

Taxonomic Paper

A new toad species of the genus *Brachytarsophrys*Tian & Hu, 1983 (Anura, Megophryidae) from Guizhou Province, China

Shize Li[‡], Jing Liu[‡], Guiping Yang[§], Gang Wei[§], Haijun Su^l

‡ Moutai Institute, Renhuai, China§ Guiyang College, Guiyang, China| Guizhou University, Guiyang, China

Corresponding author: Gang Wei (wg198553@126.com), Haijun Su (hjsu@gzu.edu.cn)

Academic editor: Truong Nguyen

Received: 31 Dec 2021 | Accepted: 17 Feb 2022 | Published: 18 Feb 2022

 $\hbox{Citation: Li S, Liu J, Yang G, Wei G, Su H (2022) A new toad species of the genus \textit{Brachytarsophrys} Tian \& Hu, \\$

1983 (Anura, Megophryidae) from Guizhou Province, China. Biodiversity Data Journal 10: e79984.

https://doi.org/10.3897/BDJ.10.e79984

ZooBank: urn:lsid:zoobank.org:pub:221B3424-B5CF-429A-87BB-D948A43E79B5

Abstract

Background

The toads of the genus *Brachytarsophrys* Tian & Hu, 1983 are distributed in southern China, Myanmar, Vietnam, Laos and northern Thailand. Seven species of the genus have been recognised, of which five of them are known from China so far.

New information

Brachytarsophrys qiannanensis sp. nov., a new species of the short-legged toad genus is here described from southern Guizhou Province, China. Diagnostic characters of the new species are illustrated and comparisons with its congeners are provided. Its validity is also affirmed by its distinct mitochondrial gene sequence divergence with all congeners and its monophyly recovered in the mitochondrial gene-based phylogenetic analyses.

Keywords

Megophryidae, new taxon, phylogenetic analysis, morphology

Introduction

The short-legged toad genus *Brachytarsophrys* Tian & Hu, 1983 occurs widely in southern China, Myanmar, Vietnam, Laos and northern Thailand (Frost 2022). The generic taxonomy of this group has been controversial for a long time. Some studies (Dubois 1987, Duellman 1993, Dubois and Ohler 1998, Xu 2005, Mahony et al. 2017) considered this group as a subgenus of *Megophrys* Kuhl & Van Hasselt, 1822, but most recent studies still retained it as a genus mainly based on its distinct morphology, special ecological traits and the independent phylogenetic position (e.g. Fei et al. 1990, Ye and Fei 1992, Zhao and Adler 1993, Rao and Yang 1997, Fei et al. 2009, Pyron and Wiens 2011, Fei and Ye 2016, Chen et al. 2017, Li et al. 2020, Luo et al. 2021, Lyu et al. 2021, Tapley et al. 2021, Frost 2022).

The genus *Brachytarsophrys* currently contains seven species and Li et al. (2020) suggested they are divided into two groups, namely the *Brachytarsophrys carinense* group and the *Brachytarsophrys feae* group. The *Brachytarsophrys carinense* group contains *B. carinens* (Boulenger, 1889) and *B. intermedia* (Smith, 1921) and the *Brachytarsophrys feae* group contains *B. feae* (Boulenger, 1887), *B. chuannanensis* Fei, Ye and Huang, 2001, *B. platyparietus* Rao and Yang, 1997, *B. popei* Zhao, Yang, Chen, Chen & Wang, 2014 and *B. orientalis* Li, Lyu, Wang & Wang, 2020. Some phylogenetic studies indicated that the species diversity of *Brachytarsophrys* has been underestimated (e.g. Li et al. 2020, Lyu et al. 2021).

During field surveys in Libo County, Qiannan Autonomous Prefecture, Guizhou Province, China in 2021, we collected a series of *Brachytarsophrys* toads. Molecular phylogenetic analyses and morphological comparisons supported it as an undescribed species of *Brachytarsophrys*. We describe it herein as a new species.

Materials and methods

One adult male and three adult females of the *Brachytasophrys* sp. were collected from Libo County (**LB**), Guizhou Province, China (see Suppl. material 1, Fig. 1). In the field, the toads were euthanised using isoflurane and the specimens were fixed in 75 % ethanol. Tissue samples were taken and preserved separately in 99 % ethanol prior to fixation. The specimens were deposited in Chengdu Institute of Biology (**CIB**), Chinese Academy of Sciences (**CAS**).

All adult specimens of the *Brachytasophrys* sp. were measured. The terminology and methods followed Fei and Ye (2016) and Li et al. (2020). Measurements were taken with a dial caliper to 0.1 mm. Sixteen morphometric characters of adult specimens were

measured: eye diameter (ED, distance from the anterior corner to the posterior corner of the eye); foot length (FL, distance from distal end of shank to the tip of Toe IV); head length (HDL, distance from the tip of the snout to the articulation of jaw); maximum head width (HDW, greatest width between the left and right articulations of jaw); internasal distance (IND, minimum distance between the inner margins of the external nares); interorbital distance (IOD, minimum distance between the inner edges of the upper eyelids); length of lower arm and hand (LAL, distance from the elbow to the distal end of the Finger IV); lower arm width (LW, maximum width of the lower arm); snout-vent length (SVL, distance from the tip of the snout to the posterior edge of the vent); snout length (SL, distance from the tip of the snout to the anterior corner of the eye); length of foot and tarsus (TFL, distance from the tibiotarsal articulation to the end of the Toe IV); thigh length (THL, distance from vent to knee); tibia length (TL, from the outer surface of the flexed knee to the heel); maximal tibia width (TW); maximal tympanum diameter (TYD); upper eyelid width (UEW, greatest width of the upper eyelid margins measured perpendicular to the anterior-posterior axis). The location of the web on the phalange articulation was designated as follows: - (distal part of phalange articulation); none (middle part of phalange articulation); + (proximal part of phalange articulation); ++ (lower part of phalange articulation) followed the protocol described by Savage (1975) and Li et al. (2020).

Sex was determined by secondary sexual characters, i.e. the presence of vocal sac and nuptial pads/spines in male (Fei and Ye 2016).

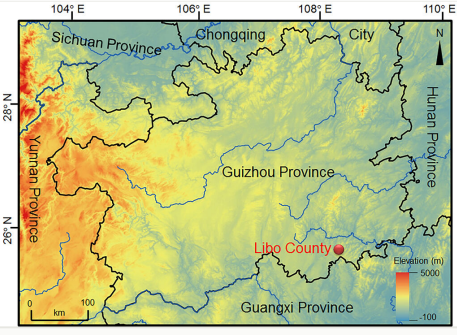


Figure 1. doi
Geographical location of the type locality of *Brachytarsophrys qiannanensis* sp. nov., Libo County, Guizhou Province, China.

