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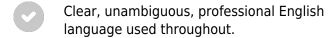
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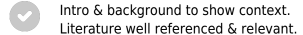
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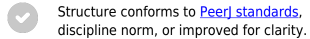
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A new species of freshwater crab (Crustacea: Brachyura: Pseudothelphusidae) from a naturally isolated orographic forest enclave within the semiarid Caatinga in Ceará, northeastern Brazil

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A new species of freshwater crab, Fredius ibiapaba, is described and illustrated from a midaltitude forested patch in Ipú (Ibiapaba plateau, Ceará, northeastern Brazil), between 635 to 782 m. The new species can be separated from its congeners by the morphology of its first gonopod: proximal half remarkably swollen, sloping abruptly downwards anteriorly to a nearly right-angular shoulder; mesial lobe much smaller than cephalic spine; cephalic lobe moderately developed; auxiliary lobe lip, delimiting field of apical spines, protruded all the way to distal margin of auxiliary lobe. Comparative 16S rDNA sequencing used to infer the phylogenetic placement of Fredius ibiapaba n. sp. revealed that it is the sister taxon of F. reflexifrons, a species which occurs allopatrically in the Amazon and Atlantic basin's lowlands (< 100 m). Fredius ibiapaba n. sp. and F. reflexifrons are highly dependent upon humidity and most probably were once part of an ancestral population living in a wide humid territory. Shrinking humid forests during several dry periods of the Tertiary and Quaternary likely have resulted in the fragmentation of the ancestral humid area and hence of the ancestral crab population. Fredius reflexifrons evolved and spread in a lowland, humid river basin (Amazon and Atlantic basins), whilst F. ibiapaba n. sp. evolved isolated on the top of a humid plateau. The two species are now separated by a vast intervening area occupied by the semiarid Caatinga.

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2 3 4 5	naturally isolated orographic forest enclave within the semiarid Caatinga in Ceará, northeastern Brazil
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29	
30	Abstract



63

31 A new species of freshwater crab, Fredius ibiapaba, is described and illustrated from a mid-32 altitude forested patch in Ipú (Ibiapaba plateau, Ceará, northeastern Brazil), between 635 to 782 33 m. The new species can be separated from its congeners by the morphology of its first 34 gonopod: proximal half remarkably swollen, sloping abruptly downwards anteriorly to a nearly 35 right-angular shoulder; mesial lobe much smaller than cephalic spine; cephalic lobe moderately 36 developed; auxiliary lobe lip, delimiting field of apical spines, protruded all the way to distal 37 margin of auxiliary lobe. Comparative 16S rDNA sequencing used to infer the phylogenetic 38 placement of Fredius ibiapaba n. sp. revealed that it is the sister taxon of F. reflexifrons, a 39 species which occurs allopatrically in the Amazon and Atlantic basin's lowlands (< 100 m). 40 Fredius ibiapaba n. sp. and F. reflexifrons are highly dependent upon humidity and most 41 probably were once part of an ancestral population living in a wide humid territory. Shrinking 42 humid forests during several dry periods of the Tertiary and Quaternary likely have resulted in 43 the fragmentation of the ancestral humid area and hence of the ancestral crab population. 44 Fredius reflexifrons evolved and spread in a lowland, humid river basin (Amazon and Atlantic 45 basins), whilst F. ibiapaba n. sp. evolved isolated on the top of a humid plateau. The two 46 species are now separated by a vast intervening area occupied by the semiarid Caatinga. 47 Introduction 48 49 Cumulative evidences from many independent sources argue in favor of the mid-altitude 50 forested patches in northeastern Brazil being remnants of a once much larger humid forest, 51 connected to both the Amazonian and Atlantic rainforests during the moister periods (e.g., 52 Andrade-Lima, 1982; Cartelle & Hartwig, 1996; de Vivo, 1997; Ab'Saber, 2000; Auler et al., 53 2004; Carnaval & Bates, 2007; Carmignotto, 2012; and references therein). These humid forest 54 refuges (Figure 1A–D), naturally isolated by the vast surrounding semiarid Caatinga (Figure 1F, 55 G), are indeed known to harbor many woody plant and animal species (fossil and Recent) 56 that are also found or are closely related to species occurring allopatrically in the Amazonian 57 and Atlantic rainforests. 58 Here we describe and illustrate a new species of a freshwater pseudothelphusid crab, 59 Fredius ibiapaba n. sp., from a mid-altitude forest enclave in Ipú (Ibiapaba plateau, Ceará, 60 northeastern Brazil), between 665 to 782 m (Figure 1A–D). Evidences from a phylogenetic 61 analysis using 16S rDNA are presented for a sister taxon relationship between Fredius ibiapaba

n. sp. and F. reflexifrons (Ortmann, 1897), a species occurring allopatrically in the Amazonian

humid lowlands. Previous hypothesis on the phylogenetic relationships of F. reflexifrons and the



possible evolutionary scenario that led to the emergence of the sister taxons *Fredius ibiapaba* n. sp. and *F. reflexifrons* are discussed.

Materials & Methods

Procedures with material examined

The specimens were collected using license permission from the Sistema de Autorização e Informação em Biodiversidade (SISBIO #29615) of the Brazilian Ministry of Environment (MMA). The studied specimens are deposited in the collections of the INPA (Instituto Nacional de Pesquisas da Amazônia, Manaus) and MZUSP (Museu de Zoologia, Universidade de São Paulo, Brazil). Measurements: cl, carapace length, taken along the carapace axis to its posterior median margin, in millimeters (mm). Dates are written in the format day.month.year, with months in lower-case Roman numerals. Abbreviations are as follows: G1, G2, first and second gonopods, respectively. Mxp3, third maxilliped. The terminal y used in the description of the G1 is referred in the figure 2.

Molecular data analysis

DNA extraction, amplification and sequencing: Muscle tissue samples were obtained from the pereopods or abdomen of *Fredius ibiapaba* n. sp., *F. buritizatilis* Magalhães, Sanches, Pileggi & Mantelatto, 2014, and *Prionothelphusa eliasi* Rodriguez, 1980. At the Laboratório de Biologia Molecular da Universidade Estadual do Sudoeste da Bahia- LBM/UESB a small region of the 16S rDNA gene was extracted with Wizard® Genomic DNA Purification Kit (Promega), amplified in a 12,5 µl final volume reaction with 2,5 mM de MgCl2 (Invitrogen), 0,05 mM de dNTP (Invitrogen), buffer 1x (Invitrogen – 10xPCR Buffer: 200mM Tris-HCl (pH 8.4), 500mM KCl), 1U de taq platinum (Invitrogen) and 0,3µM of each primer. The PCR conditions were: one cycle at 94°C, 60 sec; five cycles at 94°C, 60 sec; 45°C, 40 sec and 72°C, 60 sec; and 35 cycles at 94°C, 60 sec; 51°C, 40 sec and 72°C; 60 sec; a final extension of five minutes at 72°C



was performed. The primers used were 16Sar (5'-CCGGTCTGAACTCAGATCACGT-3') and 16Sbr (5'-CGCCTGTTTATCAAAAACAT-3') (Palumbi et al., 1991). PCR products were purified using a polietilenoglicol (PEG) 20% and sequenced in an ABI Prism 3100 Genetic Analyzer® (Applied Biosystems) at the Departamento de Tecnologia da Universidade Estadual Paulista "Júlio de Mesquita Filho", Jaboticabal. Sequencing reaction was performed with Big Dye v3.1 (Applied Biosystems), prepared with 4,75 µl ultrapure water, 1,5 µl BigDye 5x buffer, 0,75 µl BigDye terminator Mix, 2 µl primer (0,8 pmol) and 1 µl of Purified PCR product. Sequence conditions were: one minute at 96°C; 35 cycles of 15 sec at 96°C; 15 sec at 50°C and 2 minutes at 60°C. Both, forward and reverse sequence strands were obtained and the consensus generated by the software BioEdit 7.0.5 (Hall, 2005). The identities of the final sequences were confirmed with a BLAST (Basic Local Alignment Search Tool) on GenBank database. Additional comparative sequences were retrieved from GenBank (Table 1).

Phylogenetic analyses: Substitution saturation in 16S rDNA was tested using the saturation index implemented in DAMBE 5 (Xia, 2013). The sequences were grouped and edit in BioEdit and aligned using the ClustalW interface (Thompson et al. 1994). *Prionothelphusa eliasi* (Pseudothelphusidae) and *Trichodactylus dentatus* H. Milne Edwards, 1853 (Trichodactylidae) were chosen as outgroups. The best-fit model HKY + G was selected using jModeltest 2.1.7 (Darriba et al., 2012). This model was used to generate Maximum Likelihood gene trees in MEGA 6.06 (Tamura et al. 2013). Branch support values were calculated using bootstrap analyses with 1,000 replicates (Felsenstein, 1985). Only nodes with bootstrap support greater than 50 are shown on the phylogenetic tree. Nucleotide divergence estimated from pairwise distance was calculated in MEGA 6.06 with the same best-fit model (Table 2).

