A new genus of miniaturized and pug-nosed gecko from South America (Sphaerodactylidae: Gekkota)

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Sphaerodactyl geckos comprise five genera distributed across Central and South America. We estimated phylogenetic relationships among sphaerodactyl genera using both separate and combined analyses of seven nuclear genes. Relationships among genera were incongruent at different loci and phylogenies were characterized by short, in some cases zero-length, internal branches and poor phylogenetic support at most nodes. We recovered a polyphyletic Coleodactylus, with Coleodactylus amazonicus being deeply divergent from the remaining Coleodactylus species sampled. The C. amazonicus lineage possessed unique codon deletions in the genes PTPN12 and RBMX while the remaining Coleodactylus species had unique codon deletions in RAG1. Topology tests could not reject a monophyletic Coleodactylus, but we show that short internal branch lengths decreased the accuracy of topology tests because there were not enough data along these short branches to support one phylogenetic hypothesis over another. Morphological data corroborated results of the molecular phylogeny, with Coleodactylus exhibiting substantial morphological heterogeneity. We identified a suite of unique craniofacial features that differentiate C. amazonicus not only from other Coleodactylus species, but also from all other geckos. We describe this novel sphaerodactyl lineage as a new genus, Chatogekko gen. nov. We present a detailed osteology of Chatogekko, characterizing osteological correlates of miniaturization that provide a framework for future studies in sphaerodactyl systematics and biology.


INTRODUCTION

Sphaerodactyl geckos (Sphaerodactylini: Sphaerodactylidae) are a species-rich group of Neotropical lizards. They comprise more than 10% of gecko species with more than 150 described species in five genera: Coleodactylus, Lepidoblepharis, Gonatodes, Pseudogonatodes, and Sphaerodactylus (Kluge, 1995; 2001; Gamble et al., 2008a; Uetz, 2010). Sphaerodactyl geckos are distributed across Central and South America and the Caribbean, including several Pacific continental and oceanic islands, e.g. Gorgona, Cocos (Vanzolini, 1968a; Harris, 1982; Harris & Kluge, 1984; Kluge, 1995). Most sphaerodactyl gecko species are active during the day and the clade is thought to

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be secondarily diurnal, having evolved from a nocturnal ancestor (Werner, 1969; Underwood, 1970; Kluge, 1995; Röl & Henkel, 2002). They are generally small; some Sphaerodactylus species are among the smallest known amniotes, averaging only 16 mm snout–vent length (SVL) (Thomas, 1965; MacLean, 1985; Hedges & Thomas, 2001), and the largest forms do not exceed 65 mm SVL (Rivas & Schargel, 2008).

The genus Coleodactylus is distributed in northeastern South America (Kluge, 1995) and consists of five described species: C. amazonicus (Andersson, 1918), C. brachystoma (Amaral, 1935), C. meridionalis (Boulenger, 1888), C. natalensis Freire, 1999, and C. septentrionalis Vanzolini, 1980. Coleodactylus has historically been defined by the structure of the ungual sheath, the scales covering the claw, being composed of five asymmetrical scales (Parker, 1926; Vanzolini, 1957; Kluge, 1995). Coleodactylus amazonicus differs from its congeners in having an ungual sheath possessing only four asymmetrical scales, a reduction caused by the loss of the medialmost dorsal scale (Andersson, 1918; Parker, 1926; Vanzolini, 1957; Avila-Pires, 1995). Coleodactylus amazonicus also has keeled dorsal scales, while all other members of the genus have smooth scales (Vanzolini, 1957; Avila-Pires, 1995). These morphological differences cast doubt on the diagnostic reliability of the ungual sheath and other characters for the genus and/or on the allocation of C. amazonicus to Coleodactylus.

Molecular data mirror the morphological differences among Coleodactylus species. Recent molecular phylogenies recovered two deeply divergent lineages in Coleodactylus, with one clade consisting of C. amazonicus and the other made up of the remaining Coleodactylus species, the ‘C. meridionalis group’ (Geurgas, Rodrigues & Moritz, 2008; Geurgas & Rodrigues, 2010; Gamble et al., 2011). These results were not translated into a revised taxonomy however because of poor nodal support, e.g. bootstrap values because of poor nodal support, e.g. bootstrap values and Bayesian posterior probabilities, for these relationships from the molecular data. Additionally, topology tests that constrained a monophyletic Coleodactylus sensu lato (s.l.) failed to reject the hypothesis that C. amazonicus forms a clade with the remaining Coleodactylus species (Geurgas et al., 2008; Gamble et al., 2011). Non-tree-based molecular evidence supports the distinction between C. amazonicus and the remaining Coleodactylus species. Two separate deletions of 18 and 6 bp in the RAG1 gene occur in species of the C. meridionalis group, but not in C. amazonicus (Gamble et al., 2008a, 2011; Geurgas & Rodrigues, 2010). Rare genomic events such as codon deletions and insertions (indels) are relatively homoplasy-free characters and can provide strong evidence of evolutionary history (van Dijk et al., 1999; Rokas & Holland, 2000; Simmons, Ochoterena & Carr, 2001).

The sum of available data calls into question the monophyly of Coleodactylus. Coleodactylus amazonicus is morphologically distinct from the remaining Coleodactylus species. Molecular data present a mixed picture of Coleodactylus relationships and sphaerodactyl phylogeny as a whole but, like the morphological data, cast doubt on Coleodactylus monophyly. We gathered new molecular and morphological data to address these issues. Our objectives were to: test the monophyly of Coleodactylus using a multigene molecular dataset and specifically address the failure of previous topology tests to support two distinct Coleodactylus lineages; review the morphology of C. amazonicus as a means to diagnose deeply divergent clades within Coleodactylus s.l.; and characterize osteological correlates of miniaturization in Coleodactylus s.l.

**MATERIAL AND METHODS**

**TAXON SAMPLING AND MOLECULAR DATA**

We assembled a nuclear gene dataset that included multiple species from each of the currently recognized genera of the New World Sphaerodactylini: Coleodactylus s.l., Gonatodes, Lepidoblepharis, Pseudogonatodes, and Sphaerodactylus. We included several Old World members of Sphaerodactylidae as outgroups, including Sauraodactylus brosseti, Pristurus carteri, and two species of Teratoscincus (i.e. T. microlipsis and T. przewalskii). Phylogenies were rooted with the gekkonid Hemidactylus platyurus. Locality data and GenBank accession numbers for sampled taxa are listed in Table S2 of the Supporting Information.

