

A new species of the *Cyrtodactylus brevipalmatus* group (Squamata, Gekkonidae) from the uplands of western Thailand

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Abstract

An integrative systematic analysis recovered a new species of the *Cyrtodactylus brevipalmatus* group from the uplands of Thong Pha Phum National Park, Kanchanaburi Province in western Thailand. *Cyrtodactylus thongphaphumensis* sp. nov. is deeply embedded within the *brevipalmatus* group, bearing an uncorrected pairwise sequence divergence of 7.6–22.3% from all other species based on a 1,386 base pair segment of the mitochondrial NADH dehydrogenase subunit 2 gene (ND2) and adjacent tRNAs. It is diagnosable from all other species in the *brevipalmatus* group by statistically significant mean differences in meristic and normalized morphometric characters as well as differences in categorical morphology. A multiple factor analysis recovered its unique and non-overlapping placement in morphospace as statistically significantly different from that of all other species in the *brevipalmatus* group. The description of this new species contributes to a growing body of literature underscoring the high degree of herpetological diversity and endemism across the sky-island archipelagos of upland montane tropical forest habitats in Thailand, which like all other upland tropical landscapes, are becoming some of the most imperiled ecosystems on the planet.

Keywords

Bent-toed gecko, genetics, Indochina, integrative taxonomy, montane forests, morphology

Introduction

The gekkonid genus *Cyrtodactylus* Gray, 1827 contains well over 350 named and unnamed species and constitutes the third largest vertebrate genus on the planet (Grismer et al. 2021a, b; Uetz et al. 2022). To date, its extensive distribution extends across at least eight biogeographic regions and crosses a number of well-established biogeographic barriers from South Asia to western Melanesia (Grismer et al. 2022a). The ecological plasticity, phylogenetic relationships, and geographic distribution among, and within its 32 geographically circumscribed monophyletic species groups, are indicative of its ability to disperse across ephemeral seaways, major river systems, basins, mountain ranges, and land bridges, followed by extensive in situ diversification within specific geographic areas (Grismer et al. 2020, 2021a, b, 2022a).

Within Indochina and northern Sundaland, the *Cyrtodactylus brevipalmatus* group is one of the most ecologically and morphologically specialized groups within *Cyrtodactylus* (sec. Grismer et al. 2020, 2021a, b). All members bear a similar morphology, behavior, and color pattern adapted to an arboreal life style (Grismer et al. 2022b). The latest phylogenetic taxonomic treatment of the group (Grismer et al. 2022c) described four new species from Thailand, resulting in ten described and potentially as many undescribed populations needing further study. One of these undescribed populations, *C. sp.* 9 from Thong Pha Phum National Park, Kanchanaburi Province in western Thailand (Fig. 1), was first recognized on the basis of molecular phylogenetic evidence from a single specimen (Chomdej et al. 2021). We collected and sequenced eight additional specimens which corroborate the results of Chomdej et al. (2021) in that all eight specimens plus the specimen of Chomdej et al. (2021) form a monophyletic lineage deeply nested within the *brevipalmatus* group (Grismer et al. 2022c). Univariate and multivariate analyses of the eight new specimens recovered statistically significant morphological and morphospatial differences from all other members of the group which unequivocally indicate that it requires species-level recognition (Grismer et al. 2022c). As such, it is described herein.

Materials and methods

Genetic data

Methods for DNA extraction, sequencing, and editing followed Grismer et al. (2021c) and resulted in a 1,386 base pair segment of the mitochondrial NADH dehydrogenase subunit 2 gene (ND2) and adjacent tRNAs. All material examined is listed in Grismer et al. (2022c: table 1) along with GenBank accession numbers.

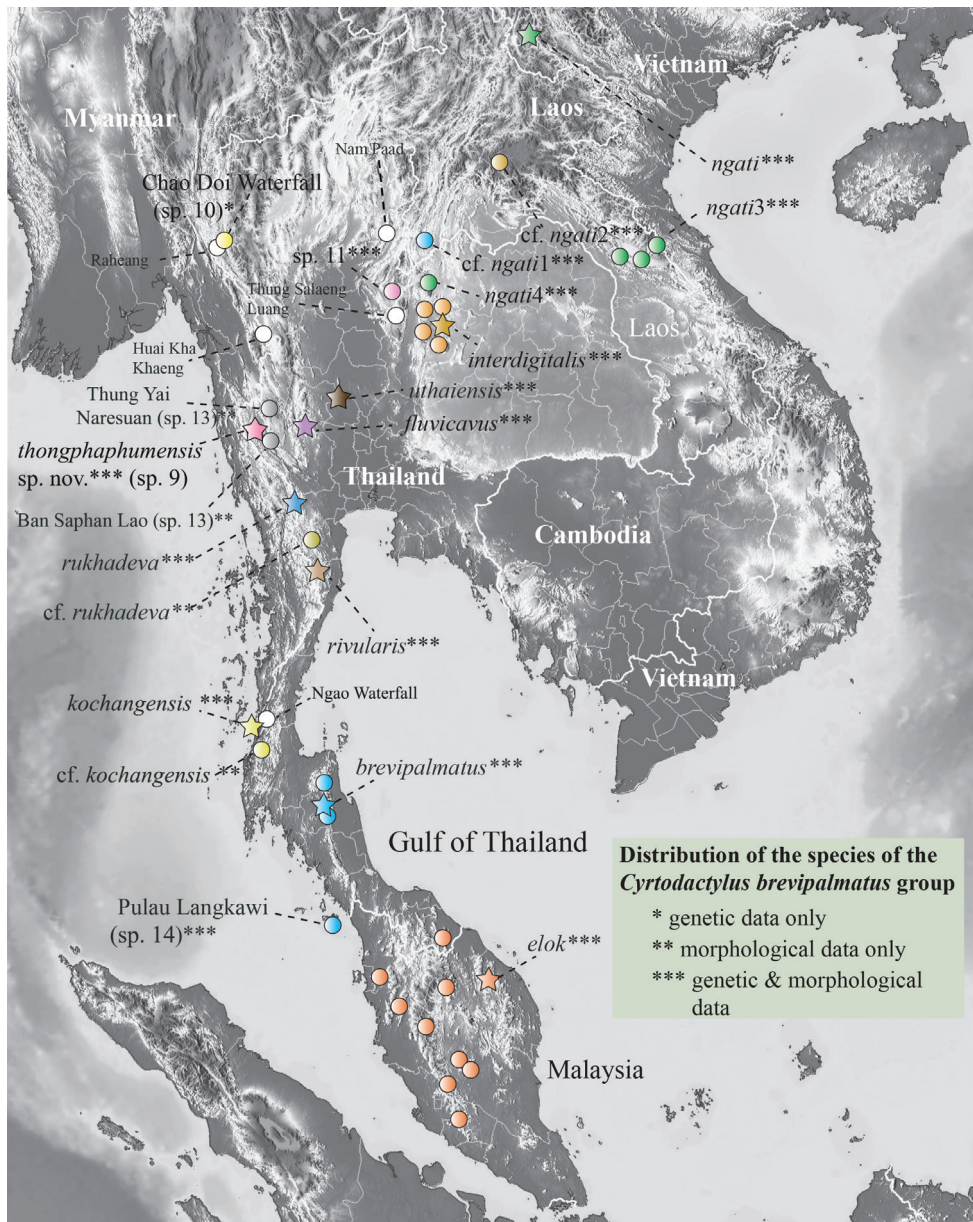


Figure 1. Distribution of nominal species and unnamed populations of the *Cyrtodactylus brevipalmatus* group. Stars denote type localities. White circles are literature localities from which specimens were not examined and remain unidentified. Locality data for all material examined is in Grismer et al. (2022c: table 1).

Morphological data

The morphological data taken included 17 meristic, 18 normalized morphometric, and eight categorical characters (Grismer et al. 2022c) (Table 1). Normalization of the morphometric characters followed the method of Chan and Grismer (2022).

Table 1. Descriptions of morphometric, meristic, and categorical characters.