Registration of nomenclatural act

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116	and hence the new names contained in the electronic version are effectively published under
117	that Code from the electronic edition alone. This published work and the nomenclatural acts it
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119	ZooBank LSIDs (Life Science Identifiers) can be resolved and the associated information
120	viewed through any standard web browser by appending the LSID to the prefix
121	http://zoobank.org/. The LSID for this publication is: [urn:lsid:zoobank.org:pub:0925982D-7441-
122	120 4256-9856-A553987956A6]. The online version of this work is archived and available from
123	the following digital repositories: PeerJ, PubMed Central and CLOCKSS.
124 125 126	Results Pseudothelphusidae Ortmann, 1893
127	Fredius Pretzmann, 1967
128	Fredius ibiapaba n. sp. (Figures 3A-E; 4A-C; 5A, C; 6A-D; 7A-E)
129	
130	Type material. Holotype, male cl 36 mm, cw 53mm (MZUSP 39710), Sítio Caranguejo, Ipú,
131	Ceará, 04°18'50" S, 40°44'47"W, 729 m, xii.2017. Paratypes: male cl 34 mm, cw 48 mm
132	(MZUSP 39169), same data as holotype; female cl 35 mm, cw 49 mm (MZUSP 39171), Sítio
133	Gameleira, Ipú, Ceará, 04°17'17" S, 40°44'44"W, 665 m, 5.i.2018; male cl 32 mm, cw 48 mm
134	(MZUSP 39167), Sítio Santa Cruz, Ipú, Ceará, 04°19'40" S, 40°45'09"W, 782 m, 10.x.2014;
135	female cl 31 mm, cw 44 mm (MZUSP 39168), Sítio Santa Cruz, Ipú, Ceará, 04°19'40" S,
136	40°45'09"W, 782 m, 23.iv.2015; male cl 41.2 mm, cw 62.6 mm (MZUSP 39742), Sítio Ipuçaba,
137	Ipú, Ceará, 27.x11.2017.
138	
139	Comparative material. Fredius fittkaui (Bott, 1967): male, cl 47.1 mm, cw 66.9 mm (MZUSP
140	24497), Guyana, Potaro-Siparuni, Rio Kuribrong, 05°22'35"N, 59°33'4"W, P. Bernardo & B.
141	Newman coll., 28.ix.2010. Fredius reflexifrons (Ortmann, 1897): Peru: male, cl 31 mm, cw 42.5



142	mm (MZUSP 6389), Rio Apiacu, DepartamentoLoreto, Boris Malkin coll., 15-25.iv.1966. Brazil:
143	male cl 37 mm, cw 52 m (MZUSP 19922), Amapá, Serra do Navio, Serra do Veado, Projeto
144	Diversitas Neotropica, M. Tavares coll. 7.v.1994. 2 males, cl 57.7 mm, cw 42 mm and cl 73.8
145	mm, cw 53 mm (MZUSP 13178), Amapá, Rio Jari, montante, Cachoeira Santo Antônio, M. Jegú
146	& J. Zuanon coll., 9-26.vi.1981. 1 macho (INPA 583), Amapá, Serra do Navio/ Serra do Veado,
147	07.v.1994. 1 macho (INPA 2125), Amapá, município de Laranjal, 16.i.2012. 1 macho (INPA
148	889), Amazonas, Manaus, Reserva do Km 41, 02°26'56"S, 59°46'13"W. 1 macho (INPA 368),
149	Amazonas, Manaus, Reserva Ducke, 22.ii.1986. 1 macho (INPA 850), Amazonas, Manaus,
150	11.vii.2001. 1 macho (INPA 852), Amazonas, Iranduba, Sítio Anaíra, 03°10'39"S, 60°07'39"W,
151	12.ix.1999. 1 macho (INPA 1254), Pará, Santarém, Comunidade Santa Rosa. 1 macho (INPA
152	851), Pará, Rio do Peixe Boi, 01°11'30"S, 47°18'54"W, E. Matos and A. Henriques Jr coll.,
153	03.iii.1995. 1 macho (INPA 1512), Pará, Bragança, Rio Chumucuí, S. Alves coll., 12.xi.2004.
154	Fredius denticulatus (H. Milne Edwards, 1853): male cl 45 mm, cw 62 mm (MZUSP 16294), Rio
155	Amapari, Serra do Navio, AP, Projeto Diversitas Neotropica, n°151, M. Tavares coll.,
156	30.iv.1994, C. Magalhães det. 16.ii.1996.
157	
158	Type locality. Sítio Caranguejo, Ipú, Ceará, 04°18'50"S, 40°44'47"W, 729 m.
159	Distribution. Only recorded from the type locality to date.
160	
161	Etymology. The specific name is a noum in apposition taken from the Tupi language word for
162	plateau, "yby'ababa", ibiapaba.
163	
164	Diagnosis. G1 robust, proximal half remarkably swollen, sloping abruptly downwards anteriorly
165	to a nearly right-angular shoulder; mesial lobe much smaller than cephalic spine; cephalic lobe
166	moderately developed; auxiliary lobe lip, delimiting field of apical spines, protruded all the way to
167	distal margin of auxiliary lobe.

Description of the holotype. Carapace transversally ovate (Figure 3A), widest at middle (cw/cl, 1.51); dorsal surface smooth, slightly convex, regions ill-defined. Gastric pits minute, very close to each other. Cervical grooves shallow, nearly straight, poorly indicated, distal ends reaching to anterolateral margin. Front deflexed, almost straight in dorsal view, entire, marked with row of very small papillae; front lower border carinate, with an almost indistinct sinus medially in frontal view; postfrontal lobules obsolete; median groove between postfrontal lobules faint. Upper orbital margin with row of very faint papillae; lower margin minutely denticulate; exorbital angle marked by obtuse tooth, followed posteriorly by faint notch. Ce pace anterolateral margin semicircular in outline, fringed by minute denticles; posterolateral margins almost straight, strongly convergent, smooth. Epistomial margin with minute papillae; epistomial tooth broadly triangular, deflexed (Figure 3C). Suborbital and subhepatic regions of carapace smooth; pterygostomial region densely pubescent around mouthparts (Figures 3B, C).

Mxp3 palp slender, long, reaching slightly beyond articulation of merus and ischium when folded. Merus markedly operculiform. Posterior half of mesial margin of merus and mesial margin of ischium with conical teeth (Figure 3B). Exopod short, 0.28 times length of lateral margin of ischium, devoid of flagellum. Efferent branchial channel opening subcircular (Figure 3B).

Chelipeds moderately heterochelous, right cheliped larger than left one (Figure 3E, F). Major cheliped merus subtriangular in cross-section; lateral surface smooth, irregular row of small tubercles of different sizes along dorsal surface; with irregular longitudinal row of small tubercles; mesial surface smooth, slightly concave to fit lateral sides of carapace; mesial lower margin with row of conical teeth slightly increasing in size distally; lateral lower margin with row of small teeth. Carpus smooth dorsally; mesial margin with row of small, irregular teeth and strong, acute spine about midlength of margin. Palm moderately swollen, smooth on lateral and mesial sides, with minute papillae on dorsal and ventral rounded faces. Dactylus regenerated.





