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# A NEW SPECIES OF OSTEOCEPHALUS STEINDACHNER, 1862 (ANURA, HYLIDAE), FROM BRAZILIAN AMAZONIA

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ABSTRACT. Treefrogs of the genus Osteocephalus have been the focus of several taxonomic and phylogenetic studies, especially in the last two decades. These recent studies have shown that the diversity of this charismatic Amazonian genus is still largely underestimated. Through the evidence of morphological and molecular data, we describe a new species of the Osteocephalus alboguttatus species group from the Purus-Madeira Interfluve, southwestern Brazilian Amazonia. The new species differs from other Osteocephalus by having a small body size (snout–vent length 32.1–44.1 mm), skin texture non-sexually dimorphic, dorsum smooth with a few scattered small tubercles, vocal sac single and subgular, frontoparietal ridges not externally visible, and a dark tan brown iris with lighter vermiculation. The rapid increase in the number of new frog species described from the Purus-Madeira Interfluve highlight the importance of sampling poorly explored and remote areas in Amazonia, as well as the value of supporting taxonomic research to accelerate species documentation in face of the biodiversity crisis.

KEY WORDS: Amazonas; Arapixi Extractive Reserve; bamboo-dominated forests; Osteocephalus alboguttatus species group; Purus-Madeira Interfluve; taxonomy

#### INTRODUCTION

The genus Osteocephalus Steindachner, 1862 (Amphibia, Anura, Hylidae), harbors 28 treefrog species distributed across Amazonian lowland rainforests and Andean foothills (Jungfer et al., 2013; AmphibiaWeb, 2021). The first attempt to recognize the diversity in this genus was made by Goin (1961), who provided a key to the genera of hylid frogs, with diagnosis to Osteocephalus, including skin texture, shape of vocal sac in males, and cranial osteology. Under this

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combination were recognized five species: Osteocephalus britti (Melin, 1941), Osteocephalus buckleyi (Boulenger, 1882), Osteocephalus leprieurii (Duméril and Bibron, 1841), Osteocephalus pearsoni (Gaige, 1929), and Osteocephalus taurinus Steindachner, 1862. This author also included Osteocephalus planiceps Cope, 1874, in the synonymy of O. leprieurii.

Trueb and Duellman (1971) were the first to review the taxonomy of this genus extensively and provided relevant baselines for the advancement of associated knowledge. However, the ''conservative approach'' in which their taxonomic decisions were based also led to a series of misinterpretations. For example, they transferred O. planiceps (that was in the synonymy of O. leprieurii sensu Goin, 1961) and Osteocephalus vilarsi (Melin, 1941) to the synonymy of O. taurinus. They also synonymized Osteocephalus festae (Peracca, 1904), Osteocephalus cabrerai (Cochran and Goin, 1970), and Osteocephalus carri (Cochran and Goin, 1970) with O. buckleyi, resulting in a considerable reduction of the species content of Osteocephalus at that time. Subsequent advances in taxonomy of the genus included the description of Osteocephalus subtilis by Martins and Cardoso (1987), the expansion of generic concept by Duellman and Hoogmoed (1992), the description of Osteocephalus oophagus by Jungfer and Schiesari (1995), and the resurrection of O. cabrerai and O. planiceps from the synonymy of O. buckleyi and O. taurinus, respectively (Duellman and Mendelson, 1995). In the last few decades, the aggregation of new morphological data and advances in analytical capacity led to a massive increase in the taxonomic knowledge of Osteocephalus, with the description of half of its currently known species diversity (e.g., Ron and Pramuk, 1999; Jungfer et al., 2000; Jungfer and Lehr, 2001; Jungfer and Hödl, 2002; Lynch, 2002; Moravec et al., 2009;

Jungfer, 2010, 2011; Ron et al., 2012; Jungfer et al., 2013, 2016; Duellman, 2019; Chasiluisa et al., 2020).

Following the advances in taxonomic knowledge, recent studies addressed the phylogenetic relationships within Osteocephalus (Moravec et al., 2009; Ron et al., 2010, 2012; Salerno et al., 2012; Jungfer et al., 2013; Ferrão et al., 2019; Chasiluisa et al., 2020). The most comprehensive phylogenetic revision revealed that Osteocephalus was in fact paraphyletic (Jungfer et al., 2013). To solve this issue, some species formerly attributed to Osteocephalus were transferred to the genus Tepuihyla Ayarzagüena, Señaris, and Gorzula, 1993, and to the newly erected genus Dryaderces (Jungfer et al., 2013). Furthermore, five species groups were defined by Jungfer et al. (2013) on the basis of molecular data: Osteocephalus alboguttatus (Boulenger, 1882), O. buckleyi, O. leprieurii, O. planiceps and O. taurinus species groups.

