# Systematics, ecomorphology, cryptic speciation and biogeography of the lizard genus Tytthoscincus Linkem, Diesmos \& Brown (Squamata: Scincidae) from the skyisland archipelago of Peninsular Malaysia 

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#### Abstract

An integrative taxonomic analysis reveals a remarkable degree of cryptic speciation between sympatric, specialized, leaf-litter species in an upland clade of the scincid genus Tytthoscincus across the sky-island archipelago of Peninsular Malaysia. Bayesian inference and maximum likelihood phylogenies based on the mitochondrial gene ND2 indicate that this clade is composed of a lineage of limb, digit and tympanum-reduced, semi-fossorial species and a lineage of leaf-litter generalist bearing longer, more robust limbs and digits, and well-developed tympanae. Behavioural differences inferred from differing ecomorphologies form the basis of hypotheses proposed to account for resource partitioning at Fraser's Hill, Pahang between the sympatric leaf-litter generalist T. bukitensis and the new semi-fossorial species T. kakikecil sp. nov. However, this hypothesis does not account for the extreme case of cryptic speciation and syntopy between T. bukitensis and a new leaf-litter generalist T. martae sp. nov. at Genting Highlands, Pahang. The phylogenetic analyses also indicate there is an additional new, upland, semi-fossorial species, T. jaripendek sp. nov., from Cameron Highlands and a new lowland, leaf-litter generalist, Tytthoscincus sp., from Hulu Terengganu, Terengganu. The specific identities of the new upland populations are corroborated by morphological analyses using combinations of principal component analysis, discriminant analysis of principal components, analysis of variance, Tukey HSD (honest significant difference) tests and Welch two-way $t$-tests. The Hulu Terengganu specimen was too damaged to provide a reliable description. A biogeographic scenario inferred from a time-calibrated BEAST analysis suggests that speciation in the upland clade has been ongoing since the Upper Pliocene following repeated episodes of secondary contact associated with glacier-driven climatic oscillations generating upland forest expansion and retraction. These analyses highlight the increasing levels of herpetological diversity and endemism being discovered in the montane regions of Peninsular Malaysia and bring to light the understudied ecological complexity of Peninsular Malaysia's leaf-litter ecosystems and the growing need for its conservation.


ADDITIONAL KEYWORDS: cryptic species - ecomorphology - leaf-litter - Peninsular Malaysia - semi-fossorial - Tytthoscincus.

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## INTRODUCTION

The sky-island archipelago of Peninsular Malaysia extends across a series of mountain ranges and islands and is emerging as one of the most herpetologically rich regions on the entire Thai-Malay Peninsula. This fragmented, upland ecosystem of habitat islands above 800 m in elevation is composed of hill dipterocarp and mossy forests that have yielded 38 new species of amphibians and reptiles from five different mountain systems in the last 10 years of exploration (Grismer, Grismer \& Youmans, 2004; Grismer, 2006a, b, 2007, 2008; Grismer et al., 2008; Wood et al., 2008, 2009; Chan et al., 2009, 2014; Grismer, Norhayati \& Chan, 2009a; Grismer, Wood, Grismer, 2009c; Matsui et al., 2009; Grismer et al., 2010a, b, 2011, 2012, 2013a, b, 2014b, c, 2015a, b, 2016b; Grismer \& Chan, 2010; Loredo et al., 2013; Matsui, Belabut \& Ahmad, 2014; Grismer \& Quah, 2015; Cobos et al., 2016; Davis et al., 2016). Not only has this growing diversity increased levels of local upland endemism, it is beginning to reveal complex patterns of closely related co-distributed species showing remarkable degrees of syntopy (Grismer et al., 2013b, 2015a; Loredo et al., 2013; Grismer \& Quah, 2015). Most notable in this regard are microhabitat-restricted leaf-litter taxa such as the Reed Snakes Macrocalamus Günther and Calamaria Boie (Quah et al., in preparation) and the diminutive Forest Skinks Tytthoscincus Linkem, Diesmos, \& Brown. The latter comprise a lineage of at least 13 small [snout-vent length (SVL) $<45 \mathrm{~mm}$ ], forest floordwelling species that collectively extend from Sulawesi across Wallace's Line to the Philippines, Borneo, Java and Peninsular Malaysia (Linkem, Diesmos \& Brown, 2011; Grismer et al., 2016a). Additional molecular evidence will no doubt demonstrate that many of the diminutive skinks in Borneo, Sumatra and Thailand, currently in the genus Sphenomorphus, should be transferred into Tytthoscincus.

In general, Tytthoscincus have relatively elongate bodies bearing short limbs and digits that are inferred here to be adaptations for locomoting in leaf-litter (Van Damme, Aerts \& Vanhooydonck, 1998, 2002; Melville \& Swain, 2000; Kohlsdorf, Garland \& Navas, 2001; Herrel, Meyers \& Vanhooydonck, 2002; Siler \& Brown, 2011). Anecdotal observations in Peninsular Malaysia indicate that Tytthoscincus forage within the cover of leaf-litter during the day but are occasionally seen on the surface during periods of overcast and light precipitation (Grismer, 2011). For their size, these skinks are surprisingly swift - moving in and out of rocks and root systems into which they irretrievably retreat at the slightest provocation. Species of Tytthoscincus are not lizards one comes across easily and considerable effort (raking leaves, turning rocks and logs, planting pitfall traps, etc.) is usually necessary for the collection
of only a few samples. Their secretive nature and ability to escape capture have contributed greatly to their low numbers in collections and a general lack of knowledge concerning their natural history. This, coupled with their relatively uniform morphology and nondescript, unremarkable, dull-brown coloration, often makes it difficult to establish species boundaries using traditional morphological characters (Grismer, 2006b, 2007, 2008; Grismer et al., 2009a, b).

Recent molecular studies (Grismer et al., 2016a; Karin, Bauer \& Das, 2016) have demonstrated that Tytthoscincus is more diverse than its morphology generally indicates. Grismer et al. (2016a) noted that on the Thai-Malay Peninsula, there are two sequentially related monophyletic lineages: a lowland swamp-dwelling clade composed of two species from Peninsular Malaysia and Singapore and an endemic montane clade composed of nine upland species with insular derivatives. New molecular and morphological data presented herein based on collections of additional material from new localities in Peninsular Malaysia and from other known localities are used here to elucidate the phylogenetic and ecological complexity within the upland clade. We also attempt to establish a stable, phylogenetic classification by disentangling a currently erroneous taxonomy based solely on morphology (Grismer et al., 2009a, b). Additionally, ecomorphological evidence will be presented that we propose contributes to a newly discovered high degree of sympatry (as many as three species at some localities) within this genus based on three new upland species discovered from the Banjaran (=mountain range) Titiwangsa and one new lowland species from the Tembat Forest Reserve, Hulu Terengganu, Terengganu. Lastly, we have expanded the molecular data set to demonstrate that Indonesian species of Sphenomorphus Fitzinger previously suspected of belonging to the genus Tytthoscincus (Linkem, 2013; Karin et al., 2016) do so indeed and their formal transference is accommodated here.

## MATERIAL AND METHODS

## Phylogenetic analysis

A data set composed of 1026 bp of the mitochondrial gene NADH dehydrogenase subunit 2 gene (ND2) was constructed from 36 individuals comprising all nine species of Tytthoscincus that comprised the ingroup plus 15 individuals from 11 other species of Tytthoscincus that, along with one specimen of Larutia seribuatensis, composed the outgroups based on Linkem et al. (2011; Table 1) and Linkem (2013). Total genomic DNA was isolated from liver or skeletal muscle from specimens stored in $95 \%$ ethanol using the Qiagen DNeasy tissue kit (Valencia, CA, USA). The ND2 gene was amplified using a double-stranded polymerase

Table 1. GenBank accession numbers for ND2 sequences and catalogue numbers of the voucher specimens used in this study

| Species | Voucher | Locality | GenBank no. |
| :---: | :---: | :---: | :---: |
| Larutia seribuatensis | LSUHC 5168 | West Malaysia, Johor, Pulau Tuali | HQ906968 |
| Tytthoscincus aesculeticolus (paratype) | FMNH 239839 | East Malaysia, Sabah Sipitang District | JF498209 |
| T. aesculeticolus | SP 06913 | East Malaysia, Sabah Sipitang District | JF498208 |
| T. atrigularis | KU 315055 | Philippines, Zamboanga City 'Province' Pasaonca Natural Park | JF497971 |
| T. atrigularis | KU 315060 | Philippines, Zamboanga City 'Province' Pasaonca Natural Park | JF497972 |
| T. batupanggah | CAS 259189 | East Malaysia, Sarawak, Gunung Penrissen | KU587723 |
| T. batupanggah | ZRC 2.7123 | East Malaysia, Sarawak, Gunung Penrissen | KU587725 |
| T. hallieri | FMNH 230184 | East Malaysia, Sabah Lahad Datu District | JF497973 |
| T. leproauricularis | LSUHC 7860 | East Malaysia, Sarawak, Gunung Penrissen | KU587724 |
| T. panchorensis (holotype) | LSUHC 4801 | West Malaysia, Penang, Bukit Panchor State Park | KU872741 |
| T. parvus | RMB 4707 | Indonesia, Sulawesi | JF498213 |
| T. sananus | JAM 8829 | Indonesia, Sanana Island | KY989394 |
| T. sibuensis | LSUHC 5583 | West Malaysia, Johor, Pulau Sibu | KU872743 |
| T. cf. sibuensis | JAM 7233 | Singapore, Nee Soon Swamp | KY989395 |
| T. temmincki | BS 10263 | Indonesia, Java | KY989396 |
| T. textus | JAM 7497 | Indonesia, Sulawesi | KY989397 |
| Upland clade |  |  |  |
| T. bukitensis | LSUHC 12750 | West Malaysia, Pahang, Genting Highlands, Awana Rd. | KY989398 |
| T. bukitensis | LSUHC 12237 | West Malaysia, Pahang, Genting Highlands, Awana Rd. | KY989399 |
| T. bukitensis (holotype) | ZRC 2.6245 | West Malaysia, Pahang, Fraser's Hill, Bishops's Trail | KY989400 |
| T. bukitensis | LSUHC 12686 | West Malaysia, Pahang, Fraser's Hill, Jeriau | KY989401 |
| T. bukitensis | LSUHC 12687 | West Malaysia, Pahang, Fraser's Hill, Jeriau | KY989402 |
| T. bukitensis | LSUHC 12226 | West Malaysia, Pahang, Fraser's Hill, Jeriau | KY989403 |
| T. bukitensis | LSUHC 11771 | West Malaysia, Pahang, Fraser's Hill, Telecom Loop | KY989404 |
| T. butleri | LSUHC 9206 | West Malaysia, Perak, Bukit Larut | KU872732 |
| T. butleri | LSUHC 12429 | West Malaysia, Perak, Bukit Larut | KY989405 |
| T. butleri | LSUHC 12432 | West Malaysia, Perak, Bukit Larut | KY989406 |
| T. butleri | LSUHC 12433 | West Malaysia, Perak, Bukit Larut | KY989407 |
| T. butleri | LSUHC 9204 | West Malaysia, Perak, Bukit Larut | KU872731 |
| T. butleri | LSUHC 12516 | West Malaysia, Penang, Penang Island, Penang Hill | KY989408 |
| T. butleri | LSUHC 12071 | West Malaysia, Penang, Penang Island, Penang Hill | KU872733 |
| T. butleri | LSUHC 11275 | West Malaysia, Penang, Penang Island, Penang Hill | KY989410 |
| T. butleri | LSUHC 11552 | West Malaysia, Kedah, Pulau Langkawi, Gunung Raya | KU872737 |
| T. butleri (holotype of T. langkawiensis) | LSUHC 6790 | West Malaysia, Kedah, Pulau Langkawi, Gunung Raya | KU872736 |
| T. butleri | LSUHC 11784 | West Malaysia, Kedah, Pulau Langkawi, Gunung Raya | KY989411 |
| T. butleri | LSUHC 11785 | West Malaysia, Kedah, Pulau Langkawi, Gunung Raya | KU872738 |
| T. butleri | LSUHC 11786 | West Malaysia, Kedah, Pulau Langkawi, Gunung Raya | KU872739 |
| T. butleri | LSUHC 9206 | West Malaysia, Kedah, Pulau Langkawi, Gunung Raya | KU872732 |
| T. ishaki | LSUHC 5165 | West Malaysia, Pahang, Pulau Tioman, Gunung Kajang | KY989413 |