Abbreviations	Characters
Morphometric characters	
SVL	snout-vent length, taken from the tip of the snout to the vent
TL	tail length, taken from the vent to the tip of the tail—original or partially regenerated
TW	tail width, taken at the base of the tail immediately posterior to the postcloacal swelling
HumL	humeral length, taken from the proximal end of the humerus at its insertion point in the glenoid fossa to the distal margin of the elbow while flexed 90°
ForL	forearm length, taken on the ventral surface from the posterior margin of the elbow while flexed 90° to the inflection of the flexed wrist
FemL	femur length, taken from the proximal end of the femur at its insertion point in the acetabulum to the distal margin of the knee while flexed 90°
TibL	tibia length, taken on the ventral surface from the posterior margin of the knee while flexed 90° to the base of the heel
AG	axilla to groin length, taken from the posterior margin of the forelimb at its insertion point on the body to the anterior margin of the hind limb at its insertion point on the body
HL	head length, the distance from the posterior margin of the retroarticular process of the lower jaw to the tip of the snout
HW	head width, measured at the angle of the jaws
HD	head depth, the maximum height of head measured from the occiput to base of the lower jaw posterior to the eyes
ED	eye diameter, the greatest horizontal diameter of the eye-ball
EE	eye to ear distance, measured from the anterior edge of the ear opening to the posterior edge of the bony orbit
ES	eye to snout distance or snout length, measured from anteriomost margin of the bony orbit to the tip of snout
EN	eye to nostril distance, measured from the anterior margin of the bony orbit to the posterior margin of the external nares
IO	interorbital distance, measured between the dorsomedial-most edges of the bony orbits
IN	internarial distance, measured between the external nares across the rostrum
EL	ear length, greatest oblique length across the auditory meatus.
Meristic characters	
SL	supralabial scales, counted from the largest scale at the corner of the mouth or posterior to the eye, to the rostral scale
IL	infralabial scales, counted from termination of enlarged scales at the corner of the mouth to the mental scale
PVT	paravertebral tubercles between the limb insertions, counted in a straight line immediately left of the vertebral column
LRT	longitudinal rows of body tubercles, counted transversely across the body midway between the limb insertions from one ventrolateral body fold to the other
VS	longitudinal rows of ventral scales, counted transversely across the abdomen midway between limb insertions from one ventrolateral fold to the other
VSM	transverse rows of ventral scales, counted along the midline of the body from the postmentals to just anterior to the cloacal opening, stopping where the scales become granular
TL4E	expanded subdigital lamellae on the fourth toe proximal to the digital inflection, counted from the base of the first phalanx where it contacts the body of the foot to the largest scale on the digital inflection—the large contiguous scales on the palmar and plantar surfaces were not counted
TL4U	small, generally unmodified subdigital lamellae distal to the digital inflection on the fourth toe, counted from the digital inflection to the claw including the claw sheath
TL4T	total number of subdigital lamellae beneath the fourth toe, $TL4E + TL4U = TL4T$
FL4E	number of expanded subdigital lamellae on the fourth finger proximal to the digital inflection, counted the same way as with TL4E
FL4U	small generally unmodified subdigital lamellae distal to the digital inflection on the fourth finger, counted the same way as with TL4U
FL4T	total number of subdigital lamellae beneath the fourth toe, $FL4E + FL4U = FL4T$
FS	enlarged femoral scales, counted from each thigh and combined as a single metric
PCS	enlarged precloacal scales, counted as a single metric
PP	number of precloacal pores in males, counted as a single metric
FP	femoral pores in males, counted from each thigh and combined as a single metric
BB	number of dark body bands, counted from between the dark band on the nape and the hind limb insertions on the body
Categorical characters	
FKT	tubercles on the flanks (present or absent)
SC1	slightly enlarged medial subcaudals (present or absent)
SC2	single distinctly enlarged, unmodified, row of medial subcaudal scales (present or absent)
SC3	enlarged medial subcaudals intermittent, medially furrowed, posteriorly emarginated (yes or no)
DCT	dorsolateral caudal tubercles (small or large)
VLF1	DCT forming a ventrolateral caudal fringe (narrow or wide)
VLF2	ventrolateral caudal fringe scales generally homogenous or not (yes or no)
TLcross	cross-section of the tail (round or square)

Phylogenetic analyses

Following Grismer et al. (2022c), an input file implemented in BEAUti (Bayesian Evolutionary Analysis Utility) v. 2.4.6 was run in BEAST (Bayesian Evolutionary Analysis Sampling Trees) v. 2.4.6 (Drummond et al. 2012) on CIPRES (Cyberinfrastructure for Phylogenetic Research; Miller et al. 2010) in order to generate a BEAST phylogeny, employing a lognormal relaxed clock with unlinked site models and linked trees and clock models. bModelTest (Bouckaert and Drummond 2017), implemented in BEAST, was used to numerically integrate over the uncertainty of substitution models while simultaneously estimating phylogeny using Markov chain Monte Carlo (MCMC). MCMC chains were run using a Yule prior for 40,000,000 million generations and logged every 4,000 generations. The BEAST log file was visualized in Tracer v. 1.7.0 (Rambaut et al. 2018) to ensure effective sample sizes (ESS) were well-above 200 for all parameters. A maximum clade credibility tree using mean heights at the nodes was generated using TreeAnnotator v. 1.8.0 (Rambaut and Drummond 2013) with a burn-in of 1,000 trees (10%). Nodes with Bayesian posterior probabilities (BPP) of 0.95 and above were considered strongly supported (Huelsenbeck et al. 2001; Wilcox et al. 2002). Uncorrected pairwise sequence divergences were calculated in MEGA 11 (Tamura et al. 2021) using the complete deletion option to remove gaps and missing data from the alignment prior to analysis.

Statistical analyses

All statistical analyses were conducted using R Core Team (2018). A Levene's test for the normalized morphometric and meristic characters was conducted to test for equal variances across all groups. Characters with equal variances ($F \geq 0.05$) were analyzed by an analysis of variance (ANOVA) and TukeyHSD post hoc test. Those with unequal variances ($F < 0.05$) were subjected to Welch's F-test and Games-Howell *post hoc* test.

Morphospatial clustering and positioning among the species was analyzed using multiple factor analysis (MFA) on a concatenated data set comprised of 38 characters including non-metric categorical characters which cannot be used in a principal component analysis (Suppl. material 1). The MFA was implemented using the mfa() command in the R package FactorMineR (Husson et al. 2017) and visualized using the Factoextra package (Kassambara and Mundt 2017). A non-parametric permutation multivariate analysis of variance (PERMANOVA) from the vegan package 2.5–3 in R (Oksanen et al. 2020) was used to determine the statistical significance of centroid locations and group clustering. The analysis used a Euclidean (dis)similarity matrix with 50,000 permutations based on the loadings of the first four dimensions recovered from the MFA. The highly morphologically derived *Cyrtodactylus elok* was not included so as to prevent biasing the morphospatial relationships among the other species (see Grismer et al. 2022b).

Results

Phylogenetic analysis

The BEAST analysis recovered the Thong Pha Phum population as being deeply embedded within the *brevipalmatus* group and the strongly supported (1.00) sister lineage to two sister groups composed of (1) *C. interdigitalis*, *C. uthaiensis*, and *C. sp. 11* and (2) *C. cf. ngati1*, *C. cf. ngati2*, *C. ngati3*, *C. ngati4*, and *C. ngati* (Fig. 2). The uncorrected pairwise sequence divergence between the Thong Pha Phum population and all other species of the *brevipalmatus* group ranges from 7.6–22.3%. (Table 2).

Statistical analyses

The ANOVA and TukeyHSD *post hoc* and Welch's F-test and Games-Howell *post hoc* tests of the adjusted morphometric and meristic characters were consistent with the phylogenetic and pairwise distance data in recovering a number of sta-

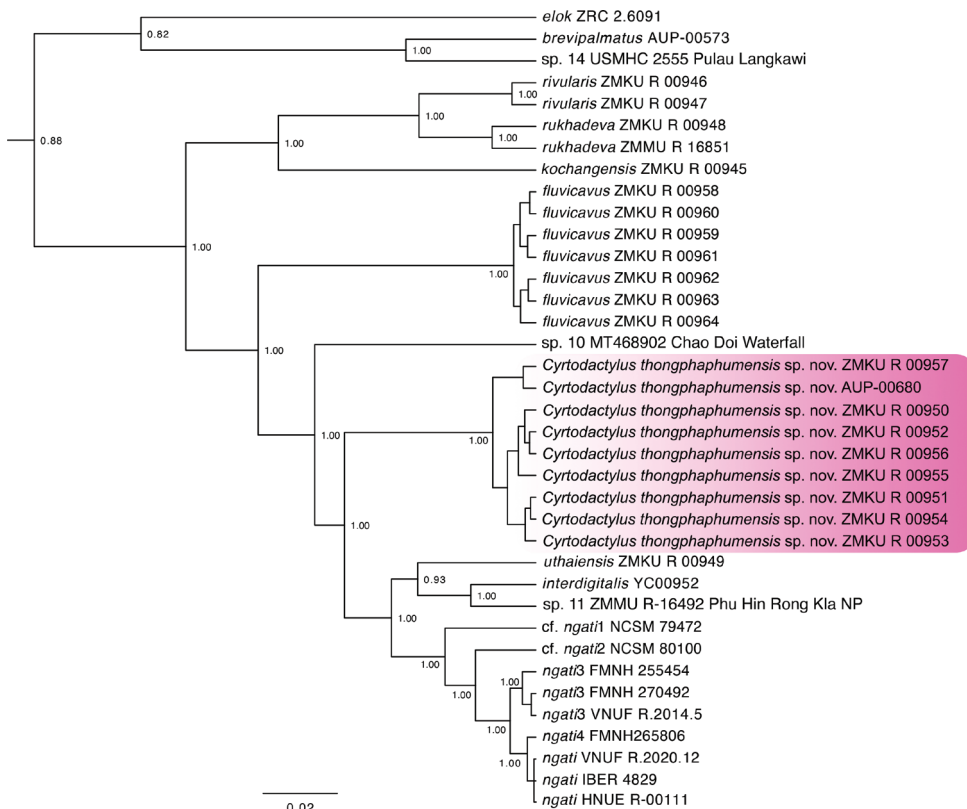


Figure 2. Maximum clade credibility BEAST phylogeny of the *Cyrtodactylus brevipalmatus* group highlighting the new species described herein. Bayesian posterior probabilities (BPP) are listed at the nodes.

Table 2. Mean (minimum–maximum) percentages of uncorrected pairwise sequence divergence (*p*-distances) among the putative species of the *Cyrtodactylus brevipalmatus* group based on 1,386 base pairs of mitochondrial NADH dehydrogenase subunit 2 gene (ND2) and adjacent tRNAs. Intraspecific p-distance are in bold font. n/a = data not applicable.

