1	The Araguaia river is an important biogeographical divide for
2	didelphid marsupials in central Brazil
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22 **Running title:** Araguaia river as a dispersal barrier for marsupials

1 Abstract

2 The Riverine Barrier model suggests that rivers may play a significant role in separating 3 widespread organisms into isolated populations. In this study we used a comparative 4 approach to investigate the phylogeography of six didelphid marsupial species in central 5 Brazil. Specifically, we evaluate the role of the mid-Araguaia River in differentiating 6 populations and estimate divergence time among lineages to assess the timing of 7 differentiation of these species, using mitochondrial DNA sequence data. The six 8 didelphid marsupials revealed different intraspecific genetic patterns and structure. The 9 three larger and more generalist species, Didelphis albiventris, Didelphis marsupialis 10 and *Philander opossum*, showed connectivity across the Araguaia River. In contrast the 11 genetic structure of the three smaller and specialist species, Gracilinanus agilis, 12 Marmosa (Marmosa) murina and, Marmosa (Micoureus) demerarae was shaped by the 13 mid-Araguaia. Moreover, the split of eastern and western bank populations of the two 14 latter species is consistent with the age of Araguaia River sediments formation. We 15 hypothesize that the role of the Araguaia as a riverine barrier is linked to the level of 16 ecological specialization among the six didelphid species and differences in their ability 17 to cross rivers or disperse through the associated habitat types.

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19 Keywords Amazonia-Cerrado ecotone, cytochrome *b*, Didelphidae, Gallery Forests,
20 Pleistocene

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1 Introduction

2 The Riverine Barrier model was first proposed by Wallace in 1852, suggesting that 3 rivers may have a significant role in separating widespread organisms into isolated 4 populations (Wallace 1852). Several vertebrate groups were extensively sampled in a 5 pioneer study in the Juruá River in Amazonia aiming to examine the riverine effects on 6 patterns of both population and species differentiation and ecological distribution 7 (Patton et al. 2000). This and several other successive studies that investigated the 8 potential role of Neotropical rivers as putative barriers reached different conclusions 9 (e.g., Ayres and Clutton-Brock 1992; Gascon et al. 2000; Matocq et al. 2000; Bates et 10 al. 2004; Faria et al. 2013a; Maldonado-Coelho et al. 2013). The lack of congruence 11 among species and rivers has been related with the species vagility and also with the 12 drainage system geography and formation history (Moritz et al. 2000; Antonelli et al. 13 2010; Leite and Rogers 2013).

14 The Araguaia River lies in the transitional area between Cerrado and Amazonia. 15 With a length of 2,110 km and reaching 3-6 km of width in its middle portion, it 16 represents the largest river of the wet-dry tropics of Brazil (Latrubesse and Stevaux 17 2002). This river is classified as an anabranching river, consisting of channels separated 18 by vegetated semi-permanent alluvial islands, with low sinuosities (Latrubesse 2008). 19 Like other rivers in the Brazilian Shield it represents a relatively stationary geographical 20 feature, which is expected to limit gene flow between forest-dwelling populations from 21 opposite banks, contrary to meandering rivers in the western region of the Amazon 22 Basin (e.g. Aleixo 2004; Bates et al. 2004; Antonelli et al. 2010; Maldonado-Coelho et 23 al. 2013). Therefore, the Araguaia River provides an excellent natural setting to test 24 biogeographic diversification hypotheses such as the riverine barrier model. Previous 25 studies reached different conclusions, showing that Araguaia might have been a barrier to two *Rhipidomys* rodent sister species (Rocha et al. 2011), but not playing an important role in the intraspecific differentiation of the rodents *Hylaeamys megacephalus* and *Oecomys* aff. *roberti* (Rocha et al. 2014). Thus, it is essential to understand to what extent these findings are consistent across different animal taxa, i.e. in a general biogeographic framework, and why different species are responding differently to the same geographic feature.

7 The family Didelphidae is among the oldest of extant mammal families (Jansa et 8 al. 2013). Species of this diverse group are common in forest and open habitat 9 communities in the Neotropics (Gardner 2008). Currently, 11 species are recognized in 10 the ecotonal area between Cerrado and Amazonia (Lacher and Alho 2001; Bezerra et al. 11 2009; Rocha et al. 2011). In this study we considered the six most common species of 12 didelphid marsupials observed in the mid-Araguaia River basin (Rocha et al. 2011): 13 Didelphis albiventris, Didelphis marsupialis, Philander opossum, Gracilinanus agilis, Marmosa (Marmosa) murina and Marmosa (Micoureus) demerarae (for recent 14 15 taxonomic revision see Voss et al. 2014). Although these six species are widely 16 distributed in lowland Neotropical forests (Gardner 2008), few studies have focused on 17 their phylogeography in central Brazil (exceptions include Costa 2003; Faria et al. 18 2013a, 2013b), and none in the ecotonal area between Cerrado and Amazonia, which 19 remains poorly studied overall (but see Lacher and Alho 2001; Bezerra et al. 2009; 20 Rocha et al. 2011).

Here, mitochondrial DNA (mtDNA) sequence data was used to investigate the role of the Araguaia River in differentiating populations of six didelphid marsupials in central Brazil. Inferring the evolutionary and demographic history of species solely on mitochondrial DNA can be misleading, due to its strictly maternal inheritance (Ballard 2004), but exploratory analyses with this marker are important to provide basic

1 knowledge of putative barriers and geomorphological events (e.g., Patton et al. 2000; 2 Costa 2003). Furthermore, molecular data combined with geographical distribution 3 information have been useful to understand the evolutionary history of species and to 4 explicitly test hypotheses of biogeographical events (e.g., Patton et al. 2000; Costa 5 2003; Antonelli et al. 2010; Nicolas et al. 2011). Additionally, comparative 6 phylogeography, using multiple co-distributed taxa, allows a better understanding of the 7 relationships between landscape formation and biotic diversification (Arbogast and 8 Kenagy 2001).