Table 1. Continued
$\left.\begin{array}{llll}\hline \text { Species } & \text { Voucher } & \text { Locality } & \text { GenBank no. } \\ \hline \text { T. ishaki } & \text { LSUHC 6150 } & \begin{array}{c}\text { West Malaysia, Pahang, Pulau Tioman, Gunung } \\ \text { Kajang }\end{array} & \text { Forthcoming } \\ \text { T. ishaki } & \text { LSUHC 6151 } & \begin{array}{c}\text { West Malaysia, Pahang, Pulau Tioman, Gunung } \\ \text { Kajang }\end{array} & \text { Forthcoming } \\ \text { T. jaripendek sp. nov. } & \text { LSUHC 11681 } & \begin{array}{c}\text { West Malaysia, Pahang, Cameron Highlands, } \\ \text { (holotype) }\end{array} & \text { Robinson Falls }\end{array}\right]$ KU872745
chain reaction (PCR) under the following conditions: $1.0 \mu \mathrm{~L}$ genomic DNA ( $\sim 10-33 \mu \mathrm{~g}), 1.0 \mu \mathrm{~L}(10 \mu \mathrm{M})$ light strand primer ( $5^{\prime}$-AAGCAGTTGGGCCCATACC-3'; Macey et al., 1997), $1.0 \mu \mathrm{~L}(10 \mu \mathrm{M})$ heavy strand primer (5'-AACCAAACCCAACTACGAAAAAT-3'; Macey et al., 1997), $1.0 \mu \mathrm{~L}$ dinucleotide pairs ( $1.5 \mu \mathrm{M}$ ), $2.0 \mu \mathrm{~L}$ $5 \times$ buffer $(1.5 \mu \mathrm{M}), 1.0 \mu \mathrm{~L} \mathrm{MgCl} 10 \times$ buffer ( $1.5 \mu \mathrm{M}$ ), $0.18 \mu \mathrm{~L}$ Taq polymerase ( $5 \mathrm{U} / \mu \mathrm{L}$ ) and $7.5 \mu \mathrm{~L} \mathrm{H} \mathrm{H}_{2} \mathrm{O}$. PCR reactions were run on an Eppendorf Mastercycler gradient thermocycler under the following conditions: initial denaturation at $95{ }^{\circ} \mathrm{C}$ for 2 min , followed by a second denaturation at $95{ }^{\circ} \mathrm{C}$ for 35 s , annealing at $52{ }^{\circ} \mathrm{C}$ for 35 s , followed by a cycle extension at $72{ }^{\circ} \mathrm{C}$ for 35 s , for 33 cycles. All PCR products were visualized on a $1 \%$ agarose gel electrophoresis. Successfully amplified PCR products were vacuum purified using MANU 30 PCR Millipore plates and purified products were resuspended in sterile molecular grade water. Purified PCR products were sequenced using the ABI Big-Dye Terminator v3.1 Cycle Sequencing Kit in an ABI GeneAmp PCR 9700 thermal cycler. Cycle sequencing reactions were purified with Sephadex G-50 Fine (GE Healthcare) and sequenced on an ABI 3730xl DNA Analyzer at the BYU DNA Sequencing Center. All sequences were edited in Geneious v6.1.8 (Drummond et al., 2011) and aligned by eye. Mesquite v3.0.2 (Maddison \& Maddison, 2015) was used to check for premature stop codons and to ensure the correct amino acid read frame.
The phylogenetic analysis applied two model-based methods: maximum likelihood (ML) and Bayesian inference (BI). The Bayesian information criterion
(BIC) implemented in IQ-TREE (Nguyen et al., 2015) calculated TVM+I+G4, HKY+G4 and TPM3+G4 to be the best-fit models of evolution for the first, second and third codon positions, respectively (Kalyaanamoorthy et al., 2017). ML analyses implemented in IQ-TREE employed 1000 bootstrap pseudoreplicates via the ultrafast bootstrap approximation algorithm (Minh, Nguyen \& von Haeseler, 2013). A codon-partitioned Bayesian analysis was carried out in MrBayes v3.2.6 (Huelsenbeck \& Ronquist, 2001; Ronquist \& Huelsenbeck, 2003) on CIPRES (Cyberinfrastructure for Phylogenetic Research; Miller, Pfeiffer \& Schwartz, 2010) using default priors. Models of evolution used were approximated from those calculated for the ML analysis and employed GTR $+\mathrm{I}+\Gamma$ for the first and third codon positions and HKY $+\Gamma$ for the second codon position. Two simultaneous runs were performed with four chains per run (three hot and one cold) using default priors. The simulation ran for four million generations, was sampled every 4000 generations from the Markov Chain Monte Carlo (MCMC) and halted after the average standard deviation split frequency was below 0.01 . The first $25 \%$ of the trees were discarded as burn-in using the sumt function. Convergence of all parameters and stationarity distributions were verified and checked in Tracer v1.6 (Rambaut et al., 2014). Nodes having ultrafast ML bootstrap values of 95 and BI posterior probabilities of 0.95 and above were considered well supported (Huelsenbeck et al., 2001; Wilcox et al., 2002; Nguyen et al., 2015). Bootstrap values and posterior probabilities between 80 and 90 were considered moderately supported.

A time-calibrated BI analysis was implemented in BEAUti v1.8.0 (Bayesian Evolutionary Analysis Utility) and run on BEAST v1.8.0 (Bayesian Evolutionary Analysis Sampling Trees; Drummond et al., 2012) employing an uncorrelated lognormal relaxed clock with unlinked substitution and clock models and monophyly unchecked for Taxon Sets. MCMC chains were run using a Species: Coalescence tree prior for 20 million generations and logged every 20000 generations. It has been demonstrated that the third codon position is susceptible to substitution saturation (Zamudio, Jones \& Ward, 1997; Carranza et al., 2000; Grismer et al., 2015a) and could contribute to overestimating node ages. However, Grismer et al. (2015a) noted that although third codon position saturation was evident in their study, it was not a significant factor in estimating node ages across various codon and gene partition schemes and was consistent with similar node age estimates using nuclear genes on the same taxa (Heinicke et al., 2011). The run was calibrated to a $1.47 \%$ mutation rate following mean estimates by Chapple et al. (2012) for New Zealand skinks (Oligsoma) and used a lognormal prior distribution and a mean in real space with an SD of 0.05 for the ucld.mean of each codon. A maximum clade credibility tree using mean heights at the nodes was generated using TreeAnnotator v1.8.0 (Rambaut \& Drummond, 2013) with $10 \%$ of the tress used as a burnin. The BEAST log file was visualized and checked in Tracer v1.6.0 (Rambaut et al., 2014) to ensure effective sample sizes (ESS) were above 200 for all parameters. Because ESS values for the 'prior' and 'posterior' were below 200 and the remaining parameters were considerably higher than 200 in a preliminary run using a GTR $+\Gamma$ substitution model, a re-run using the less complex HKY model was utilized because low ESS values may indicate overparameterized substitution models (Grummer, Bryson \& Reeder, 2014). Importantly, however, the divergence time estimates did not differ significantly between the GTR $+\Gamma$ and the HKY models.

## MORPHOLOGICAL ANALYSIS

All measurements were made with Mitutoyo digital calipers to the nearest 0.1 mm by LLG. Scale counts were made on the right side of the body when possible with a Nikon SMZ 1500 dissecting microscope. Scale terminology is adapted from Grismer et al. (2009a, b), $\operatorname{Lim}$ (1998) and Taylor (1935) and head scales are illustrated in Figure 1. Measurements and scale counts used are SVL measured from the tip of the rostral scale to the vent; head length (HL) measured from the anterior margin of the ear opening to the tip of the rostral scale; tympanum diameter (TD) measured from the anterior to the posterior margin of the auditory


Figure 1. Generalized illustration of head terminology of Tytthoscincus adapted from Grismer (2009a, b), Lim (1998) and Taylor (1935). cs, chinshield; fn, frontonasal; il, infralabial; lo, loreal; m, mental; n, nasal; p, parietal; pf, prefrontal; pm, postmental; po, preorbital; psl, postsupralabial; pt, primary temporal; r, rostral; sc, superciliary; sl, supralabial; so, supraorbital; st, secondary temporal. *Subocular.
meatus; axilla-groin length (AXG) measured from the anterior margin of the hind limb at its insertion point of the body to the posterior margin of the forelimb measured at its insertion point on the body with the upper limb sections held at right angles to the body; forelimb length (FL) measured from the insertion of the forelimb on the body to the base of the third toe with the brachium held at a right angle to the body and hind limb length (HDL) measured from the insertion of the hind limb on the body to the base of the fourth toe the thigh held at a right angle to the body. Ratiometric data (TD/HL, HL/SVL, AXG/SVL, FL/SVL and HDL/SVL) from adults (SVL $\geq 30 \mathrm{~mm}$ determined by the possession eggs or through dissection) were generated from raw measurements.

Scale counts taken were midbody scale rows (MB) counted as the number of longitudinal scale rows encircling the body at a point midway between the limb insertions; paravertebral scale rows (PV) counted as the number of scales in a line from the parietal scales to a point on the dorsum opposite the vent and ventral scale rows (VS) counted as a row of scales between the postmental and the edge of the cloacal plate. Other standard counts include superciliaries [given as two numbers, the first accounting for the anterior smaller scales followed by a second number accounting for the much larger, posteriorly projecting scales preceded by the notation of "' (e.g. 8`1)], supraoculars, loreals, supralabials, infralabials and third finger and fourth toe subdigital lamellae (3FL and 4TL, respectively). Additional characters examined were the degree of contact between the parietals and supraoculars, enlargement of posterior superciliary scales, degree of contact between the prefrontal scales, presence or absence of enlarged precloacal and pectoral scales, texture of
subdigital lamellae and the degree of development of the tympanum. Colour pattern characters examined were the degree of dark, dorsolateral striping and overall ground colour.

Analyses of variance (ANOVAs) were performed to ascertain if statistically significant mean differences ( $P<0.05$ ) existed among the populations for the scale count and adjusted mensural data (see below). ANOVAs having $P$ values less than 0.05 indicating that statistical differences existed were subjected to a Tukey honest significant difference (HSD) test to ascertain which population pairs differed significantly. Boxplots were generated for all characters in order to visualize trends in the data and to identify outliers.
Principal component analysis (PCA) and discriminant analysis of principal components (DAPC) were used to determine (1) the degree to which the morphological variation coincided with species boundaries delimited by the molecular phylogenetic and univariate analyses, (2) to ascertain which category of characters - discrete (scale counts) or continuous (mensural) - performed best at delimiting species and (3) whether sympatric and syntopic populations have different ecomorphologies that are separated in morphospace. PCA, implemented by the prcomp command in $R$ v3.2.1 (R Core Team, 2015), searches for the best overall low-dimensional representation of significant morphological variation in the data. Characters used in the PCAs and DAPCs were discrete meristic data from the scale counts MB, PV, VS, 3FL and 4TL and the continuous mensural data from TD, HL, AXG, FL and HDL. To remove the effects of body size from the latter, we used the following equation: $X_{\text {adj }}=X-$ $\beta\left(\mathrm{SVL}-\mathrm{SVL}_{\text {mean }}\right)$, where $X_{\text {adj }}=$ adjusted value, $X=$ measured value, $\beta=$ unstandardized regression coefficient for each OTU (operational taxonomic unit), SVL $=$ measured SVL and SVL ${ }_{\text {mean }}=$ overall average SVL of all OTUs (Thorpe, 1975, 1983; Turan, 1999; Lleonart, Salat, \& Torres, 2000). All PCA data were natural logtransformed prior to analysis and scaled to their SD in order to normalize their distribution so as to ensure characters with very large and very low values did not over-leverage the results owing to intervariable nonlinearity and to transform meristic and mensural data into comparable units for analysis.

To characterize clustering and separation in morphospace, a DAPC was performed to search for linear combinations of morphological variables having the greatest between-group variance and the smallest within-group variance (Jombart, Devillard \& Balloux, 2010). DAPC relies on log-transformed data from the PCA as a prior step to ensure that variables analyzed are not correlated and number fewer than the sample size. Principal components (PCs) with eigenvalues greater than 1 were retained for the DAPC according to the criterion of Kaiser (1960). Separate PCA and DAPC
analyses were performed on the scale count (discrete) and mensural (continuous) data in order to ascertain which, if any category of characters, performed better at delimiting species with respect to the phylogenetic analyses. Total evidence PCA and DAPC analyses were also performed using a concatenated data set to illustrate an unbiased morphospatial relationship of the OTUs. All statistical analyses were performed using the platform R v3.2.1 (R Core Team, 2015).

Two specimens from Genting Highlands (FMNH 199945, Ulu Kali, Pahang and LSUHC 12072, Awana Road, Pahang) and one specimen from Telom Valley, Cameron Highlands (ZRC 2.5944) from which we were unable to obtain DNA sequence data did not differ discretely from some of the other species in the montane clade. In order to place them taxonomically as conferre (i.e. cf. = compared), each specimen was run separately in a concatenated PCA and DAPC. Membership probabilities were calculated based on posteriors retained from the discriminant functions of the DAPC (Jombart \& Collins, 2015) using the round (head(dapc1\$posterior), 3) command in R v3.2.1 ( R Core Team, 2015).

Specimens examined are listed in the Appendix. Museum acronyms follow Sabaj-Pérez (2010) except for the following non-standard acronyms for the Department of Wildlife and National Parks collection (DWNP), PERHILITAN, Pahang, Malaysia; La Sierra University Herpetological Collection (LSUHC), La Sierra University, Riverside, CA, USA; Langkawi Research Center of Universiti Kebangsaan Malaysia (LRCUKM) on Langkawi Island, Kedah, Malaysia; Rafe M. Brown field series (RMB), University of Kansas, Lawrence, KS, USA and Jimmy A. McGuire field series (JAM), University of California, Berkeley, CA, USA.

## RESULTS

## MOLECULAR ANALYSES

The ML and the BI analyses produced trees with non-conflicting topologies and the ML topology is presented here with Bayesian posterior probabilities and ultrafast ML bootstrap values placed at the nodes, respectively (Fig. 2). The molecular analyses corroborate some of the earlier morphological hypotheses of species designation (Grismer, 2006b; Grismer et al., 2009a, b) by indicating that Tytthoscincus ishaki Grismer, T. perhentianensis Grismer, Wood, \& Grismer and T. temengorensis Grismer, Norhayati, \& Chan are not nested within other species. The analyses similarly corroborate the species designation of T. butleri (Boulenger) from Bukit Larut, Perak but indicate that it is paraphyletic with respect to T. langkawiensis Grismer from the Langkawi Archipelago, Kedah


Figure 2. Maximum likelihood phylogram depicting the relationships of species within the genus Tytthoscincus with Bayesian posterior probabilities and ultrafast ML bootstrap values, respectively, labelled at the nodes. Black circles denote time-calibrated nodes labelled in millions of years before present.
(Grismer, 2008) and a newly discovered population of Tytthoscincus from Pulau Pinang (Penang Island, Malaysia). However, these three allopatric populations have an uncorrected pairwise sequence divergence no greater than $2.04 \%$ among them and correspondingly extremely short branch lengths (Fig. 2). Considering this, and that there are no discrete morphological differences nor statistically different mean values of any character separating these populations from one another (Table 2) and that they completely overlap in the PCA and DAPC analyses (Fig. 3), the Penang Island population is considered conspecific with $T$.
butleri and T. langkawiensis Grismer, 2008 becomes a junior synonym of T. butleri (Boulenger, 1912) by way of nomenclatural priority.