We extracted genomic DNA from tissues using the DNeasy Blood and Tissue kit (Qiagen, Valencia, CA, USA) and used PCR to amplify gene fragments of seven nuclear loci for sequencing. Six loci were protein-coding regions: recombination-activating gene 1 (RAG1); recombination-activating gene 2 (RAG2); oocyte-maturation factor MOS (C-MOS); acetylcholinergic receptor M4 (ACM4 or CHRMA4); phosducin (PDC); and protein tyrosine phosphatase, non-receptor type 12 (PTPN12). The seventh locus included intron 8 (in Gallus) and flanking exon regions of RNA binding motif protein, X-linked (RBMX). Primers are listed in Table S1 of the Supporting Information. We purified PCR products using Exonuclease I and Shrimp Alkaline Phosphatase (Hanke & Wink, 1994). Big Dye sequencing was conducted at the BioMedical Genomics Center, University of Minnesota. Sequences were assembled and checked for accuracy using Sequencher 4.8 (Gene Codes, Ann Arbor, MI, USA). We translated protein-
coding genes to amino acids using MacClade 4.08 (Maddison & Maddison, 1992) to confirm codon alignment and gap placement. We aligned RBMX sequences initially using T-Coffee (Notredame, Higgins & Heringa, 2000) and subsequently fine-tuned the alignment by hand.

**PHYLOGENETIC ANALYSES**

We conducted several phylogenetic analyses of the nuclear dataset. The seven loci were concatenated to conduct partitioned maximum likelihood (ML) analysis. We also analysed each locus separately. All ML analyses were conducted using RAxML 7.2.6 (Stamatakis, 2006). The concatenated ML analysis consisted of 19 partitions, with data partitioned by gene and by codon, except the intron RBMX, which consisted of a single partition. ML analyses of individual protein-coding loci also partitioned data by codon. All ML partitions utilized the GTR + Gamma model of sequence evolution and nodal support was estimated with 1000 bootstrap replicates (Felsenstein, 1985).

We conducted Bayesian analyses of the nuclear dataset using MrBayes 3.1.2 (Huelsenbeck & Ronquist, 2001; Ronquist & Huelsenbeck, 2003). All analyses used a neighbour-joining tree as a starting topology. Analyses of the individual genes involved two independent runs, each consisting of four parallel Markov chain Monte Carlo (MCMC) chains per run for five million generations and sampled every 1000 generations. Each Bayesian analysis of the individual genes utilized a model of molecular evolution as determined by Akaike’s information criterion (AIC) in jModeltest (Posada, 2008). Analyses of the concatenated dataset partitioned data by codon with a separate partition for RBMX. Each partition utilized a model of molecular evolution as determined by AIC with model parameters estimated independently using the unlink option. The concatenated analysis involved two independent runs, each consisting of six parallel MCMC chains per run for ten million generations and sampled every 1000 generations. We assessed convergence and stationarity in all Bayesian analyses by plotting likelihood values in Tracer 1.5 (Rambaut & Drummond, 2007) as well as plotting split frequencies between independent runs using AWTY (Nylander et al., 2008).

Poor phylogenetic resolution among sphaerodactyl genera in the concatenated analyses and incongruence among individual gene trees (see Results) motivated us to explore additional means of estimating phylogenetic relationships that could incorporate the sometimes diverse histories of individual genes. The probability of incomplete lineage sorting is increased when there are short internal branches (Maddison, 1997; Rosenberg & Tao, 2008), so we used two methods to estimate species trees that can accommodate individual gene genealogies. The first method, MDC (minimized deep coalescence), used individual gene trees to find a species tree that minimized the number of deep coalescent events across all loci (Maddison, 1997; Maddison & Knowles, 2006). The second method, BCA (Bayesian concordance analysis), estimated the species tree possessed by the plurality of clades recovered from individual loci, the concordance tree, and also estimated the proportion of loci that shared a specific clade with the concordance tree, the concordance factor (Ane et al., 2007; Baum, 2007).

We estimated the MDC tree using Mesquite 2.73 (Maddison & Maddison, 2008). This method required that ‘species’ be identified a priori and that individuals or taxa from the analyses of separate loci be assigned to each of these ‘species’. Because we were interested in relationships among sphaerodactyl genera, we treated genera as ‘species’ in the MDC analysis. We accommodated phylogenetic uncertainty associated with the reconstruction of the individual gene trees using the Augist Mesquite module (Oliver, 2008). We estimated 1000 MDC trees with each search randomly sampling from the posterior distribution of trees from the Bayesian analyses of each of the nuclear loci. We used the subtree pruning and regrafting heuristic search algorithm with a maximum of 100 equally parsimonious trees saved at each search. Tree weights were stored for each search in the event multiple equally parsimonious MDC trees were found. The MDC species tree was calculated as a 50% majority-rule consensus tree with bipartition frequencies providing a measure of nodal support.

We estimated the BCA tree using BUCKy 1.4.0 (Ane et al., 2007). We conducted three separate analyses, each with a different a priori discordance level among gene trees, which was controlled by the variable $\alpha$ (Ane et al., 2007). Setting $\alpha = 0$, for example, imposes a single species tree on all of the loci, while at the other extreme setting $\alpha = \infty$ forces each locus to have its own independent history. We used an interactive web-based tool (http://www.stat.wisc.edu/~ane/bucky/prior.html) to calculate $\alpha$ values for our data. Each value for $\alpha$ placed a different prior on the number of possible species trees: $\alpha = 0.1$ placed a high prior on one distinct tree; $\alpha = 1.0$ placed a high prior on 2–3 species trees; and $\alpha = 10$ placed a high prior on 5–6 species trees. All BUCKy analyses were run for 10 000 000 generations following a 10% burn-in.

**HYPOTHESIS TESTING**

We tested the monophyly of Coleodactylus s.l. using two different methods. We implemented the
likelihood-based Shimodaira–Hasegawa (SH) test (Shimodaira & Hasegawa, 1999), which compared the constrained topology, a monophyletic Coleodactylus s.l., with the unconstrained ML tree. Per-site log likelihoods were estimated in RAxML 7.2.6 (Stamatakis, 2006) and P-values were calculated using CONSEL (Shimodaira & Hasegawa, 2001). We also tested alternative phylogenetic hypotheses in a Bayesian framework. We used the filter option in PAUP* 4.0b10 (Swofford, 2002) to calculate the posterior probability of a monophyletic Coleodactylus s.l. in the posterior distribution of trees from the MrBayes analyses. We tested the monophyly of Coleodactylus s.l. using the concatenated nuclear gene dataset and each locus separately.

Short internal branches connected the six sphaerodactyl genera in both the concatenated trees as well as individual gene trees (see Results). These short internal branches not only increased the likelihood of incomplete lineage sorting, as mentioned above, but the limited number of character changes along these extremely short branches could make it difficult to adequately compare alternative hypotheses using the SH test. Some of these internal branches were so short as to have effectively zero branch length. These phenomena could explain why previous attempts to test Coleodactylus monophyly failed to adequately distinguish among competing hypotheses (Geurgas et al., 2008; Gamble et al., 2011). We examined our ML trees for the presence of zero-length branches using a likelihood ratio test with the ‘describe trees’ function in PAUP* 4.0b10 (Swofford, 2002). Briefly, the likelihood of the best tree was compared with the likelihood of the same tree but with a single branch collapsed to zero using the likelihood ratio test. Each of the four branches connecting the six sphaerodactyl genera was sequentially tested in this manner. A significant result meant the branch length was significantly different from zero. Significance levels were Bonferroni-corrected for the number of intergeneric branches.