The *Brachytasophrys* sp. was also compared with all other *Brachytarsophrys* species, based on morphological characteristics. Comparative morphological data were obtained from literature for *B. carinense* (Boulenger 1889, Li et al. 2020), *B. chuannanensis* (Fei and Ye 2001, Li et al. 2020), *B. feae* (Boulenger 1887, Li et al. 2020), *B. intermedia* (Smith 1921, Li et al. 2020), *B. orientalis* (Li et al. 2020), *B. platyparietus* (Rao and Yang 1997, Li et al. 2020) and *B. popei* (Zhao et al. 2014, Li et al. 2020).

The advertisement calls were recorded in the field on 6 August 2021 in Libo County, Qiannan Autonomous Prefecture, Guizhou Province, China. It was recorded in the stream at ambient air temperature of 18.0°C and air humidity of 80%. SONY PCM-D50 digital sound recorder was used to record within about 50 cm of the calling individual. The sound files in wave format were resampled at 48 kHz with sampling depth 24 bits. PRAAT 6.0.27 (Boersma 2001) was used to obtain the sonograms and waveforms (window length = 0.005s). Raven pro 1.5 software (Bioacoustics Research Program 2013) was used to quantify the acoustic properties (window size = 256 points, fast Fourier transform, Hanning window). Terminology of advertisement call analyses and description followed Köhler et al. (2017). Ambient temperature was taken by a digital hygrothermograph.

Four specimens of the Brachytasophrys sp. were included in the molecular analyses (Table 1). Total DNA was extracted using a standard phenol-chloroform extraction protocol (Sambrook et al. 1989). Three fragments of the mitochondrial 16S rRNA (16S), cytochromeoxidase subunit I (COI) and cytochrome b (Cytb) genes were amplified. For 16S, the primers P7 (5'-CGCCTGTTTACCAAAAACAT-3') and P8 (5'-CCGGTCTG AACTCAGATCACGT-3') were used following Simon et al. (1994); for COI, Chmf4 (5'-TYTCWACWAAYCAYAAAGAYATCGG-3') and Chmr4 (5'-ACYTCRGGRTGRCCRAAR AATCA-3') were used following Che et al. (2012) and for Cytb, PFGlu14140L (5'-GAAAAACCACTGTTGTHHYTCAACTA-3') and PFThr15310 (5'-CGGYTTA CAAGACCGRTGCTTT-3') were used following Zhang et al. (2013). Gene fragments were amplified under the following conditions: an initial denaturing step at 95 °C for 4 min; 36 cycles of denaturing at 95 °C for 30 s, annealing at 54 °C (for 16S)/49 °C (for COI)/50 °C (for Cytb) for 40 s and extending at 72 °C for 70 s. Sequencing was conducted using an ABI3730 automated DNA sequencer in Shanghai DNA BioTechnologies Co., Ltd. (Shanghai, China). New sequences were deposited in GenBank (for GenBank accession numbers see Table 1).

Table 1. Information for samples used in molecular phylogenetic analyses in this study (/ = not available).						
ID	Species	Localities	Voucher	GenBank accession number		
				16S	COI	Cytb
1	Brachytarsophrys qiannanensis sp. nov.	China: Libo County, Qiannan Autonomous Prefecture, Guizhou	CIB LB20210806053	OK104099	OK104052	OK127913

ID	Species	Localities	Voucher	GenBank accession number		
				16S	COI	Cytb
2	Brachytarsophrys qiannanensis sp. nov.	China: Libo County, Qiannan Autonomous Prefecture, Guizhou	CIB LB20210806054	OK104100	OK104053	OK127914
3	Brachytarsophrys qiannanensis sp. nov.	China: Libo County, Qiannan Autonomous Prefecture, Guizhou	CIB LB20210806055	OK104101	OK104054	OK127915
4	Brachytarsophrys qiannanensis sp. nov.	China: Libo County, Qiannan Autonomous Prefecture, Guizhou	CIB LB20210806056	OK104102	OK104055	OK127916
5	Brachytarsophrys orientalis	China: Jiulianshan Nature Reserve, Longnan County, Jiangxi	SYS a004225	1	MT162625	MT162650
6	Brachytarsophrys orientalis	China: Jiulianshan Nature Reserve, Longnan County, Jiangxi	SYS a004228	1	MT162628	MT162653
7	Brachytarsophrys orientalis	China: Jiulianshan Nature Reserve, Longnan County, Jiangxi	SYS a004226	1	MT162626	MT162651
8	Brachytarsophrys orientalis	China: Jiulianshan Nature Reserve, Longnan County, Jiangxi	SYS a004486	1	MT162629	MT162654
9	Brachytarsophrys orientalis	China: Jiulianshan Nature Reserve, Longnan County, Jiangxi	SYS a005451	1	MT162632	MT162655
10	Brachytarsophrys orientalis	China: Jiulianshan Nature Reserve, Longnan County, Jiangxi	SYS a004227	1	MT162627	MT162652
11	Brachytarsophrys orientalis	China: Huboliao Nature Reserve, Nanjing County, Fujian	SYS a003340	1	MT162624	MT162649
12	Brachytarsophrys orientalis	China: Gutian Township, Shanghang County, Fujian	SYS a003249	1	MT162623	MT162648
13	Brachytarsophrys popei	China: Taoyuandong Nature Reserve, Yanling County, Hunan	SYS a001864	KM504256	MH406361	MH407191