9 The distribution of haplotypes across the river holds information about the 10 population processes that have shaped the history of each species (e.g., Patton et al. 11 2000; Nicolas et al. 2011). Patton et al. (2000) formulated three different hypotheses for 12 riverine divergence, which included primary diversification, secondary contact and 13 dispersal. The first hypothesis postulates a complete barrier formed by the river to an existing taxon range, resulting in a reciprocally monophyly of sister clades from 14 15 opposite banks. In the secondary contact hypothesis, the river is only a common contact 16 zone, and despite of the observed monophyly, haplotypes from opposite banks are not 17 sister clades. The third case comprises dispersal events from an established population 18 to the opposite riverbank, resulting in paraphyletic relationships. Moreover, low 19 differentiation will be revealed by a certain degree of haplotype sharing across the river 20 (see Patton et al. 2000 for further explanations). To frame the phylogenetic predictions 21 of the river barrier hypothesis we included in the analysis samples of didelphid 22 marsupial species from other geographical areas, including the Atlantic Forest, Pantanal and Amazonia. We also estimated divergence time among lineages to assess the timing 23 24 of differentiation and to assess the potential impact of external events (e.g. 25 geomorphological) on population differentiation (Leite and Rogers 2013). If the mid-

Araguaia River represents a barrier to populations of didelphid species, it is expected 1 2 that the diversification between populations from opposite river banks coincides with 3 the formation of the river during the Pleistocene, roughly 240,000 years ago (Latrubesse 4 and Stevaux 2002; Valente and Latrubesse 2012). This means that for older divergence 5 events, the river is unlikely to be the primary agent of divergence, although it could 6 nonetheless be acting as barrier to gene flow (e.g., Maldonado-Coelho et al. 2013). 7 Finally, we also explored the species demographic history to detect signatures of 8 population expansion and/or bottlenecks. If the mid-Araguaia River was a primary 9 barrier in population divergence, one should not find evidence of population expansion 10 by the river (Moritz et al. 2000, Leite and Rogers 2013). On the other hand, if there is signature of historical population fluctuations, the Araguaia River could be just a 11 secondary barrier as populations could have been isolated by other geographical 12 processes (i.e. refuges; Moritz et al. 2000, Leite and Rogers 2013). 13

14

15 Methods

16 Sample collection

17 This work focused on six didelphid marsupials: D. albiventris (n = 15), D. marsupialis 18 (n = 50), G. agilis (n = 108), M. murina (n = 80), M. demerarae (n = 35) and P. 19 opossum (n = 54) from central Brazil. Samples were collected in several localities in the 20 both margins of the mid-Araguaia River, two islands within this river and also in its 21 tributaries, Javaés and Coco Rivers (Figure 1). Sampling was carried out between June 22 2007 and November 2008, using a standardized trapping protocol to sample small 23 nonvolant mammals in upland and floodplain gallery forests (for detailed sampling at 24 mid-Araguaia see Rocha et al. 2011). Tissue samples of the mid-Araguaia River have been deposited at Universidade Federal do Espírito Santo (UFES), Vitória, Brazil. 25

Additional samples from central Brazil were obtained via museum tissue collection or kindly shared by colleagues (for details on biological material and sequences used, including sampling localities for each species see Table S1 and Gazetteer). The central Brazilian region treated in this work encompasses mainly samples from the ecotonal region between Cerrado and Amazonia, but also samples from Cerrado (Figure 1).

Additionally, we included sequences of didelphid marsupial species from other
geographical areas, including the Atlantic Forest, Pantanal and Amazonia (Figure 1),
available in GenBank: *D. albiventris* (n = 5), *D. marsupialis* (n = 8), *G. agilis* (n = 26), *M. murina* (n = 23), *M. demerarae* (n = 20) and *P. opossum* (n = 6), in order to
investigate the geographical structure and phylogeographic affinities of central Brazil.
All sequences used in this study were deposited in GenBank (Table S1).

12

13 **DNA extraction and amplification**

14 DNA was extracted from liver or ear tissue preserved in ethanol using salt-extraction 15 method (Bruford et al. 1992). Cytochrome b (mt-Cytb) fragments with 801 base pairs 16 (bp) were amplified by polymerase chain reaction using the primers MVZ05 and 17 MVZ16 (Smith and Patton 1993). PCR reactions were performed in a 25 µl total 18 volume, including 2.5 μ l of 10 × PCR buffer, 1.0 μ l of MgCl₂ (50 mM), 0.5 μ l of dNTP 19 mixture (10mM), 0.3 µl of Platinum® Tag DNA polymerase (Invitrogen), 0.3 µl of each 20 primer (10 µM), and 1 µl of DNA template (100 ng/µl). Amplifications were performed with the following PCR profile: initial denaturation at 94 °C (5 min), followed by 39 21 22 cycles with denaturation at 94 °C (30 s), annealing at 48 °C (45 s), polymerization at 72 23 °C (45 s), and a final extension at 72 °C (5 min). Mitochondrial fragments were purified 24 using ExoSap-IT® (USB Corporation) and sequenced using an automatic sequencer 25 ABI 3130-XL (Perkin Elmer, Applied Biosystems, Foster City, California), with the above-listed primers. Sequence alignment was performed using CLUSTALW algorithm
 implemented in MEGA 6 (Tamura et al. 2013). Electropherograms and sequences were
 manually checked and edited in MEGA 6.

4

5 **Phylogeographic relationships**

6 In order to investigate the geographical distribution and phylogeographic affinities of 7 six didelphid marsupials, and specifically to test if the mid-Araguaia River has a 8 significant role in structuring populations of those didelphid species in central Brazil, 9 relationships amongst haplotypes were estimated through Bayesian inference (BI) 10 performed in the MrBayes v.3.1.2 (Ronquist and Huelsenbeck 2003). For this analysis 11 we used only one individual per haplotype representing populations from central Brazil, 12 but also haplotypes from the Atlantic Forest, Pantanal and other regions in the 13 Amazonia were included in posterior analyses. Haplotypes from closely related species 14 were used as outgroups. The best model of nucleotide substitution was selected in 15 MrModeltest (Nylander 2004). Haplotype trees were sampled every 100 of 10^7 16 generations until Markov chain became stationary, i.e. when standard deviation of split 17 frequencies was below 0.01. A 50 % majority rule consensus tree was obtained after 18 "burn-in" of 25 % of the sample points to generate Bayesian posterior probabilities 19 (BPP).

Pairwise sequence distances (%) within and between the main geographic clades
were estimated using the Kimura 2-parameter (K2p) model (Kimura 1980) implemented
in MEGA 6.

23 Median-joining (MJ) networks were constructed for central Brazilian samples 24 only (Table S1), using NETWORK v.4.6 (Bandelt et al. 1999). Only variable nucleotide 25 sites were included in MJ analyses. For the purposes of this analysis, we categorized two areas as comprising all populations from west and east of the Araguaia River.
Haplotypes from each bank of the Araguaia River were identified with different colours
(white – eastern bank, EAR; black – western bank, WAR) to illustrate the riverine
barrier or put in evidence the haplotype sharing across the river. For *D. marsupialis*,
samples from islands of the Araguaia River were also available and we categorized a
third area for this species, identifying samples from those islands with grey colour.

