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The first molecular phylogeny of Chilodontidae (Teleostei: Ostariophysi: Characiformes) reveals cryptic biodiversity and taxonomic uncertainty

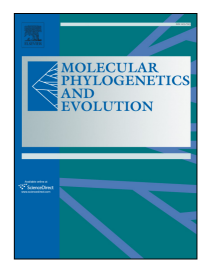
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1	The first molecular phylogeny of Chilodontidae (Teleostei: Ostariophysi:
2	Characiformes) reveals cryptic biodiversity and taxonomic uncertainty
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18	Running Title: Molecular phylogeny of Chilodontidae
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Abstract

20

Chilodontidae is a small family of eight described characiform species popularly 21 known as headstanders. These small to moderately sized fishes are well known to 22 aquarists, who prize their striking spotted pigmentation and unusual behaviors, and to 23 systematists, who have revised both chilodontid genera in recent memory and studied 24 25 their phylogenetic relationships using a comprehensive morphological dataset. However, no molecular phylogeny for the family has ever been proposed. Here, we 26 reconstruct phylogenetic relationships for all eight known chilodontid species using 27 three mitochondrial and two nuclear loci. Results largely agree with the previous 28 morphological hypothesis, and confirm the monophyly of the family as well as its 29 30 included genera, *Caenotropus* and *Chilodus*. The molecular topology differs slightly from the morphological hypothesis by placing *Caenotropus maculosus* rather than C. 31 mestomorgmatos as the sister to the remaining three congeners, and by reconstructing 32 33 the Curimatidae as the closest outgroup family, rather than the Anostomidae. However, the topologies supported by the morphological data were only slightly less likely and 34 could not be rejected via Shimodaira-Hasegawa tests. Within Chilodus, two described 35 species with distinctive pigmentation (C. fritillus and C. zunevei) appear embedded 36 within the broad distributed C. punctatus clade, suggesting the presence of cryptic taxa 37 with polymorphic pigmentation within the present concept of *C. punctatus*. Future work 38 39 should combine morphological and molecular data to revisit the taxonomy and systematics of *Chilodus* and determine species limits within the *C. punctatus*-group 40 41 sensu lato.

42

43 Keywords: Amazon, Freshwater fishes, Headstanders, Multilocus analysis, Systematics.

44

45 1. Introduction

46

40	
47	The 275 species in the fish superfamily Anostomoidea of the order
48	Characiformes (Eschmeyer and Fong, 2013), are widely distributed in Central and
49	South American freshwater ecosystems from Costa Rica to Argentina (Vari, 1983) and
50	in northern South America occur in river systems to both sides of the Andean
51	Cordilleras. Anostomoidea includes the families Anostomidae, Chilodontidae,
52	Curimatidae and Prochilodontidae, that together form a major recognized monophyletic
53	assemblage (Vari, 1983; Buckup, 1998; Sidlauskas, 2008) supported by
54	synapomorphies related to modifications to the gill arches, musculature and dentition.
55	The major clades within the superorder are defined by multiple derived features of those
56	body systems, plus the neurocranium, jaws, connective tissues, pectoral girdle and axial
57	skeleton (Vari, 1983; Vari, 1989; Vari et al., 1995; Castro and Vari, 2004; Sidlauskas
58	and Vari, 2008).
59	Species of Chilodontidae (Fig. 1), one of the four families in the Anostomoidea,
60	adopt a typical head-down orientation while swimming and resting (Vari et al., 1995),
61	an orientation unusual within Characiformes, and are consequently popularly known as
62	headstanders. Given this unusual orientation, small to moderate body sizes and striking
63	pigmentation patterns, chilodontids are well known among aquarists (Isbrücker and
64	Nijssen, 1988). Members of Chilodontidae are broadly distributed in the Amazon and
65	Orinoco river basins, the coastal rivers draining the Guianas, and the Rio Parnaíba basin
66	in northeastern Brazil (Vari and Raredon, 2003; Vari et al., 2009) where they feed on a
67	combination of small invertebrates, sponges and detritus (Goulding et al., 1988; Vari

and Raredon, 2003). The family is composed by eight recognized species, six of which

69 were previously studied in taxonomic reviews of *Chilodus* (Isbrücker and Nijssen,

1988) and *Caenotropus* (Vari et al., 1995). Two additional species were described in

subsequent decades: *Chilodus fritillus* by Vari and Ortega (1997) and *Caenotropus*

schizodon by Scharcansky and Lucena (2007).

Vari (1983) and Vari et al. (1995) proposed a series of synapomorphies for 73 Chilodontidae, as well as for its two included genera, *Caenotropus* and *Chilodus*. In the 74 latter publication, Vari et al. (1995) proposed intrageneric phylogenetic relationships for 75 *Caenotropus* on the basis of 10 morphological characters. Scharcansky and Lucena 76 77 (2007) more recently expanded that analysis to address the phylogenetic placement for *C. schizodon* described in the same publication. No set of phylogenetic relationships 78 within *Chilodus* has ever been proposed. Using a multilocus molecular dataset, Oliveira 79 et al. (2011) corroborated the monophyly of Anostomoidea and of Chilodontidae, 80 though their analysis included only two chilodontid species (*Caenotropus labyrinthicus* 81 82 and Chilodus punctatus). No more detailed molecular hypothesis of phylogenetic relationships within Chilodontidae is available. 83 Herein, we present the first phylogenetic analysis including all eight species in 84 the Chilodontidae. Our aims were to test the monophyly of Chilodontidae (sensu Vari, 85 1983) and both genera in the family and to infer their interspecific relationships using a 86 model-based phylogenetic analysis of molecular data. We also discuss the 87 biogeographic distribution of the family, apparent morphological convergences, and the 88 possibility of unrecognized cryptic species within the context of the molecular results. 89 90

91 2. Material and Methods

92 2.1. Taxon sampling

At least one specimen of all eight species of Chilodontidae (Table 1) wasincluded in the analysis. We sampled broadly across the distributional range of

Caenotropus labyrinthicus and *Chilodus punctatus*, which are the most common species 95 96 of chilodontids and among the most widespread of the species that have been the subject of recent analysis among all New World characiforms. The map in Fig. 2 97 illustrates the sampling localities for the ingroup taxa and was prepared using the 98 Quantum GIS 1.7.1 and Cartographer module of Mesquite (Maddison and Maddison, 99 2013). We included several species of each of the other three anostomoid families 100 (Anostomidae, Curimatidae and Prochilodontidae) as outgroups, as well as one species 101 102 in Hemiodontidae that was used to root the tree (Table 1). Tissues were preserved in 95% ethanol or a saturated DMSO/NaCl solution. Voucher specimens were formalin-103 fixed, alcohol-preserved and deposited in collections (Table 1; abbreviations follow 104 http://www.asih.org/codons.pdf). 105 106 2.2. DNA extraction and sequencing 107 Total DNA was extracted from ethanol-fixed muscle tissue with a DNeasy 108 Tissue kit (Qiagen Inc.; www.qiagen.com) following the instructions of the 109 110 manufacturer, or following a modified NaCl extraction protocol adapted from Lopera-Barrero et al. (2008). Partial sequences of the genes 16S rRNA (16S, 608 bp), 111 cytochrome oxidase C subunit 1 (COI, 633 bp) and cytochrome B (Cytb, 985 bp) were 112 amplified using one round of polymerase chain reaction (PCR). Additionally, we 113 114 obtained sequences of the myosin heavy chain 6 gene (Myh6, 704 bp), and 115 recombination activating gene 1 (Rag1, 1210 bp) through nested-PCR following the 116 procedures detailed in Oliveira et al. (2011). PCR amplifications were performed in 117 12.5 µl reactions containing 9.075 µl of double-distilled water, 1.25 µl 5 x reaction buffer, 0.375 µl MgCl₂, 0.25 µl dNTP mix at 8 mM, 0.25 µl of each primer at 10 µM 118 119 (list of primers in Table 2), 0.05 µl Platinum Taq DNA polymerase enzyme (Invitrogen;

120	www.invitrogen.com) and 1.0 μl genomic DNA (10-50 ng). The amplification cycles
121	consisted of an initial denaturation (4 min at 95°C) followed by 28 cycles of chain
122	denaturation (30s at 95°C), primer hybridization (30-60s at 52-54°C), and nucleotide
123	extension (30-60s at 72°C). All PCR products were visually identified in a 1% agarose
124	gel. Samples were cleaned using ExoSAP (Hanke and Wink, 1994) and subsequently
125	sequenced using dye terminators (Big Dye [™] Terminator v 3.1 Cycle Sequencing Ready
126	Reaction Kit, Applied Biosystems; www.appliedbiosystems.com) purified again
127	through ethanol precipitation and loaded onto an automatic sequencer ABI 3130-
128	Genetic Analyzer (Applied Biosystems) at either the Universidade Estadual Paulista,
129	Botucatu, São Paulo, Brazil, or Oregon State University, Corvallis, Oregon, USA.
130	Consensus sequences were assembled and edited in BioEdit 7.0.9.0 (Hall, 1999) and
131	Geneious 6.1 (Biomatters, 2013). Where uncertainty of nucleotide identity was detected,
132	IUPAC ambiguity codes were applied.