ID	Species	Localities	Voucher	GenBank accession number		
				16S	COI	Cytb
14	Brachytarsophrys popei	China: Taoyuandong Nature Reserve, Yanling County, Hunan	SYS a001865	KM504257	MT162620	MT162645
15	Brachytarsophrys popei	China: Taoyuandong Nature Reserve, Yanling County, Hunan	SYS a001866	KM504258	MT162621	MT162646
16	Brachytarsophrys popei	China: Jinggang Shan, Jiangxi	SYS a004209	MK524124	MK524155	1
17	Brachytarsophrys platyparietus	China: Duodihe, Dayao county, Yunnan	SYS a005919	1	MT162633	MT162656
18	Brachytarsophrys platyparietus	China: Mt. Jinzhong, Longlin County, Guangxi	SYS a002236	1	MT162622	MT162647
19	Brachytarsophrys platyparietus	China: Mt. Fanjing, Tongren City, Guizhou	YPX43968	1	MT162644	MT162667
20	Brachytarsophrys platyparietus	China: Mt. Mopan, Xinping County, Yunnan	SYS a007774	1	MT162634	MT162657
21	Brachytarsophrys platyparietus	China: Mt. Mopan, Xinping County, Yunnan	SYS a007775	1	MT162635	MT162658
22	Brachytarsophrys platyparietus	China: Mt. Mopan, Xinping County, Yunnan	SYS a007776	1	MT162636	MT162659
23	Brachytarsophrys platyparietus	China: Mt. Mopan, Xinping County, Yunnan	SYS a007777	1	MT162637	MT162660
24	Brachytarsophrys platyparietus	China: Yilong Township, Shiping County, Yunnan	SYS a007790	1	MT162638	MT162661
25	Brachytarsophrys platyparietus	China: Yumen Township, Yanbian County, Sichuan	SYS a007853	1	MT162639	MT162662
26	Brachytarsophrys feae	China: Jingdong County, Yunnan	SYS a003912	MH406899	MH406362	MH407192
27	Brachytarsophrys feae	China: Jingdong County, Yunnan	SYS a003913	1	MH406363	MH407193
28	Brachytarsophrys feae	China: Huangcaoling, Yunnan	KIZ046706	KX811810	KX812056	1
29	Brachytarsophrys chuannanensis	China: Zihuai Township, Hejiang County, Sichuan	SYS a004926	MH406901	MT162630	1

ID	Species	Localities	Voucher	GenBank accession number		
				16S	COI	Cytb
30	Brachytarsophrys carinense	Thailand: Doi Chiang Dao, Chiang Mai	K3001	KR827713	KR087626	1
31	Brachytarsophrys carinense	Thailand: Omkoi, Chiang Mai	KIZ024170	1	MT162640	MT162663
32	Brachytarsophrys carinense	Thailand: Mae Surin NP., Mae Hong Son	KIZ024429	1	MT162641	MT162664
33	Brachytarsophrys carinense	Thailand: Thong Pha Phum, Kanchanaburi	KIZ024640	1	MT162642	MT162665
34	Brachytarsophrys intermedia	Vietnam: Krong Pa, Gia Lai	ROM 23794	1	MT162643	MT162666
35	Atympanophrys shapingensis	1	CIBSC2011102004	JX458090	JX458090	JX458090
36	Panophrys omeimontis	China: Hongya County, Sichuan	MO-HY130601	KP728257	KP728257	KP728257

For molecular analyses, the available sequence data for Brachytarsophrys were downloaded from GenBank (Table 1), primarily from previous studies (Chen et al. 2017, Li et al. 2020). For phylogenetic analyses, corresponding sequences of one Atympanophrys shapingensis Liu 1950 and one Panophrys omeimontis Liu 1950 were also downloaded (Table 1) and used as outgroups according to Chen et al. (2017). Sequences were assembled and aligned using the Clustalw module in BioEdit v.7.0.9.0 (Hall 1999) with default settings. For phylogenetic analyses of mitochondrial DNA, the dataset was concatenated with 16S, COI and Cytb gene sequences. To avoid under- or overparameterisation (Lemmon and Moriarty 2004, McGuire et al. 2007), the best partition scheme and the best evolutionary model for each partition were chosen for the phylogenetic analyses using PARTITIONFINDER v. 1.1.1 (Robert et al. 2012). In this analysis, 16S gene and each codon position of protein-coding genes were defined and Bayesian Inference Criteria was used. As a result, the analysis suggested that the best partition scheme is 16S gene/each codon position of protein-coding genes and selected GTR + G + I model as the best model for each partition. Phylogenetic analyses were conducted using Maximum Likelihood (ML) and Bayesian Inference (BI) methods, implemented in PhyML v. 3.0 (Guindon et al. 2010) and MrBayes v. 3.12 (Ronquist and Huelsenbeck 2003), respectively. For the ML tree, branch supports were drawn from 10,000 non-parametric bootstrap replicates. In BI, two runs each with four Markov chains were simultaneously run for 50 million generations with sampling every 1,000 generations. The first 25% trees were removed as the "burn-in" stage followed by calculations of Bayesian posterior probabilities and the 50% majority-rule consensus of the post burn-in trees sampled at stationarity, bootstrap supports (BS) and Bayesian Posterior Probabilities (BPP) are shown at the nodes. Finally, mean genetic distance between samples in this

study, based on uncorrected p-distance model, was estimated using MEGA v. 6.06 (Tamura et al. 2013) with the pairwise deletion setting for the Gap/Missing Data.

Data resources

All the sequences in this study were retrieved from GenBank and the accession numbers of the newly-determined sequences in this study are shown in Table 1.

Taxon treatment

Brachytarsophrys qiannanensis Li, Liu, Yang, Wei, & Su, sp. n.

ZooBank 92662CAA-955E-41B2-B991-8905E6E65FFA

Materials

Holotype:

a. scientificName: Brachytarsophrys qiannanensis; kingdom: Animalia; phylum: Chordata; class: Amphibia; order: Anura; family: Megophryidae; genus: Brachytarsophrys; higherGeography: South-western China; country: China; stateProvince: Guizhou Province; county: Libo County; municipality: Qiannan Autonomous Prefecture; locality: Changniu Village; verbatimElevation: 1190; verbatimCoordinates: 25.572492°N, 108.274189°E; georeferenceSources: georeferenceSources; eventDate: 06/08/2021; individualID: CIB LB20210806054; individualCount: 1; sex: male; lifeStage: adult; recordedBy: Jing Liu; identifiedBy: Shize Li; type: Even

Paratype:

a. scientificName: Brachytarsophrys qiannanensis; kingdom: Animalia; phylum: Chordata; class: Amphibia; order: Anura; family: Megophryidae; subgenus: Brachytarsophrys; higherGeography: South-western China; country: China; stateProvince: Guizhou Province; county: Libo County; municipality: Qiannan Autonomous Prefecture; locality: Changniu Village; verbatimCoordinates: 25.572492°N, 108.274189°E; georeferenceSources: Google Earth; eventDate: 06/08/2021; individualID: CIB LB20210806055-57; individualCount: 3; sex: 3 females; lifeStage: adul; recordedBy: Jing Liu; identifiedBy: Shize Li; type: Even

Diagnosis

Morphometric measurements for specimens examined are given in Table 2 and Suppl. material 1. See Fig. 2A-E for dorsal and ventral view of body, dorsal and ventral view of hand and ventral view of foot.

