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A new species of *Brasilotyphlus* (Gymnophiona: Siphonopidae) and a contribution to the knowledge of the relationship between *Microcaecilia* and *Brasilotyphlus*

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Abstract

A third species of *Brasilotyphlus*, a siphonopid caecilian, is described based on six specimens from two twin mountains in Roraima state, northern Brazil. *Brasilotyphlus dubium* **sp. nov.** differs from all other congeners in having a combination of 123–129 primary annuli and 9–16 secondary annular grooves. The first molecular data were generated and analyzed for *Brasilotyphlus*, and the genus was recovered as monophyletic and nested within a paraphyletic *Microcaecilia*. The extent of genetic and taxon sampling, and levels of phylogenetic support are not considered sufficient enough to place *Brasilotyphlus* in the synonymy of *Microcaecilia*.

Key words: Brazil, caecilians, phylogeny, taxonomy

Introduction

Brasilotyphlus Taylor 1968 is a Brazilian endemic genus of caecilian amphibian (Gymnophiona), first described to accommodate *Gymnopsis braziliensis* Dunn 1945 from Manaus municipality, Amazonas. Currently, *B. braziliensis* (Dunn 1945) is known only from its type locality and Reserva do Gavião (02°25' S, 59°50' W), approximately 70 km from Manaus (Maciel & Hoogmoed 2011). Four decades after Taylor's work, a second species of the genus, *B. guarantanus* Maciel, Mott and Hoogmoed 2009, was described from Guarantã do Norte municipality in the state of Mato Grosso and Parauapebas municipality, state of Pará, Brazil (Maciel *et al.* 2009). In the most recent comprehensive taxonomic overview of Gymnophiona, *Brasilotyphlus* was diagnosed by a unique combination of morphological characters within Siphonopidae of a diastema between the vomerine and palatine teeth and eyes covered by bone (Wilkinson *et al.* 2011). Wilkinson & Nussbaum (2006), Maciel *et al.* (2009) and Maciel & Hoogmoed (2011) argued that *Brasilotyphlus* is very similar to *Microcaecilia*, the most species-rich genus of siphonopid caecilian. Considering that the diastema is the main difference between these genera, Maciel *et al.* (2009) and Maciel & Hoogmoed (2011) questioned whether the two genera are valid entities or should be considered synonymous. However, Wilkinson *et al.* (2013) highlighted that the difference between *Microcaecilia* and *Brasilotyphlus* is more complex, because the diastema in *Brasilotyphlus* is large and associated with three other characters: palatine series extending posteriorly slightly further than premaxillary-maxillary series; semicircular

vomerine series; and relatively posterior choanae. One approach to testing the reciprocal monophyly of these two genera would be to use molecular data, however such data are not available for *Brasilotyphlus*.

During two expeditions to the Serra do Apiaú and Serra da Maroquinha, two isolated granitic massifs in Mucajaí municipality, state of Roraima, northern Brazil, some of us (AF, PMSN and SMS) and colleagues found five specimens of an undescribed species of a siphonopid caecilian. Subsequently, an additional specimen deposited in the herpetological collection of the Museu Paraense Emílio Goeldi (Coleção Herpetológica Osvaldo Rodrigues da Cunha, MPEG), Belém, Pará, Brazil, was also identified as the same putative species. Here we describe this as a third species of *Brasilotyphlus*, from c. 620 km NW from the type locality of *B. braziliensis* and 1,500 km NW from the type locality of *B. guarantanus* (Figure 1). Moreover, we provide the first molecular assessment of the systematics of *Brasilotyphlus*.