133

134 2.3. Alignment and phylogenetic analyses

Consensus sequences of each gene for each individual were independently 135 aligned using the Muscle program (Edgar, 2004) under default parameters. The 136 resulting alignments were inspected by eye for obvious misalignments that were then 137 corrected manually. GenBank accession numbers appear in Table 1 and the matrix was 138 139 deposited in TreeBase (http://treebase.org) under number 14605. To evaluate the occurrence of substitution saturation, the index of substitution saturation (Iss) as 140 described by Xia et al. (2003) and Xia and Lemey (2009) was estimated using Dambe 141 142 5.3.38 (Xia, 2013). The nucleotide frequencies were computed in MEGA 5.0 (Tamura et al., 2011). 143

144	Maximum likelihood (ML) analyses were generated in a partitioned (13
145	partitions, Table 3) RAxML (Stamatakis, 2006) analysis using the CIPRES web server
146	(Miller et al., 2010). Random starting trees were used for ML tree search and all other
147	parameters were set to default values. All ML analyses were performed under GTR+G
148	since RAxML only applies this model (Stamatakis et al., 2008). The robustness of the
149	topology was investigated using 1000 bootstrap pseudoreplicates.
150	Maximum parsimony (MP) analysis was performed using PAUP* 4.0b10
151	(Swofford, 2003). Heuristic searches were performed with minimally 1000 random
152	addition replicates and TBR branch swapping. All characters were unordered and all
153	transformation series were equally weighted. Branches with maximum length of zero
154	were collapsed. Gaps were treated as missing data. The resulting topologies were
155	statistically tested with the bootstrap method (Felsenstein, 1985) using 1000
156	pseudoreplicates.
157	We inferred a Bayesian topology (BI) with a partitioned matrix using MrBayes
158	3.1.2 (Huelsenbeck and Ronquist, 2001; Ronquist and Huelsenback, 2003) via the
159	CIPRES web portal (Miller et al., 2010). Schema ranging from 1 to 13 partitions was
160	tested following Li et al. (2007) under the Akaike information criterion (AIC) as
161	detailed in Table 3 and in Additional file 1. Parameters were estimated using ModelTest
162	3.6 (see Posada and Crandall, 1998 for model symbols) implemented in PAUP* 4.0b10
163	(Swofford, 2003) on each partition (Table 3). We performed two runs of four
164	independent MCMC chains with 10 million replicates each, sampling one tree every one
165	thousand generations. The distributions of log likelihood scores were examined using
166	Tracer 1.5 (Rambaut and Drummond, 2007) in order to determine stationarity and
167	decide if extra runs were required to achieve convergence. The first one million

168	generations (10%) were discarded as burn-in, and the remaining trees were used to
169	construct a 50% majority rule consensus tree in PAUP* (Swofford, 2003).
170	To test the degree of support for the resulting molecular phylogenies versus the
171	previously published morphological hypothesis, we compared the maximum likelihood
172	unconstrained tree to the maximum likelihood trees generated under two different
173	constraint trees in RAxML. In the first, we constrained the four anostomoid families to
174	conform to the arrangement proposed by Vari (1983) in which Anostomidae and
175	Chilodontidae are sister taxa, as are Curimatidae and Prochilodontidae. In the second,
176	the ingroup Chilodontidae was constrained to conform to the morphological hypothesis
177	of relationships within <i>Caenotropus</i> of Vari et al. (1995), modified by the addition of <i>C</i> .
178	schizodon as sister to C. labyrinthicus as proposed by Scharcansky and Lucena (2007).
179	Constraint trees were constructed in Mesquite (Maddison and Maddison, 2013). We
180	compared the maximum likelihood topologies inferred under these three scenarios using
181	the Shimodaira-Hasegawa test (Shimodaira and Hasegawa, 1999) as implemented in the
182	phangorn (Schliep, 2011) package in R (R Development Core Team, 2013). Within
183	phangorn, we compared the likelihood fits assuming a GTR substitution model, four
184	discrete intervals of the gamma distribution (k=4), and 10,000 bootstrap replicates. We
185	computed likelihoods and p-values with and without optimizing the rate matrix and base
186	frequencies in phangorn (Table 4, Additional files 2 and 3). We also filtered the
187	combined results from both Bayesian Markov chains to determine the percentage of
188	trees in the posterior distribution that were consistent with bipartitions present in the
189	morphological reconstruction, but not present in the best-supported molecular
190	reconstruction.
101	

191

192 3. Results

193	Most sequences from the outgroup species were previously published in the
194	phylogeny of the Characidae by Oliveira et al. (2011); however, the sequences from the
195	cytochrome oxidase C subunit I (COI) for these species were newly generated for this
196	study. The concatenated matrix from 13 outgroups and 32 specimens of the
197	Chilodontidae include 4140 bp and 1506 variable sites of which 1179 were parsimony
198	informative. The Iss index indicated no saturation in transitions or transversions in both
199	asymmetrical (Iss.cAsym) and symmetrical (Iss.cSym) topologies. Table 2 contains
200	numbers of base pairs (bp) after alignment, primer sequences, and nucleotide
201	composition for each analyzed gene. Comparisons of log likelihoods, AIC and BIC
202	values among different partitioning schemes (from 1 to 13 partitions) were tested and
203	are presented in Additional file 1.
204	Throughout the text and in Fig. 3, measures of support are indicated as a series
205	of three numbers on selected internal branches of the trees subtending labeled clades,
206	starting with posterior probabilities in Bayesian Inference (BI) analysis and followed by
207	non-parametric bootstrap percentages from maximum likelihood (ML) and parsimony
208	(MP) analyses, respectively (e.g., 1/100/100, see Fig. 3); dashes represent values lower
209	than 0.9 (B) or 50% (ML and MP) and asterisks represent nodes that have different
210	topologies in different analytical methods. Nodes without support values greater than
211	0.9 (B) and 50% (ML and MP) were collapsed.
212	The Bayesian results represent a majority rule consensus of 18002 post-burn-in
213	trees, the likelihood analysis yielded a single tree with a sum of branch lengths (SBL) of
214	1.931, and the parsimony analysis returned a single tree (TL: 4726; CI: 0.439; RI:
215	0.714). Fig. 3 shows the maximum likelihood topology, along with bootstrap and
216	posterior probabilities values from the three analyses, all of which returned very similar
217	results. Anostomidae, Chilodontidae, Curimatidae and Prochilodontidae were all

corroborated as monophyletic by all three analyses with 100% bootstrap support or 218 219 posterior probability equal to one. We obtained a well-supported clade (1/93/70)composed of the Chilodontidae, Curimatidae and Prochilodontidae. The Bayesian 220 221 consensus differs from the maximum likelihood topology only in a single relationship among the species of the outgroup Curimatidae. The clade composed by Chilodontidae 222 and Curimatidae was well-supported in BI (0.9) and ML (82) but with low support in 223 the MP reconstruction (41). Otherwise, optimal topologies from the three analyses agree 224 225 completely.

Within Chilodontidae (clade 1), the monophyly of *Caenotropus* (clade 2) was 226 also well-supported with C. maculosus (clade 3) as the sister of a clade composed by its 227 three congeners. In clade 4, the analysis placed C. mestomorgmatos as sister to clade 5 228 composed of C. labyrinthicus and C. schizodon. Clade 6 is composed by two specimens 229 230 of C. schizodon from the Rio Tapajós in the eastern portions of the Amazon basin and one specimen of C. labyrinthicus from the Río Nanay in Peru (Fig. 2), in the western 231 232 portion of that river system, thereby rendering C. labyrinthicus paraphyletic (albeit with only moderate statistical support: -/71/63). 233

Our results corroborated *Chilodus* as a monophyletic genus (clade 7). Within *Chilodus, C. gracilis* (from the Rio Negro in Brazil) appeared as monophyletic (clade 8) and as sister to clade 9 containing the other three currently recognized species of the genus (*C. fritillus, C. punctatus* and *C. zunevei*). *Chilodus punctatus*, however, was not recovered as a monophyletic group as a consequence of *C. fritillus* and *C. zunevei* nesting among the twelve sampled individuals of *C. punctatus*.