Brachytarsophrys qiannanensis sp. nov. could be distinguished from its congeners by a combination of the following morphological characters: (1) body size small (SVL 70.1 mm in male and 80.1 – 84.9 mm in females); (2) tongue pyriform, feebly notched posteriorly; (3) tibiotarsal articulation reaching to commissure of jaw when leg stretched forward; (4) toes about one third to two thirds webbed in males; (5) male with a single

subgular vocal sac and a brown nuptial pad present on the dorsal surface of the first finger.

Table 2.

Measurements of the adult specimens of *Brachytarsophrys qiannanensis* sp. nov. Units are given in mm. See abbreviations for the morphological characters in Materials and Methods section.

Measurement	Male (n = 1)	Females (n = 3)		
		Range	Mean ± SD	
SVL	70.1	80.1 – 84.9	82.5 ± 2.4	
HDL	18.9	20.6 – 24.8	22.0 ± 2.4	
HDW	32.2	34.4 – 38.8	36.5 ± 2.2	
SL	8.1	9.0 – 10.3	9.6 ± 0.7	
IND	7.8	7.8 – 8.4	8.1 ± 0.3	
IOD	9.4	9.2 – 11.3	10.5 ± 1.1	
UEW	4.1	5.4 – 6.0	5.6 ± 0.3	
ED	5.9	6.2 – 7.2	6.7 ± 0.5	
TYD	4.1	3.0 – 4.1	3.7 ± 0.6	
LAL	29.5	32.8 – 34.2	33.5 ± 0.7	
LW	6.4	6.0 - 6.5	6.3 ± 0.2	
THL	29.6	28.5 – 34.8	31.7 ± 3.2	
TL	29.4	29.4 – 31.4	30.3 ± 1.0	
TW	10.3	9.7 – 10.6	10.2 ± 0.5	
TFL	41.9	42.7 – 44.8	43.6 ± 1.1	
FL	32.6	31.8 – 32.3	32.0 ± 0.3	

Description of holotype: An adult male, SVL 70.1 mm; head enormous, extremely depressed, about 1.7 times as broad as long; snout short, rounded in dorsal view, slightly protruding beyond margin of lower jaw; canthus rostralis indistinct; loreal region very oblique, slightly concave; nostril closing to the tip of snout; tympanum not obvious; eye large, eye diameter 31 % of head length; maxillary teeth present, vomerine teeth present on two vomerine ridges; tongue pyriform, notched posteriorly.

Fore-limbs short and moderately robust, the length of lower arm and hand 42 % of SVL; fingers rather short without web, relative finger lengths: I < II < V < III; tips of digits round, feebly dilated; lateral fringes absent; metacarpal tubercle two, inner one significantly enlarged, outer one slightly enlarged.

Hind-limbs relatively short and robust, heels not meeting when thighs are positioned at right angles to the body, tibiotarsal articulation reaching to commissure of jaw when leg stretched forward; tibia length longer than thigh length; relative toe lengths I < II < V < III < IV; tips of toes round, slightly dilated; toes about one third to two thirds webbed and lateral fringe wide, the webbing formula is $I(1) - (2^+) II(2^-) - (3^-) III(2^{1/2}) - (4^-) IV(4^{++}) - (2^-) V$; inner metatarsal tubercle oval-shaped; outer metatarsal tubercle absent.



Figure 2. doi

Photos of the holotype CIB LB20210806054 of *Brachytarsophrys qiannanensis* sp. nov. in life. **A** dorsal view; **B** ventral view; **C** dorsal view of hand (insert: the nuptial pad on the dorsal surface of the first finger); **D** ventral view of hand; **E** ventral view of foot.

Dorsal skin rough, several conical tubercles scattered on flank of trunk, dorsum of body and limbs; upper eyelid with several tubercles and one enlarged to form horn; tubercles on the dorsum forming a U-marking on the anterior dorsum; a dark brown streak on dorsum of head and between the eyes; supratympanic fold distinct, from posterior corner of eye to a level above the shoulder.

Ventral surface smooth; pectoral gland distinct, closer to axilla than to mid-ventral line; rear of thigh with a small femoral gland, around which densely arranged granules forming a granular patch.

Colouration of holotype in life: Dorsal brown, a dark brown streak on dorsum of head and between the eyes; dark tubercles present on the dorsum, forming a U-marking, some dark tubercles scattered on the shoulder and posterior dorsum, flank of body scattered with some light brown tubercles; upper lip light brown; tympanic region

brown; dorsal digits with dark brown transverse bands and three transverse skin ridges on the dorsal shank and thigh; ventral surface brown-black, pectoral gland yellowish; several yellowish spots on two sides of belly; lower surface of digits purple-grey; webs, palms and soles purple-grey; tip of digits, two metacarpal tubercle and inner metatarsal tubercle grey-white; nuptial pad brown; the tubercles at upper eyelid yellowish; pupils black; iris brownish.

Colouration of holotype in preservation: Colour of dorsal surface fades to pale brown; ventral surface brown; the posterior of ventral surface of body; tip of digits, two metacarpal tubercles and inner metatarsal tubercle grey-white fades to white (Fig. 3).

Variation: Measurements of the type series are shown in Suppl. material 1. In this new species, the females had larger bodies than male (Table 2). In life, the diagnostic morphological characters of all paratypes were identical to those of the holotype (Fig. 4). However, colouration and stripe patterns differed amongst individuals. In CIB LB20210806055, the brown patches on dorsum are irregular (Fig. 4A), in the ventral surface, some dark tubercles on the throat and the tubercles on the thigh are white (Fig. 4B); in CIB LB20210806056, the brown patches forming an X-marking on the dorsum (Fig. 4C) and the colouration of ventral surface is lighter (Fig. 4D).

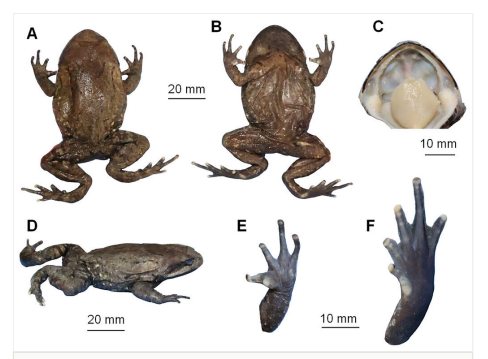


Figure 3. doi

The holotype specimen CIB LB20210806054 of *Brachytarsophrys qiannanensis* sp. nov. in preservative. **A** dorsal view; **B** ventral view; **C** view of oral cavity; **D** lateral view; **E** ventral view of hand; **F** ventral view of foot.