The molecular analyses also indicate there are two well-supported (BI 1.00/ML 100) major lineages within the upland clade of Tytthoscincus (Fig. 2). The first is composed of two semi-fossorial sister species (see below for the morphological designation of this ecomorph): the newly discovered population T. jaripendek sp. nov. (see description below) from Cameron Highlands, Pahang and a new species, T. kakikecil sp. nov. (see description below) from Fraser's Hill and Ulu Kali,

Table 2. Summary statistics and scale counts of the three allopatric populations of Tytthoscincus butleri

|  | Pulau Langkawi | Bukit Larut | Penang Island |
| :---: | :---: | :---: | :---: |
| Supralabials | 6 | 6 | 6 |
| Infralabials | 5 (10) or 6 (1) | 5 | 5 |
| Supraoculars | 4 | 4 | 4 |
| Frontoparietal contacting supraoculars | 2, 3, 4 | 2, 3, 4 | 2, 3, 4 |
| Superciliaries | 91 | 91 | 91 |
| Prefrontals in contact | Yes | Yes | Yes |
| Loreals | 1 (1) or 2 (10) | 1 (1) or 2 (7) | 2 |
| Tympanum deeply recessed and pigmented | Yes | Yes | Yes |
| Pectoral scales enlarged | No - weakly | Weakly - yes | Yes |
| Lamellae texture | Keeled | Keeled | Keeled |
| Midbody scales | 32-38 | 32-35 | 30-35 |
| Mean | 34.9 | 33.7 | 32.3 |
| SD | $\pm 1.87$ | $\pm 1.11$ | $\pm 2.51$ |
| $n$ | 11 | 7 | 3 |
| Paravertebral scales | 60-72 | 65-72 | 64-75 |
| Mean | 68.5 | 69.2 | 70.3 |
| SD | $\pm 3.24$ | $\pm 2.28$ | $\pm 5.69$ |
| $n$ | 11 | 7 | 3 |
| Ventral scales | 63-72 | 63-69 | 68-73 |
| Mean | 67.6 | 66.4 | 69.67 |
| SD | $\pm 3.41$ | $\pm 2.15$ | $\pm 2.89$ |
| $n$ | 11 | 7 | 3 |
| 3rd finger lamellae | 7 | 7 | 7 |
| Mean | 7 | 7 | 7 |
| SD | $\pm 0.0$ | $\pm 0.0$ | $\pm 0.0$ |
| $n$ | 11 | 7 | 3 |
| 4th toe lamellae | 10-13 | 10-13 | 10-12 |
| Mean | 11.5 | 11.7 | 11.0 |
| SD | $\pm 0.82$ | $\pm 0.95$ | $\pm 1.00$ |
| $n$ | 11 | 7 | 3 |
| TD/HL | 0.13-0.16 | 0.13-0.16 | 0.13-0.14 |
| Mean | 0.14 | 0.14 | 0.13 |
| SD | $\pm 0.01$ | $\pm 0.01$ | $\pm 0.01$ |
| $n$ | 8 | 7 | 3 |
| HL/SVL | 0.20-0.22 | 0.19-0.22 | 0.19-0.22 |
| Mean | 0.21 | 0.21 | 0.20 |
| SD | $\pm 0.01$ | $\pm 0.01$ | $\pm 0.01$ |
| $n$ | 8 | 7 | 3 |
| AXG/SVL | 0.53-0.57 | 0.52-0.56 | 0.50-0.58 |
| Mean | 0.54 | 0.54 | 0.54 |
| SD | $\pm 0.01$ | $\pm 0.02$ | $\pm 0.04$ |
| $n$ | 8 | 7 | 3 |
| FL/SVL | 0.22-0.28 | 0.23-0.26 | 0.22-0.23 |
| Mean | 0.24 | 0.24 | 0.23 |
| SD | $\pm 0.02$ | $\pm 0.01$ | $\pm 0.01$ |
| $n$ | 8 | 7 | 3 |
| HDL/SVL | 0.33-0.40 | 0.34-0.38 | 0.33-0.37 |
| Mean | 0.35 | 0.36 | 0.35 |
| SD | $\pm 0.02$ | $\pm 0.01$ | $\pm 0.02$ |
| $n$ | 8 | 7 | 3 |
| Maximum SVL | 40.1 | 40.9 | 38.0 |

All measurements are in millimetres. AXG, axilla-groin length; FL, forelimb length; HDL, hind limb length; HL, head length; SVL, snout-vent length; TD, tympanum diameter.

PCA and DAPC of meristic characters


PCA and DAPC of mensural characters



Figure 3. Principal component analysis (PCA) and discriminant analysis of principal components (DAPC) analyses of scale count and adjusted mensural data.

Genting Highlands, Pahang. Together, these two species comprise the sister group to the second major lineage consisting of leaf-litter generalists (see below for the morphological designation of this ecomorph). This lineage is composed of T. bukitensis, a new species $T$. martae sp. nov. (see description below) from Awana Road, Genting Highlands, Pahang; T. ishaki; T. perhentianensis; T. temengorensis and T. butleri. Additionally, expanded genetic sampling demonstrates that $T$. bukitensis is not endemic to Fraser's Hill (Grismer, 2007), but extends as far south to at least Awana Road in Genting Highlands and as such, is syntopic with $T$. martae sp. nov. (Fig. 2). Lastly, the analyses demonstrate that the Tembat Forest Reserve population from

Hulu Terengganu, Terengganu represents a fourth new species, Tytthoscincus sp. nov. However, this specimen died in a pitfall trap and subsequent predation by insects left it damaged beyond description.

The expanded molecular data set confirmed that Sphenomorphus sananus (Kopstein), S. temmincki (Duméril \& Bibron) and S. textus (Müller) should be included in the genus Tytthoscincus as previously hypothesized by Linkem et al. (2011) and Linkem (2013) being that they are embedded within the phylogeny of Tytthoscincus (Fig. 2) and are thus transferred herein.

Using a limited taxon set and a ML analysis, Karin et al. (2016) demonstrated that Tytthoscincus
batupanggah Karin, Das \& Bauer and T. leproauricularis Karin, Das \& Bauer were part of a monophyletic group that included species from Sulawesi and Borneo. Increased taxon sampling from Sulawesi and Peninsular Malaysia and the additional use of a BI analysis here indicate that the group to which T. batupanggah and T. leproauricularis belong is paraphyletic with respect to the montane clade from Peninsular Malaysia. Tytthoscincus batupanggah is the sister species of a monophyletic group composed of T. atrigularis, T. hallieri and T. parvus as Karin et al. (2016) indicated but now included in that group are T. panchorensis Grismer, Muin, Wood, Anuar, \& Linkem, T. sibuensis Grismer, T. textus, T. sananus, T. cf. sibuensis and T. temmincki (Fig. 2). Karin et al. (2016) placed T. leproauricularis as the sister species to T. aesculeticola but the analyses herein show it to be the potential sister lineage to a clade containing all other Tytthoscincus although this relationship is only weakly supported (0.81/69).

## MORPHOLOGICAL ANALYSES

The PCA and DAPC for the continuous, adjusted mensural characters are largely consistent with the molecular analyses in that they indicate little to no overlap between the semi-fossorial and fossorial species (Fig. 3). PC1 and PC2 account for $84 \%$ of the total variation and load most heavily for AXG and FL along PC1 (52\% of the total variation) and for TD along PC2 (32\% of the total variation) (Table 3). Boxplots for these characters show that T. jaripendek sp. nov. and T. kakikecil sp. nov. along with T. temengorensis have relatively longer trunk lengths (AXG) than all other species; that along with T. temengorensis and T. perhentianensis, they have relatively shorter FL than all other species; and they have the relatively smallest tympanae of all species (Fig. 4). The DAPC further characterizes these results (Fig. 3). These characters are commonly correlated with a fossorial lifestyle (see below).

The PCA and DAPC for the discrete, meristic characters (scale counts) do not coincide as well with the molecular analyses as the mensural data in that T. perhentianensis clusters between the semi-fossorial species T. jaripendek sp. nov. and T. kakikecil sp. nov. (Fig. 3). PC1 and PC2 account for $76 \%$ of the total variation and load most heavily for the number of subdigital lamellae on the third finger (3FL), fourth toe (4TL) and paravertebral scales (PV) along PC1 (60\% of the total variation) and for MB along PC2 (16\% of the total variation; Table 4). The boxplots for 3TL and MB show that T. jaripendek sp. nov. and T. kakikecil sp. nov. along with T. perhentianensis have fewer lamellae (=shorter third toes) than all other species and that T. jaripendek sp. nov. and T. kakikecil sp. nov. have fewer MB than all other species (Fig. 5). These results are further characterized in the DAPC (Fig. 3). Shorter digits are commonly correlated with a fossorial lifestyle (see below).

The concatenated PCA and DAPC analyses are the preferred analyses in that they performed best overall with respect to the molecular analyses in that there was unequivocal separation between the semi-fossorial and fossorial species as well as clear separation among other combinations of the leaf-litter generalist species (Fig. 6). More precisely, the PCA analysis demonstrates that the upland clade of Tytthoscincus is composed of two non-overlapping, distinctive, ecomorphological lineages along PC1 that are referred to here as the semifossorial and the leaf-litter generalist lineages (Fig. 1). PC1 accounts for $44 \%$ of the total variation and loads most heavily for characters related to a reduction in limb length and number of digital lamellae (FL, HDL, 3 FL and 4TL; Figs 4, 5; Table 5) - characters commonly correlated with a fossorial lifestyle. Ordination of PC1 and PC2 show complete separation of the semi-fossorial species T. jaripendek sp. nov. and T. kakikecil sp. nov., the leaf-litter generalists T. perhentianensis and T. ishaki from each other and all other species. The leaf-litter generalists $T$. temengorensis clusters separately from all other species except T. butleri. The limbs of T. jaripendek

Table 3. Summary statistics and principal component scores for scaled ratiometric characters

|  | PC1 | PC2 | PC3 | PC4 | PC5 |
| :--- | ---: | ---: | ---: | ---: | ---: |
| SD | 1.606044 | 1.270911 | 0.670349 | 0.549558 | 0.232432 |
| Proportion of variance | 0.515880 | 0.323040 | 0.089870 | 0.060400 | 0.010800 |
| Cumulative proportion of variance | 0.515880 | 0.838920 | 0.928790 | 0.989200 | 1.000000 |
| Eigenvalues | 2.579378 | 1.615215 | 0.449368 | 0.302014 | 0.054025 |
| AXG | -0.518408 | -0.214519 | 0.507062 | 0.615025 | 0.223311 |
| TD | -0.163549 | -0.656511 | -0.706972 | 0.179008 | 0.101942 |
| HL | -0.451126 | -0.413177 | 0.289974 | -0.730316 | -0.091236 |
| FL | -0.541198 | 0.335782 | -0.256324 | 0.131888 | -0.715024 |
| HDL | -0.456174 | 0.489398 | -0.305438 | -0.197345 | 0.648195 |

AXG, axilla-groin length; FL, forelimb length; HDL, hind limb length; HL, head length; TD, tympanum diameter.


Figure 4. Boxplots of the ratiometric data.

Table 4. Summary statistics and principal component scores for meristic characters

|  | PC1 | PC2 | PC3 | PC4 |  |
| :--- | :--- | :--- | :--- | :--- | :--- |
| SD | 1.734939 | 0.906664 | 0.789836 | 0.588183 | 0.445139 |
| Proportion of variance | 0.602 | 0.16441 | 0.12477 | 0.06919 | 0.03963 |
| Cumulative proportion of variance | 0.602 | 0.76641 | 0.89118 | 0.96037 | 1 |
| Eigenvalues | 3.010012 | 0.822039 | 0.623841 | 0.345959 | 0.198149 |
| MB | -0.388991 | 0.634759 | -0.463629 | 0.476040 | -0.064813 |
| PV | -0.443814 | -0.567318 | 0.093480 | 0.542347 | 0.422256 |
| VS | -0.391034 | 0.340741 | 0.838640 | 0.020419 | -0.165086 |
| 3FL | -0.504837 | 0.118398 | -0.165947 | -0.661859 | 0.515292 |
| 4TL | -0.493909 | -0.380930 | -0.213200 | -0.201920 | -0.724375 |

FL, forelimb length; MB, midbody scale rows; PV, paravertebral scale rows; TL, toe subdigital lamellae; VS, ventral scale rows.
sp. nov. and T. kakikecil sp. nov. are also much less robust than those of the other species and they have extremely reduced, shallow, pigmented tympanae (Fig. 7). Varying combinations of T. bukitensis, T. butleri, T. martae sp. nov. and T. temengorensis overlap along PC1 and/or PC2.
The PC2 accounts for an additional $19 \%$ of the total variation (Fig. 6) and loads heavily for reduced tympanae (TD), shortened head (HL) and a longer trunk (AXG; Figs 4, 5; Table 5) - characters commonly correlated with a fossorial lifestyle. The PC3 accounts for an additional $10 \%$ of the total variation and loads heavily for ventral scales (VS; Table 5). The first eight PCs had eigenvalues ranging from 0.17 to 4.44 and accounted for $98 \%$ of the total variance (Table 5) and the first three of these were retained for the DAPC analysis. The DAPC analysis also shows exclusive clustering among the semi-fossorial species T. jaripendek sp. nov. and T. kakikecil sp. nov., the leaf-litter generalists T. perhentianensis and T. ishaki, as well as broad overlap among T. butleri, T. bukitensis, T. martae sp. nov. and T. temengorensis (Fig. 6).

The morphological analyses support the molecular analyses further in the specific recognition of the newly discovered populations from Cameron Highlands (T. jaripendek sp. nov.) and its sister species from Fraser's Hill and Genting Highlands (T. kakikecil sp. nov.) being that each of them is discretely and statistically diagnosable from each other and all other species of Tytthoscincus in the montane clade (Tables 6, 7; Fig. 2). Additionally, they have a $4.0 \%$ uncorrected pairwise sequence divergence between them. The other new species from Fraser's Hill and Genting Highlands (T. martae sp. nov.) is statistically diagnosable from most species in the montane clade and clusters separately from them in the concatenated PCA and DAPC. It is statistically separated from T. butleri, T. ishaki and T. perhentianensis but not T. temengorensis and T. bukitensis although it does cluster separately from $T$. temengorensis in the PCA and DAPC (Table 5; Fig. 6). The phylogenetic analysis indicates that considering T. bukitensis and T. martae sp. nov. conspecific would render the former paraphyletic (Fig. 2).

## TAXONOMY

The molecular phylogeny combined with the morphological data indicate that the sister lineages T. jaripendek sp. nov. from Cameron Highlands and T. kakikecil sp. nov. from Cameron Highlands, Fraser's Hill and Genting Highlands warrant species recognition. The molecular phylogeny also suggests that T. martae sp. nov., from Fraser's Hill and Genting Highlands, also warrants species recognition even though its morphological separation from T. bukitensis is less clear (see below).

> ORder Squamata Oppel, 1811
> Family Scincidae Gray, 1825
> Genus TytTHoscincus Linkem, Diesmos \& Brown, 2011

## Tytthoscincus Jaripendek sp. nov.

Cameron Highlands Forest Skink Mengkarung Hutan Cameron Highlands
(Fig. 8; TABLE 8)
Tytthoscincus sp. Grismer et al. 2016a: 236.
Holotype: Adult male (LSUHC 11680) collected at Robinson's Falls, Cameron Highlands, Pahang, Peninsular Malaysia (N $04^{\circ} 27.959^{\prime}$, E $101^{\circ} 23.129^{\prime}$; 1411 m in elevation) by Evan S. H. Quah and L. Lee Grismer on 26 April 2014.

Paratypes: LSUHC 11679, 11681 and 11987 all bear the same collection data as the holotype except the latter was collected on 4 September 2014.