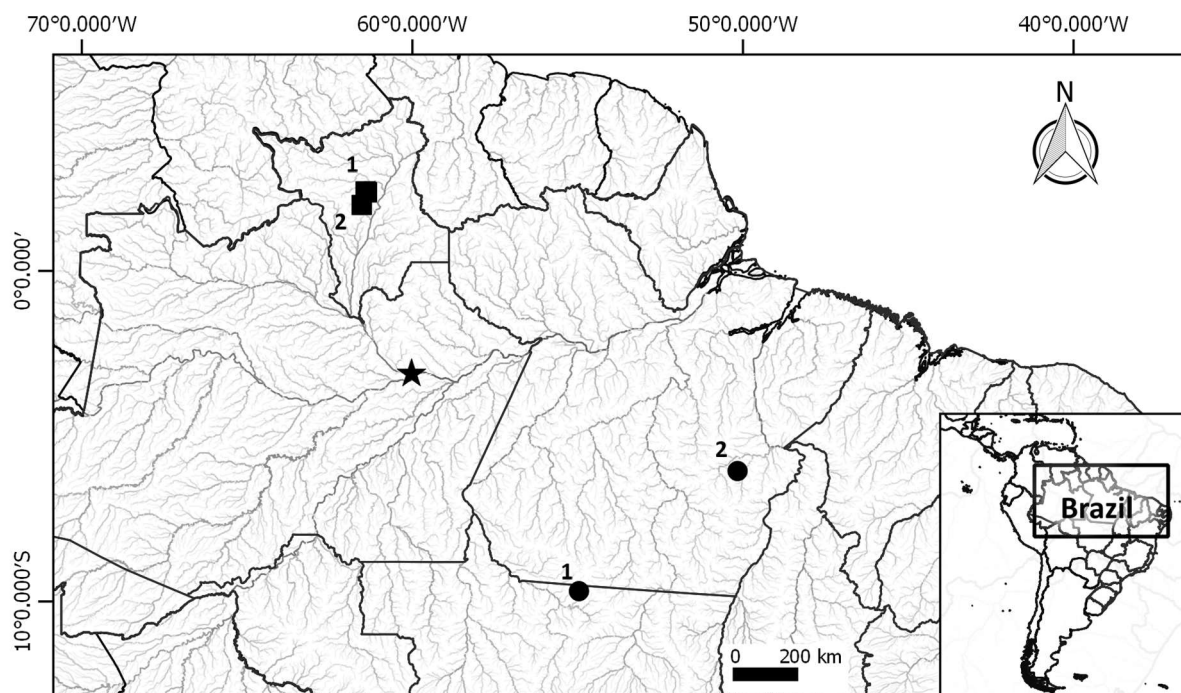


FIGURE 1. Map showing the distribution of *Brasilotyphlus*: star represents the type locality of *B. braziliensis* [Manaus, Amazonas, Brazil (Dunn 1945)]; circle 1 is the type locality of *B. guarantanus* [Municipality of Guarantã do Norte, Mato Grosso, Brazil (Maciel *et al.* 2009)]; circle 2 is the paratype locality of *B. guarantanus* [Municipality of Parauapebas, Pará, Brazil (Maciel *et al.* 2009)]; square 1 is the type locality of *Brasilotyphlus dubium* **sp. nov.** [Municipality of Mucajaí, Roraima, Brazil (this study)]; square 2 is the locality of a *Brasilotyphlus dubium* **sp. nov.** paratype [MPEG 7779; Municipality of Iracema, Roraima, Brazil (this study)].

Material and methods

Morphological data. Six collected specimens were investigated based on nine meristic and 16 morphometric characters. We used the respective abbreviations for morphological characters following Maciel & Hoogmoed (2011) and for anatomy of the nuchal region following Kamei *et al.* (2009). TL—total length; HH—head height at level of corner of mouth; HL—head length, measured from tip of snout to posterior edge of skull (felt through the skin of the dorsal surface of the collar region); HW—head width at corner of mouth; BH—body height at midbody; BW—body width at midbody; WTR—body width at five primary folds anterior to the vent; SP—distance between snout tip and anterior margin of mouth; TMM—distance between tentacle and margin of mouth; TCM—distance between tentacle and corner of mouth; TN—distance between tentacle and nostril; TT—shortest distance between opposite tentacles; NN—shortest distance between opposite nostrils; NCM—distance between nostril and corner of mouth; NMM—distance between nostril and margin of mouth; HWN—head width at level of nostrils;

AIV—primary annular grooves interrupted by vent; CSG—secondary annular grooves completely encircling body; CPA—primary annular grooves completely encircling body; PA—primary annuli; SG—secondary grooves (= number of primary annuli bearing secondary grooves); DT—dentary teeth; PMT—premaxillary-maxillary teeth; PVT—prevomerine teeth; PT—palatine teeth; NG1—first nuchal groove (between head and collars); NG2—second nuchal groove (between first and second collars); NG3—third nuchal groove (between collars and anteriormost annulus). Dermal and subdermal scales were examined following Wilkinson *et al.* (2013).

TABLE 1. Sequences of Gymnophiona used in this study. Taxa, voucher numbers and NCBI GenBank accession numbers. *from this study.