7

8 **Divergence time and gene flow**

9 To estimate the time to most recent common ancestor (t_{MRCA}) we ran a single data set 10 with 87 *mt-Cytb* sequences including haplotypes from all six species from this study 11 (Table S1) and, in order to use multiple calibration points, we included other Didelphidae sequences obtained from GenBank (Caluromys lanatus: U34664; 12 13 Chironectes minimus: AJ628363; Lutreolina crassicaudata: AJ628364; Monodelphis 14 domestica: HQ651773; Thylamys karimii: EF051700). Molecular clock was tested by 15 performing a maximum likelihood (ML) analyses in MEGA 6. The ML values of the 16 obtained topologies were compared with and without molecular clock constraints also 17 using MEGA 6. The presence of strict molecular clock for this dataset was rejected. The 18 t_{MRCA} analyses were performed in BEAST 2.1.3 (Bouckaert et al 2014) using the 19 Bayesian relaxed clock model and allowing the branch length to vary according to an 20 uncorrelated lognormal distribution (Drummond et al. 2006). We used the Yule process 21 speciation model as tree prior and the GTR+I+G evolution model. We followed the 22 phylogeny of Voss & Jansa (2009) to set monophyletic constrains. Four calibration 23 points were used: the diversification of the Didelphidae and the diversification of 24 Didelphimorphia, which were set as minimum and maximum ages (Meredith et al. 25 2011), and two minimum divergence dates within the marsupial radiation, which were

1 set assuming a log-normal distribution with standard deviation of 1.0 (Jansa et al. 2013). 2 The minimum age of the diversification of Didelphidae and Didelphimorphia was the 3 oldest unequivocal fossil belonging to clade at 11.608 Ma for both (Meredith et al. 4 2011). The maximum age of Didelphidae was based on stratigraphic bounding, which 5 encompasses two stratigraphic units without fossil records of this clade at 28.5 Ma (Meredith et al. 2011). The maximum age of Didelphimorphia was based on 6 7 phylogenetic bounding, which encompasses the age of the stem of didelphimorph at 8 65.8 (Peradectes) (Horovitz et al. 2009; Meredith et al. 2011). The two minimum 9 divergence dates within the marsupial clade were the divergence between Didelphis and 10 Philander at 3.3 Ma; and the split between Monodelphis and Marmosa at 12.1 Ma 11 (Jansa et al. 2013). We performed two independent runs of 100,000,000 generations 12 each and sampled every 10,000 generations. Analyses were performed in the CIPRES 13 Science Gateway. Convergence of the MCMC chains was verified in Tracer v.1.6 by 14 checking the effective sample size values (ESS). The resulting trees were combined 15 using LogCombiner and TreeAnotator after a burn-in of 10 %. Consensus trees were 16 visualized in FigTree 1.4.

17 The IM program (Hey and Nielsen 2004) was used to estimate the effective 18 number of migrants per generation (m), time of divergence (t) between populations from 19 either river bank of the Araguaia River, and the effective population size in both 20 populations as well as the ancestral population (qEAR, qWAR and qA, respectively). 21 For these analyses we only used samples from the immediate vicinity of the Araguaia 22 River (see Table S1). For G. agilis we did not perform this analysis since we had only a 23 single sample from the western river bank. For D. marsupialis we categorized three 24 populations (as this species inhabits both the river banks and mid-river islands) and we 25 performed all three pairwise combinations: eastern and western river banks, eastern

1 river bank and islands and western river bank and islands. We used an inheritance scale of 0.25 for the mtDNA and the HKY model of evolution. Substitution rates per 2 3 nucleotide were obtained using BEAST (as mentioned above), and transformed into 4 complete-locus substitutions (Mean = 0.000014, 95% HPD = 0.000003-0.00003). We 5 did at least five preliminary trial runs for each species in order to assess the range of prior distributions prior for theta (varying from 10 to 30), migration (varying from 2 to 6 7 20) and time estimation (varying from 6 to 20) (Hey and Nielsen 2004). We assumed 8 symmetrical gene flow (i.e. m1 = m2) for all species to avoid over-parameterizing the 9 model. We used a burn-in period of 500,000 steps and recorded results every hour. We 10 allowed the IM program to run until the lowest effective sample size (ESS) were at least 11 500 (e.g., Peters et al. 2005). We only reported results from the longest run for all 12 species (D. albiventris: 64,217,790 steps, minimum ESS = 5570; D. marsupialis: 13 10,663,380 steps, minimum ESS = 4548; *P. opossum*: 8,258,956 steps, minimum ESS = 14 5027; M. murina: 32,290,531 steps, minimum ESS = 1388; and M. demerarae: 15 51,902,032 steps, minimum ESS = 14,564). We assumed a generation time of 1 year for 16 all species.

17

18 **Demographic history**

19 Number of haplotypes (h), number of polymorphic sites (S), haplotype (Hd) and 20 nucleotide (π) diversity values for the central Brazilian samples were estimated with 21 DnaSP v.5 (Librado and Rozas 2009).

22 Deviation from neutrality in central Brazil was tested through the following

- 23 statistics based on frequency spectrum of mutations, Tajima's D (Tajima 1989) and
- 24 Ramos-Onsins and Rozas R₂ (Ramos-Onsins and Rozas 2002), and based on haplotype
- 25 frequencies, Fu's Fs (Fu 1997), using DnaSP v.5. Coalescence simulations with 1,000

- 1 replicates were applied to determine the p-value of each statistics. Significant p-values
- $2 \quad (< 0.05)$ were taken as evidence a scenario of demographic expansion.

3	Demographic history of the six didelphid marsupials was further investigated
4	using Bayesian Skylines Plots (BSP) implemented in BEAST 2.1.3. We used
5	substitution rates per nucleotide per million year obtained in the above-mentioned
6	estimates of the divergence time (Mean = 0.017 , 95% HPD = $0.004-0.035$), and prior
7	best models of nucleotide substitution selected in MrModeltest. Two independent runs
8	of 100,000,000 generations each and sampled every 10,000 generations, were
9	performed for each analysis. Convergence of the MCMC chains was verified in Tracer
10	v.1.6 by checking the effective sample size values (ESS). Skyline plots were
11	constructed using Tracer v.1.6.

12

13 **Results**

14 Didelphis albiventris

15 The BI analyses based on 14 haplotypes showed three well-supported clades for D. 16 albiventris: the Southwest clade (SWC), the Cerrado clade (CE) and the Amazonia-17 Cerrado ecotone clade (ECO) (Figure 2a). The SWC clade included haplotypes from 18 Southern Brazilian Atlantic Forest and from Bolivian Chaco domains (Figure 2a-c). 19 The ECO clade is represented mainly by haplotypes from the Amazonia-Cerrado 20 ecotone, especially from mid-Araguaia River (Figure 2). The CE clade is represented 21 mainly by haplotypes from Cerrado (Figure 2). However, haplotype HDa1 from the 22 Amazonia-Cerrado ecotone (locality 34) is closely related to the CE clade, and 23 haplotype HDa8 from Cerrado (locality 77) is closely related to the ECO clade (Figure 24 2).

1 The average genetic distance between SWC clade and CE plus ECO clades is 2 6.6 ± 0.9 %, and their t_{MRCA} dates to the Pleistocene at 0.76–2.52 Myr BP (Table 1, 3 Figure S1). The average K2p genetic distance between the two central Brazilian clades 4 (CE and ECO) is of 1.1 ± 0.3 %, and their t_{MRCA} dates to the Pleistocene at 0.24–0.98 5 Myr BP (Table 1, Figure S1).