Diagnosis: Tytthoscincus jaripendek sp. nov. can be differentiated from all other species of Tytthoscincus in the montane clade by having the combination of $8 ` 1$ superciliaries; a shallow, pigmented tympanic depression; no enlarged pectoral scales; 28-30 midbody


Figure 5. Boxplots of the scale count data.
scales (MB); 63-65 paravertebral scales; 60-62 ventral scales; keeled, subdigital lamellae; five or six subdigital lamellae on the third finger; nine or ten subdigital lamellae on the fourth toe; TD/HL $=0.11-0.13$; HDL/ SVL $=0.18-0.19 ; \mathrm{AXG} / \mathrm{SVL}=0.55-0.57 ; \mathrm{FL} / \mathrm{SVL}=$
$0.18-0.21 ; \mathrm{HDL} / \mathrm{SVL}=0.31-0.33$ and a maximum SVL of 33.9 mm (Table 6). All non-ratiometric characters are presented in all other Tytthoscincus and species of Sphenomorphus suspected of being Tytthoscincus in Grismer et al. (2016a: 237) and Karin et al. (2016: 416).


Figure 6. A, concatenated principle component analysis (PCA) of the montane clade of Tytthoscincus showing separation of the semi-fossorial species (triangles) and the leaf-litter generalists (circles) along axis PC1. B, concatenated discriminant function of principle component analysis (DAPC) emphasizing interspecific variation and complete separation of the semifossorial species from one another and the leaf-litter generalists.
Table 5. Summary statistics and principal component scores for meristic and scaled ratiometric variables

|  | PC1 | PC2 | PC3 | PC4 | PC5 | PC6 | PC7 | PC8 | PC9 |  |
| :--- | ---: | ---: | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |
| SD | 2.107388 | 1.395620 | 1.014235 | 0.888918 | 0.783016 | 0.719016 | 0.565313 | 0.411946 | 0.349538 | 0.225295 |
| Proportion of variance | 0.444110 | 0.194780 | 0.102870 | 0.079020 | 0.061310 | 0.051700 | 0.031960 | 0.016970 | 0.012220 | 0.005080 |
| Cumulative proportion | 0.444110 | 0.638880 | 0.741750 | 0.820770 | 0.882080 | 0.933780 | 0.965740 | 0.982710 | 0.994920 | 1.000000 |
| $\quad$ of variance |  |  |  |  |  |  |  |  |  |  |
| Eigenvalues | 4.441085 | 1.947756 | 1.028673 | 0.790175 | 0.613114 | 0.516984 | 0.319579 | 0.169699 | 0.122177 | 0.050758 |
| MB | -0.303549 | 0.035739 | -0.343574 | 0.549427 | -0.487490 | -0.325005 | 0.173186 | -0.318962 | 0.090920 | -0.058284 |
| PV | -0.321171 | -0.214824 | -0.172704 | -0.604050 | 0.086622 | -0.391133 | 0.210385 | -0.307255 | -0.395894 | 0.010386 |
| VS | -0.261003 | -0.081636 | -0.641692 | 0.074810 | 0.440500 | 0.471684 | -0.221627 | -0.187923 | 0.079655 | -0.023302 |
| 3TL | -0.413132 | -0.140872 | -0.019955 | 0.136672 | -0.139177 | 0.311290 | 0.363137 | 0.612392 | -0.408808 | -0.010529 |
| 4FL | -0.384368 | -0.132103 | -0.020961 | -0.433226 | -0.382836 | 0.060766 | -0.181724 | 0.224593 | 0.640385 | 0.053585 |
| TD | 0.055018 | 0.575580 | -0.391448 | -0.097530 | 0.266474 | -0.399060 | 0.203116 | 0.440121 | 0.179600 | -0.073900 |
| HL | -0.187674 | 0.574420 | -0.000883 | -0.114987 | -0.307726 | 0.094936 | -0.573638 | -0.028975 | -0.424818 | 0.086714 |
| AXG | -0.241444 | 0.482122 | 0.308350 | -0.122662 | 0.053034 | 0.376710 | 0.480371 | -0.372963 | 0.164776 | -0.239593 |
| FL | -0.406735 | 0.079774 | 0.299189 | 0.219141 | 0.358316 | -0.177021 | -0.020664 | -0.028813 | 0.105362 | 0.720126 |
| HDL | -0.395209 | -0.092507 | 0.318901 | 0.188388 | 0.313002 | -0.277125 | -0.332015 | 0.092727 | 0.021455 | -0.635613 |

AXG, axilla-groin length; FL, forelimb length; HL, head length; HDL, hind limb length; MB, midbody scale rows; PV, paravertebral scale rows; TD, tympanum diameter; TL, toe subdigital lamellae;


Figure 7. A, B, dorsal and lateral views of the anterior portion of the body and head, respectively, of the leaf-litter generalist Tytthoscincus butleri (LSUHC 9204) showing its more robust forelimbs and deeply set, well-developed tympanum. C, D, dorsal and lateral views of the anterior portion of the body and head, respectively, of the semi-fossorial Tytthoscincus kakikecil sp. nov. (LSUHC 11769) showing its more gracile forelimbs and shallow, poorly developed tympanic depression.

Description of holotype: Adult male, SVL 32.2 mm ; tail length 38.0 mm ; AXG 18.4 mm ; HL 6.1 mm ; head width 3.7 mm ; snout to anterior margin of foreleg 11.2 mm ; rostral wider than long, in broad contact with frontonasal; frontonasal wider than long; prefrontals large, broadly contacting on midline; frontal elongate, diamond-shaped, in contact with first two supraoculars; four supraoculars; frontoparietals in contact posterior to frontal, contacting second, third and fourth supraoculars anterolaterally and parietals and interparietal posteriorly; interparietal dia-mond-shaped, large, slightly projecting posteriorly, eyespot in posterior projection; parietals large, in medial contact posterior to interparietal, contacting fourth supraocular anteriorly; enlarged nuchal scales absent; nasals small, widely separated, trapezoidal, contacting rostral anteriorly, frontonasal dorsally, first loreal posteriorly, first supralabial ventrally; nostril in centre of nasal; supranasals absent; two loreals, posterior loreal taller than wide; upper and lower preocular present; lower preocular followed by a series of suboculars; eight superciliaries, posterior superciliary elongate and projecting dorsomedially; six supralabials, third, fourth and fifth below eye; two postsupralabials; two primary temporals; three secondary
temporals, uppermost not contacting parietals; lower eyelid transparent, scaly, no enlarged central window; mental twice as wide as long; single, large postmental, contacting first infralabial on each side; two enlarged pairs of chinshields posterior to postmental, anterior pair contacting medially, posterior pair widely separated posteriorly by a single scale; anterior and posterior chinshield pairs contacting first, second and third infralabials; five infralabials; external ear opening 0.8 mm wide, approximately one-half diameter of eyeball, subcircular, lacking anterior lobules; and tympanum very shallow, pigmented.

Body scales smooth, cycloid, imbricate; ventral scales slightly larger than dorsal scales; 28 longitudinal scale rows around midbody; 65 paravertebral scales (PV); 60 ventral scales (VS); slightly enlarged median precloacal scales overlapping outer precloacal scales; tail slightly compressed laterally; subcaudals slightly larger than dorsal caudals; limbs very thin, short (FL/ SVL $=0.21 ; \mathrm{HDL} / \mathrm{SVL}=0.33$ ), widely separated when adpressed; scales of dorsal surface slightly larger than those of ventral surface; palmar and plantar scales slightly raised; and digits short, scales of dorsal surfaces in a single row, subdigital lamellae bicarinate, five on third finger, ten on fourth toe.
Table 6. Scale counts, summary statistics and morphometric ratios of selected characters of the montane clade of Tytthoscincus from Peninsular Malaysia

|  | butleri | bukitensis | jaripendek sp. nov. | temengorensis | perhentianensis | ishaki | kakikecil sp. nov. | martae sp. nov. | cophias |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Supralabials | 6 | 6 | 6 | 6 | 6 | 6 | 6 | 6 | 6 |
| Infralabials | 5, 6 | 5 | 5 | 5 | 5, 6 | 5, 6 | 5, 6 | 5, 6 | 6 |
| Superciliaries |  | $8^{\prime} 1,9 {fc4a5ec94-370e-491e-8707-28225470598b} 2,10^{\prime} 2$ | 8`1 & 71 & \(9 \times 1\) & 91 & 8`1, 9`1, 9`2 | 8`1, 9`1, 10`2 | 81 |  |  |  |  |
| Loreals | 1,2 | 2 | 1, 2 | 2 | 2 | 1,2 | 1,2 | 2 | 2 |
| Tympanum | DWD | DWD | SPD | DWD | DWD | DWD | SPD | DWD | SPD |
| Pectoral scales enlarged | Yes (10), no (11) | Yes (9), no (1) | No | Yes | Yes | No | Yes | Yes | Yes |
| Lamellae texture | Keeled | Keeled | Keeled | Keeled | Smooth | Keeled | Keeled | Keeled | Keeled |
| Midbody scales | 32-38 | 31-35 | 28-30 | 30-35 | 29,30 | 30-32 | 30-33 | 31-35 | 23 |
| Mean | 34.2 | 32.5 | 29.3 | 32.5 | 29.5 | 30.7 | 32.2 | 33.3 | 23.0 |
| SD | $\pm 1.8$ | $\pm 1.3$ | $\pm 1.0$ | $\pm 3.6$ | $\pm 0.7$ | $\pm 1.1$ | $\pm 1.3$ | $\pm 1.3$ | $\pm 0.0$ |
| $n$ | 20 | 8 | 4 | 2 | 2 | 3 | 6 | 7 | 1 |
| Paravertebral scales | 60-75 | 71-74 | 63-65 | 68-70 | 65, 66 | 68-73 | 67-73 | 70-74 | 56 |
| Mean | 68.9 | 72.9 | 64.3 | 69.0 | 65.5 | 70.0 | 69.8 | 72.3 | 56.0 |
| SD | $\pm 3.4$ | $\pm 1.2$ | $\pm 1.0$ | $\pm 1.4$ | $\pm 0.7$ | $\pm 2.6$ | $\pm 2.2$ | $\pm 1.4$ | $\pm 0.0$ |
| $n$ | 20 | 8 | 4 | 2 | 2 | 3 | 6 | 7 | 1 |
| Ventral scales | 62-73 | 64-74 | 60-62 | 70, 71 | 61, 62 | 60-70 | 65-69 | 61-72 | 56 |
| Mean | 67.6 | 68.3 | 60.6 | 70.5 | 61.5 | 65.3 | 67.0 | 68.4 | 56.0 |
| SD | $\pm 3.1$ | $\pm 3.5$ | $\pm 1.0$ | $\pm 0.7$ | $\pm 0.7$ | $\pm 5.0$ | $\pm 1.5$ | $\pm 3.5$ | $\pm 0.0$ |
| $n$ | 20 | 8 | 4 | 2 | 2 | 3 | 6 | 7 | 1 |
| 3rd finger lamellae | 6-8 | 7 | 5 or 6 | 7, 8 | 6 | 7 | 6 | 7 | 5 |
| Mean | 7 | 7 | 5.1 | 7.5 | 6 | 7.0 | 6 | 7 | 5.0 |
| SD | $\pm 0.4$ | $\pm 0.0$ | $\pm 0.5$ | $\pm 0.7$ | $\pm 0.0$ | $\pm 0.0$ | $\pm 0.0$ | $\pm 0.0$ | $\pm 0.0$ |
| $n$ | 20 | 8 | 4 | 2 | 1 | 3 | 6 | 7 | 1 |
| 4 th toe lamellae | 10-13 | 12 or 13 | 9 or 10 | 11 | 10 | 11 | 10 | 12 or 13 | 10 |
| Mean | 11.6 | 12.4 | 9.5 | 11 | 10 | 11.0 | 10 | 12.7 | 10.0 |
| SD | $\pm 0.8$ | $\pm 0.5$ | $\pm 0.6$ | $\pm 0.0$ | $\pm 0.0$ | $\pm 0.0$ | $\pm 0.0$ | $\pm 0.5$ | $\pm 0.0$ |
| $n$ | 20 | 8 | 4 | 2 | 1 | 3 | 6 | 7 | 1 |
| TD/HL | 0.13-0.16 | 0.12-0.16 | 0.11-0.13 | 0.13 | 0.13-0.14 | 0.13-0.16 | 0.10-0.12 | 0.12-0.16 | 0.15 |
| Mean | 0.14 | 0.14 | 0.12 | 0.13 | 0.14 | 0.14 | 0.11 | 0.13 | 0.15 |
| SD | $\pm 0.01$ | $\pm 0.01$ | $\pm 0.01$ | $\pm 0.0$ | $\pm 0.01$ | $\pm 0.2$ | $\pm 0.01$ | $\pm 0.02$ | $\pm 0.0$ |
| $n$ | 18 | 8 | 4 | 2 | 2 | 3 | 6 | 7 | 1 |
| HL/SVL | 0.19-0.21 | 0.19-0.24 | 0.18-0.19 | 0.21-0.23 | 0.22-0.23 | 0.18 | 0.18-0.21 | 0.18-0.26 | 0.15 |
| Mean | 0.20 | 0.22 | 0.19 | 0.22 | 0.23 | 0.18 | 0.19 | 0.21 | 0.15 |
| SD | $\pm 0.01$ | $\pm 0.02$ | $\pm 0.01$ | $\pm 0.1$ | $\pm 0.1$ | $\pm 0.0$ | $\pm 0.01$ | $\pm 0.03$ | $\pm 0.0$ |
| $n$ | 18 | 8 | 4 | 2 | 2 | 3 | 6 | 7 | 1 |