Terminal taxa	Voucher	16SrRNA	Cytb
<i>Brasilotyphlus guarantanus</i> 1	MUFAL10363	MG162597*	MG162605*
<i>Brasilotyphlus guarantanus</i> 2	MUFAL10379	MG162598*	MG162606*
<i>Brasilotyphlus guarantanus</i> 3	UFMT5009	MG162599*	-
<i>Brasilotyphlus guarantanus</i> 4	UFMT5012	MG162600*	-
<i>Brasilotyphlus dubium</i> sp. nov. 1	MUFAL13638	MG162601*	MG162607*
<i>Brasilotyphlus dubium</i> sp. nov. 2	MUFAL13639	MG162602*	MG162608*
<i>Brasilotyphlus dubium</i> sp. nov. 3	MUFAL13641	MG162603*	MG162609*
<i>Boulengerula fischeri</i>	BMNH 2008.607	FR691661	FR691666
<i>Boulengerula taitanus</i>	NMK A/3112	AY954504	AY954504
<i>Caecilia gracilis</i>	BMNH:2008.747	KF540147	KF540147
<i>Caecilia tentaculata</i>	BMNH:2008.750	KF540146	KF540146
<i>Chikila fulleri</i>	BNHS 5514	NC_021369	NC_021369
<i>Crotaphatrema lamottei</i>	BMNH:2008.274	NC019596	NC019596
<i>Dermophis mexicanus</i>	No voucher	KF540150	KF540150
<i>Gegeneophis ramaswamii</i>	MW 331	AY456250	AY456250
<i>Gegeneophis seshachari</i>	BNHS:4601	HQ444015	HQ444103
<i>Geotrypetes seraphini</i> 1	MVZ 252475	GQ244469	GQ244469
<i>Geotrypetes seraphini</i> 2	BMNH 2005.2	AY954505	AY954505
<i>Gymnopsis multiplicata</i> 1	MVZ:Herp:179536	NC020139	NC020139
<i>Gymnopsis multiplicata</i> 2	MVZ 171331	GQ244471	GQ244471
<i>Ichthyophis glutinosus</i>	MW 1733	AY456251	AY456251
<i>Luetkenotyphlus brasiliensis</i>	BMNH:2005.3	KF540158	KF540158
<i>Microcaecilia dermatophaga</i>	BMNH:2008.716	KF540159	KF540159
<i>Microcaecilia</i> sp1.	MTR20960	MG162604 *	MG162610 *
<i>Microcaecilia savagei</i>	MAD 496	EU753992	EU754004
<i>Microcaecilia</i> sp2.	IWK 0128	GQ244473	GQ244473
<i>Microcaecilia</i> sp2.	No voucher	EU753993	EU754005
<i>Microcaecilia unicolor</i>	BMNH:2008.749	KF540160	KF540160
<i>Rhinatrema bivittatum</i>	BMNH 2002.6	AY456252	AY456252
<i>Rhinatrema shiv</i>	IRSNB:1991	GU566189	GU566190
<i>Schistometopum gregorii</i>	BMNH:2008.559	KF540163	KF540163
<i>Schistometopum thomense</i>	CAS 219292	GQ244476	GQ244476
<i>Scolecormorphus vittatus</i>	BMNH 2002.100	AY456253	AY456253
<i>Siphonops annulatus</i> 1	BMNH 2005.9	NC007911	NC007911
<i>Siphonops annulatus</i> 2	MVZ 162588	EU753986	EU754003
<i>Typhlonectes natans</i> 2	No voucher	NC002471	NC002471
<i>Typhlonectes natans</i> 1	FC 13178	EU753984	EU753999

Comparisons with *B. braziliensis* were made using only the literature (Taylor 1968; Maciel & Hoogmoed 2011) and with *B. guarantanus* using two specimens from Parauapebas, Pará, Brazil (Museu de História Natural da Universidade Federal de Alagoas, MUFAL 10363; 10379) and literature data (Maciel *et al.* 2009). A small ventral incision was made and all specimens were sexed by direct examination of the gonads. Specimens were classified as immature when gonads could not be clearly differentiated.

Molecular data. New molecular genetic data were generated from seven specimens including three specimens of the new species (MUFAL 13638, 13639, 13641) and four *B. guarantanus*, two from the type locality, Guarantã do Norte, Mato Grosso (Universidade Federal de Mato Grosso, UFMT 5009 and the paratype UFMT 5012), and two specimens from the paratype locality, Parauapebas, Pará, ca. 661 km from type locality (MUFAL 10363, 6° 3' 8" S, 50° 7' 40" W; and 10379, 6° 3' 29" S, 50° 11' 50" W). Total genomic DNA was extracted using a salt precipitation method (Fetzner 1999). Fragments of two mitochondrial genes (cytochrome b and 16SrRNA) were amplified and sequenced using the primers MNCN-Glu and CytB (San Mauro *et al.* 2004), and 16sar and 16sbr (Palumbi *et al.* 1991) respectively. PCR and sequencing protocols used were those developed by San Mauro *et al.* (2004) for caecilians. All sequences were edited and assembled using the default settings in Bioedit v.7.2.5 (Hall 1999), sequences of cytochrome b were additionally translated into amino acids for confirmation of alignment. To infer the phylogenetic placement of the new species and that of *B. guarantanus*, sequences of cytochrome b and 16SrRNA from 37 species of caecilians representing all ten families for which homologous sequences were available in Genbank and including all siphonopid species (accession numbers available in Table 1) were downloaded and incorporated in the data matrix. The 16SrRNA and cytochrome b alignments are deposited on TreeBASE (<http://purl.org/phylo/treebase/phyloids/study/TB2:S22971>).