Species	1. <i>C. brevipalmatus</i>	2. <i>C. cf. ngatii</i>	3. <i>C. cf. ngatii</i>	4. <i>C. elok</i>	5. <i>C. flavigaster</i>	6. <i>C. interdigitalis</i>	7. <i>C. kochianguensis</i>	8. <i>C. ngatii</i> ; C. <i>ngatii</i> 3 and C. <i>ngatii</i> 4	9. <i>C. riedelii</i>	10. <i>C. rukhadewa</i>	11. <i>C. thongphaphumensis</i>	12. <i>C. sp. nov.</i>	13. <i>C. sp. nov.</i>	14. <i>C. sp. nov.</i>	15. <i>C. utthaiensis</i>
N	1	1	1	1	1	7	1	1	7	2	2	9	1	1	1
1.	n/a														
2.	21.03	n/a													
3.	21.68	4.39	n/a												
4.	20.77	22.58	21.42	n/a											
5.	18.86 (18.84–18.97)	10.64 (10.58–10.84)	11.02 (10.97–11.23)	20.15 (20.13–20.26)	0.10 (0.00–0.26)										
6.	20.77	6.97	9.16	22.84	12.02 (12.00–12.13)	n/a									
7.	19.35	14.58	14.71	20.90	12.31 (12.26–12.31)	15.23	n/a								
8.	20.70 (20.65–20.90)	3.30 (2.84–4.00)	3.71 (3.35–4.26)	21.11 (20.90–21.42)	11.34 (11.10–11.87)	8.13 (7.74–8.65)	14.58 (14.45–14.84)	0.84 (0.00–1.55)							
9.	20.00	15.87	15.03	21.61	12.57	15.48	12.26	15.03	0.52						
10.	(19.74–20.26) (15.61–16.13)	(14.84–15.23)	(21.42–21.81)	(12.26–13.03)	(15.23–15.74)	(12.00–12.52)	(14.71–15.48)								1.55
11.	20.65 (20.13–21.16)	15.42 (14.86–16.00)	14.84–15.48 (14.86–16.13)	21.61 (21.16–22.06)	12.25 (11.61–13.03)	16.00 (15.35–16.65)	13.10 (12.52–13.68)	15.23 (14.19–16.23)							
12.	19.87	7.93 (7.74–8.00)	9.51 (9.42–9.55)	22.02 (21.81–22.32)	9.75 (9.55–9.94)	8.96 (8.77–9.03)	13.22 (13.03–13.29)	8.81 (8.13–9.68)							
13.	20.39	7.23	8.90	22.19	11.12 (11.10–11.23)	3.87	14.58								
14.	6.45	20.90	20.65	20.00	18.34 (18.32–18.45)	20.13	19.10	20.52 (20.26–20.65)	19.74 (19.48–20.00)	20.00 (19.48–20.52)	19.60 (19.48–19.87)	18.84 (19.48–20.52)	19.61 n/a		
15.	19.74	5.81	8.13	21.16	10.12 (10.06–10.32)	7.1	13.94 (6.58–7.61)	6.97 (13.68–14.19)	13.94 (13.29–14.58)	7.80 (7.61–7.87)	8.39 (7.61–7.87)	6.58 (7.61–7.87)	19.48 n/a		

Table 3. Significant *p*-values from the results of the ANOVA and Welch's *F* (*) analyses comparing the normalized morphometric and meristic characters of *Cyrtodactylus thongphaphumensis* sp. nov. to other species of the *Cyrtodactylus brevipalmatus* group. Only species with and $N > 2$ are included. No significant differences were recovered for SVL. Abbreviations are in the Materials and methods.

Morphometric characters	AG*	HumL*	ForL	FemL	TibL	HL	HW	HD*	ED*	EE*	ES	EN*	IO	EL	IN
<i>C. brevipalmatus</i>						0.01	< 0.001			< 0.001	0.03				
<i>C. fluvicavus</i>			0.0					0.007			0.013	0.023		0.007	
<i>C. interdigitalis</i>					0.00		0.007								
<i>C. ngati</i>	< 0.001						< 0.001	0.042			0.007	< 0.001		< 0.001	0.000
<i>C. ngati</i> 3	0.001					0.01	0.03	0.003	0.043		0.001	0.019	0.019		0.003
<i>C. rukhadeva</i>	0.02	0.004				0.02		0.033							
Meristic characters	SL	IL*	PVT*	LRT	VS	VSM	TL4E	TL4T	FL4E	FL4U*	FL4T*	FS	PCS*	BB*	
<i>C. brevipalmatus</i>					< 0.001	0.003				0.022			< 0.001	0.05	
<i>C. fluvicavus</i>					< 0.001	< 0.001				0.001	0.004	0.020			
<i>C. interdigitalis</i>					0.003		< 0.001	0.005		0.043		0.01		< 0.001	
<i>C. ngati</i>	0.003				0.016	0.011			0.000	< 0.001	< 0.001		< 0.001	< 0.001	
<i>C. ngati</i> 3					0.001	0.042				< 0.001	0.001				
<i>C. rukhadeva</i>	0.029				< 0.001		0.002						0.001		

Table 4. Summary statistics from the PERMANOVA analysis from the loadings of dimension 1–4 of the MFA comparing *Cyrtodactylus thongphaphumensis* sp. nov. to all other species the *Cyrtodactylus brevipalmatus* group with sample sizes > 1 . Bold fonts denote significant differences.

OTU pairs	F model	R ²	p-value	p-adjusted
<i>C. rukhadeva</i>	88.504	0.847	0.000	0.001
<i>C. cf. ngati</i> 2	56.471	0.876	0.020	1.000
<i>C. ngati</i> 3	59.321	0.868	0.006	0.324
<i>C. interdigitalis</i>	85.773	0.896	0.002	0.112
<i>C. ngati</i>	134.367	0.937	0.006	0.332
<i>C. brevipalmatus</i>	80.229	0.879	0.001	0.025
<i>C. fluvicavus</i>	55.127	0.809	0.000	0.008
<i>C. rivularis</i>	9.485	0.542	0.022	1.000
<i>C. sp. 13</i>	30.716	0.793	0.022	1.000

tistically significant differences between the Thong Pha Phum population and all other species (Table 3). Thong Pha Phum population plotted separately in the MFA with meristic data contributing 16.5% of the inertia in dimension 1, categorical morphology contributing 15.3% of the inertia in dimension 2, and normalized morphometric data contributing 13.6% of the inertia in dimension 3 (Fig. 3). The PERMANOVA analysis recovered the morphospatial position of the Thong Pha Phum population as being statistically different from *C. brevipalmatus*, *C. cf. ngati*2, *C. ngati*3, *C. ngati*, *C. fluvicavus*, *C. interdigitalis*, *C. rivularis*, *C. rukhadeva*, and *Cyrtodactylus* sp. 13 (Table 4).