Table 6. Continued

|  | butleri | bukitensis | jaripendek sp. nov. | temengorensis | perhentianensis | ishaki | kakikecil sp. nov. | martae <br> sp. nov. | cophias |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| AXG/SVL | 0.50-0.57 | 0.48-0.63 | 0.55-0.57 | 0.50-0.57 | 0.54-0.55 | 0.49-0.54 | 0.52-0.65 | 0.47-0.55 | 0.58 |
| Mean | 0.54 | 0.53 | 0.56 | 0.54 | 0.55 | 0.52 | 0.57 | 0.51 | 0.58 |
| SD | $\pm 0.02$ | $\pm 0.05$ | $\pm 0.01$ | $\pm 0.5$ | $\pm 0.1$ | $\pm 0.03$ | $\pm 0.05$ | $\pm 0.03$ | $\pm 0.0$ |
| $n$ | 16 | 8 | 4 | 2 | 2 | 3 | 6 | 7 | 1 |
| FL/SVL | 0.22-0.28 | 0.22-0.27 | 0.18-0.21 | 0.21-0.22 | 0.20-0.21 | 0.23-0.24 | 0.18-0.23 | 0.21-0.26 | 0.17 |
| Mean | 0.24 | 0.25 | 0.20 | 0.22 | 0.21 | 0.23 | 0.21 | 0.24 | 0.17 |
| SD | $\pm 0.02$ | $\pm 0.02$ | $\pm 0.02$ | $\pm 0.01$ | $\pm 0.1$ | $\pm 0.01$ | $\pm 0.02$ | $\pm 0.02$ | $\pm 0.0$ |
| $n$ | 18 | 8 | 4 | 2 | 2 | 3 | 6 | 7 | 1 |
| HDL/SVL | 0.33-0.40 | 0.34-0.40 | 0.31-0.33 | 0.32-0.38 | 0.29-0.30 | 0.34-0.35 | 0.30-0.34 | 0.33-0.38 | 0.28 |
| Mean | 0.36 | 0.37 | 0.32 | 0.35 | 0.30 | 0.35 | 0.32 | 0.35 | 0.28 |
| SD | $\pm 0.02$ | $\pm 0.02$ | $\pm 0.01$ | $\pm 0.0$ | $\pm 0.1$ | $\pm 0.01$ | $\pm 0.02$ | $\pm 0.02$ | $\pm 0.0$ |
| $n$ | 18 | 8 | 4 | 2 | 2 | 3 | 6 | 7 | 1 |
| Maximum SVL | 40.9 | 42.6 | 33.9 | 35.5 | 30.0 | 41.0 | 36.2 | 44.0 | 35.9 |




Coloration: Overall dorsal ground colour of head, body, limbs and tail dark brown; head speckled with lightcoloured markings; labials distinctly banded; anterior of flanks to posterior margin of jaws dull-orange bearing irregularly shaped, dark speckles and elongate markings; dorsum, posterior portion of flanks and tail bearing small, light-coloured spots; limbs faintly mottled; ventral surfaces dull-yellow, immaculate; lateral margin of gular region stippled with dark brown; and palmar and plantar surfaces, subtibial, subforearm and subcaudal regions heavily stippled with dark brown.

Variation: The paratypes closely approach the holotype in all aspects of coloration except for LSUHC 11987 which has considerably more gular stippling and is darker dorsally overall. Meristic variation is presented in Table 8.

Distribution: Tytthoscincus jaripendek sp. nov. is known only from the type locality at Robinson's Falls, Cameron Highlands, Pahang, Peninsular Malaysia (Fig. 9) but is likely to range more widely across the Cameron Highlands plateau.

Natural history: Tytthoscincus jaripendek sp. nov. inhabits leaf-litter in hill dipterocarp forests at 1411 $m$ in elevation (Fig. 10). All specimens were collected by raking leaves that had accumulated at the foot of an earthen bank. Specimens were collected both day and night and during periods of heavy rain.

Etymology: The specific epithet jaripendek is derived from the Malay words jari meaning finger and pendek meaning short and refers to this species' short digits as indicated by its low numbers of subdigital lamellae.

Comparisons: Tytthoscincus jaripendek sp. nov. can be differentiated from all other species of Tytthoscincus of the montane clade (Table 6), except its sister species T. kakikecil sp. nov., by having a shallow, pigmented tympanic depression vs. a well-developed, deeply recessed, non-pigmented tympanum. From T. kakikecil sp. nov., it differs discretely and statistically by having $63-65$ vs. 67-73 paravertebral scales, $60-62$ vs. 65-69 ventral scales (Tables 6, 7), and statistically significant differences in the mean number of paravertebral scales, ventral scales, 3TL and FL/SVL (Tables 6, 7). From all other species except T. perhentianensis, it differs further in having fewer midbody scales (28-30 vs. 30-38 collectively). Tytthoscincus jaripendek sp. nov. also bears statistically significant mean differences of all ten characters in varying combinations across all other species (Table 7). Meristic and body shape trends of these characters among species in the montane clade are presented in Figures 4 and 5.
Table 7. Matrix of species pairs having significantly different ( $P<0.05$ ) mean values of varying combinations of meristic and scaled ratiometric characters based on a Tukey HSD test

|  | bukitensis | butleri | ishaki | jaripendek | kakikecil | martae | perhentianensis temengorensis |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| bukitensis | * |  |  |  |  |  |  |
| butleri | MB, PV | * |  |  |  |  |  |
| ishaki | AXG, TD, HL | AXG, VMB, TD, HL | * |  |  |  |  |
| jaripendek | MB, PV, VS, 3TL, 4TL, AXG, HL, FL, HDL | MB, PV, VS, 3TL, 4TL, AXG, HL, FL, HDL | $\begin{aligned} & \text { PV, 3TL, TD, HL, } \\ & \text { FL, HDL } \end{aligned}$ | * |  |  |  |
| kakikecil | $\begin{aligned} & \text { PV, 3TL, 4TL, AXG, HL, } \\ & \text { FL, HDL } \end{aligned}$ | $\begin{aligned} & \text { 3TL, 4TL, AXG, HL, } \\ & \text { FL, HDL } \end{aligned}$ | TD, HL, HDL | PV, VS, 3TL, FL | * |  |  |
| martae |  | PV, 4TL | $\begin{aligned} & \text { 4TL, TD, AXG, HL, } \\ & \text { FL } \end{aligned}$ | MB, PV, VS, 3TL, 4TL, AXG, FL, HDL | $\begin{aligned} & \text { 3TL, 4TL, TD, } \\ & \text { HL, HDL } \end{aligned}$ | * |  |
| perhentianensis | AXG, PV, 3TL, HL | MB, 3TL, 4TL, AXG | AXG, TD, HL | 3TL, AXG, HL, FL | AXG, HL | $\begin{aligned} & \text { PV, 3TL, } \\ & \text { 4TL, } \\ & \text { AXG, TD } \end{aligned}$ |  |
| temengorensis | 3TL, HDL | 3TL | AXG, TD, HL | VS, 3TL, AXG | 3TL, AXG |  | 3TL, HL, AXG |

[^1]
# Tytthoscincus kakikecil sp. NOV. 

Fraser's Hill Forest Skink
Mengkarung Hutan Bukit Fraser
(Fig. 11; TABLE 9)
Holotype: Adult male (LSUHC 11769) collected along Richmond Road, Frasers's Hill, Pahang, Peninsular Malaysia ( $\mathrm{N} 03^{\circ} 42.590^{\prime}$, E $101^{\circ} 44.236^{\prime}$; 1271 m in elevation) by Evan S. H. Quah on 27 September 2013.

Paratypes: LSUHC 11770 and 11772 collected on the Telecom Loop Road, Fraser's Hill, Pahang, Peninsular Malaysia ( $\mathrm{N} 03^{\circ} 43.112^{\prime}$, E $101^{\circ} 45.158^{\prime}$; 1305 m in elevation) by Evan S. H. Quah and L. Lee Grismer on 27 September 2013. LSUHC 12754 collected at Ulu Kali, Genting Highlands, Pahang, Peninsular Malaysia (N $03^{\circ} 25.759^{\prime}$, E $101^{\circ} 47.328^{\prime}$; 1670 m in elevation) by Evan S. H. Quah and L. Lee Grismer on 22 June 2016.

Additional specimens examined: FMNH 19945 from Ulu Kali, Genting Highlands and ZRC 2.5944 from Telom Valley, Cameron Highlands, Pahang, Peninsular Malaysia.

Diagnosis: Tytthoscincus kakikecil sp. nov. can be differentiated from all other species of Tytthoscincus in the montane clade by having the combination of $8 `$, $9^{`} 1$ or $9^{`} 2$ superciliaries; a shallow, pigmented tympanic depression; enlarged pectoral scales; 30-33 midbody scales; 67-73 paravertebral scales; 65-69 ventral scales; keeled, subdigital lamellae; six subdigital lamellae on the third finger; ten subdigital lamellae on the fourth toe; TD/HL $=0.10-0.12 ; \mathrm{HL} / \mathrm{SVL}=0.18-$ 0.21 ; AXG/SVL $=0.52-0.65 ; \mathrm{FL} / \mathrm{SVL}=0.18-0.23$; $\mathrm{HDL} / \mathrm{SVL}=0.30-0.34$; and a maximum SVL of 36.2 mm (Table 6). All non-ratiometric characters are scored across all other Tytthoscincus and species of Sphenomorphus suspected of being Tytthoscincus in Grismer et al. (2016a: 237) and Karin et al. (2016: 416).

Description of holotype: Adult male, SVL 30.8 mm ; original tail 33.9 mm ;AXG 19.9 mm ;HL 6.2 mm ; head width 3.4 mm ; snout to anterior margin of foreleg 10.6 mm ; rostral wider than long, in broad contact with frontonasal; frontonasal wider than long; prefrontals large, in broad contact on midline; frontal elongate, diamond-shaped, in contact with first two supraoculars; four supraoculars; frontoparietals in contact posterior to frontal, contacting second, third and fourth supraoculars anterolaterally and parietals and interparietal posteriorly; frontoparietals non-overlapping; interparietal diamond-shaped, large, slightly projecting posteriorly, eyespot in posterior projection; parietals large, in medial contact posterior to interparietal, contacting fourth supraocular anteriorly; enlarged nuchal scales absent; nasals small, widely separated, trapezoidal, contacting rostral anteriorly, frontonasal dorsally, first loreal posteriorly, first supralabial


Figure 8. Holotype of Tytthoscincus jaripendek sp. nov. (LSUHC 11680) from Robinson Falls, Cameron Highlands, Pahang, Peninsular Malaysia.
ventrally; nostril in centre of nasal; supranasals absent; two loreals taller than wide, posterior loreal smallest; upper and lower preocular present; lower preocular followed by a series of suboculars; nine superciliaries, posterior superciliary elongate and projecting dorsomedially; six supralabials, third, fourth and fifth below eye; two postsupralabials; three primary temporals; three secondary temporals, uppermost not contacting parietals; lower eyelid transparent, scaly, no enlarged central window; mental twice as wide as long; single, large postmental, contacting first infralabial on each side; two enlarged pairs of chin shields posterior to postmental, anterior pair contacting medially, posterior pair widely separated posteriorly by a single scale; anterior and posterior chinshield pairs contacting first, second and third infralabials; five infralabials; external ear opening 0.6 mm less than onehalf diameter of eye, subcircular, lacking anterior lobules; tympanum very shallow, pigmented.

Body scales smooth, cycloid, imbricate; ventral scales slightly larger than dorsal scales; 33 longitudinal scale rows around midbody; 73 paravertebral scales; 65 ventral scales; slightly enlarged precloacal scales; tail slightly compressed laterally; subcaudals same size as dorsal caudals; limbs very thin, short (FL/ $\mathrm{SVL}=0.22 ; \mathrm{HDL} / \mathrm{SVL}=0.32$ ), widely separated when adpressed; scales of dorsal surface slightly larger than those of ventral surface; palmar and plantar scales slightly raised; and digits short, scales of dorsal surfaces in a single row, subdigital lamellae bicarinate, ten on fourth toe; six on third finger.

Coloration: Overall dorsal ground colour of head, body, limbs and tail brown; head speckled with lightcoloured markings; labials bearing light centres; dorsum, posterior portion of flanks and tail bearing small, light-coloured spots; limbs mottled; anterior of flanks to posterior margin of jaws dull-orange bearing irregularly shaped speckles and elongate markings; all ventral surfaces dull-yellow, immaculate; lateral margin of gular region stippled with dark brown; and palmar and plantar surfaces, subtibial, subforearm and subcaudal regions heavily stippled with dark brown.

Variation: The paratypes closely approach the holotype in all aspects of coloration except LSUHC 12754 is considerably darker in overall in dorsal coloration (Fig. 11). Meristic variation is presented in Table 9.

Distribution: Tytthoscincus kakikecil sp. nov. is known from the town of Fraser's Hill at Richmond Road and the Telecom Loop and from the upper elevations of Genting Highlands at Ulu Kali, Pahang, Peninsular Malaysia (Fig. 9).

Natural history: Tytthoscincus kakikecil sp. nov. inhabits leaf-litter in hill dipterocarp and mossy forests from at least 1305-1670 m in elevation (Fig. 12). All specimens at Fraser's Hill were collected by raking leaves that had accumulated on earthen banks along the road or by turning rocks and logs buried in leaf-litter. LSUHC 12754 from Genting Highlands was found abroad at night in a dirt parking lot next to a rubbish

Table 8. Meristic and ratiometric characters of the type series of Tytthoscincus jaripendek sp. nov.

|  | LSUHC 11679 | LSUHC 11680 | LSUHC 11681 | LSUHC 11987 |
| :---: | :---: | :---: | :---: | :---: |
|  | Paratype | Holotype | Paratype | Paratype |
| Supralabials | 6 | 6 | 6 | 6 |
| Infralabials | 5 | 5 | 5 | 5 |
| Supraoculars | 4 | 4 | 4 | 4 |
| Frontoparietal contacting supraoculars 2, 3 and 4 | Yes | Yes | Yes | Yes |
| Superciliaries | 81 | 81 | 81 | 81 |
| Prefrontals in contact | Yes | Yes | Yes | No |
| Loreals | 1 | 2 | 1 | 1 |
| Tympanum shallow, pigmented | Yes | Yes | Yes | Yes |
| Midbody scale rows | 29 | 28 | 30 | 30 |
| Dorsal scales striated | No | No | Weakly | No |
| Paravertebral scale rows | 64 | 65 | 65 | 63 |
| Ventral scale rows | 61 | 60 | 62 | 60 |
| Enlarged pectoral scales | Yes | Yes | Yes | Yes |
| Enlarged precloacal scales | Yes | Yes | Yes | Yes |
| 3rd finger lamellae | 5 | 5 | 5 | 6 |
| 4th toe lamellae | 9 | 10 | 9 | 10 |
| Lamellae texture | Keeled | Keeled | Keeled | Keeled |
| HL | 6.2 | 6.1 | 6.2 | 6.3 |
| TD | 0.8 | 0.8 | 0.7 | 0.7 |
| TD/HL | 0.12 | 0.13 | 0.12 | 0.11 |
| HL/SVL | 0.19 | 0.19 | 0.18 | 0.19 |
| AXG | 18.6 | 18.4 | 18.8 | 18.5 |
| AXG/SVL | 0.56 | 0.57 | 0.55 | 0.56 |
| FL | 6.6 | 6.6 | 6.2 | 7.0 |
| FL/SVL | 0.2 | 0.21 | 0.18 | 0.21 |
| HDL | 10.7 | 10.8 | 10.5 | 10.9 |
| HDL/SVL | 0.32 | 0.33 | 0.31 | 0.33 |
| SVL | 32.9 | 32.2 | 33.9 | 32.7 |
| Sex | Male | Male | Female | Female |

AXG, axilla-groin length; FL, forelimb length; HDL, hind limb length; HL, head length; SVL, snout-vent length; TD, tympanum diameter.
pile on the edge of a mossy forest that we presume to be its natural habitat (as opposed to the rubbish pile).