We identified the best-fitting substitution models for each gene using jModeltest 2.1.7 (Posada 2008) with the Akaike information criterion (AIC). The best-fit substitution model was TIM2+I+G for 16srRNA and TPM3uf+I+G for cytochrome b, however, because these models are not implemented in MrBayes 3.4.5 (Huelsenbeck & Ronquist 2001), the GTR+I+G model was implemented for the concatenated dataset. Bayesian analyses were conducted with a concatenated matrix using MrBayes 3.4.5 (Huelsenbeck & Ronquist 2001). In order to explore parameter space thoroughly, we implemented four incrementally heated Metropolis-coupled Markov chains Monte Carlo (MCMC), using random trees as an initial step for chains, and ran them for 20 million generations each, sampling every 1,000 generations. Because higher ESS values indicate lower standard errors, MCMC sampling was considered adequate if the effective sampling size (ESS) for each parameter was higher than 200 and if the standard deviation of the split frequencies was lower than 0.01, both checked using Tracer v1.6 (Rambaut *et al.* 2014). The first 25% of the samples were discarded as burnin. Analyses were summarized with a 50% majority rule consensus. The reliability of each node was evaluated based on the posterior probability (PP) in the Bayesian analysis.

We tested the reciprocal monophyly of *Brasilotyphlus* and *Microcaecilia* in a maximum likelihood (ML) phylogenetic framework using the Approximately Unbiased (AU) test (Shimodaira 2002). We compared the optimal maximum likelihood tree to the ML tree in which all sampled *Microcaecilia* were enforced as monophyletic. ML phylogenies were constructed using RAxML 8.2.10 (Stamatakis 2014) on the CIPRES Science Gateway (Miller *et al.* 2010). Data were partitioned by gene, and nodal support estimated using rapid bootstrapping (Stamatakis *et al.* 2008), which was stopped automatically after 102 replicates. The AU test was implemented in PAUP* v4.0a161 (Swofford 2003) with likelihood scores estimated using the GTR+G+I model and p-values generated using 10,000 RELL bootstrap replicates.

Results

Brasilotyphlus dubium sp. nov.

Figs. 2–4; Table 2

Holotype. Museu de História Natural da Universidade Federal de Alagoas (MUFAL) 13638, field number MTR 23151, a male collected by Pedro M. Sales Nunes, Antoine Fouquet and Felipe Franco Curcio, from Serra da Maroquinha (N 2° 22' 44", W 61° 22' 37"), 400 m from sea level, Mucajaí municipality, Roraima, Brazil, May 2012.

Paratypes (n = 5). MUFAL 13639, male, field number MTR 23216, collected in pitfall trap, with same collection data as the holotype; MUFAL 13640, field number SMS 873, immature, MUFAL 13641, field number SMS 920, female, and MUFAL 13642, field number SMS 940, female, collected by Sergio Marques de Souza, Pedro M. Sales Nunes and Antoine Fouquet, from Serra do Apiaú (N 2° 24' 30", W 61° 24' 54") at 140 m, 685 m and 835 m from sea level respectively, state of Roraima, Brazil, in November 2011; and MPEG 7779, female, collected by Laurie J. Vitt, from 7 km east of the Ajarani river (N 1° 59' 50", W 61° 32' 4"), Iracema municipality, Roraima, Brazil, 3 July 1993.

Identification. The new species is considered a species of *Brasilotyphlus* on the basis of it having eyes covered by bone and a diastema between the vomerine and palatine teeth with the following associated characters: palatine extends posteriorly to the series of premaxillary-maxillary teeth; a semicircular vomerine series of teeth.

Diagnosis. *Brasilotyphlus dubium* **sp. nov.** differs from *B. braziliensis* in having fewer primary and secondary annular grooves (123–129 and 9–16 vs 142–147 and 23–36, respectively), and in having premaxillary-maxillary teeth reaching the level of the posterior edge of the choanae (in *B. braziliensis* the maxillary teeth do not reach the level of the choanae). The new species differs from *B. guarantanus* in having fewer primary annuli (123–129 vs 151–170) and more secondary annular grooves (9–16 vs 0–2).



FIGURE 2. MUFAL 13638, holotype of *Brasilotyphlus dubium* **sp. nov.** from Serra da Maroquinha, Roraima, Brazil. Left column: head and anterior part of body in dorsal (above), lateral (middle) and ventral view (below). Right column: posterior part of body in dorsal (above), lateral (middle) and ventral view (below).