195	Fingers not gaping when closed, tips not crossing. Minor cheliped similar in shape.
196	Thoracic sternal suture 2/3 complete, distinct; sternal suture 3/4 interrupted, visible only
197	laterally (Figure 3B); sternal sutures 4/5and 5/6 interrupted, ending just before reaching midline
198	of thoracic sternum; sternal sutures 6/7 and 7/8 complete. Midline of thoracic sternum deeply
199	incised in sternites VII and VIII.
200	All abdominal segments free. Lateral margins of male telson slightly concave, tip rounded
201	(Figure 3B).
202	G1 postimal half remarkably swollen, sloping abruptly downwards anteriorly to a
203	nearly right-angular shoulder (Figure 4B, C). Subapical bulge moderately developed around
204	lateral and sternal sides (Figures 4B; 5A, B; 6A). Marginal suture straight (Figure 4C). Marginal
205	lobe truncate, projected distally beyond abdominal surface, junction marked by distinct
206	depression. Mesial lobe much smaller than cephalic spine, showing as triangular, acute spine,
207	pointing to abdominal direction (Figures 4A, C, 5B; C; 6A, C). Cephalic spine very strong,
208	acuminate at tip, pointing in mesial direction (Figures 4A–C, 5B; C; 6A–C). Cephalic lobe
209	prominent, truncate, tip rounded, with several very small spines along lateral, mesial and sternal
210	sides (Figures 4A, B; 6A, B). Auxiliary lobe much shorter than cephalic lobe in abdominal view,
211	separated from it by distinct depression, their junction forming lateral channel running distally in
212	almost straight direction before ending in inward curve subterminally (Figures 4A; 6A). Field of
213	apical spines large, open, flattened, elongated, ear-shaped, provided with small spinules,
214	delimited by lateral and abdominal lips of apex (Figures 4A, B; 5B; 6A, B).
215	G2 slightlinger than G1 when abdomen folded beneath cephalothorax; very slender,
216	tapering distally progressively, distal part moderately flattened, with rather dense, minute
217	spinules along sternal side.
218	

Cutting margin of dactylus and fixed finger both with larger teeth interspersed with smaller ones.



219 Remarks. Fredius ibiapaba n. sp. is herein assigned to the genus Fredius, whose diagnostic 220 characters (Rodriguez, 1982; Rodriguez & Pereira, 1992) are readily recognized in the new 221 species, namely, exopod of mxp3 short, about 0.3 times length of outer margin of ischiu E 31 222 widest at base (Figure 4B, C); marginal lobe simple, ending in an inverted cup-shaped 223 elongation at base of field of apical spines; subapical bulge covering lateral and sternal sides; 224 field of apical spines large, open, flattened, ear-shaped, with small scattered spinules at 225 proximal sternal border (Figures 4A–C; 5B; 6A, B). 226 The new species morphologically resembles *Fredius denticulatus*, *F. fittkaui*, *F.* 227 reflexifrons and F. ykaa in that the gonopod cephalic spine is much more developed than the mesial lobe (see Magalhães & Rodriguez, 2002: 679, fig. 1; 683, fig. 2, respectively; Rodriguez 228 229 & Campos, 1998: 766, fig. 2O, P) (Figure 4A, C; 5B; 6A, C), whereas other species either have 230 the gonopod cephalic spine little larger than the mesial lobe (*F. stenolobus* and *F. adpressus*), 231 or have it much shorter than the mesial lobe (e.g., F. buritizalis, F. platyacanthus, and F. estevi), 232 or have the cephalic spine and the mesial lobe similar in size (e.g., F. granulatus, F. chaffanjoni 233 (see Magalhães et al., 2014 and references therein). 234 Fredius ibiapaba n. sp. stands apart from Fredius denticulatus, F. fittkaui, F. reflexifrons 235 and F. ykaa in having the G1 proximal half remarkably swollen on the abdominal side, sloping 236 abruptly downwards anteriorly to a nearly right-angular shoulder (Figures 4B, C), whereas in the 237 latter four species the G1 shoulder is clearly more gently sloping distally (Figure 4E, F). 238 The closest morphological resemblance of Fredius ibiapaba n. sp. is with F. reflexifrons of 239 which it additionally differs (1) in having the auxiliary lobe lip, delimiting the field of apical spines, 240 protruded all the way to the distal margin of the auxiliary lobe (Figure 4A, B), whereas in F. 241 reflexifrons the lip fades away well before reaching the distal margin of the lobe (Figure 4D, E); 242 and (2) the subapical bulge markedly less swollen (Figures 5A, C) and the G1 apex much less 243 tilted so that the mesial lobe is not visible in sternal view (Figure 5A), in contrast to F. 244 reflexifrons (Figures 5B, D, respectively). Also, in F. ibiapaba n. sp. the cephalic lobe is truncate



245 (Figure 4A, 6A), whereas in *F. reflexifrons* it tapers progressively to a distinct narrower tip 246 (Figure 4D, 6D).

Fredius ibiapaba n. sp. further differs from *F. ykaa* in that the G1 shoulder is high and robust (Figures 4B, C), whilst in *F. ykaa* the G1 shoulder is remarkably lower; it can be easily further differentiated from *F. denticulatus* in that its G1 caudal lobe lacks a field of spines spirally twisted to a transverse position (viz., Rodriguez & Campos, 1998) and from *F. fittkaui* in having the G1 cephalic spine straight and sharply acuminate, whereas in *F. fittkaui* it is curved and round tipped.

Discussion

Phylogenetic analysis

The mitochondrial loci 16S was successfully amplified and sequenced for *Fredius buritizatilis*, *F. ibiapaba* n. sp., and *Prionothelphusa eliasi*. Additional sequences used were retrieved from GenBank (Table 1). Bootstrap support Lues are shown on nodes of the phylogenetic tree (Figure 8). *Fredius ibibapaba* n. sp. is very well supported (98) as a sister to *F. reflexifrons* reflecting the close morphological resemblance between the two species.

The divergence rates between *Fredius reflexifrons* and *F. ibiapaba* n. sp. (4%) is higher than between *F. estevisi* x *F. stenolobus*, *F. platyacanthus* x *F. stenolobus* and *F. platyacanthus* x *F. estevisi* all with of 2% of divergence (Table 2). Morphology and molecular data hence provide evidences for the differentiation between *F. ibiapaba* n. sp. and *F. reflexifrons*.

A survey of the pseudothelphusids described from 1840 to 2004 (Yeo et al., 2008) showed that the curve of described species is still far from being asymptotic. And indeed, new species are still being discovered either by collecting in new biomes (e.g., *F. buritizalis* from a palm swamp known as "buritizal"), or by revisiting the taxonomy of widely disjunct species for testing as to their conspecific status, such as *F. ibiapaba* n. sp. and *F. reflexifrons*.



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Zoogeographical notes

272 Fredius currently consists of 16 species (Table 3), distributed over a vast territory, which 273 encompass five main river basins (Rodriguez & Campos, 1998; Magalhães et al., 2014): (1) the 274 Orinoco River basin; (2) the Essequibo-Cuyuni River basin; (3) the Amazon River basin; (4) the 275 Madeira River basin and its tributary (Machado River); and (5) the Atlantic rivers basin, a 276 coastal drainage of small rivers in northern South American (Guyana, Suriname and French 277 Guiana) discharging directly into the Atlantic Ocean. 278 Rodriguez & Pereira (1992) performed a cladistic analysis of *Fredius* and suggested that 279 F. reflexifrons and F. adpressus were sister species. The purported clade F. reflexifrons/ F. 280 adpressus was presumably supported by three putative synapomorphies: (1) [G1] mesial lobe 281 attached to back of auricular lobe; (2) basal denticle of mesial lobe present; and (3) subapical 282 bulge well developed. 283 Later, however, Rodriguez & Campos (1998) reviewed the previous data and performed a 284 new analysis in which they decided that character 1 (mesial lobe attached to back of auricular 285 lobe) was no longer tenable and hence was eliminated from the new analysis. They also 286 realized that the basal denticle of the mesial lobe was indeed present in F. adpressus (character 287 2), but was absent in all other Fredius species. They further concluded that the subapical bulge 288 was actually "reduced" in F. adpressus and "strongly developed" in F. granulatus, F. 289 reflexifrons, F. fittkauii, and F. denticulatus, so that these latter two characters were also 290 removed from the new analysis. Therefore, the putative sister taxon relationship between F. 291 reflexifrons and F. adpressus dissolved. Rodriguez & Campos (1998) put forward, instead, the 292 hypothesis that F. reflexifrons was sister to F. fittkauii, not to F. adpressus, based on the 293 assumption that F. reflexifrons and F. fittkauii synapomorphically share the cephalic lobe distal 294 margin armed with several spinules. However, as found here, this character is more widely 295 distributed being also found in F. ibiapaba n. sp. and, therefore, cannot be used to argue for the

sister taxon relationship between *F. reflexifrons* and *F. fittkauii*.