The *O. alboguttatus* species group was recognized to accommodate three species from western Amazonia (Jungfer et al., 2013): O. alboguttatus, Osteocephalus heyeri Lynch, 2002, and O. subtilis Martins and Cardoso, 1987. Despite the absence of morphological synapomorphies supporting this species group, its members share a small body size (snout–vent length  $<$  50 mm), dark irises, and the absence of keratinized spicules on dorsum. Osteocephalus alboguttatus was firstly described as Hyla alboguttata by Boulenger (1882) and transferred to Osteocephalus more than a century later by Wiens et al. (2006). On the other hand, O. subtilis and O. heyeri were described without arguments supporting its generic allocation. Although the associated information on O. heyeri and O. subtilis is sparse, the geographic ranges of these species seem narrowly restricted (Jungfer et al., 2013). Data on reproductive behavior are also scarce for

members of this species group, but the smooth dorsum and subgular vocal sacs led Jungfer et al. (2013) to hypothesize that they are phytotelm-breeders. However, this hypothesis is still treated as uncertain (Blotto et al., 2021).

During fieldwork in southwestern Brazilian Amazonia, the first author (P.R.M.-S.) collected some specimens of Osteocephalus with morphological affinities with members of the *O. alboguttatus* species group. Nevertheless, we cannot attribute these specimens to any nominal species on the basis of their morphological distinctiveness, which supports them as representatives of a new species. In this study, we combined both morphological and molecular data to describe this new species and allocate it into the O. alboguttatus species group.

# MATERIAL AND METHODS

Three adult specimens were collected in the municipality of Sena Madureira (state of Acre) in November 2011 and Arapixi Extractive Reserve (municipality of Boca do Acre, state of Amazonas) in January 2016, both localities in southwestern Brazilian Amazonia. Specimens were euthanatized with peritoneal injection of 2% lidocaine, fixed with 10% neutral buffered formalin, and stored in 70% ethanol. Liver tissue samples were obtained before fixation and preserved in 100% ethanol. We analyzed the external morphology of the newly collected specimens and additional Osteocephalus deposited at the following zoological collections: Museu Nacional (MNRJ), Rio de Janeiro, Brazil; Universidade Federal do Acre (UFAC-RB), Rio Branco, Brazil; Museu de Zoologia da Universidade de São Paulo (MZUSP), São Paulo, Brazil; Instituto Nacional de Pesquisas da Amazônia (INPA-H), Manaus, Brazil; Coleção Zoológica Paulo Burhein, Universidade Federal do Amazonas (CZPB), Manaus, Brazil; Museum of Comparative Zoology (MCZ), Harvard University, Cambridge, Massachusetts, U.S.A., Biodiversity Museum University of Kansas (KU), Lawrence, Kansas, U.S.A. (specimens listed in the Appendix 1).

We used a digital caliper under a stereomicroscope to the nearest 0.1 mm to take 10 morphometric measurements following Watters et al. (2016): horizontal eye diameter (ED), eye–nostril distance (EN), foot length (FL), head length (HL), head width (HW), interorbital distance (IOD), snout–vent length (SVL), horizontal tympanum diameter (TD), tibia length (TL), and upper eyelid width (UEW). Color in life was analyzed and described from field notes and digital photographs. Sex was determined by the presence of vocal slits and nuptial excrescences. Webbing formulae followed the standardization of Savage and Heyer (1967) as adapted by Myers and Duellman (1982). Morphological terminology followed Watters et al. (2016).

For molecular analyses, we sequenced a 547–base pair (bp) fragment of the 16S ribosomal RNA, considered a universal barcode for Neotropical amphibians (Vences et al., 2005). Genomic DNA was extracted by standard protocols for a commercial kit (Wizard® Genomic DNA Purification Kit, Promega, Wisconsin, U.S.A.), with the target fragment amplified by polymerase chain reaction (PCR) with the primers 16Sar and 16Sbr and standard protocols (Palumbi et al., 1991). The PCR product was purified with PEG (polyethyleneglycol) 8000 and sequenced by standard protocols of the Big Dye Terminator Kit (Applied Biosystems, Waltham, Massachusetts, U.S.A.) in an ABI PRISM 3500 (Applied Biosystems) automated sequencer. We used Geneious (Biomatters, Auckland, New Zealand; Kearse et al., 2012) to check and edit the raw sequence.

The newly generated 16S sequence was included in a data set of homologous sequences downloaded from GenBank representing 48 individuals, 22 species, and all species groups of Osteocephalus. Sequences representing the closely related genera Dryaderces, Tepuihyla, Itapotihyla, and Phyllomedusa were used as outgroups. Information related to sequences used in molecular analyses are presented in Appendix 2.