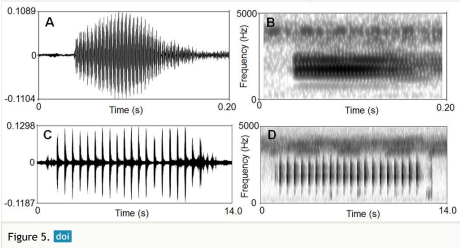


Figure 4. doi

Colour variation *Brachytarsophrys qiannanensis* sp. nov. in life **A** dorsal view of the female specimen CIB LB20210806055; **B** ventral view of the female specimen CIB LB20210806055; **C** dorsal view of the specimen the female specimen CIB LB20210806055; **D** ventral view of the female specimen CIB LB20210806056.

Secondary sexual characters: The adult male with a single subgular vocal sac and brown nuptial pad present on the dorsal surface of the first finger (Fig. 2C).

Advertisement call: The call description is based on recordings of the holotype CIB LB20210806054 (Fig. 5), calling from beneath a large stone in a streamlet and the ambient air temperature was 18.0° C. Each call consists of 16-20 (mean $17.3.5\pm2.3$, n = 3) notes. Call duration was 7690-11330 ms (mean 9068 ± 1974 , n = 3). Call interval was 10980-15670 ms (mean 13325 ± 3316 , n = 2). Each note had a duration of 129-348 ms (mean 249 ± 36 , n = 52) and the intervals between notes 180-395 ms (mean 269 ± 42 , n = 49). Amplitude modulation within the note was apparent, beginning with moderately high energy pulses, increasing slightly to a maximum by approximately mid-note and then decreasing towards the end of each note. The average dominant frequency was 1740 ± 168 Hz (1640-2330 Hz, n = 3).



Visualisation of advertisement calls of *Brachytarsophrys qiannanensis* sp. nov. **A** waveform showing one note; **B** sonogram showing one note; **C** waveform showing 20 notes of one call; **D** sonogram showing 20 notes of one call (A and B are the same note, C and D are the same call).

Comparisons: Comparative data of *Brachytarsophrys qiannanensis* sp. nov. with other seven recognised congeners of *Brachytarsophrys* are given in Suppl. material 2

Brachytarsophrys qiannanensis sp. nov. differs from *B. orientalis* by having a smaller body size , SVL 70.1 mm in male and 80.1-84.9 mm in females (vs. 76.8-82.7 mm in males, n = 7 and 88.6 mm in female); different webbing formula I (1) - (2⁺) II (2-) -(3-) III (2½) - (4-) IV (4⁺⁺) - (2-) V in male and I (1½) - (2⁺) II (2-) - (3⁺) III (2½) - (4) IV (4⁺⁺) - (2) V in female (vs. I (1½) - (2) II (1½) - (3) III (2½) - (4) IV (4) - (2) V in male and I (2) - (2⁺) II (1⅓) - (3) III (3-) - (4) IV (4⁺) - (2½) V in female); the male specimen with a brown nuptial pad present on the dorsal surface of the first finger (vs. dark brown nuptial pads present on the dorsal surface of the first two fingers)(Li et al. 2020).

Brachytarsophrys qiannanensis sp. nov. differs from *B. popei* by having tongue feebly notched posteriorly (vs. deeply notched behind); nuptial pad of the male without spines on the dorsal surface of the first finger (vs. with black nuptial spines present on the dorsal bases of the first two fingers); webbing formula I (1) - (2⁺) II (2⁻) - (3⁻) III (2½) - (4-) IV (4⁺⁺) - (2-) V in male and I (1½) - (2⁺) II (2-) - (3⁺) III (2½) - (4) IV (4⁺⁺) - (2) V in female (vs. I (1½) - (2) II (1½) - (3) III (2½) - (3½) IV (3¾) - (2) V in male and I (1½) - (2⁺) II (1½) - (3) III (2½) - (4-) IV (4-) - (2) V in female) (Zhao et al. 2014, Li et al. 2020, Suppl. material 2).

Brachytarsophrys qiannanensis sp. nov. differs from B. platyparietus by having a smaller size SVL 70.1 mm in male and 80.1-84.9 mm in three females (vs. 88.5-13.0 mm in males, n=6 and 118.5-131.0 mm in female, n=3); lateral fringes on toes narrow (vs. wide); brown nuptial pad without nuptial spines present on the dorsal surface of the first finger (vs. dark brown nuptial pads with black nuptial spines present

on the dorsal bases of the first two fingers); webbing formula I (1) - (2^+) II (2^-) - (3^-) III (2^+) - (4^-) IV (4^{++}) - (2^-) V in male and I (1^+ 2) - (2^+) II (2^-) - (3^+) III (2^+ 2) - (4^- 2) V in female (vs. I (1^+ 2)-(2^+) II (1^+ 2)-(3^+ 3) III (2^+ 3)-(3^+ 3) IV(3^+ 3)-(2^- 7) V in male and I (1^+ 2)-(2^+ 1) II (1^+ 2)-(3^+ 3) III (2^+ 3)-(4^- 7) IV (4^- 3)-(2^+ 3) V) (Rao and Yang 1997, Li et al. 2020, Suppl. material 2).

Brachytarsophrys qiannanensis sp. nov. differs from *B. feae* by having a smaller body size in male, SVL 70.1 mm (vs. 78.5-94.9 mm in males, n=5); tibiotarsal articulation reaching to commissure of jaw when leg stretched forward both in male and females (vs. reaching to shoulder in females); brown nuptial pad present on the dorsal surface of the first finger (vs. dark brown nuptial pads present on the dorsal bases of the first two fingers); webbing formula I (1) - (2⁺) II (2-) - (3-) III (2½) - (4-) IV (4⁺⁺) - (2-) V in male (VS. I (2)-(2⁺⁺) II (2-)-(3)III (2¾)-(4) IV (4)-(2¾) V) (Boulenger 1887, Li et al. 2020, Suppl. material 2).

Brachytarsophrys qiannanensis sp. nov. further differs from *B. feae* by having more notes of each call (16-20 notes vs. 4-5 notes); a longer call duration 7690–11330 ms (vs. 2256–35488 ms) and a higher dominant frequency 1640 – 2330 Hz (vs. 1378 Hz) (Wogan et al. 2004).

Brachytarsophrys qiannanensis sp. nov. differs from B. chuannanensis by having a smaller body size SVL 70.1 mm in male (vs. 91.4 - 109.4 mm in males, n = 12); the male with brown nuptial pad on dorsal surface of the first finger (vs. dark brown nuptial pads present on the dorsal bases of the first two fingers); lateral fringes on toes of male wider (one third as broad as distal toe phalanx vs. one fifth)(Fei and Ye 2001, Li et al. 2020, Suppl. material 2).