6 The MJ network was constructed with 15 samples from central Brazil (Table 2). 7 CE and ECO clade are separated by four mutations (Figure 2b). Haplotype HDa4 is the 8 ancestor of the CE clade, and it is shared by specimens from both riverbanks of the Araguaia (Figure 2b). Hypothetical ancestor of the ECO clade was not sampled (Figure 9 10 2b). The migration rate *m* estimated in IM (using ECO clade) had a long tail towards 11 higher values (Figure S2). The best estimate of divergence time t between the eastern 12 and western river banks was 0.645 Myr BP (95% HPD = 0.031-1.389 Myr BP, Table 3). 13 Eastern (qEAR), western (qWAR) and ancestral (qA) populations had similar inferred sizes (Table 3, Figure S3). Neutrality tests showed non-significant values (Table 2), not 14 15 rejecting the constant population model. BSP for the ECO clade also support a constant 16 population (Figure 3).

17

18 Didelphis marsupialis

19 The BI analyses based on 13 haplotypes showed three well-supported clades for *D.* 20 *marsupialis*: the western Amazonia clade (WAM), the Guiana Shield clade (GS) and the 21 Amazonia-Cerrado ecotone clade (ECO) (Figure 4). The WAM clade included all 22 haplotypes from Juruá River, western Amazonia, with exception of a sample from 23 locality 15 (Peru) (Figure 4). The GS clade is represented by haplotypes from the 24 Guyana (Figure 4). The ECO clade is represented mainly by haplotypes from the 25 Amazonia-Cerrado ecotone, especially from mid-Araguaia River, but also a sample from Tocantins River (Figure 4). The relation between these three clades is not resolved
 (Figure 4a), but their t_{MRCA} dates to the Pleistocene at 0.44–1.65 Myr BP (Table 1,
 Figure S1).

4 The MJ network was constructed with 50 samples of the ECO clade (Table 2). 5 Haplotype HDm1 is the ancestor of the Araguaia River samples and it is shared by 6 specimens from both riverbanks (Figure 4b). Haplotype HDm3 is also shared by 7 specimens from both riverbanks, along with specimens from the islands of the Araguaia 8 River (Figure 4b). Haplotype HDm2 was only sampled in islands, HDm4 in the eastern 9 bank, and HDm5 in the western bank of the Araguaia River (Figure 4b). The IM results 10 of the ECO clade suggested lower gene flow than D. albiventris, but still substantial 11 ranging from 0.95 (95% HPD = 0.41 to 16.95) for western river bank and islands to 12 1.61 (95% HPD = 0.71 to 19.01) for eastern and western river banks (Table 3, Figure 13 S2). Although the divergence time t did not converged well for the three pairwise 14 combinations, the best estimate of divergence time between the eastern and western 15 river banks was 0.702 Myr BP (95% HPD = 0.025-1.392 Myr BP, Table 3). The eastern 16 population had a lower inferred size (qEAR) than the ancestral population (qA) and the 17 western one (qWAR, Table 3, Figure S3). Neutrality tests showed non-significant 18 values (Table 2), not rejecting the constant population model. BSP for the ECO clade 19 also support a constant population (Figure 3).

20

21 Philander opossum

The BI analyses based on 15 haplotypes showed two well-supported clades for *P*. *opossum*: the western Amazonia clade (WAM) and the Araguaia River clade (AR) (Figure 5a). The WAM clade included all haplotypes from Juruá River, western Amazonia (Figure 5). The AR clade is represented by haplotypes from the Araguaia River, including Amazonia-Cerrado ecotone and Cerrado domains (Figure 5).
 Haplotypes from the Pantanal (PAN) formed a polytomy with those from the Araguaia
 River (AR), central Brazil clade (Figure 5).

- 4 The MJ network was constructed using 54 samples of the AR clade (Figure 5b, 5 Table 2). The haplotype HPo1 is the ancestor of the mid-Araguaia River samples and it 6 is shared by specimens from both riverbanks (Figure 5b). Haplotypes HPo6 and HPo7 7 are also shared by specimens from both riverbanks (Figure 5b). The IM results of the 8 AR clade suggested some gene flow between eastern and western river banks, with the 9 posterior distributions of migration rate m peaking at 1.74 (90% HPD = 0.615 to 9.115,10 Table3). The time of divergence t did not converged well (Figure S2), probably due to 11 the high values of gene flow which could mask a signal of population separation. The 12 eastern population had a lower inferred size (qEAR) than the ancestral population (qA) 13 and the western one (qWAR, Table 3, Figure S3). Neutrality tests showed nonsignificant values (Table 2), not rejecting the constant population model. BSP for the 14 15 AR clade also support a constant population (Figure 3).
- 16

17 Gracilinanus agilis

18 The BI analyses based on 65 haplotypes showed six well-supported clades for G. agilis: 19 the east clade (EAC), the north-eastern clade (NEC), the western clade (WCE), the 20 southern clade (SC), the central Brazil plateaux clade (CBP) and the central-east clade 21 (CEC, Figure 6a). The east clade (EAC) included almost all haplotypes from the moist 22 forest enclaves in the east of the São Francisco River, with exception of two haplotypes 23 from locality 65 (Rio de Contas, state of Bahia, Figure 6). The north-eastern clade 24 (NEC) is represented by haplotypes from the north-eastern Caatinga and Cerrado, which 25 are located in the west margin of São Francisco River (Figure 6). The western clade (WCE) included haplotypes from Amazonia-Cerrado ecotone and western Cerrado
 (Figure 6). The southern clade (SC) included haplotypes from southern Cerrado and
 Pantanal (Figure 6). The central Brazil plateaux clade (CBP) and central-east clade
 (CEC) included haplotypes from central Brazil plateaux and from central-east Cerrado,
 respectively (Figure 6)

6 The relation between most of these clades is not resolved, but their t_{MRCA} dates 7 to the Pliocene/Pleistocene at 1.15–3.32 Myr BP (node 3, Figure 6a, Table 1). The 8 average genetic distance between clades varied from 4.7 ± 0.6 % (EAC and the rest of 9 the clades) to 2.5 ± 0.4 % (WCE and CBP). The t_{MRCA} of central Brazilian clades (WCE 10 + CBP + CEC + SC) was estimated during the Pleistocene at 0.71–1.94 Myr BP (Table 11 1, Figure S1).

12 The MJ network was constructed using 56 samples of the WCE clade (Figure 13 6b). The haplotype HGa4 is centrally located relatively to samples from the eastern 14 bank of the Araguaia River (Figure 6b). The haplotype HGa11 was only sampled in the 15 western bank of the Araguaia River (Figure 6b). Neutrality tests showed non-significant 16 values (Table 2), not rejecting the constant population model. BSP for the WCE clade 17 also support a constant population (Figure 3).