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The discovery of *F. ibiapaba* n. sp. revealed that it is actually the sister group of *F.* reflexifrons, as shown by a comparative 16S rDNA sequencing used to infer the phylogenetic placement of *Fredius ibiapaba* n. sp. (Figure 8). The distribution range of *Fredius ibiapaba* n. sp. is very narrow and currently restricted to a humid enclave, a small mid-altitude forested patch in Ipú (Ceará, northeastern Brazil, Figure 1A–E), nested within the vast semiarid Caatinga domain (Figure 1F, G). The orographic forest enclaves, such as Ipú, are typically located along the slopes of plateaus, between 600 and 1100 m, hence high enough to receive rainfall of more than 1200 mm year⁻¹ of Atlantic origin (Tabarelli et al., 2004 and references therein). These enclaves are regionally known as "Brejos" (or "Brejos de altitude" or even "Brejos nordestinos") (Andrade-Lima, 1982; Silva & Casteletti, 2003; Tabarelli & Santos, 2004). Fredius ibiapaba n. sp. inhabits the mid-highlands of the Ibiapaba plateau, between about 635 to 782 m, where it digs burrows among the leaf litter, alongside little streams and water ponds inside forest stands or directly on the humid forest floor (Figure 1E). In contrast, F. reflexifrons is widely distributed in the Amazon basin's lowlands (< 100 m) from as far west as Peru (Ampyiacu River, a tributary of the Amazonas River) to as far east as the Atlantic basin (French Guiana) (Magalhães, 2003). It is found in burrows alongside the

"igarapés" (streams) or digs its burrows on the humid forest floor (Magalhães & Rodriguez, 2002).

Fredius ibiapaba n. sp. and F. reflexifrons are highly dependent upon humidity and most probably were once part of an ancestral population living in a wide humid territory. The shrinking humid forests during several dry periods of the Tertiary and Quaternary (Katzer, 1933; Andrade-Lima, 1953; Bigarella et al., 1975; Ab'Saber, 1977; Bigarella & Andrade-Lima, 1982; Andrade-Lima, 1982; Clapperton, 1993; Thomas, 2000; Haffer, 2001; Haffer & Prance, 2002) likely have resulted in the fragmentation of the ancestral humid area and hence of the ancestral crab population, which was split into two species. Fredius reflexifrons evolved and spread in a lowland, humid river basin and is now widely distributed, whilst F. ibiapaba n. sp. evolved





isolated on the top of a humid plateau (Figure 1A–E). The two species are now separated by a vast intervening area occupied by the semiarid Caatinga (Figure 1F, G).

The expansion and shrinkage of mountain, floodplain, and gallery forests, associated to complex topography are known to have fected flora and fauna (Vanzolini, 1970; Vanzolini & Williams, 1970; Vuilleumier, 1971; Andrade-Lima, 1982; Teixeira et al., 1986; Haffer, 1969; 2001; Haffer & Prance, 2002; Santos et al., 2007; Leite et al., 2016).

Andrade-Lima (1982) provided a number of examples of plant species that are now confined to the Brejos, isolated from the surrounding, widely distributed Caatinga. He found two floristic components in these refuge. The top of hills, one whose species and genera are mostly originary from the southeastern flora, lies further inland in the states of Alagoas and Rio Grande do Norte; and a second one in the humid mid highlands closer to the coast, especially between Pernambuco and the border of Ceará and Piauí states (referred to as the Pernambuco Centre by Santos et al., 2007), in which the Amazonian flora are better represented (Andrade-Lima, 1982). Santos et al. (2007) found strong bootstrap support for a close floristic relationship between the Pernambuco Centre and Amazonian localities.

It has long been known that a number of freshwater fish species inhabiting the Brejos have their closest relationships with the Amazonian Basin (Géry, 1969; Paiva, 1978; Weitzman & Weitzman, 1982; Ploeg, 1991; Vari, 1991; Menezes, 1996; Rosa & Groth, 2004). More recently, Pinheiro & Santana (2016) described a new species of freshwater crab genus *Kingsleya* Ortmann, 1897 (also a Pseudothelphusidae), from a Brejo about 750 m in Arajara, district Ceará. Previously to their discovery *Kingsleya* was known from nine species inhabiting the Amazonian lowlands (Pedraza & Tavares, 2015).

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354 355 356	References
357	Ab'Saber, A. N. (1977). Espaços ocupados pela expansão dos climas secos na América do Sul,
358	por ocasião dos períodos glaciais quaternários. Paleoclimas, (3), 1-19.
359	Ab'Saber, A. N. (2000). Spaces occupied by the expansion of bry climates in South America
360	during the Quaternary ice ages. Revista do Instituto Geológico, 21(1-2), 71-78.
361	Andrade-Lima, D. D. (1953). Notas sobre a dispersão de algumas espécies vegetais no Brasil.
362	Anais da Sociedade de Biologia de Pernambuco, 11(1), 25-49.
363	Andrade-Lima, D. (1982) Present day forest refuges in northeastern Brazil. Biological
364	diversification in the tropics (ed. by G.T. Prance), pp. 245–254. Columbia University Press,
365	New York.
366	Auler, A. S., Wang, X., Edwards, R. L., Cheng, H., Cristalli, P. S., Smart, P. L., & Richards, D. A.
367	(2004). Quaternary ecological and geomorphic changes associated with rainfall events in
368	presently semi-arid northeastern Brazil. Journal of Quaternary Science, 19(7), 693-701.
369	Bigarella, J.J. & Andrade-Lima, D. (1982). Paleoenvironmental changes in Brazil. Biological
370	diversification in the tropics (ed. by G.T. Prance), pp. 27–40. Columbia University Press,
371	New York.
372	Bigarella, J.J., Andrade-Lima, D. & Riehs, P.J. (1975). Considerações a respeito das
373	mudanc, as paleoambientais na distribuição de algumas espécies vegetais e animais no
374	Brasil. Anais da Academia Brasileira de Ciências, 47, 411–464.



375	Carmignotto, A. P., de Vivo, M. D., & Langguth, A. (2012). Mammals of the Cerrado and
376	Caatinga: distribution patterns of the tropical open biomes of Central South America.
377	Bones, clones and biomes. The history and geography of recent Neotropical mammals (BD
378	Patterson and LP Costa, eds.). University of Chicago Press, Chicago, Illinois, 307-350.
379	Carnaval, A. C., & Bates, J. M. (2007). Amphibian DNA shows marked genetic structure and
380	tracks Pleistocene climate change in northeastern Brazil. Evolution: International Journal of
381	Organic Evolution, 61(12), 2942-2957.
382	Cartelle, C., & Hartwig, W. C. (1996). A new extinct primate among the Pleistocene megafauna
383	of Bahia, Brazil. Proceedings of the National Academy of Sciences, 93(13), 6405-6409.
384	Clapperton, C. M. (1993). Nature of environmental changes in South America at the Last Glacial
385	Maximum. Palaeogeography, palaeoclimatology, palaeoecology, 101(3-4), 189-208.
386	Cumberlidge, N., Alvarez, F., & Villalobos, J. L. (2014). Results of the global conservation
387	assessment of the freshwater crabs (Brachyura, Pseudothelphusidae and
388	Trichodactylidae): The Neotropical region, with an update on diversity. ZooKeys, (457), 133.
389	Darriba, D., Taboada, G. L., Doallo, R., & Posada, D. (2012). jModelTest 2: more models, new
390	heuristics and parallel computing. Nature methods, 9(8), 772.
391	de Vivo, M. (1997). Mammalian evidence of historical change in the Caatinga semiarid
392	vegetation of northeastern Brazil. Journal of Comparative Biology, 2(1): 65-73.
393	Felsenstein, J. (1985). Confidence limits on phylogenies: an approach using the bootstrap.
394	Evolution, 39(4), 783-791.
395	Géry, J. (1969). The freshwater fishes of South América. Pp. 828-848, in: Fitkau, E.J. et al.
396	(eds.) Biogeography and ecology in South America. Dr. W. Junk, The Hage.
397	Haffer, J. (1969). Speciation in Amazonian forest birds. Science, 165:131–137.
398	Haffer, J. (2001). Hypotheses to explain the origin of species in Amazônia, p. 45–118. In: Vieira,
399	I.C.; Silva, J.M.C.; Oren D.C.; D'Incao, M.A. (orgs) Diversidsade biológica e cultural da
400	Amazônia. Belém, Museu Paraense Emílio Goeldi, 421 p.