**MORPHOLOGICAL DATA**

We examined both internal and external morphological characters from specimens of several species of Sphaerodactylidae, including exemplars from each of the currently recognized sphaerodactyl genera, to assess the monophyly of Coleodactylus s.l. (see Supporting Information). We also examined C. amazonicus specimens from several localities across its distributional range. We viewed osteological characters using a variation of a common clearing and double staining technique (Hanken & Wassersug, 1981). This method is especially useful for small animals in which dry skeletal preparation techniques are not suitable due to the potential risk of damage by the insects used to prepare them or to distortion caused by the drying and shrinkage of unossified portions of the skeleton. We modified the protocol in that we did not remove the integument from specimens, and used KOH only as a clearing reagent, without exposing specimens to enzymatic solutions of trypsin or pancreatin. Specimens were observed under a Leica MS6 dissecting microscope. Illustrations were traced with Adobe Illustrator CS3 13.0.2 directly over a series of digital photographs taken with a Nikon Coolpix 995 camera (3.1 Megapixels, 3× Optical Zoom) at different magnifications. Images were complemented with drawings made with a camera lucida.

**RESULTS**

**TAXON SAMPLING AND MOLECULAR DATA**

The nuclear gene dataset consisted of 4116 aligned base pairs from seven loci for 33 gecko taxa (Table 1).
Sequence alignment was unambiguous for protein-coding regions, but several insertion/deletions (indels) were detected in five of the genes (Table 1, Fig. 1). Indels in \textit{RAG1}, \textit{C-MOS}, and \textit{ACM4} have been commented on previously (Gamble et al., 2008a, c; Geurgas et al., 2008). Both \textit{RBMX} and \textit{PTPN12} had single codon deletions in \textit{C. amazonicus} samples. The \textit{RBMX} deletion occurred in the region analogous to exon 8 in chicken (\textit{Gallus}).

Several clades received high levels of support in both analyses, including: a clade consisting of \textit{Gonatodes} + \textit{Lepidoblepharis}; a clade consisting of \textit{Coleodactylus sensu stricto} (s.s.) + \textit{Pseudogonatodes} + \textit{Sphaerodactylus}; and \textit{Sphaerodactylini}. Generic-level sphaerodactyl clades were all well supported with the exception of \textit{Coleodactylus} s.l., which was polyphyletic with regards to other sphaerodactyl genera; for example, \textit{C. amazonicus} did not form a clade with the remaining sampled \textit{Coleodactylus} species. We recovered three clades within \textit{C. amazonicus}: one clade consisted of individuals from eastern Amazon (Pará); the second clade consisted of individuals from southwestern Amazon (Rondônia and Rio Ituxi, Amazonas); and the third clade consisted of individuals from central and northern Amazon (near Manaus, PR).
Amazonas, and Roraima). ML branch lengths among these *C. amazonicus* clades were equivalent to species-level divergences within other sphaerodactyl genera and between the species *Teratoscincus microl epis* and *T. przewalskii* (Fig. 1).

Topologies among individual gene trees were largely incongruent (Fig. 2). The only well-supported nodes in all of the analyses were nodes subtending each of the sphaerodactyl genera, although, as with the concatenated analyses, *Coleodactylus s.l.* was polyphyletic in all loci with *C. amazonicus* samples forming their own clade distinct from other sampled *Coleodactylus* species.

The MDC consensus tree (Fig. 3) was largely congruent with the concatenated ML and Bayesian trees and recovered a well-supported Sphaerodactylina clade; and a *C. amazonicus* clade. BCA with varying α levels produced identical concordance trees and concordance factors. The BCA tree (Fig. 3) was similar to the MDC consensus tree. Although it is difficult to assess what constitutes a significant concordance factor (Baum, 2007) the Sphaerodactylina clade and *Lepidoblepharis + Gonatodes* clade were the only relationships that received concordance factors exceeding 0.50.

**HYPOTHESIS TESTING**

Results of the SH tests that constrained *Coleodactylus s.l.* as monophyletic were not significant (Table 2). The Bayesian posterior probability of a monophyletic *Coleodactylus s.l.* was zero for the concatenated data and low, but not significant, for most of the individual gene analyses (Table 2).

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**Figure 2.** Cladograms for each nuclear locus and the concatenated nuclear gene dataset illustrating relationships among sphaerodactyl genera estimated using maximum likelihood. Branches with lengths not significantly different from zero are indicated with an asterisk.
We used the likelihood ratio test to determine whether branch lengths of any of the four branches connecting sphaerodactyl genera were significantly different from zero (Fig. 2). The concatenated data, RBMX, PTPN12, ACM4, and RAG1, had two of four internal branches with lengths not significantly different from zero. PDC had three of four branches not significantly different from zero. RAG2 and C-MOS had all four branches not significantly different from zero.

**MORPHOLOGICAL DATA**

We recovered several morphological traits to aid in the diagnosis and description of a new genus and provide a detailed osteology to guide future research in sphaerodactyl biology and evolution. Morphological descriptions and comparisons are explained in detail below, after we address taxonomic changes.

**TAXONOMY**

The combined morphological and molecular evidence suggested a new generic-level sphaerodactyl clade be described. We also redescribe Coleodactylus s.s. in light of our results.

**REPTILIA: SQUAMATA:**

**SPHAERODACTYLIDAE**

**CHATOGEKKO** Gamble, Daza, Collin, Vitt and Bauer, gen. nov. (Figs 5, 6)

Type species: Sphaerodactylus amazonicus (Andersson, 1918)


**Chatogekko** is distinguished from all gekkotans by the following unique combination of characters: (1) between two and four loreal scales (ls, Fig. 4A, also present in some *Sphaerodactylus*); (2) claws enclosed in an ungual sheath consisting of four scales (Parker, 1926; Vanzolini, 1957; Avila-Pires, 1995): inner supero-lateral (isl, Fig. 4B), outer supero-lateral (osl, Fig. 4B), inner infero-lateral (iil, Fig. 4C), and outer infero-lateral (oil, Fig. 4C) (ventrolaterals sensu Kluge, 1995); (3, Fig. 4D) keeled scales on dorsal body surface (Vanzolini, 1957; Avila-Pires, 1995), also present in some *Sphaerodactylus*; (4, Fig. 5A) bony external nares large and entering or approaching contact between prefrontal and nasals (as a consequence of extensive overlapping contact of maxilla and prefrontal); (5, Fig. 5A) posterior edge of premaxilla (i.e. the ascending nasal process) contacts medial process of frontal bone (Daza et al., 2008); (6, Fig. 5A) posterior edge of ascending nasal process bifurcated; (7, Fig. 5A) internasal contact absent; (8, Fig. 5A) jugal bone vestigial and limited to tip of maxilla; (9, Fig. 5A) postparietal process of parietal contacting supraoccipital and otooccipital, but not squamosal. Reduced paroccipital process located dorsally to fenestra ovalis.