Brachytarsophrys qiannanensis sp. nov. from *B. carinense* by having a smaller body size SVL 70.1 mm in male and 80.1-84.9 mm in females (vs. 91.6-123.0 mm in males, n= 4 and 124.0-168.0 mm, n= 3); by the absence of dermal ridges on dorsum (vs. present); tibiotarsal articulation reaching to commissure of jaw when leg stretched forward (vs. reaching to axilla in females and angle of mouth in males); webbing formula I (1) - (2⁺) II (2-) - (3-) III (2½) - (4-) IV (4⁺⁺) - (2-) V in male (VS. I (1½)-(2⁺⁺) II (2)-(3⁺⁺) III (3)-(4) IV(4⁺⁺)-(2½) V) (Boulenger 1889, Li et al. 2020, Suppl. material 2).

Brachytarsophrys qiannanensis sp. nov. differs from *B. intermedia* by having a smaller body size (SVL 70.1 mm in male and 80.1 - 84.9 mm in females vs. 86.0 - 103.0 mm in males, n = 7 and 92.0 mm in female) and the absence of glandular folds on dorsum (vs. present) (Smith 1921, Li et al. 2020, Suppl. material 2).

Etymology

The specific name *qiannanensis* refers to the distribution of this species, Qiannan Autonomous Prefecture, the County to where the type locality of the species belongs. We propose the common English name "Qiannan short-legged toad" and Chinese name "Qian Nan Duan Tui Chan (黔南短腿蟾)".

Distribution

Brachytarsophrys qiannanensis sp. nov. is known from the type locality, Libo County, Guizhou Province, China at elevations between 1100 – 1200 m a.s.l.

Ecology

Brachytarsophrys qiannanensis sp. nov. inhabits a mountain stream (Fig. 6) covered by evergreen broadleaf forest, there being only a small amount of water on the surface of the stream. Advertisement call of males can be heard from beneath the rocks at night and the females were frequently found near large rocks.

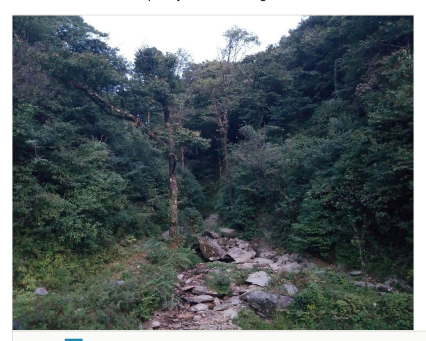


Figure 6. doi
Habitat of *Brachytarsophrys qiannanensis* sp. nov. in the type locality, Libo County, Guizhou Province, China.

Analysis

Phylogenetic analyses

Aligned sequence matrix of 16S + COI + Cytb contains 2061 bp. ML and BI trees of the mitochondrial DNA dataset presented almost consistent topology (Fig. 7). *Brachytarsophrys qiannanensis* sp. nov. was clustered into the *Brachytarsophrys* clade and grouped with *B. orientalis* and *B. popei* with high supported values (node supports in ML and BI: 90 and 1.00).

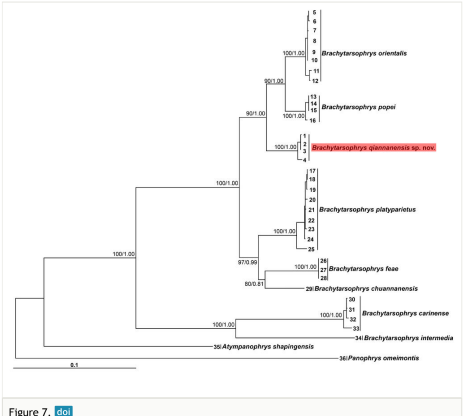


Figure 7. doi

Phylogenetic tree reconstructed using Bayesian Inference (BI) and Maximum Likelihood (ML) methods, based on 16S, COI and Cytb genes. Values at the nodes correspond to BPP and BS. Samples 1-36 refer to Table 1.

The mean genetic distances (p-distance) between Brachytarsophrys qiannanensis sp. nov. and its congeners was 2.1% on 16S (with B. popei), 4.9% on COI (with B. orientalis) and 6.6% on Cytb (with B. orientalis), these distances being much higher than those between some pairs of recognised congeners (Suppl. materials 3, 4, 5); for example, the COI pdistance was 4.3% between B. orientalis and B. popei, while the Cytb p-distance between B. orientalis and P. popei was 3.3%.

Molecular phylogenetic analyses showed that the population of Brachytarsophrys from Libo County, Qiannan Autonomous Prefecture, Guizhou Province, China is also distinct from its congeners.

Discussion

The previous morphological studies indicated that, in the genus Brachytarsophrys, only B. carinense was recorded from Leishan and Anlong Counties of Guizhou Province (e.g. Wu et al. 1986, Fei et al. 2009, Fei et al. 2012). Lyu et al. (2014) further identified the population from Fanjing Mountain, Guizhou Province as B. chuannanensis, based on morphological comparisons. Based on molecular phyogenetic analyses and morphological comparisons, Li et al. (2020) suggested that B. platyparietus should be a valid species and populations in south-western China previously recognised as B. carinense should be reidentified as B. platyparietus, including the population from Fanjing Mountain, Guizhou Province. Moreover, Li et al. (2020) suggested that the populations from the western Hunan Province and north-western part of Guangxi Province, all adjoining Guizhou Province, belonged to B. popei and B. orientalis, respectively. Accordingly, it is inferred that the population of Brachytarsophrys from Anlong County in western Guizhou Province, near Yunnan Province, may be B. platyparietus and that from Leigong Mountain in southeastern Guizhou Province, near the Hunan Province, may be B. popei or B. orientalis. In addition, the taxonomic assignments of the specimens from Fanjing Mountain, Guizhou Province are still doubtful because Li et al. (2020) reported that the specimen from Fanjing Mountain unexpectedly shared a common holotype on the COI gene with several specimens from the northern and central parts of Yunnan Province which is quite far from Fanjing Mountain in the eastern Guizhou Province. It is a pity that Li et al. (2020) did not provide more morphological and bioacoustics information of this specimen for comparisons with other species. Obviously, the taxonomic profiles of Brachytarsophrys in Guizhou Province are still unresolved and further investigations on this group should be conducted in the region.

Brachytarsophrys qiannanensis sp. nov. seems to be the smallest species (SVL 70.1 mm in male and SVL < 85 mm in females) in the genus *Brachytarsophrys*. Whether its niche characteristics promote the special morphology in this species maybe an interesting evolutionary question.