The molecular dataset was aligned by the MAFFT (multiple alignment using fast Fourier transform) online server through the E-INS-i strategy with default parameters (Katoh and Standley, 2013). With the use of the aligned dataset with 550 bp and 55 terminals, we performed phylogenetic inferences under Bayesian (BI) and maximum likelihood (ML) optimality criteria. Best adjusted model of nucleotide evolution for these inferences was determined under the Bayesian information criterion (BIC) with PartitionFinder v2.1.1 (Lanfear et al., 2017). The BI inference was conducted in MrBayes v3.2.6 (Ronquist et al., 2012) with two parallel runs of the Markov chain Monte Carlo (MCMC) algorithm with 10 million iterations. Convergence of parameters (split frequencies  $SD < 0.01$  and estimated sample size  $> 200$ ) was assessed with Tracer v1.7 (Rambaut et al., 2018), and the maximum clade credibility tree was extracted after 25% burn-in. The ML inference was conducted in the IQ-TREE (Nguyen et al., 2015) online server (Trifinopoulos et al., 2016). Branch support was estimated with 10,000 ultrafast bootstrap replications, 1,000 maximum iterations, and a minimum correlation coefficient of 0.99. Following Huelsenbeck and Ronquist (2005) and Minh et al. (2013), we consider as highly supported branches those with posterior probability values  $>0.95$  (BI) and ultrafast bootstrap values greater than 95 (ML). We used MEGA v7 (Kumar et al., 2016) to compute uncorrected pairwise

genetic distances between samples of the O. alboguttatus species group, with gaps removed with a pairwise deletion option.

#### RESULTS

Both phylogenetic inferences yielded identical topologies, with some well-supported major clades and lower supported relationships among them (Fig. 1). Most of these major clades represent the species groups proposed for Osteocephalus, corroborating the monophyletic nature of the O. alboguttatus, O. leprieurii, O. taurinus, and O. planiceps species groups. Only the O. buckleyi species group was recovered as paraphyletic. The tree topology retrieved the O. alboguttatus species group as a highly supported clade, in both BI and ML analyses. Our results also show distinct lineages attributed to O. subtilis. Two poorly supported subclades were recovered: one containing O. heyeri and the upland O. subtilis from Serra do Divisor, state of Acre, Brazil, and other containing *O. alboguttatus*, the new species, and lowland O. subtilis from its type locality (Cruzeiro do Sul, state of Acre, Brazil). The new species is recovered as sister to *O. alboguttatus* with low (BI) to moderate support (ML) (Fig. 1). The lowest interspecific pairwise genetic distance within the *O. alboguttatus* species group  $(1.5\%)$ were obtained between O. heyeri from Leticia in Colombian Amazonia and lowland O. subtilis (Table 1). This result indicates that they can even be conspecific, and their identities should be carefully reviewed and properly assessed. Regarding the new species, genetic distances range from 2.7% (compared with lowland  $O$ . *subtilis*) to 5.1% (compared with upland O. subtilis). The latter comparison also represented the highest genetic distance between lineages within the  $O.$  alboguttatus species group included in our analysis. In light of this



Figure 1. Bayesian (BI) phylogenetic tree of *Osteocephalus* and closely related genera, inferred from 547-bp fragment of the mitochondrial gene 16S. Results of a maximum likelihood (ML) analysis are included, and branch support values for both inferences are shown above and below branches (posterior probabilities for BI and ultrafast bootstraps for ML, respectively). Support values below 0.8 (BI) and 70 (ML) were omitted. The new species described in this study is highlighted in bold.



TABLE 1. Uncorrected pairwise genetic distances between Osteocephalus melanops and the most closely related taxa, from the Osteocephalus alboguttatus species group. Distances are calculated from a fragment of the mitochondrial gene 16S. Values are presented in percentages (%). The new species is highlighted in bold.

molecular distinctiveness and evidence of morphological diagnosis, we describe below the new species of Osteocephalus.

# Taxonomic account Osteocephalus melanops, sp. nov. urn:lsid:zoobank.org:act:26656D84-0E89- 42D4-AEE1-0E2546444BD4 Figures 2–5

Holotype. MNRJ 93639 (field no. PRMS 350, Figs. 2–5), an adult male collected on 23 January 2016 in the Extractive Reserve Arapixi, municipality of Boca do Acre, state of Amazonas, Brazil (08°58′29″S, 67°51′38″W; 135 m above sea level [a.s.l.]) by Paulo R. Melo-Sampaio and Josimar C. da Silva.

Paratopotype. MNRJ 93640 (field no. PRMS 377, Figs. 2–4), an adult female collected on 25 January 2016 by Paulo R. Melo-Sampaio and Josimar C. da Silva.

Paratype. UFAC-RB 5553 (Fig. 4), an adult female collected on 26 November 2011 in Ramal do Cassiriã, in front of Colônia Matão, municipality of Sena Madureira, state of Acre, Brazil (09°09′05″S, 68°32′35″W; 166 m a.s.l.), by Paulo R. Melo-Sampaio, Júnior M. L. Maciel, and Evandro Silva.

Etymology. The specific epithet melanops is derived from Greek ''melanos-'' meaning black and ''ops-'' meaning eye. The name is a reference to intense dark coloration of the species' irises.

Generic Placement. The new species is assigned to the genus Osteocephalus on the basis of molecular phylogenetic relationships and morphological similarities with other species of the genus.