Description of the holotype. Morphometric and meristic data are in Table 2. Specimen in good condition, a 3.7 mm midventral incision and slightly broken corners of the mouth. Body subcylindrical, slightly flattened dorsoventrally throughout (BW x BH 3.9 x 2.9 mm), slightly narrower anteriorly and posteriorly (WTR 3.6 mm). In dorsal view, head intermediate between U- and V-shaped. In lateral view, top of head slightly convex; upper lip

slightly concave at the anterior end and lower lips straight. Snout projects strongly beyond recessed mouth (SP 1 mm). Eyes not visible. Tentacles slightly elevated and not visible from above, closer to the corner of the mouth (TCM 1.3 mm) than to nares (TN 1.6 mm). Nares visible from above, but not from below. Teeth pointed and gently recurved. Premaxillary-maxillary teeth monocuspid, forming a series (21 teeth) with posterior maxillary teeth slightly smaller, extending to the level of the posterior edge of the choanae. Nine prevomerine and 10 palatine bicuspid teeth, with no apparent variation in size, smaller than those of the premaxillary-maxillary series and with a large diastema between them, which corresponds to a distance of approximately three tooth positions. Dentary teeth monocuspid, forming a series of 13 teeth, posterior ones slightly smaller, larger than those of premaxillary-maxillary series. Subcircular choanal apertures, separated by an approximate distance of 1.5 times the width of each choanal aperture, anterior edges approximately level with tentacles. Two collars clearly marked by three nuchal grooves (NG1, NG2, and NG3); NG1 and NG2 completely encircling the body, NG3 incomplete, curving posteriorly on the venter. In dorsal view, NG1 straight, NG2 slightly curved anteriorly, and NG3 straight; first collar smaller than second. In ventral view, NG1 curved anteriorly, NG2 straight and NG3 slightly incomplete and curved posteriorly; first and second collar similar in size. NG1 oblique laterally. A conspicuous transverse groove is present on the dorsolateral surface of the second collar. Following collars, 125 PAs, being 123 complete and two interrupted by the vent; First SG short, dorsally located on 112th PA; SGs complete from 119th PA. Vent with six main denticulations and some irregular subdivisions, the interdenticular creases shorter anteriorly. Dorsally, body terminus strongly convex. Distinct vertical terminal keel present. Annular scales limited to a single and incomplete row in the 92th groove of scales wider than long (e.g., 0.1 x 0.2 mm); in a single incomplete row at 107th groove (e.g., 0.1 x 0.3 mm) and in a complete row of ovate scales at 120th groove (e.g., 0.4 x 0.8 mm).