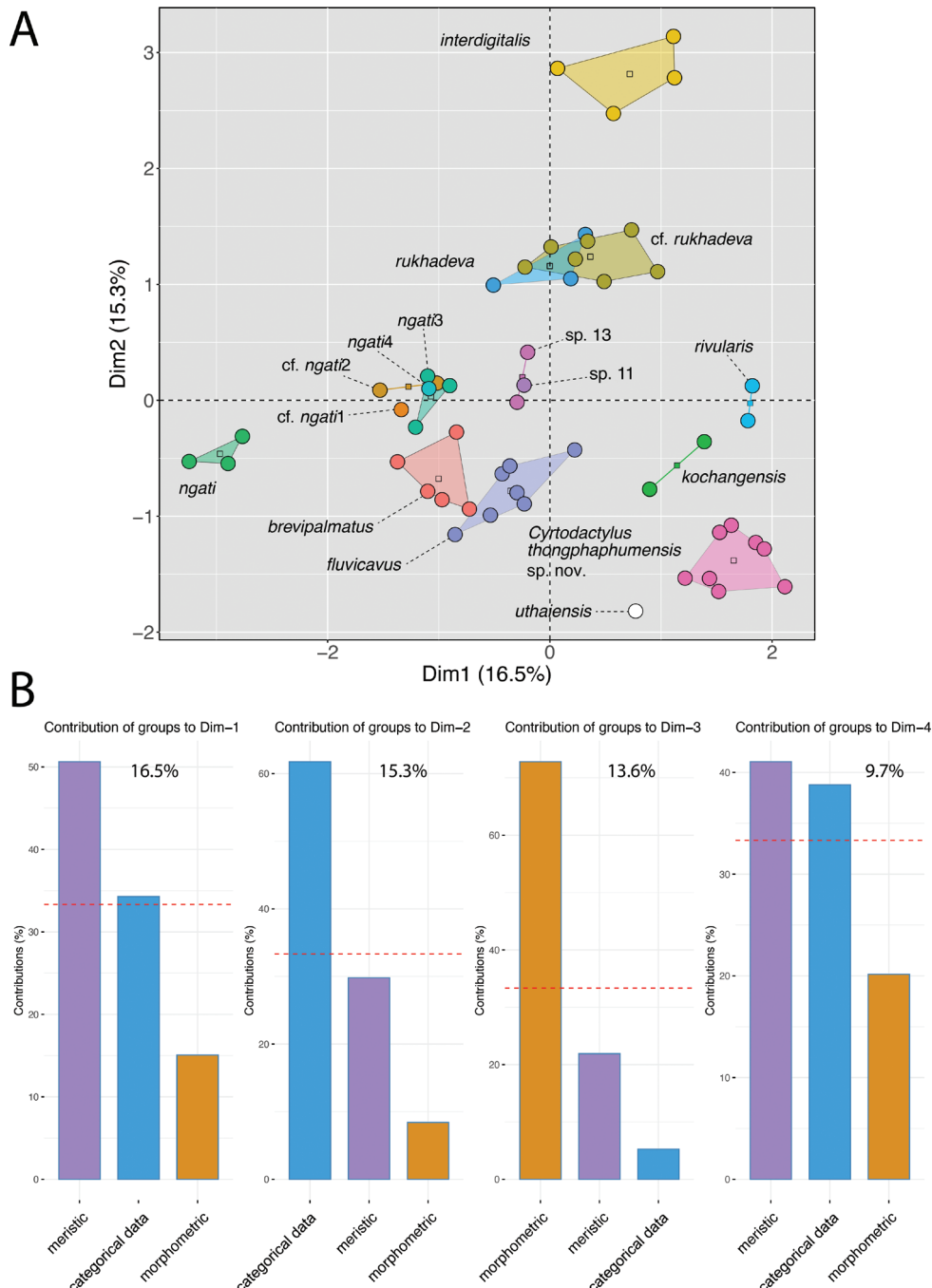


Figure 3. **A** MFA of the species-level lineages based on the BEAST phylogeny (Fig. 2) **B** Percent contributions of each data type to the inertia of dimensions 1–4 of the MFA. Percentage values on the bar graphs are the amounts of inertia for the respective dimensions.

Taxonomy

Given the phylogenetic delimitation of the Thong Pha Phum population (Fig. 2), its statistically significant diagnostic morphological differences (Table 3), its statistically significant diagnostic placement in morphospace (Fig. 3, Table 4), and its notable difference in pairwise sequence divergence from all other species (Table 2), we describe it below as new species.

Cyrtodactylus thongphaphumensis sp. nov.

<https://zoobank.org/4BB0E9B3-1BFF-49BC-BF77-79BF8CC95D27>

Suggested Common Name: Thong Pha Phum Bent-toed Gecko

Figs 4, 5

Cyrtodactylus sp. 9 Chomdej et al. 2021: 2; Grismer et al. 2022b: 248; Grismer et al. 2022c: 115.

Type material. **Holotype.** Adult male ZMKU R 00953 from Thong Pha Phum National Park, Pilok Subdistrict, Thong Pha Phum District, Kanchanaburi Province, Thailand (14.69339°N, 98.40534°E, 914 m a.s.l.), collected by Korkhwan Termprayoon, Akrachai Aksornneam, Natee Ampai, and Siriporn Yodthong on 8 April 2019.

Paratypes. Adult males ZMKU R 00951, ZMKU R 00954 and ZMKU R 00956 and adult females ZMKU R 00950, ZMKU R 00952, ZMKU R 00955, and ZMKU R 00957 bear the same collection data as the holotype.

Diagnosis. *Cyrtodactylus thongphaphumensis* sp. nov. can be separated from all other species of the *brevipalmatus* group by the combination of having 12–14 supralabials, 8–10 infralabials, 30–36 paravertebral tubercles, 19–21 rows of longitudinally arranged tubercles, 30–34 longitudinal rows of ventrals, 150–173 transverse rows of ventrals, 8–10 expanded subdigital lamellae on the fourth toe, 11–14 unexpanded subdigital lamellae on the fourth toe, 20–24 total subdigital lamellae on the fourth toe; seven or eight expanded subdigital lamellae on the fourth finger, 10–12 unexpanded subdigital lamellae on the fourth finger, 18–20 total subdigital lamellae on the fourth finger; 12–16 total number of enlarged femoral scales, 12–16 total number of femoral pores in males; 15 precloacal pores in males; 15–17 enlarged precloacals; enlarged femorals and enlarged precloacals not continuous; proximal femorals smaller than distal femorals; small tubercles on forelimbs and flanks; large dorsolateral caudal tubercles and wide ventrolateral caudal fringe; ventrolateral caudal fringe composed scales of different size; tail square in cross-section; maximum SVL 76.6 mm; 3–5 dark transverse body bands (Table 5).

Description of holotype (Fig. 4). Adult male SVL 73.2 mm; head moderate in length (HL/SVL 0.27), width (HW/HL 0.70), depth (HD/HL 0.39), distinct from neck, triangular in dorsal profile; lores concave slightly anteriorly, weakly inflated posteriorly; prefrontal region concave; canthus rostralis rounded; snout elongate (ES/HL 0.40), rounded in dorsal profile; eye large (ED/HL 0.25); ear opening horizontally

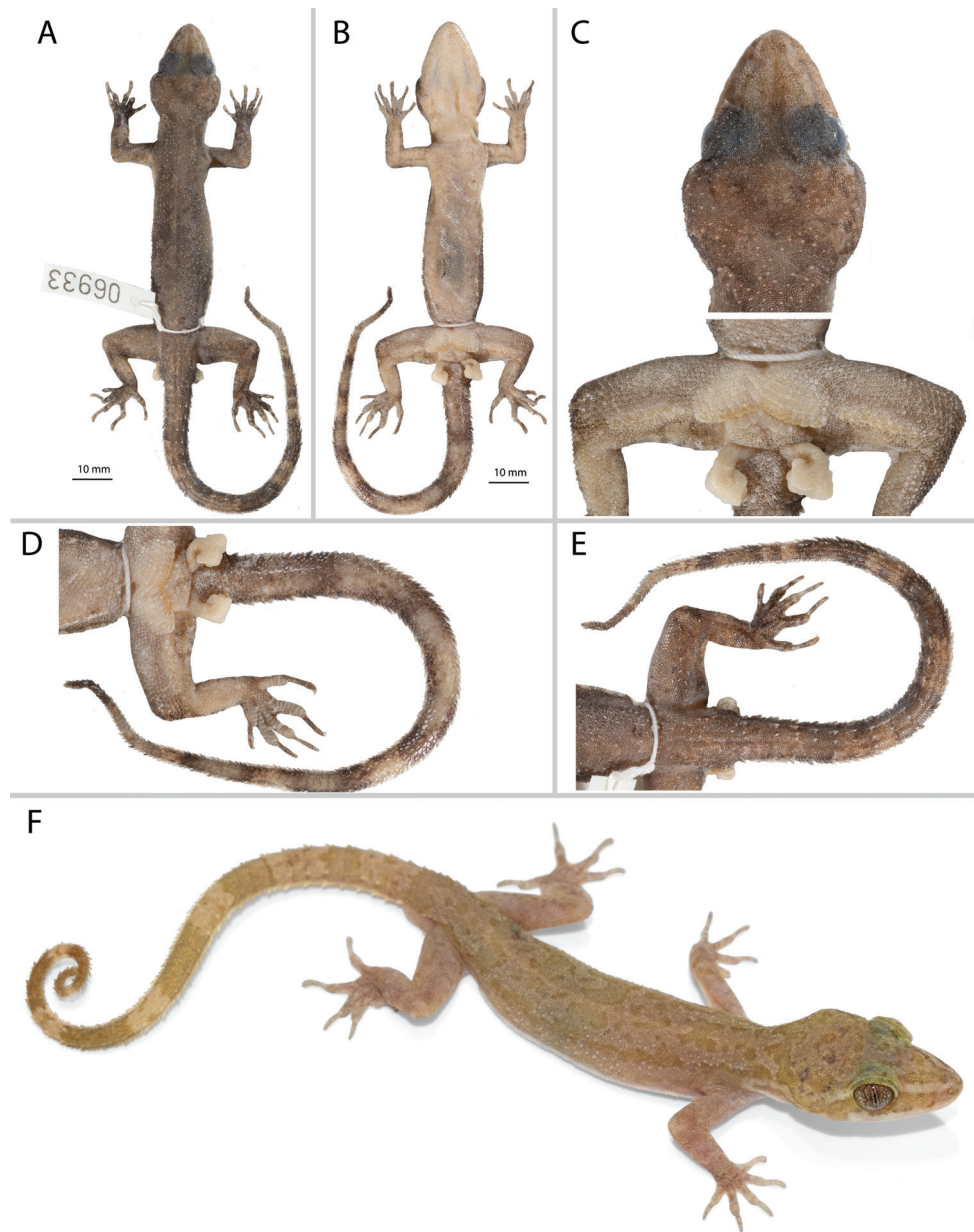


Figure 4. Adult male holotype of *Cyrtodactylus thongphaphumensis* sp. nov. ZMKU R 00953 (field no. AA 06933) from Thong Pha Phum National Park, Piloc Subdistrict, Thong Pha Phum District, Kanchanaburi Province, Thailand. **A** dorsal view **B** ventral view **C** dorsal view of head and ventral view of pelvic region **D** dorsal view of tail and **E** ventral view of tail in preservative **F** holotype in life.

elliptical, small; eye to ear distance greater than diameter of eye; rostral rectangular, divided by a dorsal furrow, bordered posteriorly by large left and right supranasals and one small azygous internasal, bordered laterally by first supralabials; external nares bor-

dered anteriorly by rostral, dorsally by large supranasal, posteriorly by two unequally sized smaller postnasals, bordered ventrally by first supralabial; 14R/14L rectangular supralabials, second through eighth supralabials nearly same size as first, then tapering below eye; 10R/10L infralabials tapering smoothly to just below and slightly past posterior margin of eye; scales of rostrum and lores flat to slightly domed, larger than granular scales on top of head and occiput; scales of occiput intermixed with distinct, small tubercles; superciliaries subrectangular, largest anterodorsally; mental triangular, bordered laterally by first infralabials and posteriorly by large left and right trapezoidal postmentals contacting medially for 45% of their length posterior to mental; one row of enlarged, square to rectangular sublabials extending posteriorly to sixth(L) and fifth(R) infralabial; gular and throat scales small, granular, grading posteriorly into slightly larger, flatter, smooth, imbricate, pectoral and ventral scales.