- Haffer, J., & Prance, G. T. (2002). Impulsos climáticos da evolução na Amazônia durante o

 Cenozóico: sobre a teoria dos Refúgios da diferenciação biótica. Estudos avançados,
- 403 16(46), 175-206.
- Hall, T. (2005). Bioedit biological sequence alignment editor. Version 7.0.4. Ibis Therapeutics,
- 405 Carlsbad.
- 406 Katzer, F. (1933) Geologia do Estado do Pará. Boletim do Museu Paraense Emílio Goeldi de
- 407 História Natural e Etnografia, 9, 1–270.
- 408 Leite, Y.L.R., Costa, L.P., Loss, A.C., Rocha, R.G., Batalha-Filho, H., Bastos, A.C., Quaresma,
- V.S., Fagundes, V., Paresque, R., Passamani, M., Pardini, R. (2016). Neotropical forest
- expansion during the last glacial period challenges refuge hypothesis. Proceedings of the
- 411 National Academy of Sciences, 113(4), 1008-1013.
- 412 Magalhães, C. (2003). Brachyura: Pseudothelphusidae Trichodactylidae, p. 143–297. In: Melo,
- 413 G.A.S. de (ed.) In: Melo, G.A.S. (Ed.), Manual de Identificação dos Crustáceos Decápodos
- de Água Doce Brasileiros. São Paulo, Edições Loyola. 429 p.
- 415 Magalhães, C. (2009). A new species of freshwater crab of the genus *Fredius* Pretzmann, 1967
- 416 from the middle Amazon River basin, Brazil (Crustacea: Decapoda: Pseudothelphusidae).
- 417 Proceedings of the Biological Society of Washington, 122(1), 81-86.
- 418 Magalhães, C. & Rodriguez, G. (2002). The systematic and biogeographycal status of *Fredius*
- 419 reflexifrons (Ortmann, 1897) and Fredius fittkauii (Bott, 1967) (Crustacea:
- 420 Brachyura:Pseudothelphusidae) from the Amazon and Atlantic Guianas River basins. Acta
- 421 Amazonica, 32(4): 677-689.
- 422 Magalhães, C., Abrunhosa, F. A., Pereira, M. D. O., & Melo, M. A. (2005). New records of
- 423 Fredius denticulatus (H. Milne-Edwards, 1853) and F. reflexifrons (Ortmann, 1897), and the
- 424 eastern limits of the distribution of pseudothelphusid crabs (Crustacea: Decapoda) in Brazil.
- 425 Acta Amazonica, 35(1), 93-96.



120	magainaes, C., & Pereira, G. (2007). Assessment of the decapod crustacean diversity in the
27	Guayana Shield region aiming at conservation decisions. Biota Neotropica, 7(2), 111-124.
28	Magalhães, C.; Sanches, V.Q.A.; Pileggi, L.G.; Mantelatto, F.L. (2014). Morphological and
29	molecular characterization of a new species of Fredius (Decapoda, Pseudothelphusidae)
30	from Rondônia, southern Amazonia, Brazil, p. 101–114. In: Yeo, D.C.J.; Cumberlidge, N.;
31	Klaus, S. (eds.) Advances in freshwater decapod systematics and biology. Crustacean
32	Monographs 19. Leiden, Brill. 296 p.
33	Menezes, N.A. (1996). Methods for assessing freshwater fish diversity. Pp. 289-295 in: Bicudo,
34	C.E. de M. & N.A. Menezes (eds.), Biodiversity in Brazil: a first approach. CNPq, São
35	Paulo.
36	Mora-Day, J., Magalhães, C., & El Souki, M. (2009). Lista Sistemática de los
37	macroinvertebrados Colectados Durante el RAP Alto Cuyuní 2008, Estado Bolívar,
38	Venezuela. In Evaluación Rápida de la Biodiversidad de los Ecosistemas Acuáticos de la
39	Cuenca Alta del Río Cuyuní, Guayana Venezolana. Conservation International.
40	Paiva, M. P. (1978). Ictiofauna e as grandes represas brasileiras. Revista Dae, 38(116), 49-57.
41	Palumbi, SR, Martin A, Romano S, McMillan WO, Stice L, Grabowski G. (1991). The Simple
42	Fool's Guide to PCR, Version 2. University of Hawaii Zoology Department, Honolulu, 45 pp
43	Pedraza, M. & Tavares, M. (2015). A new species of freshwater crab of the genus Kingsleya
44	Ortmann, 1897 (Crustacea: Brachyura: Pseudothelphusidae) from Amazonia, Brazil.
45	Zootaxa, 4032(4): 444-450.
46	Pinheiro, A.P. & Santana, W. (2016). A new and endagered species of Kingsleyia Ortmann,
47	1897 (Crustacea: Decapoda: Brachyura: Pseudothelphusidae) from Ceará, northeastern
48	Brazil. Zootaxa, 4171(2): 365-372.
49	Ploeg, A. (1991). Revision of the South American cichlid genus Crenicichla Heckel, 1840, with
50	descriptions of fifteen new species groups, phylogeny and biogeography (Pisces,



451	Perciformes, Cichlidae) (Doctoral dissertation, PhD Thesis, Universiteit van Amsterdam,
452	Netherland).
453	Rodriguez, G. (1982). Les crabes d'eau douce d'Amerique. Famille des Pseudothelphusidae.
454	Faune Tropicale, 22. ORSTOM, Paris. 224 p.
455	Rodriguez, G., & Pereira, G. (1992). New species, cladistic relationships, and biogeography of
456	the genus Fredius (Decapoda: Brachyura: Pseudothelphusidae) from South America.
457	Journal of Crustacean Biology, 12(2), 298-311.
458	Rodriguez, G., & Campos, M. R. (1998). A cladistic revision of the genus <i>Fredius</i> (Crustacea:
459	Decapoda: Pseudothelphusidae) and its significance to the biogeography of the Guianan
460	lowlands of South America. Journal of Natural History, 32(5), 763-775.
461	Rosa, R. S., & Groth, F. (2004). Ictiofauna dos ecossistemas de brejos de altitude de
462	Pernambuco e Paraíba. Brejos de Altitude em Pernambuco e Paraíba: História Natural,
463	Ecologia e Conservação. Série Biodiversidade, 9, 201-210.
464	Santos, A. M. M., Cavalcanti, D. R., Silva, J. M. C. D., & Tabarelli, M. (2007). Biogeographical
465	relationships among tropical forests in north-eastern Brazil. Journal of Biogeography, 34(3),
466	437-446.
467	Silva, J.M.C. & Casteletti, C.H.M. (2003) Status of the biodiversity of the Atlantic Forest of
468	Brazil. The Atlantic Forest of South America: biodiversity status, threats, and outlook (ed. by
469	C. Galindo-Leal and I.G. Câmara), pp. 43–59. Center for Applied Biodiversity Science and
470	Island Press, Washington, DC.
471	Smalley, A. E. (1964). The river crabs of Costa Rica, and the subfamilies of the
472	Pseudothelphusidae. Tulane studies in Zoology, 12(1), 5-13.
473	Suárez, H. (2015). Six new species of freshwater crabs from Pantepui, Venezuela (Crustacea:
474	Decapoda: Pseudothelphusidae). Anartia, 25, 64-94 [2013].



475 Tabarelli, M., & Santos, A. M. M. (2004). Uma breve descrição sobre a história natural dos 476 brejos nordestinos. Brejos de Altitude em Pernambuco e Paraíba, História Natural, 477 Ecologia e Conservação, 9, 17-24. 478 Tamura, K., Stecher, G., Peterson, D., Filipski, A., & Kumar, S. (2013). MEGA6: Molecular 479 Evolutionary Genetics Analysis version 6.0. Molecular biology and evolution, 30(12), 2725– 480 2729. doi:10.1093/molbev/mst197 481 Teixeira, D.M., Nacinovic, J.B. & Tavares, M.S. (1986) Notes on some birds of northeastern 482 Brazil. Bulletin of the British Ornithologists' Club, 106, 70–74. 483 Thomas, M. F. (2000). Late Quaternary environmental changes and the alluvial record in humid 484 tropical environments. Quaternary International, 72(1), 23-36. Thompson, J. D., Higgins, D. G., & Gibson, T. J. (1994). CLUSTAL W: improving the sensitivity 485 486 of progressive multiple sequence alignment through sequence weighting, position-specific 487 gap penalties and weight matrix choice. Nucleic acids research, 22(22), 4673-4680. 488 Vanzolini, P.E. (1970). Zoologia sistemática, geografia e a origem das espécies. Instituto de 489 Geografia, Universidade de São Paulo, 56 p. (Série Teses e Monografias, 3). 490 Vanzolini, P.E. & Williams, E.E. (1970). South American anoles: Geographic differentiation and 491 evolution of the Anolis chrysolepis species group (Sauria, Iguanidae). Arquivos de Zoologia, 492 19: 1–298. 493 Vari, R. P. (1991). Systematics of the neotropical characiform genus Steindachnerina Fowler 494 (Pisces: Ostariophysi). Smithsonian Contributions to Zoology, 507, 1-118. 495 Vuilleumier, B. S. (1971). Pleistocene changes in the fauna and flora of South America. 496 Science, 173(3999), 771-780. 497 Weitzman, S.H. & Weitzman, M. (1982). Biogeography and evolutionary diversification in the 498 Neotropical freshwater fishes, with comments on the refuge theory. Pp. 403-422, in: 499 Prance, G.T. (ed.) Biological Diversification in the Tropics. Columbia University Press, New 500 York.