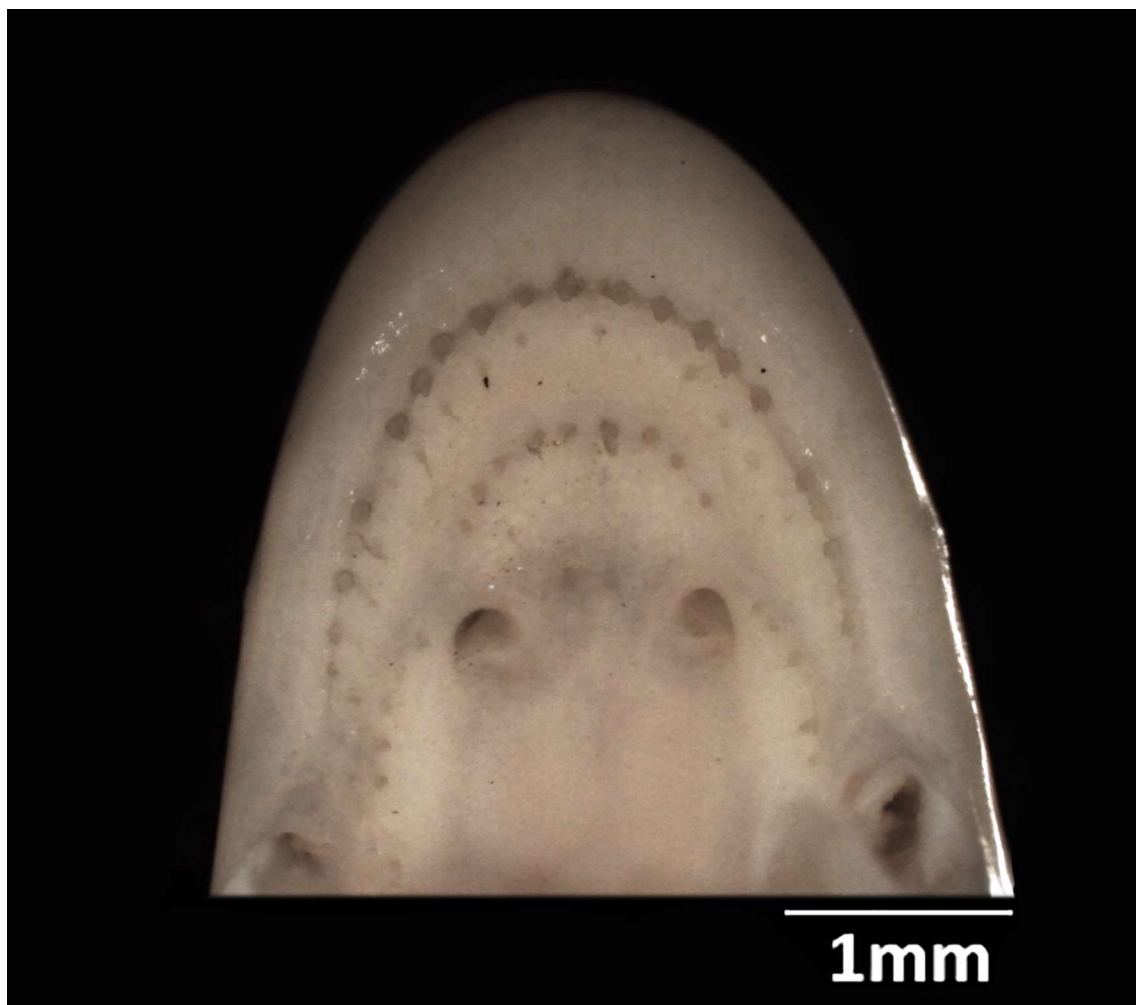


FIGURE 3. Palatal view of *Brasilotyphlus dubium* sp. nov. (holotype) showing disposition of tooth rows, diastema and choanae. Scale bars = 1 mm.

Coloration: In life, body pale purple; head pinkish. Venter and lateral surfaces, areas surrounding vent, nostrils and tentacles paler than dorsum. In preservative, body brownish; paler anteriorly than posteriorly in dorsal and ventral view. Ventral and lateral surfaces slightly paler than dorsum along the entire body. Areas surrounding the vent, nostrils, tentacles and lips less pigmented, as is the ventral surface anterior to the second collar.

TABLE 2. Morphometric (in mm) and meristic data for the type series of *Brasilotyphlus dubium* **sp. nov.** [data from this study (MUFAL 13638-13642 and MPEG 7779), *B. guarantanus* [data from this study (MUFAL 10363 and 10379) and from Maciel & Hoogmoed (2011)] and *B. braziliensis* [from Taylor (1968) and Maciel & Hoogmoed (2011)]. * = Holotype. Abbreviations are given in text.

	<i>B. dubium</i> sp. nov.						<i>B. guarantanus</i>			<i>B. braziliensis</i>
	MUFAL 13638*	MUFAL 13639	MUFAL 13642	MUFAL 13641	MUFAL 13640	MPEG 7779	MUFAL 10363	MUFAL 10379	range	range
TL	16.8	18.1	18.2	14.2	11.2	13.7	32.8	23.6	164–305	147–260
HW	3.5	3.5	3.2	2.8	2.7	2.8	3.9	3.6	2.5–4.2	2.7–5.1
HWN	1.9	1.9	1.6	1.4	1.3	1.3	2.1	2	–	–
HL	4.9	5.1	4.7	4.6	4.2	4.6	5.1	5.3	3.8–6.2	4.5
HH	2.5	2.7	2.6	2.4	2.2	1.9	3.1	2.7	1.5–3.4	2
SP	1	1	1	1.1	0.9	1.1	1.0	1.2	0.8–1.5	1.2
BW	3.9	4.3	3.6	3.3	3	3.5	5.7	4	3.3–4.9	3–6
BH	2.9	3.6	3.2	2.4	2.3	2.5	4	3.3	2.7–4.3	2.8
WTR	3.6	3.7	3.6	3	3	3.2	4.8	3.4	2.9–4.7	3.1
NN	1.3	1.5	1.2	1.2	1.1	0.9	1.8	1.6	0.9–1.5	1.2
NMM	0.9	1	0.8	0.8	0.7	1	1.1	1.2	0.6–1.2	0.7
NCM	3.2	3.3	3.3	3	2.1	3	–	–	–	–
TN	1.6	1.6	1.6	1.4	1.2	1.5	2.0	1.9	1.3–2.4	1.5–2.6
TT	3.2	3.2	2.7	2.4	2.3	2.3	–	–	2.2–3.4	2.6
TMM	0.3	0.3	0.3	0.3	0.3	0.3	0.4	0.4	0.2–0.5	0.2
TCM	1.3	1.4	1.7	1.6	1.1	1.5	1.5	1.6	0.8–1.6	0.8
PMT	21	22	23	22	22	19	19	18	17–24	15–25
PVT	9	8	9	8	8	7	11	8	8–13	8–12
PT	10	10	12	10	10	12	12	14	8–14	10–14
DT	13	18	15	14	16	16	16	17	13–21	13–18
PA (CPA)	125 (123)	128 (126)	123 (121)	129 (126)	128 (125)	125 (121)	162 (160)	150 (147)	151–170	142–147
SG (CSG)	13 (6)	9 (3)	15 (6)	16 (5)	9 (3)	14 (2)	0	0	0–2	23–36 (4–7)
AIV	2	2	2	3	3	3	2	3	0–3	2