Body relatively short (AG/SVL 0.46) with well-defined ventrolateral folds; dorsal scales small, granular interspersed with larger, conical, semi-regularly arranged, weakly keeled tubercles; tubercles extend from occipital region onto base of tail and slightly beyond as paravertebral rows; smaller tubercles extend anteriorly onto nape and occiput, diminishing in size anteriorly; approximately 20 longitudinal rows of tubercles at midbody; approximately 34 paravertebral tubercles; tubercles on flanks; 34 longitudinal rows of flat, imbricate, ventral scales much larger than dorsal scales; 166 transverse rows of ventral scales; 15 large, pore-bearing, precloacal scales; no deep precloacal groove or depression; and two rows of enlarged post-precloacal scales on midline.

Forelimbs moderate in stature, relatively short (ForL/SVL 0.13); granular scales of forearm larger than those on body, interspersed with large flat tubercles; palmar scales rounded, slightly raised; digits well-developed, relatively short, inflected at basal interphalangeal joints; digits narrower distal to inflections; subdigital lamellae wide, transversely expanded proximal to joint inflections, narrower transverse lamellae distal to joint inflections; claws well-developed, claw base sheathed by a dorsal and ventral scale; 8R/8L expanded and 11R/11L unexpanded lamellae beneath the fourth finger; hind limbs larger and thicker than forelimbs, moderate in length (TibL/SVL 0.14), covered dorsally by granular scales interspersed with moderately sized, conical tubercles dorsally and posteriorly and anteriorly by flat, slightly larger, subimbricate scales; ventral scales of thigh flat, imbricate, larger than dorsals; subtibial scales flat, imbricate; one row of 6R/8L enlarged pore-bearing femoral scales not continuous with enlarged pore-bearing precloacal scales, terminating distally at knee; 7R/8L enlarged femoral scales; proximal femoral scales smaller than distal femorals, the former forming an abrupt union with much smaller, rounded, ventral scales of posteroventral margin of thigh; plantar scales flat, subimbricate; digits relatively long, well-developed, inflected at basal interphalangeal joints; 8R/8L wide, transversely expanded subdigital lamellae on fourth toe proximal to joint inflection extending onto sole, and 12R/12L unexpanded lamellae beneath the fourth toe distal to joint inflection; and claws well-developed, claw base sheathed by a dorsal and ventral scale.

Tail original, 94.6 mm long (TL/SVL 1.29), 5.0 mm in width at base, tapering to a point; nearly square in cross-section; dorsal scales flat, intermixed with tubercles



Figure 5. Paratypes of *Cyrtodactylus thongphaphumensis* sp. nov. in preservative from Thong Pha Phum National Park, Pilok Subdistrict, Thong Pha Phum District, Kanchanaburi Province, Thailand.

forming paravertebral rows anteriorly and larger tubercles forming dorsolateral longitudinal rows; large, posteriorly directed, semi-spinose tubercles forming wide ventrolateral caudal fringe; larger scales of ventrolateral fringe occur at regular intervals; medial subcaudals enlarged but not paired, an enlarged single medial subcaudal longitudinal row absent; subcaudals, larger than dorsal caudals; base of tail bearing hemipenal swellings; 3R/3L conical postcloacal tubercles at base of hemipenal swellings; and postcloacal scales flat, imbricate.

Coloration in life (Fig. 4). Ground color of the head body, limbs, and tail dull yellow; diffuse darker mottling on the top of the head; wider, pale-brown pre- and postorbital stripe extends from external nares to angle of jaw; whitish canthal and postorbital stripe dorsal to pale-brown pre- and postorbital stripe; faint, pale brown, nuchal band bearing two posteriorly directed projections; paired dark-brown paravertebral blotches on nape; four wide, irregularly shaped and broken transverse body bands edged in slightly

pale brown between limb insertions; band interspaces bearing irregularly shaped scattered pale-brown markings; very faint pale-brown speckling on limbs and digits; seven wide pale-brown caudal bands separated by seven paler colored bands; posterior five pale-brown caudal bands encircle tail; ventral surfaces of body and limbs beige, generally immaculate, subcaudal region generally darker; iris orange-gold in color bearing black vermiculations.

Variation (Fig. 5, Table 5). Individuals of the type series are very similar in overall coloration and pattern. TL and TW of complete original tails (ZMKU R 00951–00952, ZMKU R 00954, ZMKU R 00957) are 80.1–94.7 mm (mean 89.1 ± 6.5 mm; $N = 4$) and 4.2–4.9 mm (mean 4.7 ± 0.3 ; $N = 4$), respectively. ZMKU R 00956 has a short, partially regenerated tail which lacks banding (TL 27.7 mm, TW 5.1 mm). Similarly, the posterior sections of the tails in ZMKU R 00950 (TL 75.5 mm, TW 5.0 mm) and ZMKU R 00955 (TL 73.3 mm, TW 4.7 mm) are regenerated. Specimens ZMKU R 00950, ZMKU R 00952, and ZMKU R 00954 have three as opposed to four body bands in the holotype and ZMKU R 00955 has five body bands. Raw morphometric and meristic differences within and among all species of the *brevipalmatus* group are listed in Table 5.

Distribution. *Cyrtodactylus thongphaphumensis* sp. nov. is currently known only from the type locality at Thong Pha Phum National Park, Pilok Subdistrict, Thong Pha Phum District, Kanchanaburi Province, Thailand (Fig. 1).

Etymology. The specific epithet *thongphaphumensis* is in reference to the type locality of Thong Pha Phum National Park.

Comparisons. *Cyrtodactylus thongphaphumensis* sp. nov. is the sister species to a clade composed of eight lineages in the phylogenetic sequence of *C. uthaiensis*, sp. 11, *C. interdigitalis*, *C. cf. ngati1*, *C. cf. ngati2*, *C. ngati3*, and the sister lineages *C. ngati4* and *C. ngati* (Fig. 2). *Cyrtodactylus thongphaphumensis* sp. nov. differs from those lineages by an uncorrected pairwise sequence divergence of 7.6–9.7% and from all members of the *brevipalmatus* group by 7.6–22.3% (Table 2). It differs discretely from *C. elok* by having as opposed to lacking paravertebral tubercles, femoral and precloacal pores, and by having 19–21 as opposed to 4–7 longitudinal rows of tubercles. It differs from *C. brevipalmatus*, *C. fluvicavus*, *C. interdigitalis*, *C. ngati*, *C. ngati3*, and *C. rukhadeva* in having statistically significant different mean values of combinations of the morphometric characters of AG, HumL, ForL, TibL, HL, HW, HD, EE, ES, EN, EL, and IN (Table 3). It differs further from those same species in having statistically significant different mean values of combinations of the meristic characters SL, PVT, LRT, VS, VSM, TL4T, FL4E, FL4U, FL4T, FS, PCS, and BB (Table 3). Discrete differences between *Cyrtodactylus thongphaphumensis* sp. nov. and other putative species and populations are presented in Table 5.

Natural history. All individuals were found in hill evergreen forest at 914 m elevation (Fig. 6). Specimens ($N = 8$) were collected at night (1900–2100 h) during the dry season (April) on tree trunks (62.5%; $N = 5$), on a building (12.5%; $N = 1$), and the ground (25.0%; $N = 2$) with a temperature of 27.0 °C and relative humidity of 71.1%. The holotype (ZMKU R 00953) and four paratypes (ZMKU R 00950, ZMKU R 00954, ZMKU R 00956–00957) were found on tree trunks ≤ 160 cm above ground level. One specimen (ZMKU R 00951) was found on a building. Two

Table 5. Sex and raw meristic, categorical, and morphometric data used in the analyses from specimens in the *Cyrtodactylus brevipalmatus* group. Abbreviations:

R/L = right/left; / = data unavailable.