18

19 Marmosa murina

The BI analyses based on 36 haplotypes showed five well-supported clades for *M. murina*: the Atlantic Forest clade (AF), the Tapajós River clade (TR), the Araguaia-Tocantins interfluve clade (ATI), the western Araguaia River clade (WAR) and the eastern Araguaia River clade (EAR) (Figure 7a). Haplotypes from the Guiana Shield (GS) and northern Amazonia (NAM) form a polytomy with the remaining haplotypes (Figure 7a). 1 The monophyly of each clade from both river banks of the Araguaia River 2 (WAR and EAR) and from the Araguaia-Tocantins interfluve (ATI) is supported with 3 high Bayesian posterior probabilities (BPP = 1.00, Figure 7a). The t_{MRCA} of these clades 4 dates to the Pleistocene at 0.39–1.32 Myr BP (Table 1), and the average genetic distance 5 between them is 1.9 ± 0.4 %.

6 The MJ network was constructed using 80 samples of WAR, EAR and ATI 7 clades (Figure 7b, Table 2). Our results also suggested that ancestors of these clades 8 have not been sampled yet and haplotypes are exclusive from each river bank (WAR 9 and EAR) (Figure 7b). Haplotypes from Araguaia-Tocantins interfluves (ATI) are 10 divergent and separated by 8 mutations, resulting in a MJ network with large branches, 11 and with no intermediary haplotypes (Figure 7b). The IM results of WAR and EAR 12 clades suggested very low gene flow between eastern and western river banks, with the 13 posterior distributions of migration rate *m* peaking at 0.001 (95% HPD = 0.001 to 0.123, 14 Table 3, Figure S2). The posterior distribution of time of divergence t peaked at 6.31 (95% HPD = 3.21 to 19.01, Figure S2), which converted to time in years suggests that 15 the populations split occurred about 0.451 Myr BP (95% HPD = 0.229-1.378 Myr BP, 16 17 Table 3). The eastern population had a lower inferred size (qEAR) than the ancestral 18 population (qA) and the western one (qWAR, Table 3, Figure S3). Neutrality tests 19 showed non-significant values (Table 2), not rejecting the constant population model. BSP for the EAR and WAR clades shows a slightly population decline to the present 20 21 (Figure 3).

22

23 Marmosa demerarae

24 The BI analyses based on 37 haplotypes showed six well-supported clades for M.

25 demerarae: the western Amazonia clade (WAM), the northern Amazonia clade (NAM),

1 the Guiana Shield clade (GS), the Amazonia-Cerrado ecotone clade (ECO), the eastern 2 Tocantins River clade (EToR) and the Atlantic Forest clade (AF) (Figure 8a). The 3 WAM clade is a divergent clade $(5.7 \pm 0.7 \%)$, including samples from southern 4 Amazon River to the Teles Pires River (Figure 8c). The AF clade is represented by 5 almost all haplotypes from the Atlantic Forest domain, with exception of a haplotype from locality 60 (São José de Lages, state of Alagoas) in the northern Atlantic Forest 6 7 (Figure 8). This haplotype is the only one located west of the São Francisco River, 8 while remaining samples from the Atlantic Forest are located east of this river (Figure 9 8c). The ECO clade includes haplotypes from the Amazonia-Cerrado ecotone, mainly 10 from both Araguaia riverbanks but also a haplotypes from the Xingu River (Figure 8c). 11 The EToR clade includes haplotypes from the eastern bank of the lower Tocantins River 12 in the Amazonia-Cerrado ecotone (HMd15 and HMd16) and in the Cerrado (HMd17). 13 Haplotype HMd12 from the western bank of the lower Tocantins River (locality 30) is 14 very distant $(4.2 \pm 0.7 \%)$ from the rest of the samples of central Brazil, and formed a 15 polytomy with AF, ECO and EToR clades (Figure 8).

16 The MJ network was constructed using 40 samples of ECO and EToR clades 17 (Figure 8b, Table 2). Haplotypes HMd1, HMd3-HMd5 and HMd7 from the eastern 18 bank of the mid-Araguaia River (EAR), have as ancestor the haplotype HMd14 from 19 the western bank of the lower Araguaia River (locality 28, Figure 8b-c). The ETOR 20 clade and the haplotype HMd12 are separated by 25 and 28 mutations, respectively, 21 from the rest of the samples, which result in a MJ network with very large branches, and 22 with no intermediary haplotypes (Figure 8b). The IM results suggested very limited 23 gene flow between eastern and western river banks, with the posterior distributions of migration rate m peaking at 0.047 (95% HPD = 0.007 to 0.641, Table 3, Figure S2). 24 25 Although the divergence time t did not converged well, there was a peak at 1.88 (95%)

1	HPD = 0.693 to 5.859, Figure S2), which converted to time in years suggests that the
2	populations started diverging about 0.134 Myr BP (95% HPD = 0.050-0.419 Myr BP,
3	Table 3). The eastern population had a lower inferred size $(qEAR)$ than the ancestral
4	population (qA) and the western one ($qWAR$, Table 3, Figure S3). Neutrality tests
5	showed non-significant values (Table 2), not rejecting the constant population model.
6	BSP for the ECO clade also support a constant population (Figure 3).
7	
8	Discussion
9	Comparative phylogeography and species ecology
10	The six didelphid marsupials analyzed in this work revealed different patterns of
11	differentiation across the mid-Araguaia River. The three larger species, Didelphis
12	albiventris (body size: 265–363 mm), D. marsupialis (310–460 mm) and P. opossum
13	(180–260 mm), revealed extensive intraspecific haplotype sharing across the river
14	(Figure 2b, 4b and 5b). Consistent with this haplotype sharing, the IM results also
15	confirmed gene flow among Araguaia river banks for the two latter species (Table 3).
16	In contrast, the three smaller marsupial species (Gracilinanus agilis, Marmosa
17	murina and M. demerarae, body sizes: 75–108, 98–155 and 110–190 mm, respectively)
18	showed no haplotype sharing across the mid-Araguaia River. For G. agilis we did not
19	observe haplotype sharing across the river (Figure 6b), but the geographical gap
20	between our samples in this region makes the role of the river as a barrier less obvious.
21	Therefore, only increasing sampling effort in the western river bank will help to proper
22	access the role of the Araguaia River in the differentiation of this species. For M.
23	demerarae, although the paraphyletic relationship was not supported in the Bayesian
24	analysis (Figure 8a), the observed ancestral relationships in the MJ networks (Figure 8b)
25	revealed that the western bank population was most likely the origin from which the

1 eastern bank was colonized. This pattern is also supported by the observation that for M. 2 demerarae, as well as for D. marsupialis, P. opossum and M. murina, the eastern 3 population had a lower inferred size (qEAR) than the ancestral population (qA) and the 4 western one (qWAR). Although tentative this pattern could be explained by the fact that 5 these four species are either typically Amazonian species (D. marsupialis and P. 6 opossum) or that occur mainly in rainforests (M. murina and M. demerarae), and the 7 western bank of the mid-Araguaia might represent a previous distribution limit of these 8 species.