Variation and additional information from paratypes. Variation in some meristics and morphometrics is summarized in Table 2. The vertical terminal keel is less distinct in MUFAL 13642. In preservative, MUFAL 13641 and 13642 have steel grey body color, but the pattern of paler regions is similar to that observed on the holotype. The first collar of MUFAL 13641 and 13642 are less robust than observed in the holotype, forming an almost straight transition line from head to body. In paratype MPEG 7779, scales begin in the 90th groove, limited to a single and incomplete row of small scales, wider than long (e.g., 0.1 x 0.3 mm); in a single incomplete row at 109th groove (e.g., 0.1 x 0.4 mm) and in a complete row of ovate scales at 122th groove (e.g., 0.5 x 0.7 mm). Except for minor details in their visibility, the nuchal grooves of all paratypes are as described for the holotype.

Etymology. The epithet *dubium* means “dubious”, reflecting our doubt whether or not *Brasilotyphlus* should be considered a synonym of *Microcaecilia*. For nomenclatural purposes, the species epithet is considered a noun in apposition.



FIGURE 4. *Brasilotyphlus dubium* **sp. nov.** (A) Specimen from the type series in life in dorsal view. (B) Habitat at type locality, Serra da Maroquinha, Roraima, Brazil. (C) Habitat at Serra do Apiaú, Roraima, Brazil.

Phylogenetic analyses. The concatenated dataset consisted of 1,682 base pairs (bp) (605 bp of 16srRNA and 1,684 bp of cytochrome b). After removal of 121 bp in the 16SrRNA fragment (31–43, 218–284, 311–354) due to ambiguous alignment, 1,563 bp were used in the analyses. The families Caeciliidae, Typhlonectidae, Herpelidae, Indotyphlidae and Siphonopidae were each recovered as monophyletic with high support (Figure 5). Typhlonectidae and Caeciliidae were recovered as sister clades. The basal split within Siphonopidae is between a clade comprising *Luetkenotyphlus brasiliensis* (Lütken 1851) and *Siphonops annulatus* (Mikan 1820) (PP = 1), and a clade comprising all sampled species of *Microcaecilia* [*M. unicolor* (Duméril 1863), *M. dermatophaga* Wilkinson, Sherratt, Starace & Gower 2013, *M. savagei* Donnelly & Wake 2013, *Microcaecilia* sp1. and *Microcaecilia* sp2.], *B. guarantanus* and *B. dubium* **sp. nov.** (PP = 1). The latter two were recovered as monophyletic (PP = 1), but nested within a paraphyletic *Microcaecilia* (*M. savagei* and *M. sp.2* are more closely related to *Brasilotyphlus* than to other *Microcaecilia*: PP = 0.86).

The maximum likelihood tree (not shown) is similar to our Bayesian phylogeny at well-supported nodes. The families Caeciliidae, Typhlonectidae, Herpelidae and Indotyphlidae were each monophyletic with high support (bootstrap value [BS] ≥ 97), and although Siphonopidae was recovered with moderate support (BS = 79), the relationships among *Brasilotyphlus* and *Microcaecilia* were identical to those recovered in the Bayesian analysis. The AU test could not reject a monophyletic *Microcaecilia* (difference $-\ln L = 1.77643$; AU test, $P = 0.2099$).