Species	<i>Cyrtodactylus thongphaphumensis</i> sp. nov.										<i>C. brevipalmatus</i>		<i>C. cf. brevipalmatus</i> (C. sp. 1.4)		<i>C. brevipalmatus</i>	
	ZMKU R 00950	ZMKU R 00951	ZMKU R 00952	ZMKU R 00953	ZMKU R 00954	ZMKU R 00955	ZMKU R 00956	ZMKU R 00957	ZMKU R 00957 paratype	ZMKU R 00957 paratype	LSUHC 1899	LSUHC 15076	LSUHC 11788	THNHM 10670	THNHM 14112	
Sex	♂	♂	♂	♂	♂	♀	♀	♂	♀	♂	♀	♀	♀	♀	♀	♀
Supralabials (SL)																
Infralabials (IL)	12	13	13	14	14	13	13	13	13	11	12	10	10	14	12	
Paravertebral tubercles (PVT)	8	8	10	10	9	10	10	9	8	10	9	11	11	11	11	
Longitudinal rows of tubercles (LRT)	32	33	34	34	36	36	36	36	30	30	39	37	38	37	37	
Ventral scales (VS)	21	19	20	20	21	21	19	19	15	15	16	17	16	16	14	
Ventral scales along middle of the body (VSM)	34	33	33	34	30	33	32	33	32	33	38	38	36	36	39	
Expanded subdigital lamellae on 4 th toe (TL4E)	173	158	166	159	159	150	169	169	176	170	182	182	154	154	160	
Unmodified subdigital lamellae on 4 th toe (TL4U)	9	10	9	8	10	8	9	9	9	7	8	9	8	8	8	
Total subdigital lamellae 4 th toe (TL4T)	12	14	13	12	13	12	11	11	13	13	11	11	11	11	12	
Expanded subdigital lamellae on 4 th finger (FL4E)	21	24	22	20	23	20	20	22	20	22	20	19	20	19	20	
Unmodified subdigital lamellae on 4 th finger (FL4U)	8	7	7	8	8	8	8	8	8	8	8	8	7	7	8	
Total subdigital lamellae 4 th finger (FL4T)	10	12	12	11	12	11	11	12	9	11	10	10	10	10	10	
Enlarged femoral scales (RL)	18	19	19	19	20	20	19	20	17	19	18	17	17	17	18	
Total enlarged femoral scales (FS)	12	16	16	15	16	15	13	16	16	16	10	11	16	16	14	
Enlarged precloacal scales (PCS)	/	16	/	14	15	/	12	/	7	/	/	/	/	/	/	
Predental pores in males (PP)	17	15	15	15	15	15	15	15	15	7	7	7	8	8	7	
Postcloacal tubercles (PCT)	/	15	/	15	15	/	15	/	7	/	/	/	/	/	/	
Body bands (BB)	2	2R/3L	3	3	2R/3L	3	2	3	2	3	2	3	3	3	3	
	3	4	3	4	3	5	4	4	4	6	3	5	5	5	5	

Species	<i>Cyrtodactylus thongphaphumensis</i> sp. nov.												<i>C. brevipalmatus</i> (C. sp. 14)	<i>C. cf. brevipalmatus</i> (C. sp. 14)
Institutional catalog number	ZMKU R.00950	ZMKU R.00951	ZMKU R.00952	ZMKU R.00953	ZMKU R.00954	ZMKU R.00955	ZMKU R.00956	ZMKU R.00957	ZMKU 1899	LSUHC 15076	LSUHC 11788	THNHM 10670	THNHM 14112	
Sex	♀	♂	♀	♂	♀	♂	♀	♂	♂	♀	♀	♀	♀	
Small tubercles on flank (FKT)														
Dorsolateral caudal tubercles (DCT)	present	present	present	present	present	present	present	present	present	present	present	present	present	present
Ventrolateral caudal fringe narrow or wide (VLF1)	large	large	large	large	large	large	large	large	large	small	small	/	small	/
Ventrolateral caudal fringe scales generally homogeneous (VLF2)	wide	wide	wide	wide	wide	wide	wide	wide	wide	narrow	narrow	/	narrow	/
Tail cross-section (TLcross)	no	no	no	no	no	no	no	no	no	no	no	no	no	no
Slightly enlarged medial subcaudals (SCI1)	square	square	square	square	square	square	square	square	square	circular	circular	circular	circular	/
Single enlarged medial subcaudal (SCI2)	present	present	present	present	present	present	present	present	present	present	present	present	present	absent
Enlarged medial subcaudals, intermittent, medially furrowed, posteriorly emarginate (SC3)	absent	absent	absent	absent	absent	absent	absent	absent	absent	absent	absent	absent	absent	/
Morphometric data														
SVL	73.1	73.5	73.7	73.2	64.4	76.6	76.6	74.2	68.8	70.8	64.1	66.0	63.8	
AG	34.8	33.9	35.4	33.6	28.5	37.1	33.2	35.1	35.7	33.4	30.1	30.0	26.5	
HumL	8.4	7.2	9.0	9.0	7.2	8.0	8.1	8.6	9.7	9.3	8.0	9.6	9.5	
ForL	9.5	9.1	9.2	9.8	9.2	10.0	8.6	9.8	9.9	9.8	8.9	8.2	8.7	
FemL	12.8	11.6	12.3	12.5	10.9	13.7	10.8	12.5	12.0	12.6	11.5	11.7	9.8	
TibL	10.5	10.1	10.6	10.6	9.9	11.1	10.0	11.4	11.6	12.2	10.5	9.7	8.2	
HL	19.9	20.9	20.1	20.0	17.6	20.4	19.3	20.0	19.3	19.3	19.0	17.9	18.2	
HW	14.5	14.3	15.7	13.9	12.8	14.7	14.4	14.1	13.2	13.8	12.3	12.3	12.0	
HD	7.8	7.7	7.9	7.7	7.0	8.2	7.8	7.6	8.0	7.6	7.6	7.3	7.0	
ED	5.0	5.1	5.0	5.0	4.8	5.6	5.3	4.9	5.2	4.5	4.3	5.3	4.4	
EE	5.9	5.9	6.0	5.9	5.3	6.1	6.0	6.0	5.7	5.9	4.9	5.7	5.7	
ES	7.9	8.5	7.9	7.9	7.3	8.2	7.9	7.9	7.4	7.6	7.0	7.0	7.2	
EN	6.0	6.1	6.0	5.8	5.4	6.1	6.0	5.9	5.7	5.4	4.9	5.3	5.4	
IO	5.4	5.5	5.8	5.5	4.9	5.7	5.6	5.3	5.4	4.7	4.7	4.2	5.2	
EL	1.1	1.5	1.5	1.2	1.2	1.0	1.2	1.3	1.0	1.4	1.1	1.3	1.0	
IN	2.3	2.4	2.2	2.0	2.0	2.3	2.2	2.2	1.7	2.1	2.3	2.1	2.2	

Species	<i>C. elok</i>						<i>C. fluvianus</i>						<i>C. interdigitatus</i>					
	Institutional catalog number	LSUHC 8238	LSUHC 12180	ZMMU R-16144	ZMKUR 00959	ZMKUR 00958	ZMKUR 00960	ZMKUR 00961	ZMKUR 00962	ZMKUR 00963	ZMKUR 00964	TINHM 20226	TINHM 20228	THNHM 20226	THNHM 20228	Paratype	Paratype	
Sex	♀	♂	♀	♂	♂	♀	♂	♀	♀	♀	♀	♀	♀	♀	♀	♀	♀	
Single enlarged medial subcaudal (SC2)	absent	absent	absent	absent	absent	absent	absent	absent	absent	absent	absent	absent	absent	absent	absent	/	/	
Enlarged medial subcaudals intermittent, medially furrowed, posteriorly emarginate (SC3)	no	no	no	no	no	no	no	no	no	no	no	no	yes	yes	/	/	/	
Morphometric data																		
SVL	80.2	78.2	84.8	78.6	72.5	72.0	69.6	68.4	76.8	65.7	78.2	81.2	74.8					
AG	39.7	37.8	41.5	36.2	33.4	32.0	30.4	35.6	30.6	38.1	34.5	33.7						
Huml.	10.2	9.1	10.1	1.7	9.1	8.8	9.0	8.0	10.0	7.5	10.1	9.8	10.2					
Forl.	11.5	11.7	11.8	10.2	10.5	10.3	10.5	10.1	11.1	8.8	10.8	10.6	10.5					
FemL	12.9	14.2	14.6	13.1	13.1	12.5	12.5	13.5	14.1	11.5	13.9	14.7	13.2					
TibL	13.5	14.0	13.8	12.3	11.3	10.6	10.2	9.9	11.2	9.4	12.3	13.1	11.9					
HL	21.8	21.6	21.9	21.7	20.1	20.5	19.7	20.1	21.2	18.6	21.3	20.8	19.9					
HW	15.6	16.1	15.9	15.1	14.0	13.4	12.9	13.0	14.9	13.0	15.4	14.0	13.4					
HD	9.6	9.8	10.4	9.8	8.5	8.1	8.3	7.9	8.1	7.8	8.3	8.4	8.6					
ED	4.8	5.0	5.7	5.0	5.0	5.0	4.9	4.7	5.1	4.5	5.3	5.3	5.5					
EE	6.4	7.1	7.0	6.8	6.5	5.9	5.7	5.8	6.1	5.4	6.5	5.8	6.2					
ES	8.6	8.7	9.5	8.6	8.5	8.3	8.2	8.1	9.2	7.3	9.3	8.3	7.8					
EN	6.0	6.2	6.5	6.2	6.5	6.2	5.9	6.1	6.6	6.5	6.5	6.0	5.5					
IO	5.7	5.4	5.4	3.9	5.5	5.4	5.3	5.1	5.6	5.0	5.6	4.8	4.7					
EL	1.9	1.4	1.5	1.4	1.4	1.5	1.7	1.4	1.8	1.6	1.8	1.3	1.3					
IN	2.7	2.6	2.5	3.1	2.3	2.4	2.5	2.3	2.3	2.3	2.6	2.1	2.2					
<i>C. cf. kochiagensis</i>																		
Species	<i>C. cf. kochiagensis</i>																	
Institutional catalog number	ZMKUR 00935	THNHM 01667	HNUE-R00111	IEBR ♂	VNUF R.2020.12	HNUE-R00112	FMINH 255454	FMINH 270493	FMINH 270492	FMINH 265806	FMINH 79472	NCSM 1-4917	ZMMU 80100	NCSM 00947	ZMKUR 80100	NCSM R 00946	<i>C. rivularis</i>	
Sex	♀	♂	♀	♀	♀	♀	♂	♂	♂	♂	♀	♀	♀	♀	♀	♀	<i>C. rivularis</i>	
Meristic data																		
Supralabials (SL)	12R/13L	12	10	10	10	10	13	13	13	10	14	9	12	13R/12L	13R/12L			
Infralabials (IL)	9R/9L	10	9	9	9	9	10	9	11	8	11	10	12	11R/10L	10R/9L			
Paravertebral tubercles (PVT)	34	29	39	40	38	40	28	27	26	27	28	32	29	34	33			
Longitudinal rows of tubercles (LRT)	14	19	18	17	22	19	18	17	19	18	24	19	20	18				
Ventral scales (VS)	35	34	38	36	35	32	37	36	33	33	36	35	34	37				
Ventral scales along middle of the body (VSM)	172	159	168	164	178	158	159	166	158	164	166	165	160	166				