9 Haplotypes of *M. murina* on opposite riverbanks formed monophyletic clades 10 that are genetically distant from each other (Figure 7a). Although reciprocal monophyly, 11 which would confirm the formation of a complete barrier to an existing taxon (Patton et 12 al. 2000), was not observed, we hypothesized that this barrier could have been formed 13 by steps. Indeed, the Araguaia River is acting as barrier to gene flow of populations on 14 opposite river banks as indicated by IM results of *M. murina* (and also *M. demerarae*) 15 Table 3). Additionally, IM results also suggest that eastern and western populations of 16 Araguaia River most likely split about 0.451 Myr BP (0.229–1.379 Myr BP) for M. 17 *murina*, and about 0.134 Myr BP (95% HPD = 0.050–0.419 Myr BP) for *M. demerarae*, during the Pleistocene. This divergence estimates encompass the age of Araguaia River 18 19 sediments formation (roughly 0.240 Myr BP, Valente and Latrubesse 2012). Therefore, 20 we hypothesize that the Araguaia might have formed a barrier to populations of M. 21 *murina* in the mid-Araguaia (WAR and EAR) and later to the populations in the lower 22 Araguaia (ATI), but the two populations in the eastern Araguaia River (EAR and ATI) 23 have also limited gene flow among them (see Other phylogeographic patterns to further 24 discussion on ATI divergence).

1 Moreover, it should be noted that the Araguaia River marks the transition 2 between two different biomes: the Amazonia and the Cerrado. Therefore it is possible 3 that the differentiation observed across the river and in this region could be the result of 4 local adaptation to two different biomes, with reduced effective gene flow due to 5 reduced fitness of admixed individuals (e.g., Almeida et al. 2007, Miranda et al. 2007, 2009). If genetic differentiation was caused by adaptation to different biomes we would 6 7 expect sister clades related with adjacent but distinct habitats (Moritz et al. 2000; Smith 8 et al. 2005). However, limited difference in habitat between the two riverbanks allows 9 us to discard this hypothesis in explaining the intraspecific genetic structure of M. 10 *murina* and *M. demerarae*. Additionally, a signature of demographic stability detected 11 for didelphid species along the Araguaia River do not support the diversification in 12 refuges. Although the importance of Pleistocene climatic oscillations as a driver of 13 population divergence in central Brazil is still in debate, the scenario of limited or no demographic expansion has also been reported to small mammals in the western 14 15 Amazonia, contradicting the refugia model of diversification (Matocq et al. 2000; Lessa 16 et al. 2003). On the other hand, rivers of the central Brazilian plateux have been shown 17 to be stronger historical barriers in promoting differentiation (Maldonado-Coelho et al. 18 2013), as corroborated by our results.

Differences in phylogeography among species are likely to be linked to differences in the life history (e.g., mating system and reproduction success) and ecological traits (e.g., locomotion mode and habitat preferences) (Matocq et al. 2000; Rocha et al. 2014). Although body size separates the two groups of didelphid species, this morphological feature is not likely to be the main cause of the observed genetic differences in the mid-Araguaia River. Small rodents (body weight ranging from 22 to 103 g) showed distinct patterns of genetic differentiation in this river (Rocha et al. 2011,

1 2014). In addition to body size, habitat preferences and overall distribution differ among 2 didelphid species (Patton et al. 2000; Gardner 2008; Rocha et al. 2011). These 3 ecological differences probably have a great impact on their genetic structure. Marmosa 4 *murina* is known to be associated with upland forests and avoids gallery forests that 5 flood (Patton et al. 2000; Ramos Pereira et al. 2013). Marmosa demerarae and G. agilis 6 are not upland forest specialists, since they also use seasonally flooded forests to some 7 extent (Rocha et al. 2011; Ramos Pereira et al. 2013). These arboreal murine mouse 8 opossums search the understory and forest floor for invertebrate prey (Emmons and 9 Feer 1997; Gardner 2008); a foraging strategy impossible to accomplish during the 10 flooded season, which may explain their preference for dry areas (Ramos Pereira 2013). 11 Contrastingly, all of D. albiventris, D. marsupialis and P. opossum are well distributed 12 both in upland and seasonally flooded forests (Rocha et al. 2011), which have probably 13 enhanced cross-river gene flow. Indeed, previous authors have pointed out that upland forest specialists are more prone to exhibit riverine diversification than specialists of 14 15 floodplain forests (Moritz et al. 2000; Patton et al. 2000; Rocha et al. 2014), being the 16 latter capable of colonizing river islands and crossing rivers due life-history attributes 17 such as high dispersal rates (Aleixo 2006).

18

19 The importance of river geomorphology in biogeography

River basin geomorphology may be an important factor determining the importance of the river as a phylogeographic barrier. Meandering rivers, where the location of the river bed changes over time, as the Juruá River (Latrubesse 2008), may facilitate the exchange of individuals by repeatedly bringing vicariant populations into contact (Moritz et al. 2000; Patton et al. 2000). Contrastingly, anabranching rivers with low sinuosities, as the rivers in the Brazilian Shield, including Araguaia River (Latrubesse

1 2008), may be stronger barriers simply because they are more permanent physical 2 obstacles. These predictions were corroborated by previous studies at the Juruá River, 3 which was not a barrier for most of the sampled vertebrates (e.g., Gascon et al. 2000; 4 Matocq et al. 2000; Patton et al. 2000), and at Tocantins and Teles Pires Rivers, which 5 were found to be historical barriers in the differentiation of several bird species (Bates et 6 al. 2004; Maldonado-Coelho et al. 2013). They are also confirmed by our results at the 7 mid-Araguaia River, although we find the phylogeographical impact of the river to be 8 species-dependent (see also Rocha et al. 2011, 2014; Faria et al. 2013b). Our 9 comparative results suggest that semi-permanent alluvial islands characteristic from 10 anabranching rivers may be important stepping-stones facilitating the gene flow of large 11 didelphids, as well as small rodents Rocha et al. 2014), which inhabit seasonally 12 flooded forests. Indeed, haplotype sharing and gene flow among populations of D. 13 *marsupialis* of both Araguaia river banks and of the islands was confirmed by our data.

14

15 Other phylogeographic patterns

16 Phylogeographic analysis of *M. demerarae* suggests that the dispersion of this species in 17 central Brazil occurred throughout the gallery forests of the Araguaia and Tocantins 18 River. Specifically, the observed genetic pattern revealed that specimens from the 19 eastern bank of the Tocantins River are genetically similar despite the large geographic 20 distance, while most of the specimens from the eastern and western bank of the 21 Araguaia River belong to the same cluster (Figure 8c). This finding supports previous 22 studies that have shown that these gallery forest act as dispersal corridors for rainforest 23 mammal species throughout the Cerrado (e.g., Johnson et al. 1999; Costa 2003).