In recent years, more than 20 new amphibian species have been discovered in Guizhou Province, China (Frost 2022). However, during our frequent and extensive surveys in Guizhou Province from 2016 to 2021, we only found one adult male and three adult females of *Brachytarsophrys qiannanensis* sp. nov. This perhaps indicated that the population of the species in Guizhou Province is potentially small. Hence, further surveys are needed to evaluate the population status of the species.

Acknowledgements

We are grateful to editors and reviewers for their working on the manuscript. This work was supported by National Natural Sciences Foundation of China (NSFC31960099 and 32060307), Basic research project of science and technology department of Guizhou Province (Nos. [2020] 1Y083), Science and technology support project of science and technology department of Guizhou Provincial (No. [2020] 4Y029) and Forestry Science and Technology Research Project of Guizhou Forestry Department (No. [2020]13).

Author contributions

Shize Li and Gang Wei conceived and designed the experiments, performed the experiments, analysed the data, contributed reagents/materials/analysis tools, preparedfigures and/or tables, authored or reviewed drafts of the paper, approved the final draft; Jing Liu and Haijun Su performed the experiments, analysed the data, approved the final draft; Guiping Yang performed the experiments, analysed the data and approved the final draft.

References

- Bioacoustics Research Program (2013) Raven Pro: Interactive sound analysis software.
 Version1.5. http://www.birds.cornell.edu/raven
- Boersma P (2001) PRAAT, a system for doing phonetics by computer. Glot International 5: 341-345.
- Boulenger G (1887) Description of a new frog of the genus Megalophrys. Annali del Museo Civico di Storia Naturale di Genova 4: 512-513.
- Boulenger G (1889) Description of a new batrachian of the genus *Leptobrachium*, obtained by M. L. Fea in the Karens Mountains, Burma. Annali del Museo Civico di Storia Naturale di Genova 7: 748-750.
- Che J, Chen H, Yang J, Jin J, Jiang K, Yuan Z, Murphy R, Zhang Y (2012) Universal COI primers for DNA barcoding amphibians. Molecular Ecology Resource 12: 247-258. https://doi.org/10.1111/j.1755-0998.2011.03090.x
- Chen J, Zhou W, Nikolay A, Poyarkov J, Stuart B, Brown R, Lathrop A, Wang Y, Yuan Z, Jiang K, Hou M, Chen H, Suwannapoom C, Nguyen S, Duong T, Papenfuss T, Murphy R, Zhang Y, Che J (2017) A novel multilocus phylogenetic estimation reveals unrecognized diversity in Asia toads, genus *Megophrys* sensu lato (Anura: Megophryidae). Molecular Phylogenetics and Evolution 106: 28-43. https://doi.org/10.1016/j.ympev.2016.09.004
- Dubois A (1987) Miscellanea taxinomica batrachologica. Alytes 5: 7-95.
- Dubois A, Ohler A (1998) A new species of Leptobrachium (Vibrissaphora) from northern Vietnam, with a review of the taxonomy of the genus *Leptobrachium* (Pelobatidae, Megophyinae). Dumerilia 4 (14): 1-32.
- Duellman W (1993) Amphibian species of the world. Additions and corrections.
 University of Kansas, Museum of Natural History 21: 1-372.
- Fei L, Ye C, Huang Y (1990) Key to Chinese amphibians. Publishing House for Scientific and Technological, Chongqing, 344 pp.
- Fei L, Ye C (2001) The Color Handbook of the Amphibians of Sichuan. Chinese Forestry Press, Beijing, 138 pp.
- Fei L, Hu S, Ye C, Huang Y (2009) Fauna Sinica. Amphibia. Volume 2. Anura. Science Press, Beijing, 345 pp.
- Fei L, Ye C, Jiang J (2012) Colored atlas of Chinese amphibians and their distributions. Sichuan Publishing House of Science and Technology, Chengdu, 20 pp.
- Fei L, Ye C (2016) Amphibians of China. 1. Science Press, Beijing, 717-727 pp.

- Frost D (2022) Amphibian species of the world: An online reference. Version 6.1.
 Electronic database. American Museum of Natural History, New York. http://research.amnh.org/herpetology/amphibia/index.html. Accessed on: 2022-2-13.
- Guindon S, Dufayard J, Lefort V, Anisimova M, Hordijk W, Gascuel O (2010) New
 algorithms and methods to estimate maximum-likelihood phylogenies: assessing the
 performance of PhyML 3.0. Systematic Biology 59 (3): 307-321. https://doi.org/10.1093/sysbio/syg010
- Hall T (1999) BIOEDIT: a user-friendly biological sequence alignment editor and analysis program for Windows 95/98/NT. Nucleic Acids Symposium 41: 95-98.
- Köhler J, Jansen M, Rodríguez A, Kok P, Toledo LF, Emmrich M, Glaw F, Haddad CFB, Rödel MO, Vences M (2017) The use of bioacoustics in anuran taxonomy: theory, terminology, methods and recommendations for best practice. Zootaxa 4251 (1): 1-124.
- Lemmon A, Moriarty E (2004) The importance of proper model assumption in Bayesian phylogenetics. Systematic Biology 53 (2): 265-277. https://doi.org/10.1080/10635 150490423520
- Liu C (1950) Amphibians of western China. Fieldiana. Zoology Memoires 2: 1-397.
- Li Y, Zhang D, Lyu Z, Wang J, Li Y, Liu Z, Chen H, Rao D, Jin Z, Zhang C, Wang Y (2020) Review of the genus *Brachytarsophrys* (Anura: Megophryidae), with revalidation of *Brachytarsophrys platyparietus* and description of a new species from China. Zoological Research 41 (2): 105-122. https://doi.org/10.24272/j.issn.2095-8137.2020.033
- Luo T, Wang Y, Wang S, Lu X, Wang W, Deng H, Zhou J (2021) A species of the genus Panophrys (Anura, Megophryidae) from southeastern Guizhou Province, China.
 ZooKeys 1047: 27-60. https://doi.org/10.3897/zookeys.1047.61097
- Lyu J, Niu K, Li S, Li C, Wei G, Xu N (2014) Brachytarsophry chuannanensis found in Fanjingshan National Nature Reserve, Guizhou Province. Chinese Journal of Zoology 49 (3): 432-434.
- Lyu Z, Zeng Z, Wang J, Liu Z, Huang Y, Li W, Wang Y (2021) Four new species of Panophrys (Anura, Megophryidae) from eastern China, with discussion on the recognition of Panophrys as a distinct genus. Zootaxa 4927 (1): 9-40. https://doi.org/10.11646/zootaxa.4927.1.2
- Mahony S, Nicole M, Biju S, Teeling E (2017) Evolutionary history of the Asian horned frogs (Megophryinae): integrative approaches to timetree dating in the absence of a fossil record. Molecular Phylogenetics and Evolution 34 (3): 744-771.
- McGuire J, Witt C, Altshuler D, Remsen J (2007) Phylogenetic systematics and biogeography of hummingbirds: Bayesian and maximum likelihood analyses of partitioned data and selection of an appropriate partitioning strategy. Systematic Biology 56 (5): 837-856. https://doi.org/10.1080/10635150701656360
- Pyron R, Wiens J (2011) A large-scale phylogeny of Amphibia including over 2800 species, and a revised classification of advanced frogs, salamanders, and caecilians. Molecular Phylogenetics and Evolution 61: 543-583. https://doi.org/10.1016/j.ympev.2011.06.012
- Rao D, Yang D (1997) The karyotypes of Megophryinae (Pelobatidae) with a discussion on their classification and phylogenetic relationships. Asian Herpetological Research 7: 93-102. https://doi.org/10.5962/bhl.part.18858