**Figure 3.** Phylogenetic relationships among sphaerodactyl genera estimated using (A) MDC (minimization of deep coalescence events) and (B) BCA (Bayesian concordance analysis). Node values on the MDC tree are bipartition frequencies from 1000 replicate analyses randomly sampled from the Bayesian posterior distributions of the individual gene trees. Node values on the BCA tree are posterior mean concordance factors.

We used the likelihood ratio test to determine whether branch lengths of any of the four branches connecting sphaerodactyl genera were significantly different from zero (Fig. 2). The concatenated data, RBMX, PTPN12, ACM4, and RAG1, had two of four internal branches with lengths not significantly different from zero. PDC had three of four branches not significantly different from zero. RAG2 and C-MOS had all four branches not significantly different from zero.
Table 2. Results of topological constraint tests comparing a monophyletic Coleodactylus s.l. with the best phylogenetic estimates for seven nuclear genes analysed individually, as well as the combined analysis. Columns show the log likelihood (lnL) of the best tree, the likelihood of the tree with a monophyletic Coleodactylus s.l., the difference in likelihood values between the best tree and the constraint tree, and the $P$-value of the SH test. The last column shows posterior probabilities of a monophyletic Coleodactylus s.l. from the Bayesian analyses.

<table>
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<th>Dataset</th>
<th>lnL of best tree</th>
<th>lnL of constraint tree</th>
<th>Difference in lnL</th>
<th>$P$</th>
<th>Posterior probability of alternative hypothesis</th>
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<tr>
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<td>-2 121.7696</td>
<td>1.92656</td>
<td>0.29</td>
<td>0.1280</td>
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<tr>
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<td>3.77983</td>
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<tr>
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<tr>
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</table>

Figure 4. Chatogekko amazonicus specimens. A, lateral view of the head showing 2–4 loreal scales (ls); B, dorsal view of the left hand showing the inner supero-lateral and outer supero-lateral (isl and osl, Fig. 5B); C, ventral view of the left hand showing the inner infero-lateral and outer infero-lateral (iil and oil, Fig. 5C); and D, keeled scales along the dorsal surface of the body. A–C, USNM 288775; D, MZUSP 91394. Scale bar = 1mm.
visible in dorsal view; (11, Fig. 5B) paroccipital process very reduced and located dorsally to foramen ovalis (instead of posterior as in other gekkotans) and not participating in quadrate articulation (paroccipital abutting); (12, Fig. 5B) palatine exceeds vomer substantially in length; (13, Fig. 10) duplicipalatinate condition; (14) a 3-bp deletion in coding region of exon 8 (in Gallus) of RBMX; and (15) a 3-bp deletion in coding region of exon 13 (in Gallus) of PTPN12.

**Distribution:** Central and eastern Amazonia, including the Brazilian states of Acre, Amazonas, Rondônia, Mato Grosso, Roraima, Pará, and Amapá; French Guiana; Guyana; Suriname; the Venezuelan state of Amazonas; and northern Bolivia (Gasc, 1990; Avila-Pires, 1995; Langstroth, 2005; Geurgas & Rodrigues, 2010).

**Natural history:** *Chatogekko* lives in the leaf litter in a variety of undisturbed lowland forested habitats (Vitt et al., 2005). These gekkos are active throughout the day although they do not bask (Hoogmoed, 1973). Diet is made up of small insects including springtails, mites and ticks, termites, homopterans, and larval insects (Hoogmoed, 1973; Ramos, 1981; Vitt et al., 2005). Females lay one egg per clutch and can produce several clutches during the year (Hoogmoed, 1973; Gasc, 1990). *Chatogekko* can be locally very abundant but appears to be negatively affected by forest fragmentation (Carvalho et al., 2008).

**Etymology:** A composite word from the Spanish and Portuguese ‘Chato’, derived from the Greek ‘Platus’, meaning ‘flat’ and referring to its pug-nosed snout; and gekko from the Malay ‘gekoq’, onomatopoetic of the call of the species *Gekko gekko* and the common name to all limbed gekkotans. A Sri Lankan origin for the word gekko, derived from the Sinhalese word ‘gego’, is also possible (de Silva & Bauer, 2008). The name is masculine.

**Species composition:** *Chatogekko amazonicus* (Andersson, 1918). In addition, the names *C. zernyi* (Wettstein, 1928) and *C. guimaraesi* (Vanzolini, 1957) are available for populations from eastern Amazonia and southwest Amazon, respectively. See Discussion for details.

**Coleodactylus Parker, 1926**

**Type species:** *Sphaerodactylus meridionalis* (Boulenger, 1888).


*Coleodactylus* is a miniaturized species complex of diurnal sphaerodactyl geckos that can be differentiated from all other gekkotans by the following unique combination of characters: (1) claws enclosed in ungual sheath consisting of five scales (Parker, 1926; Vanzolini, 1957; Avila-Pires, 1995); (2) smooth or imbricate scales on dorsal body surface (Vanzolini, 1957; Avila-Pires, 1995), present in most other gekkotans; (3) ascending nasal process separates nasals approximately one-quarter their length, one of the shortest among sphaerodactyl geckos (Daza et al., 2008); (4) proximal portion of metatarsal IV not very expanded; and (5) two separate deletions of 18 and 6 bp in exon 1 (in Gallus) of RAG1.

**Distribution:** Northern and eastern Brazil including states of Alagoas, Bahia, Ceará, Goiás, Maranhão, Mato
Species composition: Coleodactylus brachystoma (Amaral, 1935); C. meridionalis (Boulenger, 1888); C. natalensis Freire, 1999; and C. septentrionalis Vanzolini, 1980.

OSTEOMETRY

Because most characters that differentiate Chatogekko from other sphaerodactyl geckos come from osteology, a detailed description of its skeleton would be convenient for future taxonomic differentiation of sphaerodactyl taxa. Additionally, a detailed osteology provides a baseline for future morphological research aimed at diagnosing the putative Chatogekko species. We present the osteological data in a framework that highlights the extremely small size of these lizards. It has been stated that is impossible to present a unifying model of miniaturization encompassing all lizards (Rieppel, 1984a). Even so, many anatomical similarities of the cranial structure of Chatogekko are present in other miniaturized gekkotans, especially closely related sphaerodactyl genera. Because the cranial anatomy of Sphaerodactylus roosevelti has been described in detail (Daza et al., 2008), we only highlight those structures that show differences in this new genus. We do this in the context of a descriptive approach and do not intend to imply any particular character polarity. Additionally, we review the postcranium, which has been described previously (Noble, 1921), but not in great detail.

SKULL

The skull of Chatogekko is wedge shaped with a maximum width at the level of the otic capsules. It has a rounded outline in lateral view as there is a continuous curvature from the tip of the snout to the skull table. It has the shortest muzzle unit among sphaerodactyls (Fig. 5). This is especially evident in the anterorbital region, where a high degree of overlap occurs between the bones. The premaxilla has a very elongated ascending nasal process (asnp, Fig. 5A), with lateral margins that do not converge posteriorly. The last three-quarters of this process are reduced in width to a narrower projection that contacts the medial process of the frontal. In Sphaerodactylus, this process may reach the level of the frontal bone, but never contacts it directly because the nasal bones lie between them (Daza et al., 2008). The ascending nasal process is much shorter and does not reach the level of the frontal bone in Coleodactylus. The posterior projection of the ascending nasal process varies among the specimens of Chatogekko examined and may be bifurcated or assume an almost transverse orientation.