Very divergent haplotypes of *M. demerarae* and *M. murina* were recorded in the
Araguaia-Tocantins interfluve. Previously, a potentially new species of the subgenus

1 *Micoureus* was recorded circa 300 km east of this interfluve (Costa 2003), revealing the 2 importance of this region to the diversification within this subgenus. In addition to the 3 river interfluve, palynological records of the Serra dos Carajás, state of Pará, showed 4 that this region went through several alternating periods dominated by arboreal and 5 herbaceous savanna vegetations (Iriondo and Latrubesse 1994; Behling et al. 2010). 6 However previous authors fail in support bird differentiation in glacial forest refuges, 7 instead they found evidences of riverine barrier caused by the Tocantins River 8 (Maldonado-Coelho et al. 2013). Further intensive sampling in this region will help to 9 uncover the Araguaia-Tocantins interfluve contribution to the diversification of these 10 didelphid marsupials.

11 Finally, our phylogeographic analyses of G. agilis revealed highly supported 12 clades within Cerrado of central Brazil. Faria et al. (2013a) discussed the importance of 13 the Serra Geral do Goiás to the diversification of this species. Although the geological 14 origin of these mountain chains dates to the Cretaceous (Villela and Nogueira 2011), 15 before the diversification of this genus, the western clades (SC, CBP, CEC and WCE) 16 and northern-east clade (NEC) are separated by the Serra Geral do Goiás (Faria et al. 17 2014a, and also shown in the present study). Moreover, seven geographical groups 18 within the Cerrado were recognized based on its floristic composition (Ratter et al. 2003). The western clades of G. agilis are not completely congruent with the 19 20 geographical limits of the floristic groups (Ratter et al. 2003). However, vegetation 21 pattern reflects changes that occurred during the Tertiary and Quaternary periods (Ratter 22 et al. 2003), and the extant diversity of G. agilis in Cerrado dates at 1.26 Myr BP (0.71-23 1.94 Myr BP). Thus, suggesting that vegetation fluctuations occurred during this period 24 may have influenced the intraspecific genetic structure of this species (see also 25 Nascimento et al. 2013).

1

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11

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1 Figures and Tables

Figure 1. (a) Sampling areas at the mid-Araguaia River and its tributaries (left, source: Google Earth). Sampling points in white corresponds to the eastern river bank, in black to the western river bank, and in gray to the islands of the Araguaia River. (b) Map including all sampling localities of six didelphid marsupials in South America (right). Biomes are defined in different colors, and the major rivers are indicated in blue. See Gazetteer for detailed information on localities.

8

9 Figure 2. (a) Bayesian inference tree of *mt-Cytb* haplotypes of *D. albiventris*, in 10 which the GTR+I model was used. Numbers in circles represent nodes for which time 11 to most recent common ancestor are shown in Table 1; percentage values correspond 12 to pairwise genetic distances between clades; and asterisks indicate BPP ≥ 0.95 ; 13 numbers in the tip of branches are sampling localities and acronyms are haplotype 14 designations; clades are designated by ECO (Amazonia-Cerrado ecotone clade), CE 15 (Cerrado clade) and SWC (southwest clade). (b) Median Joining network of D. 16 albiventris of the ECO and CE clades, central Brazil. Length of connecting branches 17 corresponds to the nucleotide substitutions; size of the circles is proportional to the 18 number of individuals sharing each haplotype; white corresponds to the eastern 19 riverbank of the Araguaia River, and black to the western riverbank. (c) Map of 20 sampling localities of *D. albiventris*, with the representation of the clades.

21

Figure 3. Demographic scenarios of six didelphid marsupials from central Brazil.
 Black solid curves indicate changes in effective population size and grey shadows
 indicate upper and lower 95 % confidence interval.

1 Figure 4. (a) Bayesian inference tree of *mt-Cytb* haplotypes of *D. marsupialis*, in 2 which the GTR+I model was used. Numbers in circle represent nodes for which time 3 to most recent common ancestor are shown in Table 1; percentage values correspond to pairwise genetic distances between clades; and asterisks indicate BPP ≥ 0.95 ; 4 numbers in the tip of branches are sampling localities and acronyms are haplotype 5 6 designations; clades are designated by ECO (Amazonia-Cerrado ecotone), GS 7 (Guiana Shield) and WAM (western Amazonia). (b) Median Joining network of D. marsupialis of the ECO clade, central Brazil. Length of connecting branches 8 9 corresponds to the nucleotide substitutions; size of the circles is proportional to the 10 number of individuals sharing each haplotype; white corresponds to the eastern 11 riverbank of the Araguaia River, black to the western riverbank, and grey to the 12 islands of the Araguaia River. (c) Map of sampling localities of D. marsupialis, with 13 the representation of the clades.

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15 Figure 5. (a) Bayesian inference tree of *mt-Cytb* haplotypes of *P. opossum*, in which 16 the the GTR+G model was used. Numbers in circles represent nodes for which time to 17 most recent common ancestor are shown in Table 1; percentage values correspond to 18 pairwise genetic distances between clades; and asterisks indicate BPP ≥ 0.95 ; 19 numbers in the tip of branches are sampling localities and acronyms are haplotype 20 designations; clades are designated by WAM (western Amazonia), AR (Araguaia 21 River) and PAN (Pantanal). (b) Median Joining network of *P. opossum* of the AR clade, central Brazil. Length of connecting branches corresponds to the nucleotide 22 23 substitutions; size of the circles is proportional to the number of individuals sharing 24 each haplotype; white corresponds to the eastern riverbank of the Araguaia River, and black to the western riverbank. (c) Map of sampling localities of *P. opossum*, with the
representation of the clades.

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4 Figure 6. (a) Bayesian inference tree of *mt-Cytb* haplotypes of *G. agilis*, in which the GTR+I+G model was used. Numbers in circles represent nodes for which time to 5 6 most recent common ancestor are shown in Table 1; percentage values correspond to pairwise genetic distances between clades; and asterisks indicate BPP ≥ 0.95 ; 7 8 numbers in the tip of branches are sampling localities and acronyms are haplotype 9 designations; clades are designated by the SC (southern clade), CBP (central Brazil 10 Plateau clade), CEC (central-east clade), WCE (western clade), NEC (north-eastern 11 clade) and EAC (east clade). (b) Median Joining network of G. agilis of WCE clade, 12 central Brazil. Length of connecting branches corresponds to the nucleotide 13 substitutions; size of the circles is proportional to the number of individuals sharing 14 each haplotype; white corresponds to the eastern riverbank of the Araguaia River, and 15 black to the western riverbank. (c) Map of sampling localities of G. agilis, with the 16 representation of the clades.

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18 Figure 7. (a) Bayesian inference tree of *mt-Cytb* haplotypes of *M. murina*, in which 19 the HKY+G model was used. Numbers in circles represent nodes for which time to 20 most recent common ancestor are shown in Table 1; percentage values correspond to 21 pairwise genetic distances between clades; and asterisks indicate BPP ≥ 0.95 ; 22 numbers in the tip of branches are sampling localities and acronyms are haplotype 23 designations; clades are designated by EAR (eastern Araguaia River), WAR (western 24 Araguaia River), ATI (Araguaia-Tocantins interfluve), TR (Tapajos River), AF (Atlantic Forest), NAM (northern Amazonia) and GS (Guiana Shield). (b) Median 25

Joining network of *M. murina* of EAR, WAR and ATI clades, central Brazil. Length of connecting branches corresponds to the nucleotide substitutions; size of the circles is proportional to the number of individuals sharing each haplotype; white corresponds to the eastern riverbank of the Araguaia River, and black to the western riverbank. (c) Map of sampling localities of *M. murina*, with the representation of the clades.