Etymology: The specific epithet kakikecil is derived from the Malay words kaki meaning leg or foot and kecil meaning small and refers to this species' diminutive limbs.

Comparisons: Tytthoscincus kakikecil sp. nov. can be differentiated from all other species of Tytthoscincus of the montane clade (Table 6), except its sister species $T$. jaripendek sp. nov., by having a shallow, pigmented tympanic depression vs. a well-developed, deeply recessed, weakly pigmented tympanum. From T. jaripendek sp. nov., it differs by having 67-73 vs. 63-65 paravertebral scales; $65-69$ vs. $60-62$ ventral scales; having as opposed to lacking enlarged pectoral scales; and having statistically significant differences in the mean number of PV, VS, 3TL and FL/SVL (Tables 6, 7).

Tytthoscincus kakikecil sp. nov. also bears statistically significant mean differences of all ten characters in varying combinations across all other species (Table 6). Meristic and body shape trends of these characters among species in the montane clade are presented in Figures 4 and 5.

Remarks: Tytthoscincus kakikecil sp. nov. is known from separate populations at Fraser's Hill (LSUHC 11769-70, 11772) and Ulu Kali, Genting Highlands (LSUHC 12754, FMNH 19945), $\sim 38 \mathrm{~km}$ apart (Fig. 8). We were unsuccessful in obtaining DNA sequences from FMNH 19945, also from Ulu Kali. Sly (1976) considered this specimen to be T. butleri but provided no character evidence and Grismer (2011), who did not examine the specimen, considered it to be T. bukitensis based on the fact that T. butleri occurs in different mountain range and T. bukitensis occurs 38 km to the north across habitable


Figure 9. Distribution of the species of the montane clade of Tytthoscincus in Peninsular Malaysia.
terrain (multiple point measurements made from Google Earth along an elevation profile) at Fraser's Hill in the same mountain range. FMNH 19945 plotted next to a sequenced specimen of T. kakikecil sp. nov. from Ulu Kali (LSUHC 12754) in the PCAs and DAPCs. Furthermore, based on probability estimates generated from retained posteriors from the concatenated DAPC, FMNH 19945 clustered with T. kakikecil sp. nov. with a $72 \%$ probability. However, these populations (Fraser's Hill and Ulu Kali) differ discretely in their numbers of midbody scales (33 vs. 30 or 31 , respectively), TD (TD/HL = 0.10-0.11 vs. 0.12 , respectively), $\mathrm{HL}(\mathrm{HL} / \mathrm{SVL}=0.19-0.21$ vs. 0.18 , respectively) and HDL (HDL/SVL $=0.30-0.32$ vs. 0.34 , respectively). But, based on a Welch two-sample $t$-test, only the means of the HDL/SVL ratios are significantly different ( $P=0.035$ ). In the absence of molecular data, it would be tempting to consider these populations as separate species regardless of their small sample sizes - an error that has been made before with species of Tytthoscincus (e.g. Grismer, 2007, 2008). However, the molecular data estimate these two populations have only a $1.9 \%$ sequence divergence between them and they cluster
tightly together in the concatenated PCA and DAPC (Fig. 2). This shallow divergence across $\sim 38 \mathrm{~km}$ of habitable terrain is in stark contrast to the $6.2 \%$ divergence found between individuals of T. martae sp. nov. (LSUHC 12749,12751 ) and T. bukitensis (LSUHC 12750) found only a few meters apart at Awana Road, Genting Highlands (see below). This suggests gene flow still exists between these populations and the morphological differences are due to small sample sizes ( $n=3$ for Fraser's Hill and $n=2$ for Genting Highlands) and/or sampling error. We predict that data from additional specimens from both localities and the intervening regions will most probably eclipse their current morphological differences.

## Tytthoscincus martae sp. nov. <br> Hindu Temple Forest Skink <br> Mengkarung Hutan Kuil Hindu

(Fig. 13; TABLE 10)
Holotype: Adult male (LSUHC 12688) collected at the Hindu Temple 1.2 km south of the town of Fraser's Hill


Figure 10. Microhabitat of Tytthoscincus jaripendek sp. nov. at Robinson Falls, Cameron Highlands, Pahang, Peninsular Malaysia.


Figure 11. Type specimens of Tythoscincus kakikecil sp. nov. from Pahang, Peninsular Malaysia. A, paratype LSUHC 12754 from Ulu Kali, Genting Highlands. B, holotype LSUHC 11769 from Richmond Road, Fraser's Hill.
on the road to the Gap, Pahang, Peninsular Malaysia ( $\mathrm{N} 03^{\circ} 42.099^{\prime}$, $\mathrm{E} 101^{\circ} 44.090^{\prime} ; 1078 \mathrm{~m}$ in elevation) by Evan S. H. Quah, Marta S. Grismer and L. Lee Grismer on 23 March 2016.

Paratypes: LSUHC 12517-18 collected on 3 December 2015 and LSUHC 12749 and 12751 collected on 21 June 2016 from the Awana Road, Genting Highlands, Pahang, Peninsular Malaysia (N $3{ }^{\circ} 24.036^{\prime}$, E $101^{\circ} 47.212^{\prime}$; 1052

Table 9. Meristic and ratiometric characters of the type series of Tytthoscincus kakikecil sp. nov.

|  | $\begin{aligned} & \text { LSUHC } \\ & 11769 \end{aligned}$ | $\begin{aligned} & \text { LSUHC } \\ & 11770 \end{aligned}$ | $\begin{aligned} & \text { LSUHC } \\ & 11772 \end{aligned}$ | LSUHC 12754 | FMNH 19945 | ZRC 2.5944 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | Holotype | Paratype | Paratype | Paratype |  |  |
|  | Fraser's Hill | Fraser's <br> Hill | Fraser's Hill | Genting Highlands | Genting Highlands | Cameron Highlands |
| Supralabials | 6 | 6 | 6 | 6 | 6 | 6 |
| Infralabials | 5 | 5 | 5 | 5 | 5 | 5 |
| Supraoculars | 4 | 4 | 4 | 4 | 4 | 4 |
| Frontoparietal contacting supraoculars 2, 3 and 4 | Yes | Yes | Yes | Yes | Yes | Yes |
| Superciliaries | 911 | $9 \times 1$ | 911 | $9 \times 2$ | 911 | 81 |
| Prefrontals in contact | Yes | Yes | Yes | Yes | Yes | Yes |
| Loreals | 2 | 2 | 2 | 2 | 2 | 1 |
| Tympanum shallow, pigmented | Yes | Yes | Yes | Yes | Yes | / |
| Midbody scale rows | 33 | 33 | 33 | 30 | 31 | 33 |
| Dorsal scales striated | No | No | No | No | No | No |
| Paravertebral scale rows | 73 | 69 | 67 | 71 | 71 | 68 |
| Ventral scale rows | 65 | 66 | 69 | 66 | 68 | 68 |
| Enlarged pectoral scales | Yes | Yes | Yes | Yes | No | Yes |
| Enlarged precloacal scales | Yes | Yes | Yes | Yes | Yes | Yes |
| 3rd finger lamellae | 6 | 6 | 6 | 6 | 6 | 6 |
| 4th toe lamellae | 10 | 10 | 10 | 10 | 10 | 10 |
| Lamellae texture | Keeled | Keeled | Keeled | Keeled | Keeled | Keeled |
| HL | 6.2 | 6.7 | 5.1 | 5.9 | 7.1 | 6.8 |
| TD | 0.6 | 0.8 | 0.6 | 0.7 | 0.8 | 0.8 |
| TD/HL | 0.10 | 0.11 | 0.11 | 0.12 | 0.12 | 0.11 |
| HL/SVL | 0.2 | 0.19 | 0.21 | 0.18 | 0.18 | 0.19 |
| AXG | 19.9 | 21.7 | 12.6 | 17.7 | 21.89 | 19.3 |
| AXG/SVL | 0.65 | 0.60 | 0.52 | 0.54 | 0.56 | 0.54 |
| FL | 6.9 | 7.7 | 5.6 | 7.7 | 7.9 | 6.5 |
| FL/SVL | 0.22 | 0.22 | 0.23 | 0.23 | 0.2 | 0.18 |
| HDL | 9.9 | 11.0 | 6.4 | 11.3 | 13.1 | 10.1 |
| HDL/SVL | 0.32 | 0.31 | 0.3 | 0.34 | 0.34 | 0.3 |
| SVL | 30.8 | 35.9 | 24.2 | 33.0 | 39.2 | 36.2 |
| Sex | Male | Male | 1 | Male | Female | Female |

(/), data unobtainable. AXG, axilla-groin length; FL, forelimb length; HDL, hind limb length; HL, head length; SVL, snout-vent length; TD, tympanum diameter.
m in elevation) by Evan S. H. Quah and L. Lee Grismer. ZRC 3.6246 is from the same locality as the holotype and was collected by L. Lee Grismer, Perry L. Wood, Jr. and Timothy M. Youmans on 26 August 2006.

Additional specimen examined: LSUHC 12072 from the Awana Road, Genting Highlands, Pahang, Peninsular Malaysia.

Diagnosis: Tytthoscincus martae sp. nov. can be differentiated from all other species of Tytthoscincus in the montane clade except $T$. bukitensis by having the combination of $8^{`} 1,9^{`} 1$ or $10^{`} 2$ superciliaries; a deep, weakly pigmented tympanum; enlarged pectoral scales; 31-35
midbody scales; 70-74 paravertebral scales; 61-72 ventral scales; keeled, subdigital lamellae; seven subdigital lamellae on the third finger; 12 or 13 subdigital lamellae on the fourth toe; TD/HL $=0.12-0.16 ; \mathrm{HL} /$ $\mathrm{SVL}=0.18-0.26 ; \mathrm{AXG} / \mathrm{SVL}=0.47-0.55 ; \mathrm{FL} / \mathrm{SVL}=0.21-$ 0.26 ; $\mathrm{HDL} / \mathrm{SVL}=0.33-0.38$; and a maximum SVL of 44.0 mm (Table 6). All non-ratiometric characters are scored across all other Tytthoscincus and species of Sphenomorphus suspected of being Tytthoscincus in Grismer et al. (2016a: 237) and Karin et al. (2016: 416).

Description of holotype: Adult female, SVL 34.9 mm ; non-regenerated tail length 48.4 mm ; AXG 19.1 mm ;


Figure 12. Microhabitat of Tythoscincus kakikecil sp. nov. from Pahang Peninsular Malaysia. A, hill dipterocarp forests at Jeriau, Fraser's Hill (1305 m elevation). B, mossy forest at Ulu Kali, Genting Highlands ( 1670 m elevation).

HL 7.2 mm ; head width 4.5 mm ; snout to anterior margin of foreleg 12.4 mm ; rostral wider than long, in broad contact with frontonasal; frontonasal wider than long; prefrontals large, in point contact on midline; frontal elongate, diamond-shaped, in contact with first two supraoculars; four supraoculars; frontoparietals in contact posterior to frontal, contacting second, third and fourth supraoculars anterolaterally and parietals and interparietal posteriorly; interparietal diamondshaped, large, slightly projecting posteriorly, eyespot in posterior projection; parietals large, in medial contact posterior to interparietal, contacting fourth supraocular anteriorly; one slightly enlarged nuchal scale left of midline; nasals small, widely separated, trapezoidal, contacting rostral anteriorly, frontonasal dorsally, first loreal posteriorly, first supralabial ventrally; nostril in centre of nasal; supranasals absent; two loreals, posterior loreal taller than wide; upper and lower preocular present; lower preocular followed by a series of suboculars; eight superciliaries, posterior
superciliary elongate and projecting dorsomedially; seven suboculars; six supralabials, third, fourth and fifth below eye; two postsupralabials; three primary temporals; three secondary temporals, uppermost not contacting parietals; lower eyelid transparent, scaly, no enlarged central window; mental twice as wide as long; single, large postmental, contacting first infralabial on each side; two enlarged pairs of chin shields posterior to postmental, anterior pair contacting medially, posterior pair widely separated posteriorly by a single scale; anterior and posterior chinshield pairs contacting first, second and third infralabials; six infralabials; external ear opening 0.9 mm , subcircular, lacking anterior lobules; and tympanum deeply set, non-pigmented.

Body scales smooth, cycloid, imbricate; ventral scales slightly larger than dorsal scales; 31 longitudinal scale rows around midbody; 73 paravertebral scales; 69 ventral scales; slightly enlarged median precloacal scales overlapping outer precloacal scales; tail slightly compressed


Figure 13. Type specimens of Tytthoscincus martae sp. nov. from Pahang, Peninsular Malaysia. A, holotype LSUHC 12688 from the Hindu Temple, Fraser's Hill. B, paratype LSUHC 12749 from Awana Road, Genting Highlands.
laterally; subcaudals slightly larger than dorsal caudals; limbs relatively robust, short ( $\mathrm{FL} / \mathrm{SVL}=0.22$; HDL/ SVL $=0.33$ ), widely separated when adpressed; scales of dorsal surface slightly larger than those of ventral and posterior surfaces; palmar and plantar scales slightly raised; and digits moderate in length, scales on dorsal surfaces in single row, subdigital lamellae bicarinate, seven on third finger, 12 on fourth toe.

Coloration in life: Overall dorsal ground colour of head, body, limbs and tail dark brown; head speckled with light-coloured markings; supralabials three and four bearing a light central spot; infralabials weakly banded; dorsum, posterior portion of flanks and tail bearing small, light-coloured spots; limbs mottled; anterior of flanks to posterior margin of jaws dull-orange bearing irregularly shaped dark speckles; thick, dark, irregularly margined postorbital stripe extending from eye to groin, merging
with dorsal ground colour posterior to hind limbs, bordered dorsally above anterior portion of flanks by a thick, immaculate, dull-orange stripe; all ventral surfaces dull-yellow to beige, generally immaculate; distal portions of forelimbs and palmer surfaces dark; planter surface dark; subcaudal region stippled with dark brown.