Diagnosis. A small-sized species of Osteocephalus characterized by: 1) SVL of adults 32.1–44.1 mm; 2) skin texture of dorsum non–sexually dimorphic, smooth with a few scattered small tubercles; 3) skin texture of flanks shagreened; 4) canthus rostralis rounded; 5) frontoparietal ridges not externally visible; 6) supratympanic fold thin, from the posterior edge of the eye, sloping in an arch toward the arm insertion, not reaching tympanum posteroventrally; 7) webbing on inner edge of third finger slightly extending beyond penultimate subarticular tubercle; 8) distal subarticular tubercle on finger IV bifid; 9) dorsum light tan with irregular blotches; 10) throat, chest, and abdomen uniformly cream to white; 11) large cream subocular spot; 12) flanks creamy white, with variable amount of dark vermiculation; 13) vocal sac single, subgular; 14) tibiofibular bones green in preservative; 15) in life, iris dark tan brown, with variable amount of lighter vermiculation.

Description of Holotype. Adult male with SVL of 32.1 mm. Head wider than body and slightly wider than long  $(HW/HL = 1.06)$ . Head slightly concave dorsally. Canthus



Figure 2. Right hand of the male holotype MNRJ 93639 (A) and female paratopotype MNRJ 93640 (B) of Osteocephalus melanops. Scale bars  $=$  5 mm.

rostralis straight in dorsal view, slightly elevated. Loreal region concave. Snout protruding in lateral view and truncate in dorsal view. Dentigerous processes of vomers at level of choanae, separated from each other, straight, with 16 vomerine teeth (eight in each side). Supratympanic fold present, running from the eye-tympanum area to above the arm-body insertion. Tympanum conspicuous, rounded, with diameter about 72% of the eye diameter. Axillary membrane present but poorly developed. Dorsum with scattered flat tubercles, laterally smooth, shagreened anteriorly to insertion of the arm. Throat finely granular; chest shagreened; belly granular. Posterior surface of thighs finely granular. Vent opening with a very small anal flap, located at upper level of thighs. Subcloacal and proximal thigh area covered with several small, flat tubercles.

Arms granular with a longitudinal row of small tubercles throughout the ventrolateral edge of the forearm and reaching the fourth finger. Dark nuptial excrescences covering the prepollex. Subpalmar tubercle large and elliptical. A bifid palmar tubercle and four rounded outer metacarpal tubercles. Subarticular tubercles single and rounded, except the distal tubercle on finger IV (bifid).



Figure 3. Left foot of the male holotype MNRJ 93639 (A) and female paratopotype MNRJ 93640 (B) of Osteocephalus melanops. Scale bars  $=$  5 mm.

Relative length of fingers  $3 > 4 > 2 > 1$ . Finger webbing formula: I absent II  $2-3$ <sup>+</sup> III  $3^{\text{-}}-2^{1/2}$  IV (Fig. 2). Hind limbs smooth except on proximal half of the ventral surface of the thigh (granular) and a few inconspicuous tubercles on the posterior tarsus. A large elliptical inner metatarsal tubercle and a small rounded outer one. Subarticular tubercles present, single, and conical. Toe webbing formula:  $I_1^+$ -2<sup>-</sup> II  $1^+$ -1 III  $2^{-}-2^{-}$  IV  $2^{-}-1^{1/3}$  V; relative length of adpressed toes:  $4 > 5 > 3 > 2 > 1$  (Fig. 3).

Morphometric Measurements of Holotype. EN, 3.6; ED, 3.9; FL, 21.0; HL, 11.2; HW, 11.9; IOD, 3.1; SVL, 32.1; TD, 2.8; TL, 17.6; UEW, 4.1.

Color of Holotype. In life, dorsum light brown with large cream blotches posteriorly; a well-defined dark brown interorbital transversal bar (Fig. 4) and small dark brown spots in the sacral region; dorsal surfaces of forearms light brown with dark brown marks, dorsal surfaces of thighs and shanks light brown with dark brown transversal bands. Sides of head light brown with a wide



Figure 4. Color in life of Osteocephalus melanops. (A, B) Dorsal view and close-up of the iris of the male holotype MNRJ 93639; (C) lateral view of the female paratopotype MNRJ 93640; (D) lateral view of the female paratype UFAC-RB 5553. Photographs: Paulo R. Melo-Sampaio.



Figure 5. Ventral view of species from Osteocephalus alboguttatus species group. (A) Osteocephalus melanops holotype MNRJ 93639. (B) Osteocephalus alboguttatus, QCAZ 15972. (C) Osteocephalus heyeri, unvouchered. (D) Osteocephalus subtilis, unvouchered. Photographs: Paulo R. Melo-Sampaio (A); Santiago Ron—www.bioweb.bio (B); Germán Chávez (C, D).

subocular cream spot from midposteroventral border of orbit to periphery of tympanum and border of jaw. Flank cream, areolate, with darker thin reticulation (Fig. 5). Iris dark tan brown with golden vermiculation in upper half and silvery vermiculation in lower half (Fig. 4).

In preservative, dorsal and lateral surfaces light brown, posteriorly with cream blotches. Tympanum light tan brown bordered by a dark brown ring. Wide cream subocular spot, extending posteriorly to the middle border of tympanum. The well-defined brown interorbital bar is evident. Dark brown transversal bar absent in the urostyle. Ventral surface creamy white, ventrolateral surfaces of the body white. Melanophores are evident around the lower lip, fewer on the chin. Forelimb tan laterally and dorsally, with a line of creamy white tubercles ventrally. Posterior surfaces of thighs light tan to orange. Cloacal region tan. Cloacal tubercles creamy white. Green bones, visible through the skin of tibia.