Within clade 9, two lineages were recovered. The first is composed solely by *Chilodus punctatus* from the Rio Araguaia, a large river south of the mainstream
Amazon that drains a portion of the Brazilian Shield and flows into the Rio Tocantins

and through that river to the lower Rio Amazonas (Fig. 2). The second lineage (clade 9, 243 244 0.8/92/68) is composed by specimens of *Chilodus punctatus* from the remainder of the Amazon, including the Rios Juruá, Purus and Nanay in the west, the Rio Takutu at the 245 border of Brazil and Guyana (a tributary of the upper Rio Branco) in the northeast, and 246 the Rio Guamá (a southern tributary of the lower Amazon) in the east (Fig. 2). Clade 247 nine also includes C. fritillus from south-eastern Peru in the Río Madre de Diós, an 248 upper tributary to the Rio Madeira in the western Amazon and a specimen of C. zunevei 249 250 from the Atlantic slope of the Guianas (Commewijne River, Suriname) in north-eastern South America. 251 While the tree topology discussed above is that best supported by the available 252 molecular data, the maximum likelihood trees under the topological constraints based 253 on previous morphological results (Additional files 2 and 3) are only slightly less likely 254 255 (Table 4). Shimodaira-Hasegawa tests failed to reject these two alternative topologies (Table 4). Nevertheless, no sampled trees in the Bayesian posterior distributions are 256 fully congruent with the morphological topologies. 257 258 4. Discussion 259 4.1. Interfamilial relationships within Anostomoidea 260 261 Our molecular study returns the same hypothesis of close relationship among 262 263 Chilodontidae, Curimatidae and Prochilodontidae obtained by Oliveira et al. (2011). 264 That congruence is perhaps unsurprising given the similarity of loci and taxa examined in the two studies. This arrangement differs from the interfamilial hypothesis 265

266 ((Anostomidae + Chilodontidae) + (Curimatidae + Prochilodontidae)) of Vari (1983,

267 1989), which was based on synapomorphies of multiple morphological systems. The

latter result was subsequently obtained by Buckup (1998) using much of the same data. 268 269 The arrangement of families suggested by the molecular data would imply a large number of morphological convergences or reversals and indicate a much more complex 270 271 evolutionary history of these fishes than previously suspected. Of particular note would be the very distinctive shared modifications of the gill arches in the Chilodontidae and 272 Anostomidae, which include pronounced enlargement of the upper and lower 273 pharyngeal dentition, the presence of two or more pointed cusps on those teeth, a shift 274 275 in alignment of the fourth upper pharyngeal tooth plate, vertical expansion of the fifth upper pharyngeal tooth plate, a highly developed obliquees dorsalis muscle on the 276 fourth infrapharyngobranchial and cord-like ligaments joining the ectopterygoid and 277 ventral wing of the lateral ethmoid; none of which occur elsewhere in Characiformes 278 (Vari, 1983). Under the scenario implied by the molecular data, either all of these 279 280 characters are convergent, or they were present in the common ancestor of Anostomoidea and subsequently lost during the evolution of Curimatidae and 281 282 Prochilodontidae. 283 The strength of the morphological data makes it noteworthy that the currently available genetic data do not strongly reject the morphological hypothesis of 284 interfamilial relationships. Support for the novel arrangement based on molecular 285 286 evidence was inconsistent among the three methods that we employed, and support for the maximum likelihood tree that conforms to the morphological hypothesis (Additional 287 288 file 2) was similar enough to the support for the unconstrained tree that a Shimodaira-289 Hasegawa test was unable to distinguish between these hypotheses (Table 4). None of

the sampled trees in the Bayesian posterior conform exactly to the morphological

291 hypothesis of relationships among these families, but substantial variation in results

exist. The posterior distribution includes a moderate percentage (4.44%) of topologies

in which Chilodontidae appears as sister to Prochilodontidae as well as more limited
occurrence (<1%) of sister relationships between Anostomidae and Curimatidae,
Anostomidae and Prochilodontidae, and Curimatidae and Prochilodontidae. We plan to
examine these alternatives more thoroughly in an upcoming analysis with more
complete taxon and gene sampling. Such further tests are outside the scope of the
present paper, which focuses on Chilodontidae.

299

300 4.2. Monophyly of Chilodontidae and included genera

Our results support the monophyly of the family Chilodontidae, corroborating 301 the previous morphological hypothesis of Vari (1983) and Vari et al. (1995) and 302 agreeing with Greenwood et al. (1966), who in a pre-cladistic analysis were the first 303 authors to elevate chilodontids to the familial level. Subsequently, Vari (1983) proposed 304 305 the monophyly of the family Chilodontidae based on 26 synapomorphies (his characters 74 to 99) mostly related to the gill-arches, suspensorium, pectoral girdle, and anterior 306 portions of the vertebral column. In a taxonomic revision of *Caenotropus*, Vari et al. 307 308 (1995) discovered 10 additional synapomorphies (their characters 27 to 36) for the family involving a variety of osteological and scale-based characters. Oliveira et al., 309 (2011) later obtained monophyly of the two species of Chilodontidae included in their 310 molecular study. 311

Clade 2, containing the four species of *Caenotropus*, corroborates the monophyly of that genus as hypothesized by Vari et al. (1995) on the basis of a series of eight synapomorphies involving details of the bones of the infraorbital series, jaws, neurocranium and scales among others. Similarly, clade 7, comprising all species and multiple specimens of *Chilodus* from different localities, appears as a strongly supported monophyletic lineage (1/100/100). Vari et al. (1995) has previously proposed

318 monophyly of *Chilodus* based on eight synapomorphies related to modifications of the

bones of the infraorbital series, the lower jaw, the hyoid series, the pattern of the

320 laterosensory canals in the dentary, infraorbitals and neurocranium and the complete

321 loss of the third postcleithrum.

322

323 *4.3. Interspecific relationships in Caenotropus*

Vari et al. (1995) distinguished *Caenotropus mestomorgmatos* from its 324 325 congeners as a new species on the basis of the combination of various features. Two of these, the presence of three scales (versus four) above the lateral line to the dorsal fin of 326 a broad region of dusky pigmentation across most of the dorsal fin (versus no 327 pigmentation or just a distal spot of pigmentation) were unique to the species in the 328 genus and with the latter proposed as an autapomorphy for the species. Those authors 329 330 proposed that C. mestomorgmatos was, in turn, the sister species to a clade composed of C. maculosus and C. labyrinthicus and supported their proposal with three 331 synapomorphies involving different portions of the body. Scharcansky and Lucena 332 333 (2007) subsequently described C. schizodon and hypothesized a sister relationship between that new species and C. labyrinthicus. 334

Our results are incongruent with those hypotheses, and reverse the position of *C*. 335 maculosus and C. mestomorgmatos proposed by Vari et al. (1995). The results of this 336 study consequently place C. maculosus as sister to the group containing the remaining 337 338 three species in the genus (C. labyrinthicus, C. mestomorgmatos and C. schizodon) with C. mestomorgmatos as sister to a clade formed by C. labyrinthicus and C. schizodon 339 (Fig. 3). The arrangement supported by the morphological analysis is only slightly less 340 341 likely than that based on the molecular data, and the two topologies could not be 342 distinguished by the Shimodaira-Hasegawa test (Table 4). However, no trees in the

Bayesian posterior place *C. mestomorgmatos* as sister to its remaining congeners, and
the constrained maximum likelihood tree (Additional file 3) includes a very short
internode that effectively creates a polytomy among *C. mestomorgmatos, C. maculosus*and a clade containing *C. labyrinthicus* and *C. schizodon*. Thus, the molecular support
for the novel hypothesis of relationships within *Caenotropus* is relatively robust, and it
is worth exploring whether morphological support for the basal position of *C. maculosus* exists.

As pointed out by Vari et al. (1995), all species of *Chilodus* have darkly pigmented distal portions of the dorsal fin (Fig. 1, see also Isbrücker and Nijssen, 1988). Under our molecular reconstruction, *Caenotropus maculosus* shares this plesiomorphic condition, as does *C. mestomorgmatos* to a fainter extent. Their remaining congeners would then possess a synapomorphic reduction in dorsal-fin pigmentation (Fig. 1).