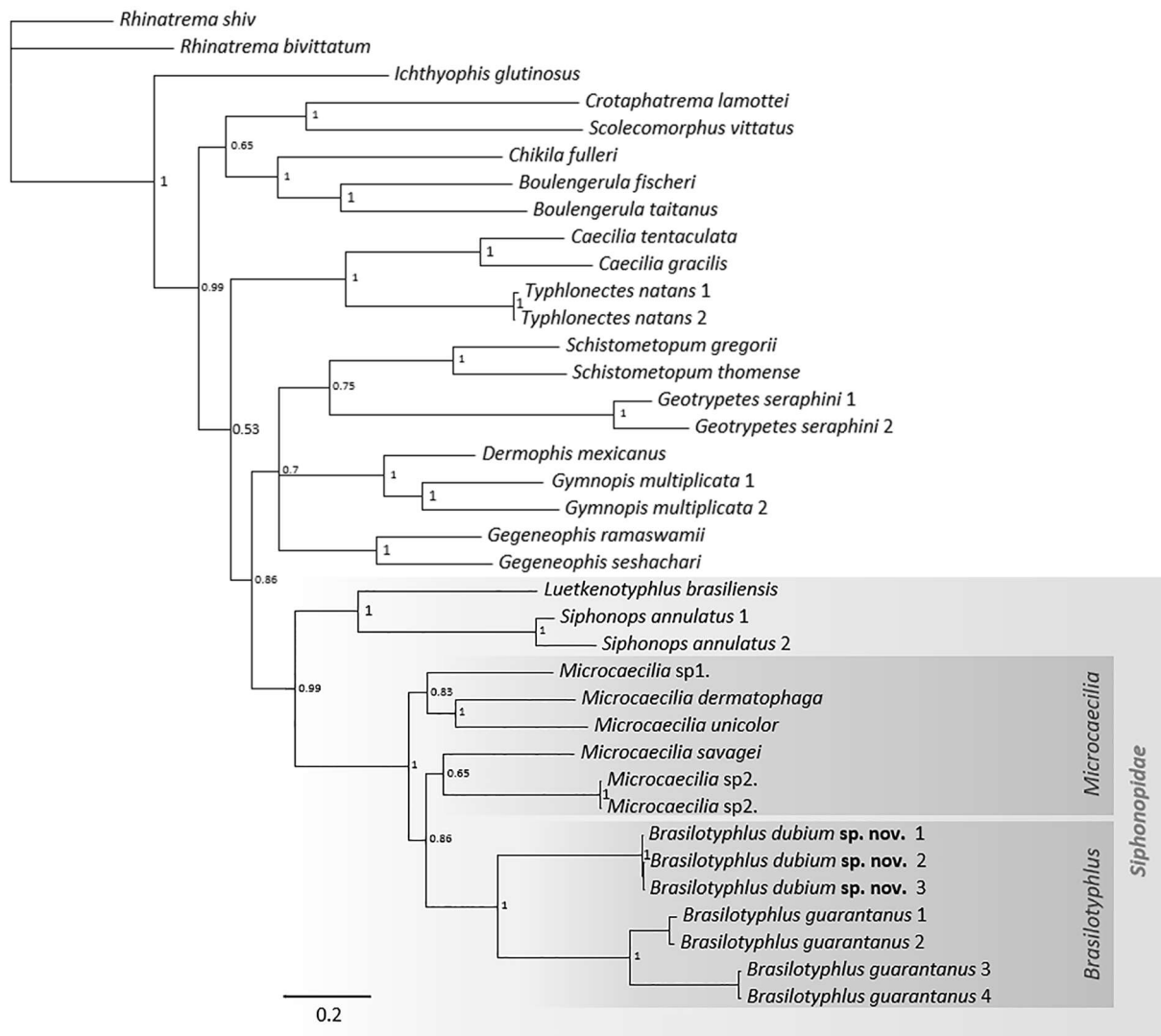


FIGURE 5. Phylogram obtained from Bayesian analysis (consensus by majority rule 50%) using 1,683 base pairs of concatenated mitochondrial genes (16SrRNA and cytochrome b). Numbers at internal branches are posterior probabilities.

Discussion

To date, *Brasilotyphlus* had not been included in any molecular phylogenetic analysis and the doubts regarding its monophyly and relationship with *Microcaecilia* were, until this study, based only on morphological data. Our mitochondrial DNA-based phylogeny includes a monophyletic *Brasilotyphlus* nested within a paraphyletic *Microcaecilia*, suggesting that *Brasilotyphlus* could be considered a synonym of *Microcaecilia* or that *Microcaecilia* should be split. However, the AU test was unable to reject the hypothesis that *Brasilotyphlus* could be considered a synonym of *Microcaecilia* and, thus, we prefer to be conservative and retain both genera because (1) phylogenetic support for the position of *Brasilotyphlus* is moderate; (2) our topology tests could not reject a monophyletic *Microcaecilia*; (3) we were not able to sample the type species of either genus; (4) we inferred phylogenetic relationships only on the basis of two mtDNA genes. We suggest that additional molecular sampling (taxa, loci and genes) should be done to clarify the relationship between these similar genera of siphonopid caecilians.

The individuals of *Brasilotyphlus dubium* **sp. nov.** from Mucajaí municipality, state of Roraima, Brazil, were collected in two adjacent mountains (Serra do Apiaú and Serra da Maroquinha), both massifs are forest covered and isolated from each other by c. 4 km of secondary forests and pasture lowlands. The individual from Iracema

municipality, state of Roraima, Brazil, was also collected in forest. Species of the genus *Brasilotyphlus* had been considered to be restricted to areas of rainforest (Rodrigues *et al.* 2004; Maciel *et al.* 2009). Therefore, we hypothesize that *B. dubium* may also be restricted to rainforest areas. However, more extensive sampling is necessary to understand the distribution and ecology of this new species.

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