The orbit in Chatogekko occupies about 32% of the skull length, which is slightly more than in other sphaerodactyls (Daza et al., 2008). As in most limbed geckos, the orbit is bounded by the postorbitofrontal, frontal, prefrontal, maxilla, and jugal (Evans, 2008; Daza & Bauer, 2010); the jugal is reduced or vestigial and contacts the tip of the posterior portion of the maxilla on the medial side. The floor of the orbit is pierced by a very large, D-shaped suborbital fenestra, which is present in all sphaerodactyls as well as the more distantly related sphaerodactylids Pristurus and Saurodactylus (Daza et al., 2008) and Euleptes (J.D.D., pers. observ.).

The rear portion of the skull is typical of miniaturized lizards (Rieppel, 1984a), which indicates how size reduction directly affects cranial structure. The basicranium is massive, being the widest part of the skull at the level of the otic capsules. The skull table is comparatively small, given that the parietals leave exposed a larger area of the basicranium. The outer margin of the basicranium (prootic, otoccipital, and supraoccipital), as a consequence, is completely visible in dorsal view. The otoccipital area is so prominent and the horizontal semicircular canal bulges to the extent that the paroccipital process, normally seen in sphaerodactyls, is totally hidden. This paroccipital process is rudimentary and plays little or no function at all in the streptostylic quadrate articulation as in other lizards (Versluys, 1912; Frazzetta, 1962; Rieppel, 1978). The quadrate is very lightly built and articulates with the basicranium in a very anterior position, just in front of the fenestra ovalis. A quadrate foramen is present but its location is more proximal than in Sphaerodactylus. The squamosal bone is minuscule, and lost in some populations of Chatogekko. When this bone is present, it barely contacts the postparietal process of the parietal and lies against the basicranium, without contacting the quadrate or wrapping around it. Another consequence of this massive basicranium is the shape of the pterygoids, which have an almost straight medial margin (i.e. not curved or sigmoidal), and create a very wide interpterygoid vacuity posteriorly. The basipterygoid process and the cartilaginous pad that covers it are very narrow in Chatogekko.

In the palate the premaxillary–vomerine fenestra is very large and irregularly shaped, and partially invaded by the maxillary lappets. The vomer is reduced in size relative to the other palatal bones, leaving a very large fenestra exochoanalis and is partially overlapped by the septomaxilla.

**JAW**

The jaw of Chatogekko is typically sphaerodactyl, very straight with an elongated dentary that extends posteriorly almost to the level of the articular surface of the craniomandibular articulation. The coronoid is low and very small, without projecting above the contour of the mandible. In lingual view, the splenial seems to be fused with the coronoid, a character that unites Pristurus with the sphaerodactyls.

**HYOID APPARATUS**

There are no major differences between the hyoid structure of Chatogekko (Fig. 7) and that of Sphaerodactylus macrolepis (Noble, 1921). In these two genera, medial or lateral projections of the hypohyal (hyoid cornu) do not exist. Among sphaerodactyls these are only present in Gonatodes. The second ceratobranchial (2 cb, Fig. 7) is comparatively shorter than in Sphaerodactylus and is oriented posteromedially. The second epibranchial (2 eb, Fig. 7) is not joined to the second ceratobranchial as in Sphaerodactylus.

**POSTCRANUM**

The postcranium of Chatogekko comprises 85% of the entire body length; the body and tail sections are subequal (Fig. 8). The vertebral column is composed of 47 vertebrae: 26 presacrals, two sacrals and 19 caudals. The presacral region comprises eight cervical, 17 thoracic and one lumbar. In the cervical region only the atlas and the axis are ribless and the remaining six cervicals bear ribs that increase in length gradually. The atlas is fused dorsally as in all sphaerodactyls except Gonatodes, which has paired elements. The 3rd and 4th cervicals have short ribs that are widened and bifurcated distally. The ribs of cervicals 5–7 are set closer to the suprascapula. The rib of the 8th cervical approaches but does not contact the sternum. The sternal ribs of the first four thoracic vertebrae are attached to the sternum directly. The 5th thoracic may be attached to the xiphisternum in specimens in which this structure is branched. The remaining thoracic vertebrae have short postxiphis-
ternal inscriptive ribs that reduce their size gradually until becoming only a small nubbin.

The longest rib is present on the 11th thoracic vertebra, after which ribs start to decrease in size until lost on the lumbar. A single lumbar vertebra does not differ in size from the posterior thoracic vertebrae. The two sacral vertebrae differ in structure. The first has expanded transverse processes that articulate with the pelvic girdle (illum) and posteriorly it is fused to the transverse processes of the second sacral, whereas the second has a short transverse process which are oriented anterolaterally. The tail is formed by 19 caudals. The pygial vertebrae have been described as those anterior caudals devoid of fracture planes (Holder, 1960). In Chatogekko, autotomy planes are visible after the 6th caudal vertebrae, but only the first three lack hemal arches. The transverse processes are elongated and oriented posteriorly on the first five caudals; these processes gradually reduce in length distally. Beyond the 6th caudal vertebrae, centrum length increases, almost doubling the length of the presacral vertebrae.

The pectoral girdle comprises suprascapulae, scapulocoracoids, epiceracoids, clavicles, interclavicle, and sternum. The suprascapula is expanded and cartilaginous. The scapular portion of the scapulocoracoid is elongated and narrow. The scapulocoracoid fenestra is closed by a projection of the cartilaginous scapular epiceracoid bar. The anterior coracid fenestra (i.e. anterior primary coracid emargination) is present, but the posterior one is absent. The clavicles are expanded medially and more or less rotated forward. They lack the clavicular fenestra, as do Lepidoblepharis (Noble, 1921; Parker, 1926) and Gonatodes. Parker (1926) also described Coleodactylus and Pseudogonatodes with no clavicular fenestra, although we found specimens of Coleodactylus and Pseudogonatodes with clavicular fenestrae, which indicates that this character is variable or polymorphic for these two genera; in Sphaerodactylus, the clavicle is invariably perforated (Noble, 1921), which we were able to corroborate in all species reviewed (see Supporting Information). The interclavicle in Chatogekko has lateral arms, but these are very broad and almost indistinct. The sternum is shield-like and well ossified.