Variation: The paratypes closely approach the holotype in all aspects of coloration except for their supralabial scales being distinctly barred (Fig. 13). The dorsum of LSUHC 12749 is slightly darker overall and the spotting less bold. The juveniles LSUHC 12072 (SVL 22.4 mm ) and 12518 (SVL 26.5 mm ) show no difference in colour pattern or pattern boldness, indicating ontogenetic changes in coloration and pattern are absent. LSUHC 12518 has a damaged neck, LSUHC 12749 has a damaged gular region and LSUHC 12751 has an incomplete tail. Meristic variation is presented in Table 10.

Table 10. Meristic and ratiometric characters of the type series of Tytthoscincus martae sp. nov.

|  | $\begin{aligned} & \text { LSUHC } \\ & 12517 \end{aligned}$ | LSUHC | LSUHC | LSUHC | LSUHC |  | $\begin{aligned} & \text { LSUHC } \\ & 12072 \end{aligned}$ |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  | 12518 | 12688 | 12749 | 12751 | 3.6246 |  |
|  | Paratype | Paratype | Holotype | Paratype | Paratype | Paratype |  |
|  | Awana | Awana | Fraser's | Awana | Awana | Fraser's | Awana |
|  | Road | Road | Hill | Road | Road | Hill | Road |
| Supralabials | 6 | 6 | 6 | 6 | 6 | 6 | 6 |
| Infralabials | 5 | 5 | 6 | 6 | 6 | 5 | 5 |
| Supraoculars | 4 | 4 | 4 | 4 | 4 | 4 | 4 |
| Frontoparietal contacting supraoculars 2, 3 and 4 | Yes | Yes | Yes | Yes | Yes | Yes | Yes |
| Superciliaries | $9 ` 1$ | 911 | $8{ }^{1} 1$ | $9 \times 2$ | 102 | 102 | 811 |
| Prefrontals in contact | Point | Yes | Point | Yes | Yes | Yes | Yes |
| Loreals | 2 | 2 | 2 | 2 | 2 | 2 | 2 |
| Tympanum deep, non-pigmented | Yes | Yes | Yes | Yes | Yes | Yes | Yes |
| Midbody scale rows | 34 | / | 31 | 35 | 33 | 33 | 34 |
| Dorsal scales striated | No | No | No | No | No | No | No |
| Paravertebral scale rows | 74 | 1 | 73 | 70 | 73 | 73 | 71 |
| Ventral scale rows | 70 | 68 | 69 | 69 | 72 | 61 | 70 |
| Enlarged pectoral scales | No | No | Yes | Yes | Yes | Yes | Weak |
| Enlarged precloacal scales | Yes | Yes | Yes | Yes | Yes | Yes | Yes |
| 3rd finger lamellae | 7 | 7 | 7 | 7 | 7 | 7 | 7 |
| 4th toe lamellae | 12 | 13 | 12 | 13 | 13 | 13 | 13 |
| Lamellae texture | Keeled | Keeled | Keeled | Keeled | Keeled | Keeled | Keeled |
| HL | 6.6 | 0.12 | 7.2 | 7.59 | 6.9 | 8.4 | 5.85 |
| TD | 0.81 | 0.23 | 0.9 | 1.21 | 0.99 | 1.02 | 0.8 |
| TD/HL | 29.76 | 6.33 | 0.12 | 0.16 | 0.52 | 0.12 | 0.14 |
| HL/SVL | 29.76 | 0.24 | 0.21 | 0.18 | 0.14 | 0.19 | 0.26 |
| AXG | 22.8 | 12.8 | 19.1 | 22.8 | 18.5 | 20.73 | 11.2 |
| AXG/SVL | 0.53 | 0.48 | 0.55 | 0.53 | 0.52 | 0.47 | 0.50 |
| FL | 29.76 | 9.22 | 7.69 | 10.93 | 0.19 | 9.28 | 5.88 |
| FL/SVL | 29.76 | 0.35 | 0.22 | 0.26 | 8.08 | 0.21 | 0.26 |
| HDL | 29.76 | 26.5 | 11.43 | 15.73 | 0.23 | 14.34 | 8.3 |
| HDL/SVL | 29.76 | 0.12 | 0.33 | 0.38 | 13.04 | 0.33 | 0.37 |
| SVL | 29.76 | 26.5 | 34.9 | 42.78 | 35.8 | 44 | 22.4 |
| Sex | / | / | Female | Female | Male | Male | / |

(/), data unobtainable. AXG, axilla-groin length; FL, forelimb length; HDL, hind limb length; HL, head length; SVL, snout-vent length; TD, tympanum diameter.

Distribution: Tytthoscincus martae sp. nov. is known from the type locality at the Hindu temple south of the town of Fraser's Hill and Awana Road at Genting Highlands, Pahang, Peninsular Malaysia (Fig. 9). However, it is likely to range further north of Fraser' Hill and south of Genting Highlands.

Natural history: Tytthoscincus martae sp. nov. inhabits leaf-litter in hill dipterocarp forests from at least $991-1239 \mathrm{~m}$ in elevation (Fig. 14). Specimens from Awana Road, Genting Highlands were collected by raking leaves that had accumulated at the foot of an earthen bank and by turning logs and rocks buried in leaf-litter. Lizards were most commonly found in areas
where moisture from the hillside had seeped down to the edge of the road. During overcast days, lizards were seen running swiftly through small rocks and root tangles on the forest floor beneath the canopy in the nearby forest. However, we do not know if these specimens were T. martae sp. nov. or T. bukitensis. At Fraser's Hill, specimens were found along the edges of seeps or small streams in damp microhabitats. ZRC 3.6246 was found beneath a log and LSUHC 12688 was found buried in the leaf-litter, both near the edges of small drainages.

Etymology: The specific epithet martae is a matronym honouring my (LLG) wife Marta S. Grismer for her long, non-wavering commitment and support of my


Figure 14. Microhabitat of Tytthoscincus martae sp. nov. from Pahang Peninsular Malaysia. A, Hindu Temple, Fraser's Hill. B, Awana Road, Genting Highlands.
work for many years, her efforts in the field on many trips across Southeast Asia and Latin America and for generating the morphological data sets for a number of our studies.

Comparisons: Tytthoscincus martae sp. nov. can be differentiated from T. jaripendek sp. nov. and T. kakikecil sp. nov. by having as opposed to lacking a deeply recessed, weakly pigmented tympanic membrane. Tytthoscincus martae sp. nov. differs discretely from T. perhentianensis and T. jaripendek sp. nov. by having $31-35$ vs. 28-30 (collectively) midbody scales and $70-74$ vs. 63-66 (collectively) paravertebral scales. It is further discretely differentiated from $T$. jaripendek sp. nov., T. perhentianensis and T. kakikecil sp. nov. by having seven vs. five or six subdigital lamellae on the third toe. It is further discretely differentiated from $T$. jaripendek sp. nov., T. perhentianensis, T. kakikecil sp. nov., T. temengorensis and T. ishaki by having 12 or 13 vs. 9-11 (collectively) subdigital lamellae on the fourth
toes. Tytthoscincus martae sp. nov. also bears statistically significant mean differences of all ten characters in varying combinations across all other species (Table 6). Numeric trends of these characters among species in the montane clade are presented in Figures 4 and 5.

Although the molecular phylogeny suggests that Tytthoscincus martae sp. nov. is a separate species being that it is not nested within any other species, it cannot be discretely nor statistically ( $P<0.05$ ) differentiated morphologically from T. bukitensis and both are sympatric at Fraser's Hill and syntopic at Awana Road, Genting Highlands. The genetic data infer that the monophyly (exclusivity) of each species is strongly supported (1.00/100 and 1.00/96, respectively; Fig. 2) and they share a modest 5.4-7.4\% uncorrected pairwise sequence divergence between them. More importantly, both the BI and ML analyses infer T. martae sp. nov. and T. bukitensis are not sister species - although support for this is moderate ( $0.89 / 77$ ) - so this conundrum cannot


Figure 15. Concatenated PCA analysis showing the placement of Tytthoscincus cophias, its tympanic depression, and overall body stature.
be solved by considering them conspecific because it would render T. bukitensis paraphyletic. Most notably, however, is that certain individuals from each species are more closely related to conspecifics from the other locality than they are to individuals of the other species found only meters apart at the same locality. For example, T. martae sp. nov. LSUHC 12749 from Awana Road is more closely related to T. martae sp. nov. LSUHC 12688 from Fraser's Hill, 38 km to the north than it is to T. bukitensis LSUHC 12750 collected within 15 min of T. martae sp. nov. LSUHC 12749 on the same day less than 5 m away in the same pile of leaf-litter (Fig. 2). Complicating this issue is that the molecular evidence demonstrates that the current paratype of T. bukitensis from Fraser's Hill (ZRC 3.6246) must be considered part of T. martae sp. nov. and is not closely related to the holotype of T. bukitensis from Fraser's Hill (ZRC 2.6245). Thus, the type series of T. bukitensis is composed of two different species that are not each other's closest relatives. Therefore, ZRC 3.6246 is removed here from the type series of T. bukitensis and transferred to the species T. martae sp. nov.

If T. bukitensis and T. martae sp. nov. were reciprocally monophyletic and occurred exclusively at different localities (i.e. they were not sympatric), we would not recognize them as separate species and posit that their
genetic distinction represents isolation by distance and individuals from the intervening regions would need to be examined and sequenced. Such is the case with the allopatric sister species Hemiphyllodactylus titiwangsaensis and H. cf. titiwangsaensis - from the same mountain range as T. bukitensis and T. martae sp. nov. - that are morphologically indistinguishable yet have an uncorrected pairwise sequence divergence between them of $12.8 \%$ (Cobos et al., 2016). However, T. bukitensis and T. martae sp. nov. are not sister species and individuals of each occur at both localities and are even syntopic at one - strongly suggesting they are separate, highly convergent, genetically exclusive lineages.

Tytthoscincus martae sp. nov. and T. bukitensis are differentiated in their $50 \%$ combined upper and lower quartile ranges of midbody scales that only narrowly approach one another at 32.9 (Fig. 5). Although the mean values of their respective counts are not statistically significantly different, this is due to a high outlying value of 35 for midbody scales in T. bukitensis (LSUHC 12750). Given that these two species are syntopic in at least one part of their range and the only way to unequivocally separate them is with a molecular analysis, we consider them cryptic species (see below). We are well aware of the potential pitfalls using a single
mitochondrial marker in cases such as these and that the use of nuclear markers might resolve the issue of potential conspecificity. However, genomic data would be the best way to test these hypotheses and plans are in progress to do so.

Specimens conferre: We were unsuccessful in obtaining DNA sequences from ZRC 2.5944 from Telom Valley, Cameron Highlands considered by Grismer (2011) to be T. bukitensis and LSUHC 12072 from the Awana Road, Genting Highlands that is newly reported here. Based on retained posteriors from a separate concatenated DAPC analyses, ZRC 2.5944 from Telom Valley clustered with T. kakikecil sp. nov. with only a $43 \%$ probability and was closest in morphospace to individuals from Fraser's Hill. Unfortunately, due to its state of preservation, we could not confirm the presence of reduced tympanae. The left tympanum looks as though it may be deeply inset but the right tympanum did not. We are also concerned with the accuracy of the mensural data. ZRC 2.5944 was collected in 1935 and the specimen is quite limp and in overall poor condition so we cannot conclude that the measurements taken from this specimen are congruous with those from freshly preserved specimens. Some scale counts (MB, PV and VS), however, are outside the range of those of $T$. jaripendek sp. nov. (the new species from Cameron Highlands) but consistent with those of T. kakikecil sp. nov. (Tables 6 and 9). We consider this specimen as $T$. cf. kakikecil sp. nov. but do not include it as part of the type series.

The retained DAPC posteriors indicate that LSUHC 12072 from the Awana Road, Genting Highlands clusters with T. martae sp. nov. with only a $59 \%$ probability which is consistent with its meristic and mensural data (Table 10). Furthermore, it plots within T. martae and outside of T. bukitensis in the concatenated PCA and the DAPC. We consider this specimen as $T$. cf. martae sp. nov. but do not include it as part of the type series.

Sphenomorphus cophias: Boulenger (1908) described Sphenomorphus cophias from a single specimen (BMNH 1946.8.3.15) collected on Gunong (=Gunung) Tahan in the Banjaran Timur at $\sim 1000 \mathrm{~m}$ in elevation and Smith (1922) reported a second specimen (which can no longer be located) from the same mountain at $\sim 1100 \mathrm{~m}$ in elevation. Both specimens are small ( 36 mm and 29 mm SVL, respectively) and Boulenger (1908) reported that the ear of BMNH 1946.8.3.15 was hidden and replaced by a depression (Fig. 15). Lim (1998) and Grismer (2011) re-examined the holotype and found that along with it being less than 45 mm SVL, it had small temporal scales that were the same size and shape as the lateral body scales; and it had short digits with toe IV being equal in length to toe

III - characters Linkem et al. (2011) used to diagnose the genus. However, based on aspects of head scalation and lower eyelid morphology, Shea (in literature, 2016) believes it may be an undescribed species of Lipina. Nonetheless, data from the holotype are presented in Table 6. If it belongs in the genus Tytthoscincus on the basis of Linkem et al. (2011), its low numbers of digital lamellae ( $3 \mathrm{FL}=5 ; 4 \mathrm{TL}=10$ ), very short limbs ( $\mathrm{FL} / \mathrm{SVL}=0.17 ; \mathrm{HDL} / \mathrm{SVL}=0.28$ ), elongate body $(\mathrm{AXG} / \mathrm{SVL}=0.58)$ and lack of a tympanum further extend the trends of these characters shown for T. jaripendek sp. nov. and T. kakikecil sp. nov. and indicate this species has a semi-fossorial ecomorphology as demonstrated by its placement in the PCA and DAPC analyses (Fig. 15). This, coupled with its upland locality in hill dipterocarp forests, suggests it may belong to the montane clade allied to the semi-fossorial lineage. However, given that there are at least two clades of distantly related species of Tytthoscincus in Peninsular Malaysia and the propensity for convergence of semi-fossorial ecomorphology in skinks in general (see below), we do not propose a formal placement in either clade in the absence of molecular data.