Variation. The female paratopotype has a darker iris and a more uniform cream color at dorsal surface of finger and toe discs (Fig. 4). The snout of the paratopotype is also more rounded than the holotype in lateral view. Morphometric measurements of the paratopotype and paratype females are as follow. MNRJ 93640: EN, 4.4; ED, 4.5; FL, 27.2; HL, 12.6; HW, 14.4; IOD, 3.8; SVL, 40.6; TD, 2.8; TL, 21.6; UEW, 4.7. UFAC-RB 5553: EN, 4.9; ED, 4.7; FL, 27.5; HL, 14.4; HW, 15.2; IOD, 3.9; SVL, 44.1; TD, 3.0; TL, 23.5; UEW, 4.8.

Comparisons. Morphological characters of the previously described species are presented in parenthesis unless otherwise stated. Osteocephalus melanops can be distinguished from members of the O. taurinus species group by having a smaller body size, with a maximum  $SVL = 44.1$  mm (minimum  $SVL =$ 64.2 mm in O. taurinus and 46.2 mm in O.

oophagus), a dark brown iris with black reticulation (iris greenish gold with bold dark brown regular radiation), frontoparietal ridges absent (present), keratinized tubercles absent in male dorsum (present) (Jungfer and Schiesari, 1995; Jungfer, 2010; Jungfer et al., 2013).

From members of the *O. leprieurii* species group, O. melanops can be distinguished by having males with single and subgular vocal sac (paired and distensible ventrolaterally to laterally in *O. leprieurii* and laterally in Osteocephalus yasuni Ron and Pramuk, 1999), keratinized tubercles absent in male dorsum (present), subdigital excrescences absent in male (present in O. leprieurii and O. yasuni), rounded canthus rostralis (distinct and angular in O. leprieurii and O. yasuni), green bones (white) and dark brown horizontal bar on iris absent (present in O. leprieurii and O. yasuni) (Ron and Pramuk, 1999; Jungfer and Hödl, 2002; Jungfer et al., 2013).

From members of the *O. planiceps* species group, O. melanops can be distinguished from Osteocephalus castaneicola Moravec, Aparicio, Guerrero-Reinhard, Calderón, Jungfer, and Gvoždík, 2009, O. planiceps, and *O. vilarsi* by having a maximum  $SVL =$ 44.1 mm (minimum SVL  $=$  47.7 mm in O. castaneicola, 48.0 mm in O. planiceps, and 47.5 mm in O. vilarsi), a dark tan brown iris (bronze with black lines radiating from the pupil), and a smooth dorsum (conspicuously tuberculated in adult males of O. vilarsi and O. planiceps), and frontoparietals not visible through skin (frontoparietals ridges well developed in O. vilarsi and O. planiceps). From species of this group with similar body sizes (Osteocephalus deridens Jungfer, Ron, Seipp, and Almendáriz, 2000, Osteocephalus fuscifacies Jungfer, Ron, Seipp, and Almendáriz, 2000, Osteocephalus leoniae Jungfer and Lehr, 2001), O. melanops can be distinguished by a smaller



Figure 6. Dorsal view of species from Osteocephalus alboguttatus species group. (A, B) Osteocephalus alboguttatus, unvouchered specimen and QCAZ 15972, respectively. (C, D) Osteocephalus heyeri, SINCHI 0727 and CZPB 1625, respectively. (E–I) Osteocephalus subtilis, unvouchered specimen (D), SINCHI 00485 (E), MTR 28225 (F), MTR 28100 (G), and CORBIDI 6176 (H), respectively; Osteocephalus cf. subtilis, an unvouchered specimen (I). Photographs: Morley Read—www.bioweb.bio (A); Santiago Ron—www.bioweb.bio (B); Germán Chávez (C, E); Alexandre Almeida (D); Pedro Peloso (F, G); Pablo Venegas (H); Jhon Jairo López Rojas (I).

female SVL, up to 44.1 mm (minimum SVL  $=$  53.2 mm in females of *O. fuscifacies* and O. leoniae), and by having a dark tan brown iris (bicolored or golden yellow) (Jungfer et al., 2000; Jungfer and Lehr, 2001; Cha´vez et al., 2008; Moravec et al., 2009; Ferrão et al., 2019).

From members of the *O. buckleyi* species group, O. melanops can be distinguished by having a single vocal sac (paired vocal sac), keratinized tubercles absent in male dorsum (present), posterior surface of thigh immaculate (with large dark brown spots), and canthus rostralis smooth (strongly tuberculate) (Ron et al., 2012; Jungfer et al., 2013, 2016). Osteocephalus melanops also has bifid subarticular tubercle on finger IV (barely bifid or single in Osteocephalus duellmani Jungfer, 2011 and single in Osteocephalus sangay Chasiluisa, Caminer, Varela-Jaramillo, and Ron, 2020) (Jungfer, 2011; Chasiluisa et al., 2020).