501	Xia, X. (2013). DAMBE5: a comprehensive software package for data analysis in molecular
502	biology and evolution. Molecular biology and evolution, 30(7), 1720-1728.
503	WoRMS (2019). Fredius cuyunis (Pretzmann, 1967). Accessed at:
504	http://www.marinespecies.org/aphia.php?p=taxdetails&id=881659 on 2019-11-19
505	Zanetti, F., Castro, P. M. D., & Magalhães, C. (2018). Freshwater crabs (Decapoda: Brachyura:
506	Pseudothelphusidae, Trichodactylidae) from the state of Roraima, Brazil: species
507	composition, distribution and new records. Nauplius, 26, 1–19.
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509	Captions for the figures and tables
510	Figure 1. (A–E) Mid-altitude, naturally isolated, humid forested patch nested within the vast
511	semiarid Caatinga domain. Sítio Caranguejo, Ipú, Ceará, 04°18'50" S, 40°44'47"W, 729 meters
512	high, type locality of <i>Fredius ibiapaba</i> n. sp. Note in (E) burrow (arrow) of <i>Fredius ibiapaba</i> n. sp.
513	among the leaf litter. (E–F) Lowland, surrounding semiarid Caatinga forest. (E) View from above
514	from Ipú. (F) Detail of a dry-stream channel.
515	
516	Figure 2. (A–B) Semi-diagrammatic view of the first male gonopod in abdominal and sternal
517	views, respectively, with the terminology used in the descriptions. CI, ceptalic lobe; cs, cephalic
518	spine; fas, field of apical spines; mal, marginal lobe; mas, marginal suture; mel, mesial lobe;
519	sab, subapical bulge.
520	
521	Figure 3. (A–D) Fredius ibiapaba n. sp., male cl 36 mm, cw 53mm (MZUSP 39710). (A–B)
522	Habitus, dorso and ventral views, respectively. (C) Cephalothorax, frontal view. (D–E) Right and
523	left chelipeds in lateral view, respectively. Scales: A–E, 10 mm.
524	
525	Figure 4. (A–F) First right male gonopod (G1) in abdominal (tilted left), lateral and mesial views
526	from A–C and D–F, respectively. (A–C) Fredius ibiapaba n. sp., male cl 36 mm, cw 53mm
527	(MZUSP 39710). (D–F) Fredius reflexifrons (Ortmann, 1897), male cl 73.8 mm, cw 53 mm
528	(MZUSP 13178). Note in (B, C) the G1 remarkably swollen, sloping abruptly downwards
529	anteriorly to a nearly right-angular shoulder (arrow), and in (E, F) the G1 shoulder clearly more
530	gently sloping distally (arrow).
531	
532	Figure 5. (A–D) First right male gonopod (G1) in sternal and apical views from A to B and C to
533	D, respectively. (A, C) Fredius ibiapaba n. sp., male cl 36 mm, cw 53mm (MZUSP 39710). (B,
534	D) Fredius reflexifrons (Ortmann, 1897), male cl 73.8 mm, cw 53 mm (MZUSP 13178). Note in

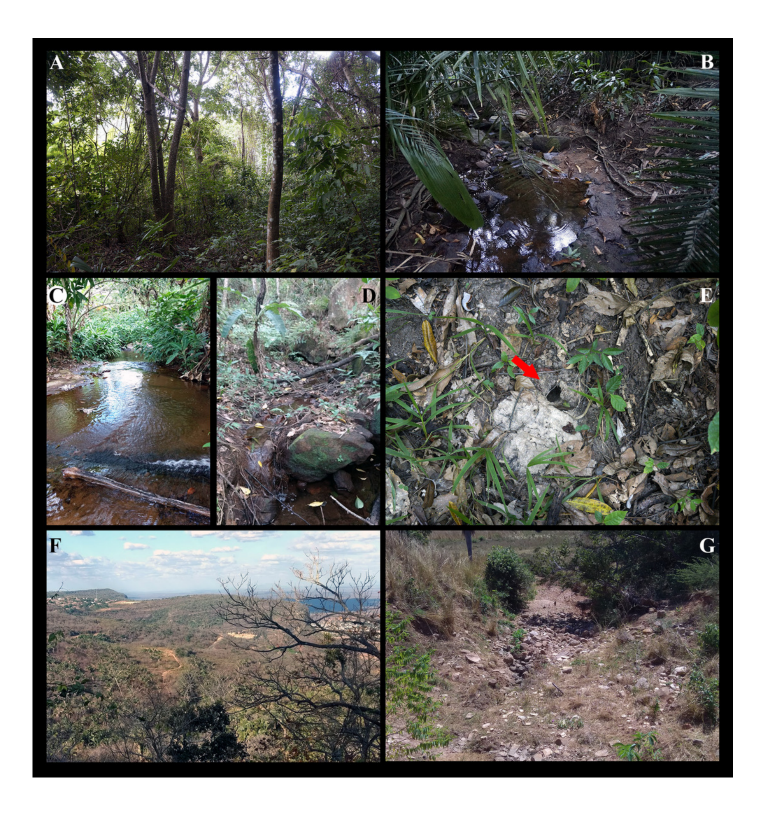


35	(A) and (C) the G1 apex much less tilted so that the mesial lobe is not visible in sternal view
536	(arrow), and the subapical bulge markedly less swollen (arrow), respectively. Note the opposite
537	in (B) and (D).
538	
539	Figure 6. (A–H) First right male gonopod (G1) in sternal, lateral, mesial, and abdominal views
540	from A–D and E–H, respectively. (A–D) Fredius ibiapaba n. sp., male cl 36 mm, cw 53mm
541	(MZUSP 39710). (E–H) Fredius reflexifrons (Ortmann, 1897), male cl 73.8 mm, cw 53 mm
542	(MZUSP 13178).
543	
544	Figure 7. (A–E) Fredius ibiapaba n. sp., paratype, male cl 41.2 mm, cw 62.6 mm (MZUSP
545	39742). Scanning electron microscopy of the first right male gonopod in mesial (tilted right),
546	sternal, apical, lateral, and mesial views. Scales: A–E, 1 mm.
547	
548	Figure 8. Phylogeny inferred from the partial mitochondrial DNA sequence of the 16S rDNA
549	gene. Note the sister taxon relationship between Fredius ibiapaba n. sp. and F. reflexifrons
550	(Ortmann, 1897).
551	
552	Table 1 – Species of <i>Fredius</i> Pretzmann, 1967, <i>Prionothelphusa</i> Rodriguez, 1980 and
553	Trichodactylus Latreille, 1828 used in the phylogenetic analyses, with respective sample locality
554	and GenBank accession number.
555	
556	Table 2 – Pairwise distance matrix from the portion of the mitochondrial 16S rRNA based on
557	~560bp.
558	
	Table 3. Geographic and altitudinal distributions for the species of <i>Fredius</i> Pretzmann, 1967.



Sítio Caranguejo, Ipú, Ceará, 04°18'50" S, 40°44'47"W, 729 meters high, type locality of *Fredius ibiapaba* n. sp.

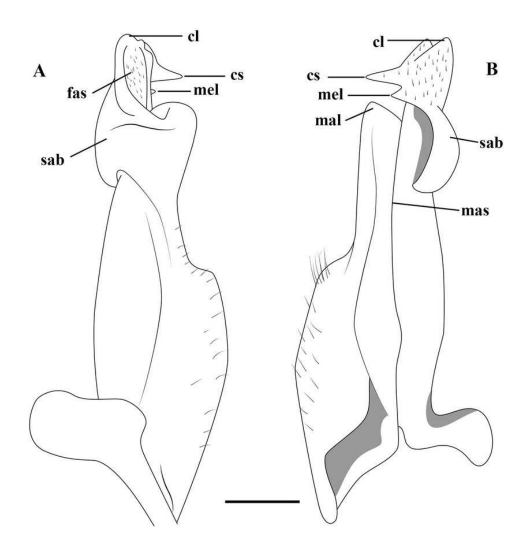
(A–E) Mid-altitude, naturally isolated, humid forested patch nested within the vast semiarid Caatinga domain. Note in (E) burrow (arrow) of *Fredius ibiapaba* n. sp. among the leaf litter. (E–F) Lowland, surrounding semiarid Caatinga forest. (E) View from above from Ipú. (F) Detail of a dry-stream channel.