## DISCUSSION

## ECOMORPHOLOGY

Within tropical ecosystems, leaf-litter is one of the most ubiquitous microhabitats in which a number of amphibians and reptiles have become adapted (Fauth, Crother \& Slowinski, 1989; Vitt \& Caldwell, 1994; Vonesh, 2001; Whitfield et al., 2007). In Peninsular Malaysia, this is particularly true for lizards where at least 35 species are considered leaf-litter inhabitants (Grismer, 2011; Grismer et al., 2011; Grismer \& Quah, 2015). As more species are described from montane regions in Peninsular Malaysia, it is becoming clear that the ecological relationships among them is far more complex than previously considered (Grismer, 2007; Grismer \& Quah, 2015) and this is especially true for closely related, syntopic, leaf-litter species (Grismer \& Quah, 2015). The fact that multiple lineages of closely related, highly specialized species such as Tytthoscincus occur in close sympatry and syntopy in such a restrictive microhabitat indicates that resource partitioning must be highly regulated (Schoener, 1974; Walter, 1991; Goodman, 2007). This may also be the case for T. leproauriculatusand T. batupanggah that occur within 100 m of one another in the uplands of Borneo at Gunung Penrissen, Sarawak (Karin et al., 2016). Knowing little about Tytthoscincus life history makes it difficult to test hypotheses as to how these species partition resources. However, because correlations between morphology and microhabitat structure underscore the link between a phenotype and an
environment (e.g. Herrel, Meyers \& Vanhooydonck, 2001; Goodman, Miles \& Schwarzkopf, 2008; Losos, 2009; Gomes, Carretero \& Kaliontzopoulou, 2016), inferences can be made based on quantifiable differences in ecomorphology (Vittet al.,1997;Vanhooydonck, Van Damme \& Aerts, 2000; Herrel et al., 2001; Losos, 2009; Lee, Skinner \& Camacho, 2013).

It has been demonstrated that body elongation, limb and digital reduction, shortening of the head and closure of the ear opening are correlated phenomena that have occurred multiple times during the evolution of fossorial and semi-fossorial skinks (Greer, 1989; Schmitz et al., 2005; Brandley, Huelsenbeck \& Wiens, 2008; Siler \& Brown, 2011; Lee et al., 2013; Wu et al., 2015). For small skinks (SVL < 50 mm ), these characteristics have been shown or suggested to be adaptive for locomoting within or beneath the surface of leaf-litter or through other types of restrictive vegetation (Van Damme et al., 1998; Melville \& Swain, 2000; Kohlsdorf et al., 2001; Herrel et al., 2002; Siler \& Brown, 2011) and that various species have differing combinations of these characters in varying degrees of development (Pianka, 1969; Melville \& Swain, 2000; Brandley et al., 2008). Tytthoscincus, in general, aligns well with this ecomorphological paradigm in that all species have relatively elongate bodies, short limbs and digits, and reside in leaf-litter and two species, $T$. jaripendek sp. nov. and T. kakikecil sp. nov., extend these trends further by having relatively shorter limbs and digits (i.e., fewer lamellae), longer bodies, shorter heads and vestigial tympanae (Tables 6, 67; Figs 4, 5, 7). The concatenated PCA, DAPC and Tukey HSD test indicate that T. jaripendek sp. nov. and T. kakikecil sp. nov. - who have varying combinations of these characters - cluster separately from all other species (Fig. 6) and differ statistically (Table 7) from various combinations of species in many characters including those associated with digit and limb reduction (3TL, 4FL, FL, HDL), elongation of the body (AXG) and shortening of the head (HL).

As to how the ecomorphological differences of the semi-fossorial species T. kakikecil sp. nov. may allow it to function differently in leaf-litter at Fraser's Hill where it is sympatric and probably syntopic with the leaf-litter generalists T. bukitensis is speculative. Melville \& Swain (2000), Bergmann \& Irschick (2010) and Lee et al. (2013) noted that shorter-bodied skinks were faster and spent more time on the surface, whereas elongate, limb-reduced species spent more time below the surface and were possibly more cryptic and substrate-specific. Although T. kakikecil sp. nov. and T. bukitensis can be 'raked up' from the same pile of leaves, we hypothesize that they may be spending different amounts of time foraging at different levels in the leaf-litter mass which can be densely packed and as thick as 0.5 m . At both Fraser's Hill and

Awana Road, we often see T. bukitensis or T. martae sp. nov. (or both) above ground moving through and across the surface of the leaf-litter but have never observed T. kakikecil sp. nov. (nor have we seen T. jaripendek sp. nov.) above ground. It is tempting to posit that these ecomorphological differences evolved as a result of competition, forcing these species into different microhabitats (e.g. Vitt et al., 1997; Herrel et al., 2001) and may have even been the initial driving force behind the speciation of the two major ecomorphological lineages within Tytthoscincus (Fig. 2). If competition was or is the driving force for the evolution of these two ecomorphologies, then we hypothesize we should find a yet undiscovered, leaf-litter generalist co-occurring with T. jaripendek sp. nov. at Cameron Highlands.

## CRYPTIC SPECIES

Ironically, many studies employing integrative taxonomic analyses that purport to reveal 'cryptic species' go on to provide traditional, discrete, morphological diagnoses for those 'cryptic species' using taxon-specific characters that have traditionally been used to diagnose other species within those groups - diagnoses that stand alone from the molecular evidence and were generated retrospectively following prudent morphological analyses (e.g. Wood et al., 2009; Grismer et al., 2013a, b; Loredo et al., 2013). So are these species truly 'cryptic'? The definition of cryptic species ranges broadly from two different species erroneously classified under the same name (Bickford et al., 2006) to species that are morphologically indistinguishable but well delineated on the basis of genetic markers (Jöger \& Schroll, 2013). Most herpetologists would probably consider Tytthoscincus species to be 'cryptic' in that they are small, elongate, little brown skinks that move rapidly in and out of the leaf-litter and are hard to tell apart. However, as demonstrated here and elsewhere (Grismer, 2006b, 2007; Grismer et al., 2009a; Linkem et al., 2011; Karin et al., 2016), with judicious examination, many of these species are easily differentiated from one another solely on the basis of morphological characters that have been useful in skink taxonomy for over a century (e.g. Boulenger, 1887, 1912). However, this is not the case with T. bukitensis and T. martae sp. nov., that live in the same piles of leaves and are not unequivocally distinguishable morphologically but are genetically exclusive and not sister species (Fig. 2). We lean toward the view of Jöger \& Schroll (2013) and consider cryptic species to be genetically exclusive lineages that are not discretely or statistically diagnosable from one another on the basis of morphology and thus consider T. bukitensis and T. martae sp. nov. to be cryptic species with respect to one another. They are not cryptic with respect to other species of Tytthoscincus in that they are discretely
and/or statistically diagnosable from them (Tables 6, 7). As noted above, differences in ecomorphology can be used to entertain hypotheses as to how the syntopic semi-fossorial and leaf-litter generalists may be partitioning resources. However, this simplistic explanation does not account for the morphological and ecological similarity between these syntopic, cryptic species. Understanding the ecological relationships between T. bukitensis and T. martae sp. nov. will contribute a great deal to a general understanding of the complex nature of leaf-litter ecology. Ecological studies of these species are currently being designed.

## BIOGEOGRAPHY IN A SKY-ISLAND ARCHIPELAGO

Understanding the phylogenetic relationships among unrelated, co-distributed species across a common landscape and the phylogeographic structure within other species across those same landscapes provides a relative, temporal framework necessary to interpret how those lineages evolved. Time-calibrating those relationships allows researchers to distinguish between old (pre-Pleistocene) and recent cladogenic events among sympatric populations (Bell et al., 2012; Grismer et al., 2015a). Time-calibrated phylogeographic and phylogenetic studies have demonstrated that speciation in tropical, montane forests is not solely the result of oscillating, climatic events of the Late Pleistocene as posited by many (Morley, 2000, 2012; Bird, Taylor \& Hunt, 2005; Outlaw \& Voelker, 2008; Reddy, 2008; Cannon, Morley \& Bush, 2009; Cannon, 2012) but have emerged along a significantly longer time continuum and that some upland species have existed in isolated refugia as paleoendemics since the Miocene (Vences et al., 2009; Tolley et al., 2011; Bell et al., 2012; Grismer et al. 2015a). In comparison, other species distributed across the same landscapes may remain essentially morphologically and genetically identical (Loredo et al., 2013; Grismer et al., 2014a, 2015b). In the sky-island archipelago of Peninsular Malaysia, Grismer et al. (2013b, 2016c) noted that the deep genetic divergences within co-distributed clades of Slender Geckos Hemiphyllodactylus and Stream Toads Ansonia, respectively, dated from the Late Eocene to the Late Pliocene ( $\sim 35.5-2.7 \mathrm{Mya}$ ). These deep divergences stand in contrast to those of co-distributed species of snakes (Asthenodipsas vertebralis, A. lasgalenesis and Lycodon butleri) whose allopatric populations bear less than $1.0 \% \mathrm{mtDNA}$ sequence divergence among them across distances as great as 210 km (Loredo et al., 2013; Grismer et al., 2014a). Loredo et al. (2013) and Grismer et al. (2014a, 2015b) hypothesized that being vagile and not micro-habitat-restricted enabled allopatric populations to migrate downslope with advancing montane forests
during cyclical, glacial maxima (Cannon et al., 2009; Woodruff, 2010; Cannon, 2012) and establish repeated episodes of secondary contact. Whereas small, non-vagile (Hemiphyllodactylus) and microhabitat-restricted (Ansonia) species remained in place and diverged along their own genetic trajectories.

Node age estimates from a time-calibrated BEAST analysis plotted on the ML topology (Fig. 2) illustrate that within the upland clade of Tytthoscincus, both relatively old and recent cladogenic events have contributed to the diversity within this lineage. The separation between the semi-fossorial and leaf-litter generalist clades occurred during the Upper Pliocene (mean of 4.8 Mya; 6.8-2.8 HPD [highest posterior density]) and speciation within the leaf-litter generalists began much more recently in the Upper Pleistocene (mean of $1.7 \mathrm{Mya} ; 2.1-1.0 \mathrm{HPD}$ ). Separation between the three allopatric, phylogeographically discordant populations of T. butleri - that have an uncorrected pairwise sequence divergence of less than $1.0 \%$ across 203 km of insular and continental mountain tops - was a very recent Pleistocene event (mean of 0.60 Mya; 0.65-0.22 HPD), reminiscent of the snakes noted above. In the semi-fossorial lineage, the separation between T. jaripendek sp. nov. and T. kakikecil sp. nov., which share a $3.6 \%$ sequence divergence between them, was also a recent Pleistocene event (mean of 0.90 Mya ; 1.14-0.23 HPD). Thus, unlike other co-distributed lineages, the cladogenic history of Tytthoscincus extends across the Plio-Pleistocene boundary. The deep split during the Upper Pliocene resulting in the evolution of the different ecomorphologies is in contrast to the younger, Pleistocene cladogenic events that happened in rapid succession (T. bukitensis, (T. martae sp. nov. (T. ishaki, (T. perhentianensis, (Tytthoscincus sp. nov., (T. temengorensis, (T. butleri))))))) from approximately a mean of 1.7-0.06 Mya. The latter events were most probably the result of cyclical ecological changes generated by rapid, ongoing glacier-driven, climate oscillations (Cannon et al., 2009; Woodruff, 2010; Cannon, 2012). The phylogenetic and phylogeographic structure of 21 additional co-distributed species (Grismer et al., in preparation) support the hypothesis of Grismer et al. (2015b) in that evolution and diversification of skyisland archipelago lineages in Peninsular Malaysia happened along a continuum from the Upper Eocene well into the Pleistocene and that speciation or lack thereof has been influenced by life history and dispersal capabilities (Loredo et al., 2013; Grismer et al., 2014a, 2015b).

## Conservation

Globally, upland ecosystems in tropical rainforests are renowned for their high levels of endemism (Blackburn \& Measey, 2009; Bell et al., 2010, 2012; Grismer et al.,

2010a, b; Grismer, 2011; Tolley et al., 2011) and the sky-island archipelago of Peninsular Malaysia is no exception. It has been noted that the montane regions of Sundaland have served as refuges for upland, range-restricted species during drier interglacial periods since at least the early Miocene ( 23 Mya; de Bruyn et al., 2013; Grismer et al., 2016c) and thus, the foundations for these species' complex ecological relationships have probably been developing over considerable periods of time. We posit that understanding the current ecological relationships of microhabitat-specific, range-restricted, refugial species will contribute greatly to a general understanding of rainforest ecosystems and that such knowledge should underlie the platform upon which effective and efficient forest management and conservation programs are conceived.

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## APPENDIX

The following specimens were examined.
Tytthoscincus bukitensis: West Malaysia, Pahang, Fraser's Hill, Telecom Loop (LSUHC 11655, 11771), Jeriau (LSUHC 12226, 12686-87), Bishops's Trail [ZRC 2.6245 (holotype)]. West Malaysia, Pahang, Genting Highlands, Awana Road (LSUHC 12237, 12750).

Tytthoscincus butleri: West Malaysia: Kedah, Pulau Langkawi, Gunung Raya; LSUHC 6790, 11152, 1178487, 11840-41, LRCUKM 143, 148. West Malaysia: Kedah, Pulau Singa Besar; DWNP 3032. West Malaysia: Perak, Bukit Larut; LSUHC 9043, 9204-06, 12429, 12432-33. West Malaysia: Penang, Penang Island, Air Hitam Dam (LSUHC 11275, 11802) and Penang Hill (LSUHC 12071).

Tytthoscincus ishaki: West Malaysia: Pahang, Pulau Tioman, Gunung Kajang LSUHC 4429, 12582, ZRC 2.6157 (holotype), 2.6158-59 (paratypes).

Tytthoscincus jaripendek sp. nov.: West Malaysia: Pahang, Cameron Highlands, Robinson Falls LSUHC 11679, 11681, 11987 (paratypes), 11680 (holotype).

Tytthoscincus kakikecil sp. nov.: West Malaysia: Pahang, Fraser's Hill, Richmond Road [LSUHC 11769 (holotype)], Telecom Loop (LSUHC 11770, 11772). West Malaysia: Pahang, Genting Highlands, Ulu Kali LSUHC 12754, FMNH 19945. West Malaysia: Pahang, Cameron Highlands, Telom Valley ZRC 2.5944.

Tytthoscincus martae sp. nov.: West Malaysia: Pahang, Fraser's Hill, Hindu Temple LSUHC 12688 (holotype), ZRC 3.6246 (paratype). West Malaysia, Pahang, Genting Highlands, Awana Road LSUHC 12517-18, 12749, 12751 (paratypes), 12072.

Tythoscincus perhentianensis: West Malaysia: Terengganu, Pulau Perhentian Besar LSUHC 8075 (holotype).

Tytthoscincus temengorensis: West Malaysia: Perak, Royal Belum Park DWNP 5018 (holotype). West Malaysia: Perak, Temengor Forest Reserve, Gibbon Trail LSUHC 5650 (paratype).


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