From the closest related species of the O. alboguttatus species group (Figs. 5, 6), O. melanops can be distinguished by having a tan brown dorsum with cream blotches (light brown dorsum with small blackish dots in O. alboguttatus, brown dorsum with transversal bars in O. subtilis), a large cream subocular spot (subocular spot dirty tan in O. heyeri), a uniform whitish-cream venter (venter whitish with brown spots in  $O.$  alboguttatus, cream with dark brown reticulation in *O. heyeri*), flanks and hidden dorsal surfaces of thighs cream with darker reticulation, without



Figure 7. Geographical distribution of the *Osteocephalus alboguttatus* species group. Data compiled from literature (summarized by Frost, 2021) and museum specimens (GBIF online database; GBIF.org, 2021).

small rounded white spots (cream with numerous small rounded white spots in O. alboguttatus, brownish and usually with black-edged white spots in O. heyeri, uniformly bluish in *O. subtilis*), thin supratympanic folds (well-developed in O. subtilis), a reticulated iris (uniformly black iris in O. subtilis), and a smaller body size, with maximum  $SVL = 32.1$  mm in male (minimum  $SVL = 35$  mm in O. *subtilis* males) (Boulenger, 1882; Martins and Cardoso, 1987; Lynch, 2002).

Geographic Distribution and Habitat. Osteocephalus melanops is only known from the type locality and a second nearby site, both at the east bank of the Purus River, southwestern Brazilian Amazonia (Fig. 7). In this region, *O. melanops* is sympatric with other four species of Osteocephalus: O. castaneicola, Osteocephalus helenae (Ruthven, 1919), O. leprieurii, and O. taurinus (Fig. 8). Individuals of the new species were found from 0.5 to 2 m height on the vegetation of a bamboo-dominated forest both in terra firme (nonflooded) and várzea (seasonally flooded) habitats always close to small streams. Type locality is a site with low anthropogenic disturbance, since it is a conservation unit for sustainable use, where activities such as hunting and plant extractivism have little effect on frog fauna, and small clearing and controlled burning for subsistence plantations play minor roles.

### DISCUSSION

Twenty-nine species of Osteocephalus are currently recognized (AmphibiaWeb, 2021;



Figure 8. Sympatric species of Osteocephalus at Extractive Reserve Arapixi, municipality of Boca do Acre, state of Amazonas, Brazil. (A) Osteocephalus leprieurii (field no. PRMS 0312). (B) Osteocephalus taurinus (field no. PRMS 0155). (C) Osteocephalus helenae (field no. PRMS 0261). (D) Osteocephalus castaneicola (field no. PRMS 0082).

present study, but see Frost, 2021), of which nine are known to occur in southwestern Brazilian Amazonia: O. buckleyi, O. cabrerai, O. castaneicola, O. deridens, O. leprieurii, O. melanops, O. planiceps, O. subtilis, and O. taurinus (Souza, 2009; Jungfer et al., 2013; Ferrão et al., 2019; Melo-Sampaio et al., 2021a; this study). Nevertheless, the description of O. melanops and the existence of distinct lineages of Osteocephalus in this region (Jungfer et al., 2013; Vacher et al., 2020; present study) demonstrates that the diversity of the genus is still underestimated. Although Jungfer et al. (2013) stated that the discovery of candidate new species within the O. alboguttatus species group was unlikely, the description of O. melanops and the existence of two highly distinct lineages associated with *O. subtilis* evidence that the

knowledge on real diversity of this species group is still incipient.

Increasing the resolution of Osteocephalus diversity also depends on the constant review of historically published data (e.g., Ferrão et al., 2019; Ortiz et al., 2020). While conducting this research, we came across with a mistake published in the original description of O. heyeri (Lynch, 2002) that, despite having already been briefly corrected by Jungfer (2010), deserves to be highlighted again to prevent future misinterpretations. On the basis of external morphology of specimens depicted in Lynch (2002, fig. 1), the lower right photograph clearly depicts the *O. cabrerai* specimen (ICN jdl22733), and not an O. heyeri as stated (Jungfer, 2010). Such a mistake raises doubts about the actual voucher correspondence for the

other three depicted specimens of O. heyeri. However, we suggest that a simple inversion occurred between two photographs (upper left vs. lower right), and the specimen depicted on the upper left should correspond to the O. heyeri female holotype (ICN 46916), and the remaining two specimens should be correctly referenced. Moreover, on the basis of morphological affinities such as uniform dark iris and flanks lacking welldefined spots, we hypothesize that the specimen depicted in the upper right of this figure most likely represents an O. subtilis instead of an O. heyeri. The recent records of O. subtilis occurring close to the known distribution of O. heyeri in the Javari River basin (López and Ortega, 2013; Chávez et al., 2016) reinforce that  $O$ . heyeri and  $O$ . subtilis must be sympatric at least in part of their distributions (see Fig. 7), and errors in their taxonomic attributions may be occurring more widely.