(A–B) Semi-diagrammatic view of the first male gonopod in abdominal and sternal views, respectively, with the terminology used in the descriptions. Cl, cephalic lobe; cs, cephalic spine; fas, field of apical spines; mal, marginal lobe; mas, marginal sutur





Fredius ibiapaba n. sp., male cl 36 mm, cw 53mm (MZUSP 39710).

(A-B) Habitus, dorso and ventral views, respectively. (C) Cephalothorax, frontal view. (D-E) Right and left chelipeds in lateral view, respectively. Scales: A-E, 10 mm.





First right male gonopod (G1) in abdominal (tilted left), lateral and mesial views from A–C and D–F, respectively.

(A-C) Fredius ibiapaba n. sp., male cl 36 mm, cw 53mm (MZUSP 39710). (D-F) Fredius reflexifrons (Ortmann, 1897), male cl 73.8 mm, cw 53 mm (MZUSP 13178). Note in (B, C) the G1 remarkably swollen, sloping abruptly downwards anteriorly to a nearly right-angular shoulder (arrow), and in (E, F) the G1 shoulder clearly more gently sloping distally (arrow).



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First right male gonopod (G1) in sternal and apical views from A to B and C to D, respectively.

(A, C) Fredius ibiapaba n. sp., male cl 36 mm, cw 53mm (MZUSP 39710). (B, D) Fredius reflexifrons (Ortmann, 1897), male cl 73.8 mm, cw 53 mm (MZUSP 13178). Note in (A) and (C) the G1 apex much less tilted so that the mesial lobe is not visible in sternal view (arrow), and the subapical bulge markedly less swollen (arrow), respectively. Note the opposite in (B) and (D).

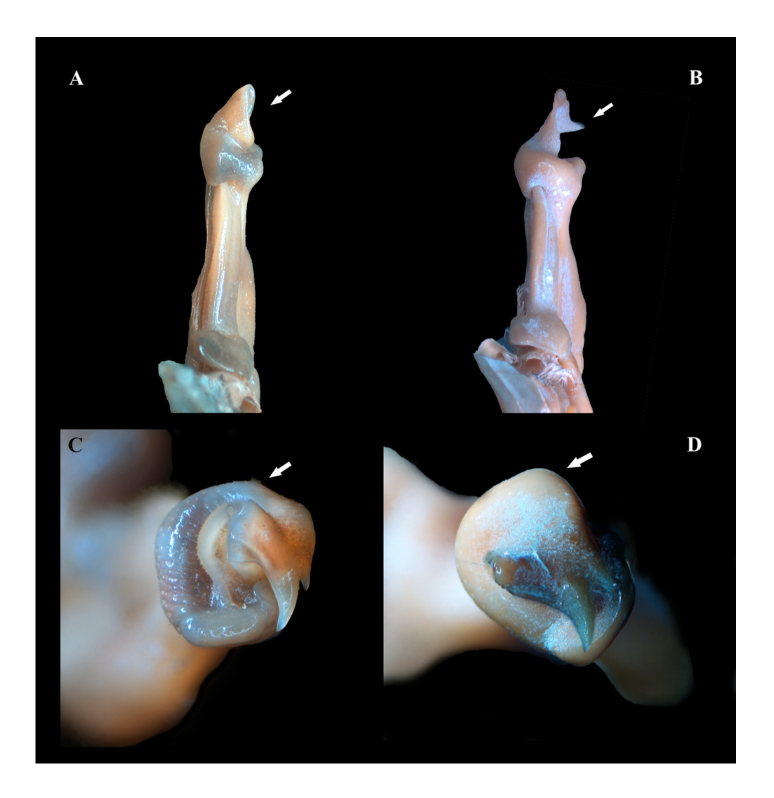




Figure 6

First right male gonopod (G1) in sternal, lateral, mesial, and abdominal views from A-D and E-H, respectively.

(A–D) *Fredius ibiapaba* n. sp., male cl 36 mm, cw 53mm (MZUSP 39710). (E–H) *Fredius reflexifrons* (Ortmann, 1897), male cl 73.8 mm, cw 53 mm (MZUSP 13178).

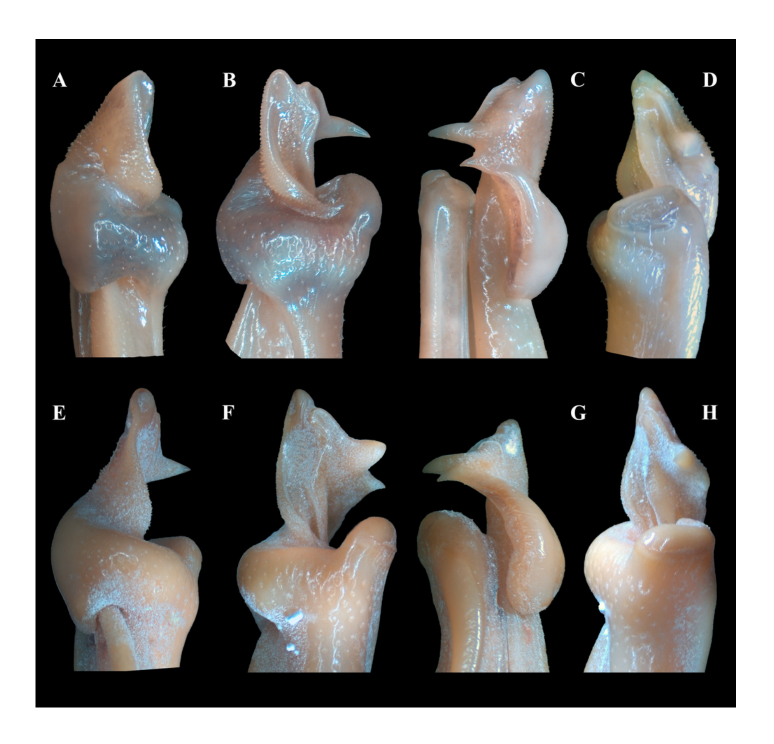


Figure 7

Fredius ibiapaba n. sp., paratype, male cl 41.2 mm, cw 62.6 mm (MZUSP 39742). Scanning electron microscopy of the first right male gonopod.

(A) mesial (tilted right), (B) sternal, (C) apical, (D) lateral, and (E) mesial views. Scales: A-E, 1 mm.

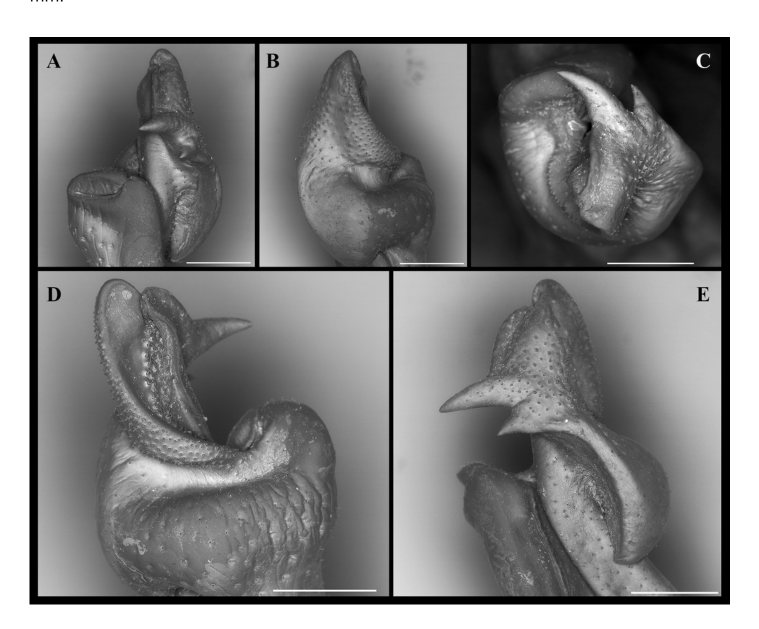




Table 1(on next page)

Phylogeny inferred from the partial mitochondrial DNA sequence of the 16S rDNA gene. Note the sister taxon relationship between *Fredius ibiapaba* n. sp. and *F. reflexifrons* (Ortmann, 1897).