The pelvic girdle is formed by the fusion of the ilium, ischium, and pubis. The ischium and pubis are in close contact with their fellows, but not fused. The ischiopubic fenestra is large and compressed anteroposteriorly. In Chatogekko the ilium is constricted dorsal to the acetabulum and extends dorsally as a rod-like process. The ischium is wider than the pubis, and the metischial processes are widely separated. The hypoischium is absent. The pubic symphysis is slender and capped by a small epipubic cartilage. In all sphaerodactyls, the pectineal process is large and ventrally directed. This is a highly diagnostic feature, mentioned by Noble (1921) as a difference between the African ‘Gonatodes dickersoni’ (now Cnemaspis dickersoni) and the Neotropical sphaerodactyls. The rounded obturator foramen for the course of nerves lies at the boundary between the ischium and pubis. This foramen is present in all limbed gekkotans and lost in pygopodids.

The limbs are short and stout, but most typical elements of the gecko appendicular skeleton (Russell, 1972; Fabrezi, Abdala & Oliveri, 2007; Russell & Bauer, 2008) are present. One variation that occurs in sphaerodactyls is the increase in number of sesamoids on the proximal epiphyseal end of the radius with respect to other lizards. These elements have been described for a few lizards, for instance Sphaerodactylus klauberi and the xantusiid Lepidophyma gaigeae (Jerez, Mangione & Abdala, 2010). In Chatogekko and Coleodactylus there are three of these elements between the radius and the humerus (Fig. 8). This number is variable among other sphaerodactyls; for example, Pseudogonatodes and
Sphaerodactylus have two, and Lepidoblepharis and Gonatodes only one. Pseudogonatodes, Coleodactylus, and Chatogekko also have sesamoids dorsal to the metacarpal–phalange articulation (Figs 5, 7). These ossifications appear sporadically in Lepidoblepharis, but not in Gonatodes.

The phalangeal formulae of the manus and pes of sphaerodactyls are typically 2-3-4-5-3 and 2-3-4-5-4 (Table 3), respectively, which are primitive for squamates (Greer, 1992). One phalanx in the fourth manual digit of Pseudogonatodes, Coleodactylus, and Chatogekko and the fourth pedal digit of Pseudogonatodes are lost (Table 3, Fig. 9) The identity of the phalanx lost is hard to determine without developmental series, but it is likely that it is either the ultimate or penultimate phalanx.

**DISCUSSION**

**PHYLOGENY**

Phylogenetic analyses of the combined nuclear gene data, using both a concatenation approach and a gene tree approach, recovered three lineages of spha-
Gonatodes + Sphaerodactylus + Coleodactylus clade. Other published molecular phylogenies have consistently recovered the Lepidoblepharis + Gonatodes clade, but have failed to recover the other clades with strong support (Gamble et al., 2008a, 2011; Geurgas et al., 2008). The difficulty in recovering these clades is probably due to the short internal branches linking genera at the base of the sphaerodactyl clade. Short internal branches are a signature of rapid cladogenesis, indicating that divergences among sphaerodactyl genera occurred in a relatively short time frame (Gamble et al., 2008a, 2011). Short internal branches can also hamper phylogenetic reconstruction (Jackman, Larson, de Queiroz & Losos, 1999; Slowinski, 2001; Poe & Chubb, 2004). Indeed, our failure to reject the hypothesis that several of those internal branches had lengths not significantly different from zero suggests hard polytomies in the molecular data (Maddison, 1989; Slowinski, 2001). One possible cause of zero-length branches is insufficient data (Poe & Chubb, 2004). This may play some role in our results as our three loci with the least amount of data, namely RAG2, C-MOS, and PDC, had either three or four of the four branches connecting sphaerodactyl genera with branch lengths not significantly different from zero. The remaining loci had more data, sometimes substantially so, and possessed only two of four branches with lengths not significantly different from zero. This was also the case with the concatenated dataset. Close examination of which branches were statistically indistinguishable from zero shows some similarities among the loci with more data (ACM4, RAG1, PTPN12, and RBMX) and the concatenated dataset (Fig. 2). The branch leading to the Gonatodes + Lepidoblepharis clade, for example, was always significantly different from zero, while the branch connecting Chatogekko with its sister taxon (which was not consistent and changed from tree to tree) was always not significantly different from zero. These similarities among the longer single-gene datasets and their concordance with the concatenated dataset indicate we had enough data for those loci. It is therefore likely that two of the four branches connecting sphaerodactyl genera actually possessed zero branch lengths. These were, in the concatenated nuclear gene dataset, the branch connecting Chatogekko to its sister taxon and the branch connecting Coleodactylus to the Pseudogonatodes + Sphaerodactylus clade. The presence of a hard polytomy in the data has serious implications for our hypothesis testing. Our topology tests were probably unable to distinguish among alternative phylogenetic hypotheses because there were very little data or, in the cases of branches with zero lengths, no data supporting any one phylogenetic hypothesis over the other. This is a difficult situation for testing phylogenetic hypotheses because the lack of data means that essentially any alternative hypotheses involving these short, zero-length branches will not be rejected. The only way of evaluating alternative hypotheses when this occurs is to look to other sources of data. In our case, we had indels and morphological data providing strong evidence that Chatogekko is distinct from Coleodactylus s.s.

Polytomies in gene trees do not automatically translate to hard polytomies in the underlying species trees (Slowinski, 2001; Poe & Chubb, 2004) and the recovery of a bifurcating sphaerodactyl phylogeny is not an impossible task. We show here that rare genomic events such as indels can be used to provide diagnostic characters for sphaerodactyl clades at multiple hierarchical levels. Deletions unique to Chatogekko in RBMX and PTPN12 and unique RAG1 deletions in Coleodactylus provide strong evidence that they are two separate lineages. Indels are considered relatively homoplasy-free characters and have proven useful in diagnosing numerous vertebrate clades (van Dijk et al., 1999; Ericson, Johansson & Parsons, 2000; de Jong et al., 2003; Townsend et al., 2004; Gamble et al., 2008b). Decreasing costs for high-throughput sequencing will make the identification and collection of this sort of data, e.g. indels or LINE/SINE insertions, easier and could prove useful in further untangling the phylogenetic relationships among sphaerodactyl geckos.

**Taxonomy**

We used molecular phylogenetic analyses to identify Chatogekko as a distinct lineage of sphaerodactyl gecko and, with a thorough examination of morphology, provided a suite of diagnostic characters for that lineage. Although some of the characters used to diagnose Chatogekko have been known for a long time, e.g. Coleodactylus has smooth dorsal scales and an ungual sheath composed of five scales while Chatogekko has keeled dorsal scales and an ungual sheath composed of four scales (Vanzolini, 1957, 1968a, b), most of our synapomorphies are new.

The discovery of generic polyphyly resulting from well-sampled phylogenetic analyses is relatively common (Lanyon, 1994; Campbell et al., 2005; Amaral et al., 2006). This problem has been particularly pervasive in geckos where digital morphology, a character suite prone to homoplasy, has played an historically important role in defining genera (Bauer, Good & Branch, 1997; Russell & Bauer, 2002). The classification of sphaerodactyl genera has been similarly dependent on digital morphology (Vanzolini, 1957; Kluge, 1995) and the historical
clustering of Chatogekko with Coleodactylus s.s. was done primarily because of superficial similarities in the ungual sheath (Vanzolini, 1957). By looking beyond the digits, we were able to uncover many morphological characters unique to Chatogekko, strengthening the argument for a taxonomic change.