Species		C.	C. d.f.	C. <i>ngati</i>		C. <i>ngati3</i>		C. <i>ngati4</i>		C. cf. <i>ngati2</i>		C. cf. <i>ngati1</i>		C. <i>ngati</i>	
Institutional catalog number	00945	<i>kohangensis</i>	<i>kohangensis</i>	HNUHM-00111	IEBR-4829	HNUF-R.2000.12	HNUF-R.00112	FMMNH-255454	FMMNH-270493	FMMNH-265806	NCSM-79472	ZMMU-R.14917	NCSM-80100	ZMKU-R.00947	ZMKU-R.9R9L
Sex	♀	♂	♂	♀	♀	♀	♀	♂	♂	♂	♀	♀	♀	♀	♀
Expanded subdigital lamellae on 4 th toe (TL4E)	9R/8L	8	8	10	9	9	10	10	8	10	9	8	10	9R/9L	♀
Unmodified subdigital lamellae on 4 th toe (TL4U)	12R/11L	13	11	10	11	10	11	11	11	11	12	10	10	13R/13L	♂
Total subdigital lamellae 4 th toe (TL4T)	21R/19L	21	13	16	17	16	21	19	21	21	18	20	22R/22L	21R/13L	
Expanded subdigital lamellae on 4 th finger (FL4E)	8R/8L	8	6	6	7	6	8	8	8	8	7	9	8R/8L	8R/8L	
Unmodified subdigital lamellae on 4 th finger (FL4U)	10R/10L	12	9	9	9	9	10	10	10	10	9	10	11R/10L	12R/12L	
Total subdigital lamellae 4 th finger (TL4T)	18R/18L	20	15	15	18	15	18	18	18	18	17	16	19	19R/18L	20R/20L
Enlarged femoral scales (RFM)	6R/6L	7R/7L	10R/0L	9R/8L	10R/9L	8R/7L	8R/9L	9R/9L	8R/8L	9R/8L	7R/8L	7R/8L	8R/8L	6R/8L	
Total enlarged femoral scales (FS)	12	14	20	17	19	17	16	17	18	16	17	15	16	17	14
Total femoral pores in males (FP)	/	14	14	/	/	/	/	14	15	13	/	/	/	/	/
Enlarged predoctal scales (PCS)	12	16	13	13	13	13	13	13	13	13	12	13	13	15	15
Predactal pores in males (PP)	/	16	/	/	/	/	/	13	13	13	/	/	/	/	/
Postdactal tubercles (PCT)	1R/1L	3	3	2	1	2	0	0	0	0	2	3	4	2R/2L	3R/3L
Body bands (BB)	5	5	6	6	6	6	6	3	4	3	3	3	3	3	4
Categorical data															
Small tubercles on flank (FT)	present	present	present	present	present	present	present	present	present	present	present	present	present	present	present
Dorsolateral caudal tubercles (DCT)	large	large	small	small	small	narrow	narrow	narrow	narrow	narrow	narrow	narrow	narrow	small	large
Entrolateral caudal fringe narrow or wide (VLF)	wide	wide	narrow	narrow	narrow	narrow	narrow	narrow	narrow	narrow	narrow	narrow	narrow	wide	wide
Entrolateral caudal fringe scales generally homogenous (MLF2)	no	no	no	no	no	no	no	yes	yes	yes	yes	yes	yes	yes	yes
Tail cross-section (TLcross)	square	/	circular	circular	circular	circular	circular	circular	circular	circular	circular	circular	circular	square	square
Slightly enlarged medial subcaudals (SC1)	present	present	present	present	present	present	present	/	present	present	present	present	present	absent	absent
Single enlarged medial subcaudal (SC2)	absent	absent	absent	absent	absent	absent	absent	/	absent	absent	absent	absent	absent	present	present
Enlarged medial subcaudals intermittently, medially furrowed, posteriorly emarginate (SC3)	no	no	no	no	no	no	no	/	no	no	no	no	no	no	no
Morphometric data															
SVL	60.1	70.2	66.5	68.1	69.3	46.6	83.6	70.2	74.1	73.8	78.0	87.1	77.7	73.9	68.1

Species	C. kochangensis	C. f. kochangensis	C. negati	C. negati	C. negati	C. negati	C. negati4	C. cf. negati	C. cf. negati2	C. rivularis					
Institutional catalog number	ZMUKR 00935	HNUE- R00111	HNUE- R00112	IEBR #829	VNUF R 2020.12	HNUE- R00112	FMNH 255454	FMNH 270493	FMNH 270492	FMNH 265806	FMNH 79472	ZMMU R-14917	NCSM 80100	ZMKUR 00947	ZMKU R 00946
Sex	♀	♂	♀	♀	♀	♀	♀	♂	♂	♂	♀	♀	♀	♀	♀
AG	29.0	31.5	28.8	29.8	30.2	19.7	41.3	35.4	37.0	31.3	38.2	41.9	36.8	34.8	33.2
Huml.	6.5	10.2	7.9	8.1	8.5	5.6	8.6	8.7	8.6	6.9	8.7	11.5	9.2	8.1	7.6
Fod.	7.6	8.6	9.2	10.0	10.1	6.5	10.2	9.3	10.4	10.0	10.3	10.4	10.7	9.7	9.1
FemL	10.4	12.1	11.5	11.5	11.5	7.6	13.7	12.7	13.0	13.1	13.1	15.2	14.2	11.4	10.4
TibL	8.4	11.8	10.8	11.1	11.8	7.8	12.5	11.8	11.2	11.1	12.8	12.6	12.7	11.2	10.3
HL	17.3	18.3	20.1	20.4	20.7	16.1	21.7	20.6	20.3	20.7	21.2	22.1	21.4	20.3	19.3
HW	11.6	12.1	12.6	12.0	11.8	8.8	13.8	12.5	13.0	12.3	12.7	14.8	13.5	14.9	13.7
HD	6.5	7.8	7.4	7.2	6.6	5.1	9.2	8.4	9.1	7.6	8.3	8.7	9.2	8.2	8.2
ED	4.2	5.2	3.8	4.1	3.4	2.6	4.9	4.9	4.9	4.8	6.5	4.6	6.0	5.8	5.6
EE	5.0	4.9	5.8	5.5	5.9	4.4	6.9	6.1	6.2	5.7	5.3	6.5	6.2	6.5	6.2
ES	6.9	7.5	7.5	7.6	6.9	5.0	9.0	8.3	8.3	8.2	8.7	8.8	8.4	8.3	7.9
EN	5.2	5.5	6.7	6.3	6.2	4.5	6.5	6.2	6.1	6.2	6.2	6.6	6.0	6.1	5.8
IO	4.2	4.0	5.6	5.4	5.6	4.2	6.6	5.6	5.4	5.1	4.9	3.5	5.7	5.8	5.5
EL	1.0	1.3	0.8	0.8	0.7	0.3	1.3	1.1	1.2	1.0	1.5	1.2	0.9	1.1	1.1
IN	1.9	2.2	2.8	2.6	2.6	2.0	2.8	2.5	2.5	2.3	2.7	2.2	2.5	2.3	2.0
<i>C. rukhadava</i>															
Species	ZMMU R-16851	ZMUKR R-16852	THNHM 00948	THNHM 24622	THNHM 24838	THNHM 03251	THNHM 03252	THNHM 03253	THNHM 03254	THNHM 01807	THNHM R-16492	THNHM 00104	THNHM 27821	THNHM 00104	<i>C.</i> sp. 13
Sex	♂	♀	♂	♂	♀	♂	♂	♂	♂	♂	♂	♂	♀	♀	♂
Meristic data															
Supralabials (SL)	11	9	14	11	13	13	11	12	13	12	11	12	15	13R/15L	
Infralabials (IL)	10	11	9	10	10	10	10	10	11	10	9	10	11	10R/11L	
Paravertebral tubercles (PVT)	27	30	26	28	27	27	30	30	26	30	33	33	29	33	
Longitudinal rows of tubercles (LRT)	19	20	19	18	19	18	18	19	19	19	18	18	20	17	
Ventral scales (VS)	34	43	38	38	36	37	37	39	35	34	35	37	36	36	
Ventral scales along middle of the body (VSM)	154	165	162	158	157	159	168	160	161	160	159	165	159		
Expanded subdigital lamellae on 4 th toe (TL4E)	9	9	9	8	9	9	10	9	10	10	9	9	7	8R/ (broken)L	
Unmodified subdigital lamellae on 4 th toe (TL4U)	11	11	12	11	13	12	12	15	13	13	10	12	12	12R/ (broken)L	
Total subdigital lamellae on 4 th toe (TL4T)	20	18	21	19	22	21	22	14	23	23	19	21	19	20	
Expanded subdigital lamellae on 4 th finger (FL4E)	9	8	8	7	8	8	8	8	8	10	8	8	8	7R/7L	