In fact, misidentifications appear to be even more pervasive in the taxonomic history of the *O. alboguttatus* species group. We analyzed the preserved paratype of O. heyeri (KU 220885), and on the basis of external morphology, we also suggest that it should not be conspecific with this species because of the presence of conspicuous dark blotches on its dorsum. This hypothesis is supported by the red-striped iris this specimen had in life, according to the photograph depicted in Duellman (2019) (in which the voucher number is incorrectly referenced as KU 220285). Such a characteristic is not known for members of the O. alboguttatus species group, and we suggest that specimen KU 220885 should be in fact a member of the O. planiceps species group. Therefore, all these arguments point to a composite type series of *O. heveri* and that this may also be occurring for other members of the O. alboguttatus species group, but such hypotheses still need to be tested after broader analyses of their type material.

New hypotheses and increased resolution on systematics and taxonomy of the O. alboguttatus species group are also largely hampered by sampling biases. Methods usually applied in anuran inventories are known to underestimate the diversity of species with natural low abundances and low-frequency calls, which seems to be the case for members of this species group (Cardoso and Vielliard, 1990). Another hampering factor is the pervasive misidentification led by phenotypic conservatism among species in this group (Jungfer et al., 2013; this study), especially when collection and posterior examination of specimens are not feasible (López and Ortega, 2013). Additionally, the lack of diagnostic characteristics in preserved specimens, especially the coloration patterns, also prevents the unveiling of the species diversity in this group. Therefore, we highlight that the documentation of coloration in life is crucial for the advancement of the taxonomic knowledge related to this species group.

The description of *O. melanops* expands the distribution of the  $O.$  alboguttatus species group to the Purus-Madeira Interfluve, where recent samplings also have unveiled several new species of amphibians (e.g., Melo-Sampaio et al., 2013, 2018; Ferrão et al., 2016, 2017, 2018a, b, 2020a, b; Lima et al., 2020). The increasing number of species descriptions in this interfluve is a result of two main factors: the crescent (but still low) representativeness of Amazon-based taxonomists, and the increase of inventories in poorly sampled areas (Ferrão et al., 2016; Melo-Sampaio et al., 2020, 2021b). Alarmingly, conservation of this highly biodiverse region is severely threatened by habitat loss led by illegal deforestation and wildfires (Barni et al., 2015; Brando et al., 2020; Silveira et al., 2020; Souza et al., 2020; MeloSampaio et al., 2021c). Although several protected areas exist in southwestern Amazonia (e.g., Arapixi Extractive Reserve and National Forest of Iquiri), the increase in small-patch deforestation of protected areas in this region limits the effective conservation of its biodiversity (Kalamandeen et al., 2018). A number of conservation strategies are necessary to improve the effectiveness of protected areas and support the delimitation of new protected areas in this region. We highlight the need for a massive investment in training new taxonomists and funding taxonomic studies in poorly sampled areas to accelerate both the documentation and description of Amazonian biodiversity.

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## Appendix 1. Examined specimens. Symbols:  $* =$ photograph.

Osteocephalus castaneicola  $(n = 37)$ . BRAZIL. ACRE: Feijó (UFAC-RB 4437, 4443, 4493); Mâncio Lima (UFAC-RB 1662, 1864, 1966); Manoel Urbano (UFAC-RB 4517, 4518); Marechal Thaumaturgo (UFAC-RB 2197, 2198, 2468, 2652, 2654); Sena Madureira (UFAC-RB 4706, 4726); Senador Guiomard (MNRJ 91672, UFAC-RB 4586, 4587, 4588, 4589 4590, 4591, 5197, 5652, 5653, 5753); Porto Acre (UFAC-RB 5756, 5757). AMAZONAS: Boca do Acre (UFAC-RB 4149, PRMS 0082, 0087, 0101, 0122, 0358). PERU: LORETO: Contamana (UFAC-RB 3598, 3599, 3605). Osteocephalus helenae  $(n = 33)$ . BRAZIL: ACRE: Senador Guiomard (UFAC-RB 4709, 4710, 4743– 4750, 5748). AMAZONAS: Boca do Acre (PRMS 0105, 0165–0184, 0261, 0308). Osteocephalus heyeri (n  $=$  1). BRAZIL: AMAZONAS: Rio Japurá (CZPB 1625). Osteocephalus leprieurii  $(n = 9)$ . BRAZIL: ACRE: Bujari (UFAC-RB 5730–5734). AMAZONAS: Boca do Acre (PRMS 0143, 0158, 0312, 0355). Osteocephalus oophagus ( $n = 5$ ). BRAZIL. AMAZONAS: Manaus, Reserva Ducke (INPA-H 36007, 36902, 37088, 37095, 37168). Osteocephalus planiceps  $(n = 28)$ . ECUADOR. NAPO: Puerto Misahaulli at La Cruz Blanca (MCZ-A 111188, 111190), Limoncocha (MCZ-A 98000, 98019, QCAZ 63543), Laguna Taracoa (MCZ-A 97755). ORELLANA: Yasuni National Park (QCAZ 14844\*, 20797\*, 40987\*, 51085\*, 55257\*, 55297–55298\*, 55364\*, 55378\*, 55380\*, 64119–64121\*, 64125– 64131\*), Napo River (QCAZ 43891\*, 44420\*). Osteocephalus subtilis  $(n=10)$ . BRAZIL: ACRE: Cruzeiro do Sul (MZUSP 60561, holotype); Mâncio Lima (UFAC-RB 1457–1463, 2741). PERU: UCAYALI: Ojo de Contaya (UFAC-RB 3565). Osteocephalus gr. alboguttatus ( $n = 4$ ). BRAZIL: AMAZONAS: Ipixuna, rio Juruá (INPA-H 2681, 2688, 2690, 2826). Osteocephalus taurinus  $(n = 11)$ . BRAZIL. ACRE: Sena Madureira (UFAC-RB 4728, 5559, 5735, 5736); Rio Branco (UFAC-RB 5752). AMAZONAS: Boca do Acre (PRMS 0155, 0236, 238); Careiro Castanho (INPA-H 25874, 25875, 25881). Osteocephalus vilarsi  $(n = 19)$ . BRAZIL. AMAZONAS: São Gabriel da Cachoeira, Missão Taracuá (INPA-H 40458, 40461,40470, GNM 488\* [holotype]); Novo Airão, Rio Negro Sustainable Development Reserve (INPA-H 40452, 40454, 40456, 40459, 40460, 40462, 40465, 40468), Jau National Park ´ (INPA-H 40455, 40463, 40464, 40467, 40466, 40472, 40473). Osteocephalus yasuni ( $n = 4$ ). BRAZIL. ACRE: Manuel Urbano (UFAC-RB 4519). ECUADOR. NA-PO: Yasuní Scientific Research Station (QCAZ 11336\* [holotype], QCAZ 10879\* [paratopotype], QCAZ 11329\* [paratopotype]).