Our results also suggest the possibility of cryptic diversity within the present 355 concept of Caenotropus maculosus. A deep split occurs within Clade 3, separating a 356 specimen from the Corantijn River basin of western Suriname from two others that 357 originated in the Marowijne River system of eastern Suriname and western French 358 359 Guiana. Within the intervening drainages, C. maculosus is only known from the 360 Suriname River which lies close to the Marowijne basin (Sidlauskas and Vari, 2012; 361 Mol et al., 2012). The evidence suggests the presence of two genetically isolated 362 populations. Additional studies should investigate whether these merit species status.

Our results corroborate the close relationship between *Caenotropus labyrinthicus* and *C. schizodon* (clade 5) as proposed by Scharcansky and Lucena
(2007). These species share the three character states originally discovered by Vari et al.
(1995) as autapomorphies for *C. labyrinthicus* and later proposed as synapomorphies

for that species and C. schizodon by Scharcansky and Lucena (2007). Caenotropus 367 368 *labyrinthicus* resolved as paraphyletic in our analyses (clade 5) due to a single individual of that species from the western Amazon clustering with two individuals of 369 C. schizodon of the eastern Amazon (clade 6). This relationship received only moderate 370 statistical support (-71/63), and a substantial fraction (22.4%) of the trees in the 371 posterior distribution include a monophyletic C. labyrinthicus. The apparent paraphyly 372 of that species in the best-supported tree may represent an artifact of locus selection or 373 374 model choice, or gene-tree conflict resulting from incomplete lineage sorting. Alternatively, it could reflect the presence of multiple species within the present concept 375 of C. labyrinthicus, or indicate that C. labyrinthicus and C. schizodon are morphotypes 376 of the same species. The samples of C. schizodon used herein originated in the Rio 377 Tapajós, a drainage which includes the type locality of that species. However, several 378 379 individuals of C. labyrinthicus with bifid premaxillary teeth, the purported distinguishing feature of *C. schizodon*, have been found in other portions of the Amazon 380 381 basin including the Rio Madeira of the western Amazon (J. Zuanon, pers. com.) and Rio 382 Xingu of the Brazilian Shield (observed in this study). Studies of other populations of C. labyrinthicus and C. schizodon across the range of those species within the Amazon 383 and Tocantins basins are required to evaluate the distribution and intrapopulational 384 385 consistency of the dentition characters that putatively separate these species, and to further evaluate their reciprocal monophyly. 386

As a final unanswered question within *Caenotropus*, we note that the known distribution of *Caenotropus mestomorgmatos* is pronouncedly disjunct. One population is distributed in blackwater portions of the southern Río Orinoco of Venezuela and the adjoining upper reaches of the Rio Negro of Brazil (Vari et al., 1995), and another in the blackwater Río Nanay of northeastern Peru (Vari and Ortega, 1997). All of our

- sequenced samples are from the Peruvian population. In a future study, it would be of
- 393 great interest to determine whether these two populations have diverged enough
- 394 genetically to merit possible recognition as separate species.
- 395
- 396 *4.4. Phylogenetic relationships and comments on the taxonomy of Chilodus*

Chilodus is a well known genus among aquarists and includes four currently 397 valid species (Vari and Ortega, 1997) and we herein present the first phylogenetic 398 399 reconstruction for the genus. Within the monophyletic *Chilodus*, C. gracilis appears as the sister group of a major clade (9) including the remaining three species. 400 Relationships within clade 9 indicate that the present concept of C. punctatus is 401 paraphyletic unless C. fritillus and C. zunevei are included within C. punctatus. Within 402 this broadly defined *C. punctatus* species-complex (clade 9), we found two distinct 403 genetic lineages. The first is restricted to the Rio Araguaia, a lowland river draining into 404 the Rio Tocantins system to the southeast of the Amazon basin and which represents an 405 ecoregion with one of the highest degrees of ichthyological endemism in the Neotropics 406 407 (Albert et al., 2011). Our phylogeny shows C. punctatus from the Rio Araguaia as the sister lineage of the remaining specimens which are distributed throughout the Amazon 408 409 basin (Fig. 3). This arrangement is congruent to the area cladogram for Neotropical 410 fishes constructed by Albert and Carvalho (2011) under which a clade composed by the 411 Tocantins, Araguaia and Xingu basins is sister to a clade composed by remaining 412 drainages within the Amazon.

Although Isbrücker and Nijssen (1988) did not report specimens of *Chilodus*from the Rio Araguaia basin, Lowe-McConnell (1991) cited *Chilodus* sp. from a
tributary of the Rio das Mortes in the Araguaia drainage. Later, Vari and Ortega (1997)

described the occurrence of *C. punctatus* from many Amazonian tributaries including
several in the Rio Araguaia basin and Lucinda et al. (2007) reported *C. punctatus* in the
middle Rio Tocantins. Additional and more detailed taxonomic studies are required to
evaluate whether this lineage of *C. punctatus* from the Rio Araguaia basin merits
recognition as a distinct species.

421 The second clade within the *Chilodus punctatus* species-complex (clade 10) is 422 distributed throughout the Amazon and Guianas. For the most part, our analysis failed to resolve relationships within this clade, but it did group *Chilodus punctatus* from the 423 Rio Juruá and Río Nanay with C. fritillus from the Río Madre de Dios in Peru, a 424 tributary of the Rio Mamoré-Madeira (Figs. 2-3). These river basins are included in the 425 426 major ecoregion of the western Amazon and Mamoré-Madre de Dios as detailed by Albert et al. (2011). The close genetic similarity between C. fritillus and these western 427 C. punctatus (particularly those from the Río Nanay) is striking given the pronounced 428 separation of these regions in linear distance, but much more so in terms of distances 429 along rivers. It further suggests that the intense spotting and the absence of a midlateral 430 stripe that was used to diagnose C. fritillus (Vari and Ortega, 1997) may be a function 431 of pronounced regional variation within a chromatically plastic species. Resolution of 432 this question necessitates in-depth analysis of multiple population samples of the genus 433 from the western Amazon. 434

The remaining individuals of *C. punctatus* within clade 10 are distributed throughout the Amazon drainage, including the Takutu-Branco system in the northeast of the basin, the Rio Guamá in the east of the system and the Rio Purus in the western Amazon. Notably, the clade also includes a specimen of *C. zunevei* from the independent Atlantic-draining Commewijne River in Suriname northeast of the Amazon basin (Figs. 2-3).

If *Chilodus punctatus* is eventually split into multiple species, the lineage 441 442 marked herein as Clade 10 will be the most likely to bear the original name. The holotype of C. punctatus was described from Lake Amuku within the Rupununi 443 Savannas of the upper Essequibo River system (Müller and Troschel, 1844). This 444 locality lies about 50 linear km from the Takutu River in the upper Rio Branco 445 drainage, where two samples of *C. punctatus* used in this analysis originated. During 446 high water periods, the Rupununi Savannas (the so called Rupununi portal) connect the 447 448 Essequibo River basin (via the Rupununi River and the Takutu River) with the Rio Branco, the major tributary of the Rio Negro which is, in turn, the largest northern 449 tributary of the mainstream Amazon. This interconnection allows ichthyofaunal 450 exchange between these otherwise separate Essequibo and Amazon biogeographic 451 provinces (Hubert and Renno, 2006; Souza et al., 2012). Considering that C. punctatus 452 453 has been reported from both the Essequibo and Takutu rivers (Sidlauskas and Vari, 2012; Souza et al., 2012), it is likely that our analyzed specimens from the Takutu River 454 are genetically similar to those living in the type-locality within the upper Essequibo 455 456 basin.

The taxonomic history of *Chilodus zunevei* is complex and the results of this 457 study further contribute to the uncertainty. Chilodus zunevei was originally described as 458 a distinct species by Puyo (1945). That nominal form was soon thereafter reduced to the 459 460 subspecies level by Géry (1964) who soon thereafter synonymized it into C. punctatus 461 (Géry, 1977) only to have the species later resurrected by Isbrücker and Nijssen (1988) 462 (see discussion in Vari and Ortega, 1997). The single available specimen of C. zunevei appears as nested within C. punctatus and potentially related to specimens from near the 463 type locality of the latter species. Although, it seems possible that C. zunevei will again 464 465 become a junior synonym of C. punctatus in an eventual taxonomic revision of

466 *Chilodus*, we prefer to not make a formal taxonomic change on the basis of a single467 sequenced individual.