- Robert L, Brett C, Simon Y, Stephane G (2012) PartitionFinder: Combined selection of partitioning schemes and substitution models for phylogenetic analyses. Molecular Phylogenetics and Evolution 29 (6): 1695-1701.
- Ronquist F, Huelsenbeck J (2003) MrBayes3: Bayesian phylogenetic inference under mixed models. Bioinformatics 19 (12): 1572-1574. https://doi.org/10.1093/bioinformatics/btq180
- Sambrook J, Fritsch E, Maniatis T (1989) Molecular cloning: A laboratory manual. 2nd.
 Cold Spring Harbor Laboratory Press, New York.
- Savage J (1975) Systematics and distribution of the Mexican and Central American stream frogs related to *Eleutherodactylus* rugulosus. Copeia 2: 254-306. https://doi.org/10.2307/1442883
- Simon C, Frati F, Beckenbach A, Crespi B, Liu H, Flook P (1994) Evolution, weighting
 and phylogenetic utility of mitochondrial gene sequences and a compilation of
 conserved polymerase chain reaction primers. Annals of the Entomological Society of
 America 87 (6): 651-701. https://doi.org/10.1093/aesa/87.6.651
- Smith M (1921) New or little-known reptiles and batrachians from southern Annam (Indo-China). Proceedings of the Zoological Society of London423-440. https://doi.org/10.1111/j.1096-3642.1921.tb03271.x
- Tamura K, Stecher G, Peterson D, Fiipski A, Kumar S (2013) MEGA6: molecular evolutionary genetics analysis, version 6.0. Molecular Biology and Evolution 30: 2725-2729. https://doi.org/10.1093/molbev/mst197
- Tapley B, Cutajar T, Nguyen L, Portway C, Mahony S, Nguyen C, Harding L, Luong H, Rowley J (2021) A new potentially endangered species of *Megophrys* (Amphibia: Megophryidae) from Mount Ky Quan San, north-west Vietnam. Journal of Natural History 54 (39): 2543-2575. https://doi.org/10.1080/00222933.2020.1856952
- Wogan GOU, Lwin KS, Win H, Thin T, Shein AK, Tun H (2004) The advertisement call
 of *Brachytarsophrys feae* (Boulenger,1897) (Anura:Megophryidae). Proceedings of the
 California Academy of Sciences 55: 251-254.
- Wu L, Dong Q, Xu R (1986) Amphibians of Guizhou province. Guizhou People Press, Guiyang, 33-35 pp.
- Xu J (2005) Study on some taxonomic problems of Megophryidae. Sichuan Journal of Zoology 24: 337-339.
- Ye C, Fei L (1992) Taxonomic studies of pelobatid toads, genus Brachytarsophrys of China. Acta Herpetologica Sinica56-62.
- Zhang P, Liang D, Mao R, Hillis D, Wake D, Cannatella D (2013) Efficient sequencing of anuran mtDNAs and a mitogenomic exploration of the phylogeny and evolution of frogs. Molecular Biology and Evolution 30: 1899-1915. https://doi.org/10.1093/molbev/mst091
- Zhao E, Adler K (1993) Herpetology of China. In: Contributions to Herpetology 10 (Ed.) Society for the Study of Amphibians and Reptiles. 116 pp.
- Zhao J, Yang J, Chen G, Chen C, Wang Y (2014) Description of a new species of the genus *Brachytarsophrys* Tian and Hu, 1983 (Amphibia: Anura: Megophryidae) from southern China based on molecular and morphological data. Asian Herpetological Research 5: 150-160. https://doi.org/10.3724/SP.J.1245.2014.00150

Supplementary materials

Suppl. material 1: Measurements of the adult specimens of *Brachytarsophrys* qiannanensis sp. nov. doi

Authors: Shize Li, Jing Liu, Guiping Yang, Gang Wei, HaiJun Su

Data type: morphological

Brief description: Measurements of the adult specimens of *Brachytarsophrys qiannanensis* sp. nov. Units are in mm. See abbreviations for the morphological characters in Materials and

Methods section.

Download file (10.99 kb)

Suppl. material 2: Diagnostic characters separating the new species from other species of *Brachytarsophrys* doi

Authors: Shize Li, Jing Liu, Guiping Yang, Gang Wei, Haijun Su

Data type: morphological

Brief description: Diagnostic characters separating the new species described in this study from

other species of Brachytarsophrys (/ = not available).

Download file (11.48 kb)

Suppl. material 3: Mean uncorrected genetic p-distance of the 16S gene between samples doi

Authors: Shize Li, Jing Liu, Guiping Yang, Gang Wei, Hainjun S

Data type: genomic

Brief description: Mean uncorrected genetic p-distance of the 16S gene between samples

examined in this study.

<u>Download file</u> (29.50 kb)

Suppl. material 4: Mean uncorrected genetic p-distance of the COI gene between samples doi

Authors: Shize Li, Jing Liu, Guiping Yang, Gang Wei, HaiJun Su

Data type: genomic

Brief description: Mean uncorrected genetic p-distance of the COI gene between samples

examined in this study.

Download file (27.00 kb)

Suppl. material 5: Mean uncorrected genetic p-distance of the Cytb gene between samples doi

Authors: Shize Li, Jing Liu, Guiping Yang, Gang Wei, HaiJun Su

Data type: genomic

Brief description: Mean uncorrected genetic p-distance of the Cytb gene between samples

examined in this study.

Download file (27.00 kb)