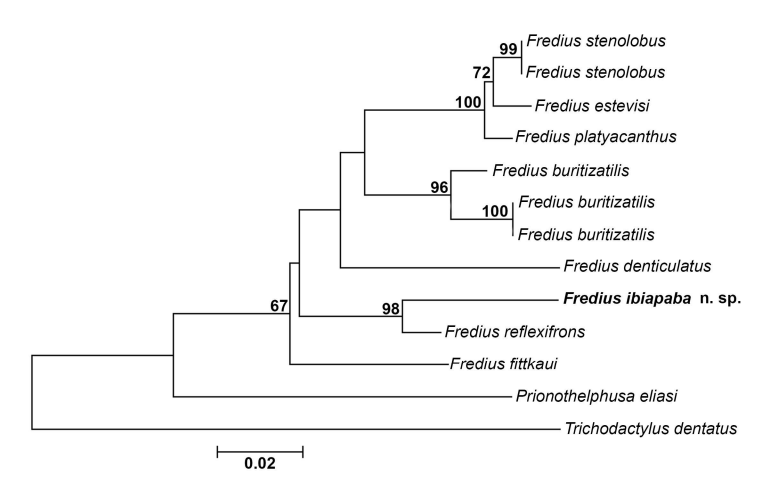




Table 2(on next page)

Species of *Fredius* Pretzmann, 1967, *Prionothelphusa* Rodriguez, 1980 and *Trichodactylus* Latreille, 1828 used in the phylogenetic analyses, with respective sample locality and GenBank accession number.



- 1 Table 1 Species of Fredius Pretzmann, 1967, Prionothelphusa Rodriguez, 1980 and
- 2 Trichodactylus Latreille, 1828 used in the phylogenetic analyses, with respective sample locality
- 3 and GenBank accession number.

Species	Locality	GenBank		
		accession numbers		
Fredius buritizatilis	Ji-Paraná, Rondônia, Brazil	JN402376		
Fredius buritizatilis	Ji-Paraná, Rondônia, Brazil	JN402377		
Fredius buritizatilis	Chupinguaia, Rondônia, Brazil	MN787136		
Fredius denticulatus	Serra do Navio, Amapá, Brazil	JN402372		
Fredius estevisi	Posto Indígena Parafuri, Roraima, Brazil	JN402379		
Fredius fittkaui	Aldeia Balawa-ú, Amazonas, Brazil	JN402373		
Fredius platyacanthus	Comunidade Paapi-ú, Roraima, Brazil	JQ414023		
<i>Fredius ibiapaba</i> n. sp.	Sítio Caranguejo, Ipu, Ceará, Brazil	MN787135		
Fredius reflexifrons	Rio Chumucuí, Bragança, Pará, Brazil	JN402378		
Fredius stenolobus	Rio Tawadu, Bolívar, Venezuela	JN402374		
Fredius stenolobus	Aldeia Palimi-ú, Rio Uraricoera, Roraima, Brazil	JN402375		
Prionothelphusa eliasi	Japurá, Vila Bittencount, Amazonas, Brazil	MN787137		
Trichodactylus dentatus	Bahia, Brazil	FM208777		



Table 3(on next page)

Pairwise distance matrix from the portion of the mitochondrial 16S rRNA based on $\sim 560 \text{bp}$.

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- 1 Table 2 Pairwise distance matrix from the portion of the mitochondrial 16S rRNA based on
- 2 ~560bp.

		1	2	3	4	5	6	7	8	9	10	11	12
1	Fredius ibiapaba n. sp.	-	-	-	-	-	-	-	-	-	-	-	-
2	Fredius reflexifrons	0,04	-	-	-	-	-	-	-	-	-	-	-
3	Fredius burutizatilis	0,10	0,07	-	-	-	-	-	-	-	-	-	-
4	Fredius buritizatilis	0,11	0,08	0,02	-	-	-	-	-	-	-	-	-
5	Fredius buritizatilis	0,11	0,08	0,02	0,00	-	-	-	-	-	-	-	-
6	Fredius denticulatus	0,12	0,09	0,08	0,08	0,08	-	-	-	-	-	-	-
7	Fredius stenolobus	0,10	0,07	0,07	0,06	0,06	0,10	-	-	-	-	-	-
8	Fredius stenolobus	0,10	0,07	0,07	0,06	0,06	0,10	0,00	-	-	-	-	-
9	Fredius estevisi	0,11	0,07	0,07	0,06	0,06	0,09	0,02	0,02	-	-	-	-
10	Fredius fittkaui	0,09	0,07	0,07	0,07	0,07	0,09	0,08	0,08	0,08	-	-	-
11	Fredius platyacanthus	0,10	0,07	0,06	0,06	0,06	0,09	0,02	0,02	0,02	0,08	-	-
12	Prionothelphusa eliassi	0,16	0,13	0,12	0,12	0,12	0,13	0,12	0,12	0,12	0,13	0,12	-
13	Trichodactylus dentatus	0,22	0,19	0,21	0,21	0,21	0,22	0,21	0,21	0,22	0,20	0,21	0,22

3



Table 4(on next page)

Geographic and altitudinal distributions for the species of *Fredius* Pretzmann, 1967.

Table 3. Geographic and altitudinal distributions for the species of *Fredius* Pretzmann, 1967.

Species Country		Environment	Altitude (m)	References	
F. ykaa Magalhães, 2009	Brazil (Amazon River basin)	Lowland streams	36 to 73	Magalhães, 2009	
<i>F. adpressus</i> Rodriguez & Pereira, 1992	Venezuela (Orinoco River basin)	Lowland streams	100	Rodriguez & Pereira, 1992	
F. beccarii (Coifmann, 1939)	Brazil, Guyana, Venezuela, Suriname (Essequibo- Cuyuni Rivers basin)	Streams (igarapés)	50 to 752	Rodriguez & Campos, 1998; Cumberlidge, Alvarez & Villalobos, 2014; Mora-Day et al., 2009; Magalhães et al., 2014; Zanetti et al. 2018	
F. buritizatilis Magalhães & Mantelatto, 2014	Brazil (Madeira River basin)	Buritizal (palm) fields	150	Magalhães et al., 2014	
F. chaffanjoni (Rathbun, 1905)	Venezuela (Orinoco River basin)	River's headwaters and mid- courses	105-300	Rodriguez & Pereira, 1992	
F. convexa (Rathbun, 1898)	Costa Rica	Highland streams	770	Smalley, 1964	
F. cuaoensis Suárez, 2015	Venezuela (Orinoco River basin)	Highland streams	950	Suárez, 2015	
F. cuyunis (Pretzmann, 1967)	British Guyana (Cuyuní River)	Lowlands	around 100	WoRMS, 2019	
F. denticulatus (H. Milne Edwards, 1853)	Brazil, Suriname, French Guiana (Amazon and	Streams (igarapés) and along river margins	70 to 400	Rodriguez & Pereira, 1992; Rodriguez &	

F. estevisi (Rodríguez, 1966)	Atlantic river basins) Brazil, Venezuela (Amazon and Atlantic	River's headwaters and streams	446 to 944	Campos, 1998; Magalhães et al., 2005; Magalhães, 2009; Cumberlidge, Alvarez & Villalobos, 2014; Magalhães et al., 2014 Mora-Day et al., 2009
F. fittkaui (Bott, 1967)	rivers basins) Brazil, Venezuela, Guyana (Amazon and Atlantic rivers basins)	Streams (iIgarapés) and along river margins	151 to 500	Rodriguez & Campos, 1998; Magalhães & Rodriguez, 2002; Cumberlidge, Alvarez & Villalobos, 2014; Magalhães et al., 2014; Zanetti et al., 2018
F. granulatus Rodriguez & Campos 1998	Colombia (Amazon River basin)	Lowlands	180 to 200	Rodriguez & Campos, 1998; Cumberlidge et al., 2014 Cumberlidge, Alvarez & Villalobos, 2014; Zanetti et al., 2018;
F. platyacanthus Rodríguez & Pereira,	Brazil, Venezuela (Atlantic	Streams (Igarapes) and	106 to 1229	Rodriguez & Pereira,

1992	rivers basin)	mountain areas		1992; Cumberlidge, Alvarez & Villalobos, 2014; Magalhães et al., 2014; Zanetti et al., 2018
F. reflexifrons (Ortmann, 1897)	Brazil, Venezuela, Suriname, French Guaiana, Peru, Guyana (Amazon and Atlantic rivers basins)	Lowland streams	37 to 200	Magalhães & Rodriguez, 2002; Magalhães et al., 2005; Cumberlidge, Alvarez & Villalobos, 2014
F. stenolobus Rodríguez & Suárez, 1994	Brazil, Venezuela (Orinoco River basin)	Streams in rocky areas	65 to 1020	Rodriguez & Campos, 1998; Magalhães & Pereira, 2007; Cumberlidge, Alvarez & Villalobos, 2014; Magalhães et al., 2014; Zanetti et al., 2018
Fredius ibiapaba n. sp.	Brazil (Orographic forest enclaves)	Burrows among the leaf litter, alongside little streams and water ponds inside forest stands or directly on the humid forest floor	665 to782	Present study