Overall, our results suggest that substantial unrecognized diversity exists within 468 the Chilodontidae and that the alpha-taxonomy of both *Caenotropus* and *Chilodus*. 469 merits revision. Such efforts should include broader geographic sampling than was 470 471 possible in the present contribution, as well as renewed attention to the patterns of color variation, squamation and tooth structure that served to diagnose the present species 472 limits within the family. Given the increased ease of integration of comprehensive 473 morphological and molecular datasets afforded by recent and ongoing ichthyological 474 collecting programs and new collaborations, it seems likely that such continent-wide 475 476 studies will soon reveal greater biodiversity than suspected previously not only among the Chilodontidae but also among other groups of Neotropical freshwater fishes. 477

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- 630
- 631
- 632 Figure Legends
- Fig. 1. Representative specimens of the eight known species in Chilodontidae. (A)

NAT

- 634 Caenotropus maculosus (live), ANSP 189147, Suriname, Litanie River, Marowijne
- 635 River basin; (B) Caenotropus mestomorgmatos (live), MZUSP 92545, Brazil, Rio
- 636 Tiquié, tributary of Rio Negro, Amazon basin; (C) Caenotropus labyrinthicus (live),
- 637 MZUSP 95908, Brazil, Rio Teles Pires, tributary of Rio Tapajós, Amazon basin; (D)
- 638 *Caenotropus schizodon* (preserved), LBP 13847, Brazil, Rio Tapajós, Amazon basin;
- 639 (E) Chilodus gracilis (preserved), LBP 6962, Brazil, Rio Negro, Amazon basin; (F)
- 640 *Chilodus punctatus* (preserved), AUM 36902, Guyana, Rupununi River, Essequibo
- 641 River basin; (G) Chilodus fritillus (live), AUM 51355, Rio Madre de Diós, a tributary
- 642 of Mamoré-Madeira system, Amazon basin; (H) Chilodus zunevei (preserved), MHNG
- 643 2608.040, French Guiana, Kaw River basin. Scale bars indicate one centimeter. Photos
- by M. Sabaj Pérez (A), F.C.T. Lima (B), J.L.O. Birindelli (C), B.F. Melo (D, E), J.
- 645 Armbruster (F), N.K. Lujan (G) and B.L. Sidlauskas (H).

646

Fig. 2. Distribution map of sampled specimens of Chilodontidae. Outgroups are notshown.

649

Fig. 3. Relationships among species of Chilodontidae and others taxa of the 650 Anostomoidea obtained by a ML partitioned analyses of the concatenated dataset. A 651 series of three numbers (e.g. 1/100/100) at each of the main nodes shows the posterior 652 653 probability for that split obtained in BI, percentage of bootstrap support obtained by ML, and percentage of bootstrap obtained by MP analysis, respectively (1000 bootstrap 654 pseudoreplicates). Dashes show values lower than 0.9 (BI) or 50% (ML, MP) and nodes 655 not supported by values higher than 0.9 (BI) or 50% (ML, MP) were collapsed. 656 Asterisks represent nodes that were not obtained by BI or MP analyses. Clades with 657

dark numbered ovals are discussed in the text.

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Additional file 1. Comparison of log likelihoods, AIC and BIC values among different partitioning schemes (from 1 to 13 partitions). For each type of analysis, the following results are shown: total number of parameters; log likelihood calculated using RAxML (L_{ML}) ; AIC values; the difference in AIC values between model i and the best model (Δi = AIC*i* – AICmin); BIC_{ML} values.

666

- 667 Additional file 2. Maximum likelihood phylogeny reconstructed in RAxML using the
- 668 interfamilial hypothesis of Vari (1983) as a topological constraint. All other settings and
- 669 parameters were identical to those used in the unconstrained analysis (Fig. 3).

670

- Additional file 3. Maximum likelihood phylogeny reconstructed in RAxML using the
- 672 intrageneric hypothesis for *Caenotropus* of Vari et al. (1995), as modified by
- 673 Scharcansky and Lucena (2007) as a topological constraint. All other settings and

MA

parameters were identical to those used in the unconstrained analysis (Fig. 3).

675



- 1 Table 1. Genus and species, voucher specimens, locality information and GenBank accession number of chilodontids and outgroup taxa used in
- 2 this study. Asterisks show sequences obtained by Oliveira et al. (2011).

C

Species	Voucher	Specimen	Locality	Coordinates	City, State	Country	16s	COI	Cytb	Myh6	Rag1
Caenotropus labyrinthicus	LBP 1828	12912	Rio Araguaia, Amazon basin	15°53'35.6"8 / 52°15'01"W	Aragarças, Goiás	Brazil	-	EU185613*	HQ289538*	-	HQ289154*
Caenotropus labyrinthicus	LBP 9216	43161	Río Apure, Orinoco basin	07°37'24.4"N / 66°24'48"W	Cabruta, Guárico	Venezuela	HQ171428*		-	HQ289136*	HQ289327*
Caenotropus labyrinthicus	LBP 3050	19138	Río Orinoco, Orinoco basin	07°38'11.6''N / 66°19'4.2''W	Caicara del Orinoco, Bolívar	Venezuela	KF562379		KF562437	KF562462	KF562485
Caenotropus labyrinthicus	OS 18770	PE10-82	Río Nanay, Amazon basin	3.751667 S / 73.287222 W	Iquitos, Loreto	Peru	KF562380	KF562408	KF562438	KF562463	KF562486
Caenotropus maculosus	MHNG 2705.038	157-13	Sipaliwini, Corantijn basin	04°38'48.3"N /57°12'53"W	Sipaliwini	Suriname	KF562381	KF562409	KF562439	KF562464	KF562487
Caenotropus maculosus	MHNG 2717.052	157-15	Tapanahony river, Marowijne basin	04°15'0"N / 54°31'33.2"W	Sipaliwini	Suriname	KF562382	KF562410	KF562440	KF562465	KF562488
Caenotropus maculosus	ANSP 189156	6895	Marowijne river basin	3° 17'24" N / 54° 4'38" W	Sipaliwini	Suriname	KF562383	KF562411	KF562441	-	KF562489
Caenotropus mestomorgmatos	ANSP 180516	PEL02- T48	Río Nanay, Amazon basin	3° 46' 45" S / 73° 22' 6" W	Iquitos, Loreto	Peru	KF562384	KF562412	KF562442	KF562466	KF562490
Caenotropus mestomorgmatos	OS 18346	PE10-67	Río Nanay, Amazon basin	3.751667 S / 73.31625 W	Iquitos, Loreto	Peru	KF562385	KF562413	KF562443	KF562467	KF562491
Caenotropus mestomorgmatos	OS 18772	PE10-93	Río Nanay, Amazon basin	3.751667 S / 73.287222 W	Iquitos, Loreto	Peru	KF562386	KF562414	KF562444	KF562468	KF562492
Caenotropus mestomorgmatos	OS 18323	PE10-139	Río Nanay, Amazon basin	3.780972 S / 73.363889W	Iquitos, Loreto	Peru	KF562387	KF562415	KF562445	-	-
Caenotropus mestomorgmatos	OS 18323	PE10-140	Río Nanay, Amazon basin	3.780972 S / 73.363889W	Iquitos, Loreto	Peru	KF562388	KF562416	KF562446	-	-
Caenotropus schizodon	LBP 13847	57304	Rio Tapajós, Amazon basin	04°16'49."S / 59°59'26.1"W	Itaituba, Pará	Brazil	KF562389	-	KF562447	-	-