Species	Institutional catalog number	<i>C. ruhobadeva</i>												<i>C. cf. ruhobadeva</i>	C. sp. 11	C. sp. 13	C. sp. 13 <i>uhainensis</i>
		ZMMU R-16851	ZMMU R-16852	ZMMU R-00948	ZMKUR 24622	THNHM 24838	THNHM 03251	THNHM 03252	THNHM 03253	THNHM 03254	THNHM 01807	ZMMU R-16492	THNHM 27821	ZMKUR 00949			
Sex	♂	♀	♀	♂	♂	♀	♂	♂	♀	♂	♂	♂	♀	♂	♀	♂	
Unmodified subdigital lamellae on 4 th finger (HL4U)	10	9	11	10	11	10	10	10	10	12	12	9	11	10	11R/11L		
Total subdigital lamellae 4 th finger (FL4T)	19	17	19	17	17	18	18	18	20	20	20	19	19	18	18R/18L		
Enlarged femoral scales (RL)	9R/8L	8R/8L	9R/8L	9R/7L	9R/9L	9R/7L	7R/7L	6R/7L	5R/8L	7R/7L	9R/8L	9R/9L	7R/10L	8R/8L			
Total enlarged femoral scales (FS)	17	16	17	18	16	14	14	13	13	14	17	18	17	17	16		
Total femoral pores in males (FP)	17	1	1	/	14	/	12	13	/	11	13	/	/	/	12		
Enlarged precloacal scales (PCS)	17	13	15	15	15	14	13	15	15	14	13	14	13	14	16	14	
Precloacal pores in males (PP)	17	/	/	15	/	14	13	/	15	14	13	/	/	/	/	14	
Postcloacal pores (PCT)	3	2	2R/3L	3	2	3	2	3	2	3	2	3	3	3	3	3R/3L	
Body bands (BB)	3	3	3	3	3	3	4	4	4	/	5	3	3	3	/	6	
Categorical data																	
Small tubercles on flank (FKT)	present	present	present	present	present	present	present	present	present	present	present	present	present	present	present	present	present
Dorsolateral caudal tubercles (DCT)	small	small	small	small	small	small	small	small	small	small	small	large	small	small	large	large	large
Ventrolateral caudal fringe narrow or wide (V/F1)	narrow	narrow	narrow	narrow	narrow	narrow	narrow	narrow	narrow	narrow	narrow	wide	narrow	narrow	narrow	wide	wide
Ventrolateral caudal fringe scales generally homogenous (VF2)	yes	yes	yes	yes	yes	yes	yes	yes	yes	yes	yes	/	yes	yes	yes	no	no
Tail cross-section (TL cross)	square	square	square	square	square	square	square	square	square	square	square	/	square	square	circular	circular	circular
Slightly enlarged medial subcaudals (SC1)	absent	absent	absent	absent	absent	absent	absent	absent	absent	absent	absent	/	present	present	present	present	present
Single enlarged medial subcaudal (SC2)	present	present	present	present	present	present	present	present	present	present	present	/	absent	absent	absent	absent	absent
Enlarged medial subcaudals intermittent, medially furred, posteriorly emarginate (SC3)	no	no	no	no	no	no	no	no	no	no	no	no	no	no	no	yes	yes
Morphometric data																	
SVL	74.9	71.7	71.6	68.3	71.8	73.6	75.3	74.7	73.2	61.5	68.1	63.7	72.9	58.1			
AG	34.6	32.6	33.9	27.3	29.9	30.9	31.3	32.2	30.3	26.2	34.6	25.8	30.6	26.6			
HmL	10.7	10.4	7.9	9.8	8.3	12.2	11.3	11.8	11.0	10.1	10.3	7.6	10.1	7.0			
ForL	8.6	7.9	9.6	8.7	8.5	9.0	10.6	9.6	9.2	7.9	8.5	8.1	9.6	8.3			
FemL	12.6	11.8	10.5	10.8	10.9	11.5	10.2	11.9	12.1	9.5	12.6	10.7	12.8	10.0			
TibL	10.1	9.3	11.2	9.7	10.7	10.9	11.7	11.3	11.1	9.1	11.4	10.1	10.2	8.4			
HL	20.2	19.2	19.7	19.7	19.9	20.8	21.3	20.8	21.5	17.9	18.4	17.6	19.9	16.1			
HW	14.6	13.4	14.0	13.1	13.9	14.9	15.0	13.1	14.1	11.8	13.1	11.9	13.8	10.9			

Species	<i>C. rukhadzeana</i>						<i>C. cf. rukhadzeana</i>						<i>C. sp. 13</i>						<i>C. nubatensis</i>	
	ZMMU R.16951	ZMMU R.16952	ZMKUR 00948	ZMMU R.16936	ZMMU R.24622	ZMMU R.24838	THNHM 03251	THNHM 03252	THNHM 03253	THNHM 03254	THNHM 01807	THNHM R.1692	THNHM 00104	THNHM 27821	THNHM R.1692	THNHM 00104	THNHM 27821	THNHM R.1692	THNHM 00104	THNHM 27821
Sex	♂	♀	♂	♀	♂	♂	♂	♂	♂	♂	♂	♀	♂	♂	♂	♂	♂	♂	♂	♂
HD	9.2	8.5	8.3	7.3	8.9	8.2	8.2	8.1	8.9	7.5	8.3	7.7	8.4	8.4	8.4	8.4	8.4	8.4	8.4	8.4
ED	4.6	4.3	5.5	4.9	5.1	5.8	5.4	5.0	5.5	4.7	4.4	4.1	5.3	5.3	4.6	4.6	4.6	4.6	4.6	4.6
EE	6.2	6.2	5.8	5.1	6.2	5.6	5.7	5.4	6.2	4.3	6.2	4.9	6.3	6.3	4.7	4.7	4.7	4.7	4.7	4.7
ES	8.3	7.7	7.9	7.4	8.1	8.4	8.8	8.1	8.6	7.3	7.7	7.2	8.0	8.0	6.4	6.4	6.4	6.4	6.4	6.4
EN	6.3	5.7	5.8	5.4	6.0	6.2	6.4	5.8	6.2	5.3	5.5	5.6	5.9	5.9	4.9	4.9	4.9	4.9	4.9	4.9
IO	3.3	3.1	5.6	4.5	4.7	5.6	5.7	5.7	5.6	4.2	2.9	4.8	6.1	6.1	4.3	4.3	4.3	4.3	4.3	4.3
EL	1.2	1.0	1.4	1.6	1.5	1.2	1.3	1.2	1.2	0.9	0.9	1.4	1.4	1.4	1.5	1.5	1.5	1.5	1.5	1.5
IN	2.2	2.1	2.1	2.0	2.2	2.4	2.5	2.4	2.3	2.0	2.3	2.1	2.3	2.3	1.8	1.8	1.8	1.8	1.8	1.8



Figure 6. Habitat of the type locality at Thong Pha Phum National Park, Pilok Subdistrict, Thong Pha Phum District, Kanchanaburi Province, Thailand.

specimens (ZMKU R 00952, ZMKU R 00955) were found on ground. At night, the new species was found to co-occur with other gekkonid lizards, *Cyrtodactylus oldhami* (Theobald, 1876), *Gekko kaengkrachanense* (Sumontha, Pauwels, Kunya, Limlikhitakorn, Ruksue, Taokratok, Ansermet & Chanhome, 2012), and *Hemidactylus garnotii* Duméril & Bibron, 1836.

Discussion

The discovery of new populations of the *Cyrtodactylus brevipalmatus* group across the archipelago of the upland sky-island habitats in Thailand will likely be commonplace with increased field work. Many such undescribed populations have already been reported and photographed on social networking platforms and these populations will be sampled and analyzed in order to ascertain their species status. Grismer et al. (2022c) pointed out that for several years many such populations went unanalyzed and were simply placed in the synonymy of either *C. brevipalmatus* or *C. interdigitalis*, only to be elevated later to species status following data-rich phylogenetic delimitation and morphological diagnostic analyses (Grismer et al. 2021c, 2022c). This current

work not only contributes to an increased understanding of the unrealized diversity within the *brevipalmatus* group, but to a growing body of literature underscoring the high degree of herpetological diversity and endemism across a sky-island archipelago of upland montane tropical forests in Thailand (see Suwannapoom et al. 2022) which like many other upland tropical landscapes, are becoming some of the most imperiled ecosystems on the planet.

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Supplementary material I

Data frame for the multiple factor analysis of the putative species of the *Cyrtodactylus brevipalmatus* group

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Data type: morphological data

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