We recovered three deeply divergent lineages within Chatogekko. These results are consistent with Geurgas & Rodrigues (2010) and Geurgas et al. (2008), who also recovered multiple species-level lineages within C. amazonicus. The geographical distribution of the three Chatogekko lineages corresponds to three described Chatogekko species, two of which are currently synonymized with C. amazonicus. Specimens from Manaus and Roraima correspond to C. amazonicus s.s., with a type locality in the central Amazon near Manaus, Amazonas, Brazil (Andersson, 1918). Specimens from Pará probably correspond to C. zernyi, with a type locality from Taperinha, Pará, Brazil, in the eastern Amazon near Santarém (Wetstein, 1928). Specimens from Rondônia and Rio Ituxi probably correspond to C. guimaraesi, with a type locality in Porto Velho, Rondônia, Brazil, in the southwestern Amazon (Vanzolini, 1957). Although our limited sampling is insufficient to resurrect C. zernyi and C. guimaraesi, the existence of available names for those clades makes such a decision reasonable and the eventual resurrection of these taxa seems inevitable. It should be noted that Geurgas & Rodrigues (2010) also recovered significant phylogenetic structure within Chatogekko amazonicus s.s and C. cf. zernyi. It is possible that splitting each species into two or more species-level lineages may be warranted although additional data would be needed to confirm this.

Coleodactylus and Chatogekko appear to be morphologically conservative and the identification of species-level lineages in both genera using morphology has historically been difficult (Moretti, 2009). Our examination of Chatogekko osteology bears this out. Even though we examined specimens from three putative Chatogekko species we could not identify morphological synapomorphies for these lineages with our data. Our results mirror other morphological analyses of Chatogekko (Vanzolini, 1968b; Avila-Pires, 1995), raising the possibility that species of Chatogekko may be morphologically cryptic. A lineage-based species concept requires that species be diagnosable and genetic evidence and the molecular synapomorphies that support each of the species-level clades within Chatogekko are sufficient to satisfy the need for diagnosability (Zink & McKitrick, 1995; de Queiroz, 1998, 2007; Sites & Marshall, 2004). That said, a thorough examination of morphological characters with a larger sample of specimens in light of the molecular phylogenetic hypothesis could be productive. Other means of identifying species, such as ecological niche modelling, cytogenetics, or multivariate morphometrics, may also prove useful (Raxworthy et al., 2007; Colli et al., 2009; Leaché et al., 2009; Oliver et al., 2009).

**Morphology**

The skull of Chatogekko exhibits interesting modifications associated with miniaturization. The extensive overlapping pattern of the premaxilla is not typical of miniaturized gekkotans (except perhaps in the pygopodid *Pletholax*), although a similar pattern is found in other miniaturized lizards. The uniqueness of the Chatogekko skull compared with other small gekkotans is not surprising, given the association between morphological novelty and miniaturization in vertebrates (Hanken, 1984). The repeated evolution of this overlapping pattern in independent lineages is simply one of several possible solutions to the problems associated with extreme size reduction and highlights the novelty often found in miniaturized taxa.

Miniaturization is often associated with paedomorphosis, the retention of juvenile traits in adult organisms (Gould, 1966; Alberch et al., 1979; Rieppel, 1996). Gekkotans possess several paedomorphic skeletal characters such as amphicoelous vertebrae (Camp, 1923; Cluge, 1967; Werner, 1971) and paired premaxilla or parietal bones (Stephenson, 1960; Cluge, 1967, 1987; Daza, 2008) although none of these skeletal changes are found exclusively in miniaturized forms. In fact, miniaturized species present a fused premaxilla and braincase bones more frequently than larger gekkotans (Daza, 2008). One character that might reflect paedomorphosis in Chatogekko is the slightly larger eyes proportional to the head (Daza et al., 2008), but this would have to be corroborated with a developmental series of different sized sphaerodactyl species.

Another interesting feature of the Chatogekko skull is the development of an incomplete secondary palate. A secondary palate is frequently listed as a very distinct structure in mammals, but is also present in some reptiles. A secondary palate is present in many fossil reptiles (Romer, 1956; Carroll, 1988; Benton, 2005), but among extant groups, this structure appears only in crocodilians, some turtles, and some lizards (Iordansky, 1973; Presch, 1976; Greer, 1977; Gaffney, 1979; Meylan et al., 2000). It has long been thought that no true secondary palate was present in lizards and the tongue was used for closing the naso-pharyngeal passages during respiration (Camp, 1923). The secondary palate in sphaerodactyls resembles that of pygopodids (Conrad, 2008) and xan-
tusiids (Malan, 1946; Savage, 1963). In sphaerodactyls, especially in Chatogekko, the secondary palate is distinctive in that the paleochoanate condition is present, but the palatine is extremely duplicipalatinate, where this bone develops a deep choanal canal formed by the vomerine process and a ventral crest of the palatine. These two structures tend to converge ventrally creating a structure that in cross-section has the shape of a ‘C’; in this sense, the palatines roof over most of the length of the choanal tubes and the ectochoanal cartilage floors the ventral surface, and extends well posteriorly so the nasal passageway opens on the posteromedial side of the palatine (ce, Fig. 10).

The amount of overlap of the premaxilla with the nasal bones, and the contact of nasals have been used as phylogenetic characters (Kluge, 1976), but it has been suggested that they should be treated as independent characters because in certain forms nasal separation can be an artefact of premaxillary overlap, even if the nasals remain in contact with one another (Daza et al., 2008). This seems to be the case in all sphaerodactyls, except in Chatogekko where there is both overlap and complete separation of the nasal bones (i.e. there is no internasal contact). A similar arrangement is present in: the pygopodid Pletholax gracilis (Rieppel, 1984b); the chameleon Rhampoholeon spectrum (Evans, 2008); the xantusiid Cricosaura typica (Savage, 1963); some miniaturized gymnophthalmids such as Bachia bicolor (Tarazona, Fabrezi & Ramirez-Pinilla, 2008), Gymnophthalmus speciosus (MacLean, 1974), Vanzosaura rubricauda (Guerra & Montero, 2009), Calyptommatus nicterus, Scriptosaura catimbau, and Nothobachia ablephara (Roscito & Rodrigues, 2011); many amphibiaenians (Montero & Gans, 2008); and to some extent in the colubrid Scaphiophis albopunctatus (Cundall & Irish, 2008). The loss of the internasal results in some substantial changes to snout configuration and to the distribution of forces; the medial laminar contact between these bones is replaced by an exclusive dorsoventral butt–lap suture with the ascending nasal process of the premaxilla. Open contact sutures are thought to work as shock absorbers or assist to allow micro-movements to dissipate forces acting between bones on the skull (Pritchard, Scott & Girgis, 1956; Jaslow, 1989), behaving in an analogous manner to the flexible material used between the slabs of concrete pavement. The loss of contact, together with the separation of nasals (dashed line in Fig. 5A), com-