Río Madre de Diós, Mamoré- Madeira system, Amazon basin Rio Negro, Amazon basin Rio Purus, Amazon basin Rio Purus, Amazon basin Rio Juruá, Amazon basin Rio Purus, Amazon basin Rio Guamá, Amazon basin Rio Araguaia, Amazon basin Rio Araguaia, Amazon basin	12.27713 S / 69.15237 W 00°00'32.1"N / 66°55'35.7"W 00°16'25.9"N / 66°38'36.5"W 00°16'25.9"N / 66°38'36.5"W 10°04'44.3" S / 67°32'33.9"W 10°04'44.3" S / 67°32'33.9"W 07°56'11.0"S / 63°27'35.3"W 01°34'00"S" / 47°09'51.4"W 15°32'25.8"S / 52°26'18.7"W	Madre de Dios São Gabriel da Cachoeira, Amazonas São Gabriel da Cachoeira, Amazonas São Gabriel da Cachoeira, Amazonas São Gabriel da Cachoeira, Amazonas Rio Branco, Acre Rio Branco, Acre Mâncio Lima, Acre Lábrea, Amazonas Ourém, Pará Barra do Garças, Mato Grosso	Peru Brazil Brazil Brazil Brazil Brazil Brazil Brazil Brazil Brazil Brazil	KF562391 KF562392 KF562393 KF562394 KF562395 KF562396 KF585008 HQ171309* KF562398 KF562399	KF562418 KF562419 KF562420 KF562421 KF562422 KF562423 KF562424 - KF562425 KF585011	- KF562449 KF562450 KF562451 KF562452 KF562453 KF585014 HQ289598* KF562455 KF562455	- KF562470 KF562471 KF562472 KF562473 KF562474 KF585017 - KF562475 KF562476	KF562495 KF562496 KF562497 KF562498 KF562498 KF562500 - HQ289211* KF562502
Rio Negro, Amazon basin Rio Negro, Amazon basin Rio Negro, Amazon basin Rio Purus, Amazon basin Rio Purus, Amazon basin Rio Juruá, Amazon basin Rio Purus, Amazon basin Rio Purus, Amazon basin Rio Guamá, Amazon basin Rio Guamá, Amazon basin Rio Araguaia, Amazon basin	00°00'32.1"N / 66°55'35.7"W 00°16'25.9"N / 66°38'36.5"W 00°16'25.9"N / 66°38'36.5"W 10°04'44.3" S / 67°32'33.9"W 10°04'44.3" S / 67°32'33.9"W 07°34'28.8" S / 72°55'24.9"W 07°56'11.0"S / 63°27'35.3"W 01°34'00"S" / 47°09'51.4"W	Cachoeira, Amazonas São Gabriel da Cachoeira, Amazonas São Gabriel da Cachoeira, Amazonas São Gabriel da Cachoeira, Amazonas Rio Branco, Acre Rio Branco, Acre Mâncio Lima, Acre Lábrea, Amazonas Ourém, Pará Barra do Garças, Mato	Brazil Brazil Brazil Brazil Brazil Brazil Brazil Brazil Brazil	KF562393 KF562394 KF562395 KF562396 KF585008 HQ171309* KF562398	KF562420 KF562421 KF562422 KF562423 KF562424 - KF562425	KF562450 KF562451 KF562452 KF562453 KF585014 HQ289598* KF562455	KF562471 KF562472 KF562473 KF562474 KF585017 - KF562475	KF562497 KF562498 KF562499 KF562500 - HQ289211* KF562502
Rio Negro, Amazon basin Rio Negro, Amazon basin Rio Purus, Amazon basin Rio Purus, Amazon basin Rio Juruá, Amazon basin Rio Purus, Amazon basin Rio Purus, Amazon basin Rio Guamá, Amazon basin Rio Araguaia, Amazon basin	00°16'25.9"N / 66°38'36.5"W 00°16'25.9"N / 66°38'36.5"W 10°04'44.3" S / 67°32'33.9"W 10°04'44.3" S / 67°32'33.9"W 07°34'28.8" S / 72°55'24.9"W 07°56'11.0"S / 63°27'35.3"W 01°34'00"S" / 47°09'51.4"W	Cachoeira, Amazonas São Gabriel da Cachoeira, Amazonas São Gabriel da Cachoeira, Amazonas Rio Branco, Acre Rio Branco, Acre Mâncio Lima, Acre Lábrea, Amazonas Ourém, Pará Barra do Garças, Mato	Brazil Brazil Brazil Brazil Brazil Brazil Brazil	KF562394 KF562395 KF562396 KF585008 HQ171309* KF562398	KF562421 KF562422 KF562423 KF562424 - KF562425	KF562451 KF562452 KF562453 KF585014 HQ289598* KF562455	KF562472 KF562473 KF562474 KF585017 - KF562475	KF562498 KF562499 KF562500 - HQ289211 [#] KF562502
Rio Negro, Amazon basin Rio Purus, Amazon basin Rio Purus, Amazon basin Rio Juruá, Amazon basin Rio Purus, Amazon basin Rio Guamá, Amazon basin Rio Araguaia, Amazon basin	00°16'25.9"N / 66°38'36.5"W 10°04'44.3" S / 67°32'33.9"W 10°04'44.3" S / 67°32'33.9"W 07°34'28.8" S / 72°55'24.9"W 07°56'11.0"S / 63°27'35.3"W 01°34'00"S" / 47°09'51.4"W	Cachoeira, Amazonas São Gabriel da Cachoeira, Amazonas Rio Branco, Acre Rio Branco, Acre Mâncio Lima, Acre Lábrea, Amazonas Ourém, Pará Barra do Garças, Mato	Brazil Brazil Brazil Brazil Brazil Brazil	KF562395 KF562396 KF585008 HQ171309* KF562398	KF562422 KF562423 KF562424 - KF562425	KF562452 KF562453 KF585014 HQ289598* KF562455	KF562473 KF562474 KF585017 - KF562475	KF562499 KF562500 - HQ289211 ³ KF562502
Rio Purus, Amazon basin Rio Purus, Amazon basin Rio Juruá, Amazon basin Rio Purus, Amazon basin Rio Guamá, Amazon basin Rio Araguaia, Amazon basin	10°04'44.3" S / 67°32'33.9"W 10°04'44.3" S / 67°32'33.9"W 07°34'28.8" S / 72°55'24.9"W 07°56'11.0"S / 63°27'35.3"W 01°34'00"S" / 47°09'51.4"W	Cachoeira, Amazonas Rio Branco, Acre Rio Branco, Acre Mâncio Lima, Acre Lábrea, Amazonas Ourém, Pará Barra do Garças, Mato	Brazil Brazil Brazil Brazil Brazil	KF562396 KF585008 HQ171309* KF562398	KF562423 KF562424 - KF562425	KF562453 KF585014 HQ289598* KF562455	KF562474 KF585017 - KF562475	KF562500 - HQ289211* KF562502
Rio Purus, Amazon basin Rio Juruá, Amazon basin Rio Purus, Amazon basin Rio Guamá, Amazon basin Rio Araguaia, Amazon basin	10°04'44.3" S / 67°32'33.9"W 07°34'28.8" S / 72°55'24.9"W 07°56'11.0"S / 63°27'35.3"W 01°34'00"S" / 47°09'51.4"W	Rio Branco, Acre Mâncio Lima, Acre Lábrea, Amazonas Ourém, Pará Barra do Garças, Mato	Brazil Brazil Brazil Brazil	KF585008 HQ171309* KF562398	KF562424 - KF562425	KF585014 HQ289598* KF562455	KF585017 - KF562475	- HQ289211* KF562502
Rio Juruá, Amazon basin Rio Purus, Amazon basin Rio Guamá, Amazon basin Rio Araguaia, Amazon basin	07°34'28.8" S / 72°55'24.9"W 07°56'11.0"S / 63°27'35.3"W 01°34'00"S" / 47°09'51.4"W	Mâncio Lima, Acre Lábrea, Amazonas Ourém, Pará Barra do Garças, Mato	Brazil Brazil Brazil	HQ171309* KF562398	- KF562425	HQ289598* KF562455	- KF562475	KF562502
Rio Purus, Amazon basin Rio Guamá, Amazon basin Rio Araguaia, Amazon basin	07°56'11.0''S / 63°27'35.3''W 01°34'00''S'' / 47°09'51.4''W	Lábrea, Amazonas Ourém, Pará Barra do Garças, Mato	Brazil Brazil	KF562398		KF562455		KF562502
Rio Guamá, Amazon basin Rio Araguaia, Amazon basin	01°34'00''S'' / 47°09'51.4''W	Ourém, Pará Barra do Garças, Mato	Brazil					
Rio Araguaia, Amazon basin		Barra do Garças, Mato		KF562399	KF585011	KF562456	KF562476	
	15°32'25.8"8 / 52°26'18.7"W		Brazil					KF562503
Rio Nanay Amazon hasin			Diuzn	KF562400	KF562426	KF562457	KF562477	KF562504
itto i tunuy, i muzon bushi	3° 52' 21" S / 73° 32' 43" W	Iquitos, Loreto	Peru	KF562401	KF562427	-	-	KF562505
Rio Nanay, Amazon basin	3.751667 S / 73.287222 W	Iquitos, Loreto	Peru	KF562402	KF562428	-	KF562479	KF562506
Rio Nanay, Amazon basin	3.751667 S / 73.287222 W	Iquitos, Loreto	Peru	KF562403	KF562429	KF562458	-	KF562507
Rio Nanay, Amazon basin	3.780972 S / 73.363889W	Iquitos, Loreto	Peru	KF562404	KF562430	-	-	-
Rio Takutu, Rio Branco, Amazon basin	03°22'55.9"N / 59°51'28.3"W	Bonfim, Roraima	Brazil	KF562405	-	KF562459	KF562482	KF562508
Rio Takutu, Rio Branco, Amazon basin	03°22'55.9"N / 59°51'28.3"W	Bonfim, Roraima	Brazil	KF562406	KF562431	KF562460	KF562483	KF562509
Commewijne river basin	05°23'47.50"N / 54°44'9.17"W	Para	Suriname	KF585009	KF585012	KF585015	KF585018	KF585020
Rio Branco, Amazon basin	02°18'02.0"N / 60°55'20.7"W	Mucajaí, Roraima	Brazil	HQ171317*	-	HQ289606*	HQ289026*	HQ289219*
Rio Tapajós, Amazon basin	08° 11'4.0" S / 55°10'47.0"W	Novo Progresso, Pará	Brazil	KF585010	KF585013	KF585016	KF585019	KF585021
	Rio Takutu, Rio Branco, Amazon basin Rio Takutu, Rio Branco, Amazon basin Commewijne river basin Rio Branco, Amazon basin	Rio Takutu, Rio Branco, Amazon basin 03°22'55.9"N / 59°51'28.3"W Rio Takutu, Rio Branco, Amazon basin 03°22'55.9"N / 59°51'28.3"W Commewijne river basin 05°23'47.50"N / 54°44'9.17"W Rio Branco, Amazon basin 02°18'02.0"N / 60°55'20.7"W Rio Tapajós, Amazon basin 08° 11'4.0" S / 55°10'47.0"W	Rio Takutu, Rio Branco, Amazon basin03°22'55.9''N / 59°51'28.3''WBonfim, RoraimaRio Takutu, Rio Branco, Amazon basin03°22'55.9''N / 59°51'28.3''WBonfim, RoraimaCommewijne river basin03°22'55.9''N / 59°51'28.3''WBonfim, RoraimaCommewijne river basin05°23'47.50''N / 54°44'9.17''WParaRio Branco, Amazon basin02°18'02.0''N / 60°55'20.7''WMucajaí, RoraimaRio Tapajós, Amazon basin08° 11'4.0'' S / 55°10'47.0''WNovo Progresso, Pará	Rio Takutu, Rio Branco, Amazon basin 03°22'55.9"N / 59°51'28.3"W Bonfim, Roraima Brazil Rio Takutu, Rio Branco, Amazon basin 03°22'55.9"N / 59°51'28.3"W Bonfim, Roraima Brazil Commewijne river basin 03°22'55.9"N / 59°51'28.3"W Bonfim, Roraima Brazil Commewijne river basin 05°23'47.50"N / 54°44'9.17"W Para Suriname Rio Branco, Amazon basin 02°18'02.0"N / 60°55'20.7"W Mucajaí, Roraima Brazil Rio Tapajós, Amazon basin 08° 11'4.0" S / 55°10'47.0"W Novo Progresso, Pará Brazil	Rio Takutu, Rio Branco, Amazon basin03°22'55.9"N / 59°51'28.3"WBonfim, RoraimaBrazilKF562405Rio Takutu, Rio Branco, Amazon basin03°22'55.9"N / 59°51'28.3"WBonfim, RoraimaBrazilKF562405Commewijne river basin03°22'55.9"N / 59°51'28.3"WBonfim, RoraimaBrazilKF562406Commewijne river basin05°23'47.50"N / 54°44'9.17"WParaSurinameKF585009Rio Branco, Amazon basin02°18'02.0"N / 60°55'20.7"WMucajaí, RoraimaBrazilHQ171317*Rio Tapajós, Amazon basin08° 11'4.0" S / 55°10'47.0"WNovo Progresso, ParáBrazilKF585010	Rio Takutu, Rio Branco, Amazon basin03°22'55.9"N / 59°51'28.3"WBonfim, RoraimaBrazilKF562405-Rio Takutu, Rio Branco, Amazon basin03°22'55.9"N / 59°51'28.3"WBonfim, RoraimaBrazilKF562406KF562431Commewijne river basin05°23'47.50"N / 59°51'28.3"WBonfim, RoraimaBrazilKF562406KF562431Commewijne river basin05°23'47.50"N / 54°44'9.17"WParaSurinameKF585009KF585012Rio Branco, Amazon basin02°18'02.0"N / 60°55'20.7"WMucajaí, RoraimaBrazilHQ171317*-Rio Tapajós, Amazon basin08° 11'4.0" S / 55°10'47.0"WNovo Progresso, ParáBrazilKF585010KF585013	Rio Takutu, Rio Branco, Amazon basin03°22'55.9"N / 59°51'28.3"WBonfim, RoraimaBrazilKF562405-KF562459Rio Takutu, Rio Branco, Amazon basin03°22'55.9"N / 59°51'28.3"WBonfim, RoraimaBrazilKF562406KF562431KF562460Commewijne river basin05°23'47.50"N / 54°44'9.17"WParaSurinameKF585009KF585012KF585015Rio Branco, Amazon basin02°18'02.0"N / 60°55'20.7"WMucajaí, RoraimaBrazilHQ171317*-HQ289606*Rio Tapajós, Amazon basin08° 11'4.0" S / 55°10'47.0"WNovo Progresso, ParáBrazilKF585010KF585013KF585016	Rio Takutu, Rio Branco, Amazon basin03°22'55.9"N / 59°51'28.3"WBonfim, RoraimaBrazilKF562405-KF562459KF562482Rio Takutu, Rio Branco, Amazon basin03°22'55.9"N / 59°51'28.3"WBonfim, RoraimaBrazilKF562406KF562431KF562460KF562483Commewijne river basin05°23'47.50"N / 54°44'9.17"WParaSurinameKF585009KF585012KF585015KF585018Rio Branco, Amazon basin02°18'02.0"N / 60°55'20.7"WMucajaí, RoraimaBrazilHQ171317*-HQ289606*HQ289026*Rio Tapajós, Amazon basin08° 11'4.0" S / 55°10'47.0"WNovo Progresso, ParáBrazilKF585010KF585013KF585016KF585019