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Appendix 2. Samples included in molecular analyses, with respective data on voucher/field numbers, localities of occurrence and GenBank accession numbers.

		Voucher/		<b>GenBank</b>
		Field		accession
<b>Species</b>	<b>Species Group</b>	number	Locality	number
Osteocephalus castaneicola	O. planiceps	NMP6d412009	Bolivia, Pando, Palmira	FJ965297
Osteocephalus castaneicola	O. planiceps	NMP6V72173	Bolivia, Pando, Nacebe	FJ965299
Osteocephalus deridens	O. planiceps	<b>OCAZ20868</b>	Ecuador, Yasuni, Orellana	JQ868501
Osteocephalus deridens	O. planiceps	KHJF006	Peru, Loreto, 28 km S Iquitos	KF002037
Osteocephalus fuscifacies	O. planiceps	<b>SMNS14194</b>	Ecuador, Napo, Jatun Sacha	KF002038
Osteocephalus leoniae	O. planiceps	KHJFTUN002	Peru, Amazonas, Santa María de Nieva	KF002056
Osteocephalus leoniae	O. planiceps	<b>KHJF112</b>	Peru. San Martín. 14 km NE Tarapoto	KF002058
Osteocephalus planiceps	O. planiceps	GGU752	Peru, Ucayali, Imiria	KF002091
Osteocephalus planiceps	O. planiceps	NMPCZE6V74913	Peru, Ucayali, Imiria	KY211978
Osteocephalus vilarsi	O. planiceps	<b>AMNHA1312546</b>	Venezuela, Amazonas, Neblina Base Camp on Rio Mawarinuma (=Rio Baria)	AY549361
Osteocephalus vilarsi	O. planiceps	MG96	Brazil, Amazonas, Missao Taracua	MK572700
Osteocephalus oophagus	$O.$ taurinus	MSH10225	Brazil, Amazonas, E. E. Anavilhanas	KF002081
Osteocephalus oophagus	$O.$ taurinus	<b>INPA-H37088</b>	Brazil, Amazonas, Reserva Ducke	MH357684
Osteocephalus taurinus	O. taurinus	APL17872	Brazil, Amazonas, Manaus, Conjunto Petros	KF002112
Osteocephalus taurinus	$O.$ taurinus	<b>INPA-H37098</b>	Brazil, Amazonas, Reserva Ducke	MH357687
Dryaderces inframaculata	outgroup	<b>INPA-H41311</b>	Brazil, Pará, middle Tapajós River	MT633109
Dryaderces pearsoni	outgroup	<b>SMNS14187</b>	Bolivia, Beni, Rurrenabaque	KF002006
Dryaderces sp.	outgroup	MTR13173	Brazil, Amazonas, Areal, Rio Abacaxis	KF002010
Tepuihyla warreni	outgroup	ROM39491	Guyana, District 7, Mt. Ayanganna	KF002185
Itapotihyla langsdorffii	outgroup	<b>MACN38643</b>	Argentina, Misiones, General Belgrano, 10 km N Bernardo de Irigoyen, Salto Andresito	AY843706
Phyllomedusa bicolor	outgroup	AJC5130	Colombia, Amazonas, Amazonas, Leticia, Reserva Natural Park	MG030713
Phyllomedusa sauvagii	outgroup	<b>MACN40002</b>	Argentina, Salta, Oran, Pichanal, Ruta Prov. 5 y Rio San Francisco	GQ366282

Appendix 2. Continued.