Figure 10. Palatal view of cleared and stained Chatogekko amazonicus specimen from Guyana (AMNH-R 132039) showing the secondary palate formed on the palatine. Abbreviations: bp, basipterygoid; bpcp, cartilaginous pad of the basipterygoid process; cc, choanal canal; ec, ectochoanal cartilage; fe, fenestra exochoanalis; pal, palatine; palvp, ventral process of the palatine; pmx-v f, premaxillary–vomer fenestra; pt, pterygoid; sof, suborbital fenestra; v, vomer; vp, vomerine process of palatine. Scale bar = 1 mm.
bined with the loss of a butt–lap joint with the maxilla (a suture present in other sphaerodactyls) suggest that the nasal bones will tend to be less stable and more inclined to move sideways. It has been demonstrated with three-dimensional finite element models that sutures relieve strain locally, but only at the expense of elevated strain in other regions (Moazen et al., 2009). Using this reasoning, a hypoth-
esized reduction in the medial strain on the nasals would have played an important part in the develop-
ment of a posterior interlocking suture with the frontal (Fig. 6). This is purely conjectural, but is
derived from the observed elaborated type of suture and comparison with a similar interlocking suture between nasals and frontal in some amphibiaenians (R. Montero, pers. comm.). The nasofrontal suture of Chatogekko is reciprocally overlapping; the nasal develops a narrow posterior process that overlaps the frontal bone, and the anterolateral process of the frontal overlaps the posterolateral surface of the nasal.

Characters from the postcranium were not diagnos-
tic for Chatogekko; nonetheless it is worthwhile com-
menting on the occurrence of perforated clavicles among sphaerodactyl geckos. The perforation was
described as variable within the gekkonid genus Cnemaspis and considered as the final stage in the
thinning process of the bone, with no phyllogenetic
significance (Smith, 1933). This statement is not
t entirely true for sphaerodactyls, where similar sized
species with comparable clavicles might have unper-
forated (e.g. Lepidoblepharis and Gonatodes) or per-
forated clavicles (e.g. Sphaerodactylus). In the latter,
perforated clavicles are present in both small and
medium-sized species, indicating that this character
might be diagnostic for the genus and have a phylo-
genetic significance at that level.

Another variable trait from the postcranium is the
phalangeal formula. These characters were used in
previous intergeneric cladistic analyses of sphaero-
dactyl geckos (Kluge, 1995). The absence of the fourth
phalangeal element in the fourth finger was one of
the characters that supported the sister relationship of Coleodactylus s.l and Pseudogonatodes; likewise,
Pseudogonatodes was differentiated from Coleodactylus s.l. by the loss of the fourth phalangeal element in
the fourth toe. A re-examination of Kluge’s (1995)
dataset showed that he scored the fourth phalangeal
element in the fourth toe (character 12) as absent in
Coleodactylus, but not in Pseudogonatodes, which is
incorrect. Reanalysis of the corrected dataset does not
produce any change in the topology (J.D.D., unpubl.
data).

We reviewed phalangeal formulae in the specimens
available and encountered a problem of homology. In
all sphaerodactyls, there is a minimum of four pha-
langes in the fourth digits of the manus and pes. Is
the element lost in Coleodactylus, Pseudogonatodes,
and Chatogekko the fourth (penultimate) phalanx and
the remaining element the fifth (ungual)? Or is the
terminal element lost and the fourth phalanx modi-
ﬁed to develop an ungual morphology? The third
phalanx of digit 4 seems to show a fusion of the third
and fourth phalanges in the manus of Pseudogonatodes,
resulting in only four phalanges in this digit. This
process is symmetrical, but in the pes there is no
sign of an ongoing fusion process. If a phalanx was
lost, we would expect to have a shorter digit, although
alternative processes such as nonossification, fusion,
and reabsorption have been discussed (Shapiro,
Shubin & Downs, 2007). To evaluate this, we mea-
sured the length of each digit; these measurements
were converted to equivalent proportions by dividing
each by the length of the shortest digit (i.e. first digit,
Fig. 11). With these values, we estimated the
increased order of digit length in both manus and pes
(Table 3). The manus in Sphaerodactylus, Pseudogo-
натodes, and Coleodactylus presented an increase
order of digit length of 1-5-2-4-3. In Lepidoblepharis
and Gonatodes, the longest digit was the fourth and
in Chatogekko the fourth digit was almost equal
to the third. The situation in Lepidoblepharis is
expected because this genus exhibits no reduction of
any kind in the fourth digit (Fig. 9A). Sphaerodactylus
exhibits similar proportions to those of Pseudogonatodes and Coleodactylus (where one phalanx is lost
or fused to another) because the second phalanx of
digit 4 is very reduced. The second digit in Gonatodes
and Chatogekko is short in comparison with the other
sphaerodactyls. In Chatogekko, digital proportions
differ from all other sphaerodactyls, as digits 2 and 5
and digits 3 and 4 become sub-equal, but the latter
are proportionally longer (Fig. 11). The pes shows a
more stable pattern; in Lepidoblepharis, the order of
increase of digit length is 1-2-3-5-4, while in the rest
of the sphaerodactyl it is 1-2-5-3-4. The only taxon
that showed element loss in the fourth finger was
Pseudogonatodes, a process that is clearly demonstra-
ated by the measurements, as the third and fourth
digits become sub-equal. Developmental data would
be necessary to corroborate fusion or loss of phalanges
in the fourth digit of the manus in Pseudogonatodes,
Coleodactylus, and Chatogekko.

CONCLUSIONS

Small size and cryptic habits have made sphaerodactyl
geckos among the most poorly studied lizard
groups. Our combined use of morphological and
molecular data led to the recognition and description
of a new genus-level lineage of sphaerodactyl gecko,
Chatogekko. Previously considered part of the genus
Coleodactylus, Chatogekko possesses a unique suite of morphological and molecular characters that distinguish it from Coleodactylus s.s. Further work with additional sampling will be necessary to uncover morphological synapomorphies for three putative Chatogekko species and other potentially undescribed taxa in the genus. Our detailed osteological data will provide a framework to move forward with that research, as well as assist more generally with the systematic research of other sphaerodactyl clades.

There are certainly many more sphaerodactyl species to be formally recognized and the use of multiple sources of data, including molecular data and morphology as done here, will be necessary to reveal the true diversity of this fascinating group of lizards.

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SUPPORTING INFORMATION

Additional Supporting information may be found in the online version of this article:

Appendix S1: Specimens used in the morphological analyses.
Table S1: Primers used in the molecular analyses.
Table S2: Specimens used in the molecular analyses.
Figure S1: Split frequencies in run1 vs. run2 for the concatenated Bayesian analysis of the nuclear gene data.
Figure S2: Partitioned Maximum Likelihood phylogeny estimated from the nuclear gene dataset. Black circles indicate nodes with bootstrap values >70.
Figure S3: Bayesian phylogenies for each of the individual loci from the nuclear gene data. Black circles indicate nodes with posterior probabilities >0.95.

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