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Leporinus altipínnis	LBP 4459	24381	Rio Negro, Amazon basin	00°40'03.1"S / 62°58'23.5"W	Barcelos, Amazonas	Brazil	HQ171321*	-	HQ289610*	HQ289030*	HQ289223*
Schizodon scotorhabdotus	LBP 3046	19130	Río Orinoco, Orinoco basin	07°38'11.6"N / 66°19'04.2"W	Caicara del Orinoco, Bolívar	Venezuela	HQ171270*	KF562432	HQ289559*	HQ288980*	HQ289177*
Schizodon vittatus	LBP 3994	23098	Rio Araguaia, Amazon baisn	11°40'9''S / 50°51'0.30''W	São Félix do Araguaia, Mato Grosso	Brazil	HQ171308*	-	HQ289597*	HQ289018*	HQ289210*
Curimatella dorsalis	LBP 3759	22034	Rio Paraguay, La Plata basin	19°34'33.7"S / 56°14'49.5"W	Aquidauana, Mato Grosso do Sul	Brazil	HQ171290*	KF562433	HQ289579*	HQ289000*	HQ289194*
Cyphocharax aspilos	LBP 6109	29560	Lago Maracaibo	09°38'53.8"N / 72°34'56.4"W	Machiques de Perijá, Zulia	Venezuela	HQ171363*		HQ289650*	HQ289071*	HQ289264*
Cyphocharax gouldingi	LBP 1537	11889	Rio Araguaia, Amazon basin	15°53'53.4"S / 52°13'00.6"W	Aragarças, Goiás	Brazil	HQ171243*	KF562434	HQ289534*	HQ288953*	HQ289150*
Potamorhina altamazonica	LBP 2571	17020	Rio Purus, Amazon basin	08°51'21.5"S / 68°42'22.6"W	Boca do Acre, Amazonas	Brazil	HQ171261*	-	HQ289552*	HQ288971*	HQ289168*
Steindachnerina brevipinna	LBP 5185	26336	Rio Paraná, La Plata basin	22°47'29''S / 53°20'58''W	Porto Rico, Paraná	Brazil	HQ171339*	-	HQ289628*	HQ289048*	HQ289241*
Hemiodus immaculatus	LBP 1725	12849	Rio Negro, Amazon basin	02°03'10.0"S / 60°06'31.7"W	Manaus, Amazonas	Brazil	HQ171246*	-	HQ289537*	HQ288956*	HQ289153*
Prochilodus reticulatus	LBP 6127	29514	Rio Catacumbo, Lago Maracaibo	09°05'08.3"N / 72°13'50.5"W	Catacumbo, Zulia	Venezuela	HQ171358*	KF562435	HQ289647*	HQ289067*	HQ289260*
Semaprochilodus laticeps	LBP 1383	12728	Río Orinoco, Orinoco basin	02°03'10.0"S / 60°06'31.7"W	Caicara del Orinoco, Bolívar	Venezuela	HQ171245*	KF562436	HQ289536*	HQ288955*	HQ289152*
3					SV.						
4											
			C								

5 Table 2. Information content and characteristics of each gene.

16S 608 COI 633 Cytb 985	1 PCR 1 PCR	16Sa-L – ACGCCTGTTTATCAAAAACAT 16Sb-H – CCGGTCTGAACTCAGATCACGT	Palumbi (1996)	0.308	0.242		
	1 PCR				0.242	0.221	0.228
Cytb 985		L6252-Asn – AAGCCGGGGGAAAGCCCCGGCAG H7271-COXI – TCCTATGTAGCCGAATGGTTCTTTT	Melo et al. (2011)	0.245	0.258	0.181	0.314
	1 PCR	LNF – GACTTGAAAAACCAYCGTTGT H08R2 – GCTTTGGGAGTTAGDGGTGGGAGTTAGAATC	Oliveira et al. (2011)	0.276	0.297	0.140	0.285
Myh6 704	1st PCR	F329 – CCGCMTGGATGATCTACAC A3R1 – ATTCTCACCACCATCCAGTTGAA	Li et al. (2007)	0.306	0.221	0.249	0.241
	2nd PCR	A3F2 – GGAGAATCARTCKGTGCTCATCA A3R2 – CTCACCACCATCCAGTTGAACAT		0.500	0.221	0.219	0.211
Rag1 1210	1st PCR	Rag1CF1 – ACCCTCCGTACTGCTGAGAA Rag1CR1 – CGTCGGAAGAGCTTGTTGCC		0.257	0.220	0.201	0.000
	2nd PCR	Rag1CF2 – TACCGCTGAGAAGGAGCTTC Rag1CR2 - TGTTGCCAGACTCATTGCCCTC	Oliveira et al. (2011)	0.257	0.238	0.281	0.222

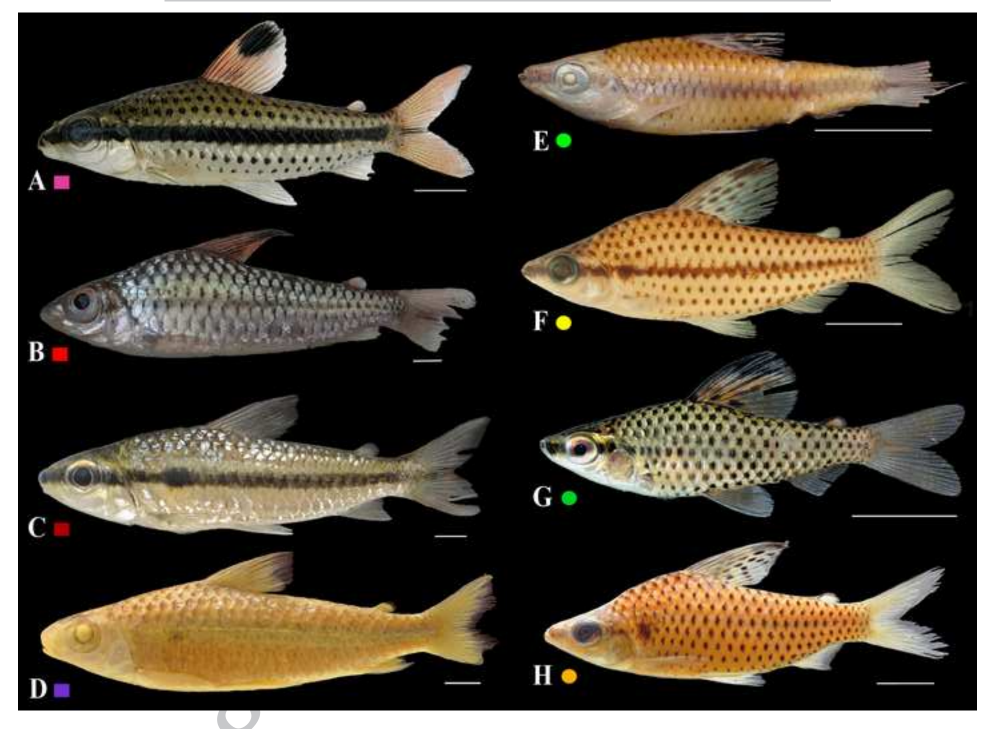
Gene and position	Partitions	Best-fit model	
16S	1 - 608	GTR+G+I	_
COI 1 st position	609 - 1241/3	TrN+G	_
COI 2 nd position	610 - 1241/3	F81	-
COI 3 rd position	611 – 1241/3	GTR+G	_
Cytb 1 st position	1242 - 2226/3	TrN+I	- 0-
Cytb 2 nd position	1243 - 2226/3	SYM+I+G	_
Cytb 3 rd position	1244 - 2226/3	TrN+I	-
Myh6 1 st position	2227 - 2930/3	GTR+G+I	- 6
Myh6 2 nd position	2228 - 2930/3	GTR+I	_
Myh6 3 rd position	2229 - 2930/3	HKY+G	-
Rag1 1 st position	2931 - 4140/3	TVM+I+G	-
Rag1 2 nd position	2932 - 4140/3	TVM+G	-
Rag1 3 rd position	2933 - 4140/3	TVM+I+G	-
			-

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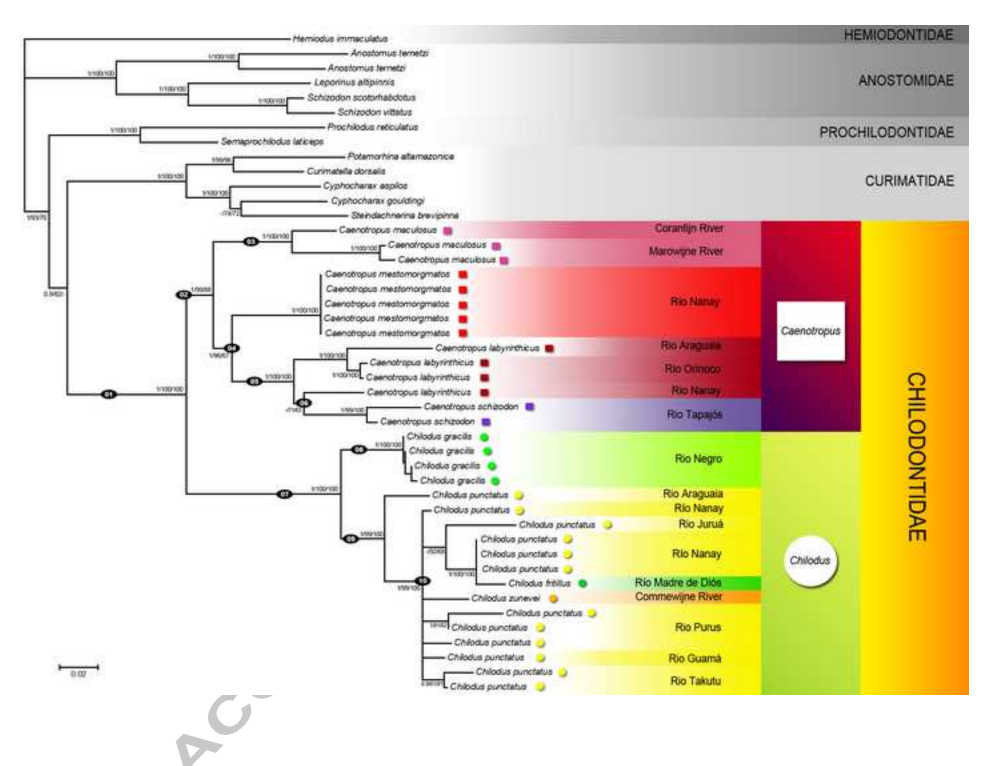
7 Table 3. Gene partitions and their models as selected by ModelTest.

Table 4. Results of Shimodaira-Hasegawa tests of alternative topologies with and without optimization of base frequencies and rate matrices in Phangorn. P-values lower than 0.05 would indicate statistical rejection of equivalence of the topologies.

	No optimiza			Optimized		
Constraint	ln L	∂L	p-value	ln L	∂L	p-value
None	-13134.40	0.00	0.7224	-12047.94	0.00	0.7317
Caenotropus (Vari et al., 1995)	-13136.61	2.21	0.4502	-12048.90	0.97	0.5345
Outgroups (Vari, 1983)	-13141.15	6.76	0.1506	-12050.72	2.78	0.3413









Highlights

The first molecular phylogeny of Chilodontidae based on all recognized species. Chilodontidae is monophyletic as well as its two genera, *Caenotropus* and *Chilodus*. *Chilodus punctatus* is paraphyletic with *C. fritillus* and *C. zunevei* nested within it. We discuss the morphological convergences and the possibility of cryptic species.

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