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8 Figure 8. (a) Bayesian inference tree of *mt-Cytb* haplotypes of *M. demerarae*, in 9 which the HKY+I+G model was used. Numbers in circles represent nodes for which 10 time to most recent common ancestor are shown in Table 1; percentage values 11 correspond to pairwise genetic distances between clades; and asterisks indicate BPP \geq 12 0.95; numbers in the tip of branches are sampling localities and acronyms are 13 haplotype designations; clades are designated by WAM (western Amazonia), NAM 14 (northern Amazonia), GS (Guiana Shield), ECO (Amazonia-Cerrado ecotone), EToR 15 (eastern Tocantins River) and AF (Atlantic Forest). (b) Median Joining network of M. 16 demerarae of the ECO (including samples from eastern Araguaia River, EAR) and 17 EToR clades, central Brazil. Length of connecting branches corresponds to the 18 nucleotide substitutions; size of the circles is proportional to the number of 19 individuals sharing each haplotype; white corresponds to the eastern riverbank of the 20 Araguaia River, and black to the western riverbank. (c) Map of sampling localities of 21 *M. demerarae*, with the representation of the clades.

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- 1 Table 1 Mean time to most recent common ancestor of the main clades of didelphid
- 2 species. Mean time and 95 % confidence intervals (95 % CI) are given in million
- 3 years before present (Myr BP). Geological time correspondent to the t_{MRCA} adapted

95 % CI

Geological Time

NodeMean1 (D. albiventris ECO / D. albiventris CE)0.56

from Cohen et al. (2013).

1 (<i>D. albiventris</i> ECO / <i>D. albiventris</i> CE) 2 (<i>D. albiventris</i> ECO + CE / <i>D.</i>	0.56	0.24–0.98	Pleistocene
albiventris SWC)	1.56	0.76-2.52	Pleistocene
1 (D. marsupialis)	0.98	0.44-1.65	Pleistocene
1 (P. opossum)	1.11	0.49–1.87	Pleistocene
1 ($G. agilis$ WCE + CBP + CEC + SC)	1.26	0.71-1.94	Pleistocene
2 (G. agilis WCE + CBP + CEC + SC / G.			
<i>agilis</i> NCE)	1.50	0.83-2.32	Pleistocene
3 (G. agilis)	2.15	1.15-3.32	Pliocene / Pleistocene
1 (M. murina EAR / M. murina WAR)	0.81	0.39-1.32	Pleistocene
2 (M. murina)	1.99	1.00-3.19	Pliocene / Pleistocene
1 (<i>M. demerarae</i> ECO + EToR + AF)	1.35	0.69–2.10	Pleistocene
2 (M. demerarae ECO + EToR + AF / M.			
demerarae GS + NAM)	1.76	0.96–2.68	Pliocene / Pleistocene
3 (M. demerarae)	2.36	1.34–3.61	Pliocene / Pleistocene

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7 **Table 2** Number of individual sequences (n), haplotypes (h), number of polymorphic

8 sites (S), haplotype (Hd) and nucleotide (π) diversity, and deviation from neutrality

9 tests (Tajima's D, Fu's Fs, and Ramos-Onsins and Rozas's R₂) of six didelphid

10 marsupial *cyt-b* sequences for central Brazil. Significance levels of observed statistics

11 are expressed as follows: *(p < 0.05), NS (not significant).

Species	n	h	S	Hd	Π	D	<mark>Fs</mark>	R ₂
D. albiventris	15	9	18	0.876	0.0062	<mark>-0.40 NS</mark>	<mark>-1.112NS</mark>	<mark>0.1246NS</mark>
D. marsupialis	50	5	6	0.642	0.0014	-0.378NS	<mark>0.253NS</mark>	<mark>0.095NS</mark>
G. agilis	108	39	94	0.951	0.0167	<mark>-1.016NS</mark>	<mark>-3.756NS</mark>	<mark>0.069NS</mark>
M. (Marmosa) murina	80	21	49	0.908	0.013	0.151NS	1.048NS	<mark>0.104NS</mark>
M. (Micoureus) demerarae	40	17	67	0.915	0.015	-0.952NS	<mark>0.734NS</mark>	0.088NS
P. opossum	54	9	11	0.723	0.0022	<mark>-0.743NS</mark>	-1.464NS	<mark>0.0783NS</mark>

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1 **Table 3** Posterior distributions of parameter estimates from IM, including the mode number of migrants per generation (*m*) between populations

2 from either Araguaia river bank, time of divergence (*t*) in millions of years before present (Myr BP), and the effective population size of both

3 populations as well as of the ancestral population (qEAR, qWAR, qIslands and qA, respectively), with 95 % highest posterior distributions

4 (HPD).

Species	Populations	<i>m</i> * (95% HPD) 3.11 (0.95–19.45)	<i>t</i> (95% HPD) 0.645 (0.031–1.389)	qEAR (95% HPD)	<i>q</i> WAR (95% HPD)	<i>q</i> Islands (95% HPD)	qA (95% HPD)	
D. albiventris	EAR/WAR			374,819 (84,849-	485,206 (79,906-	-	663,142	(38,717–
D. marsupialis	EAR/Islands	1.29 (0.54–17.71)	0.732 (0.042–1.394)	<u>1,552,824)</u> <u>60,589</u> (9,129– 200,001)	1,579,185)	20,417 (2,489–	1,594,013) 163,177	(8,133–
	WAR/Islands	0.95 (0.41–16.95)	0.717 (0.024–1.394)	290,001)	181,474 (35,625–	162,181) 29,887 (4,542–	323,533) 240,770	(13,150–
	EAR/WAR	1.61 (0.71–19.01)	0.702 (0.025–1.392)	38,214 (9,389–	454,523) 164,430 (29,479–	146,566) -	465,999) 220,331	(12,010–
P. opossum	EAR/WAR	1.74 (0.615–9.115)	0.131 (0.061–1.394)	180,152) 49,571 (16,811–	416,861) 487,524 (117,678– 841,852)	-	425,596) 407,348 820,266)	(18,535-
M. murina	EAR/WAR	0.001 (0.001-0.123)	0.451 (0.229–1.378)	138,369) 228,618 (107,940– 460,589)	841,853) 543,723 (311,752– 933,916)	-	839,266) 482,043 1,286,565)	(20,783-
M. demerarae	EAR/WAR	0.047 (0.007-0.641)	0.134 (0.050-0.419)	96,622 (30,095– 270,860)	593,992 (245,516– 1,582,395)	-	$\frac{1,280,303)}{1,500,028}$ 3,049,159)	(460,937-

5 *Reported as modes

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1 Supplementary material

Table S1 List of mitochondrial DNA haplotypes of six didelphid marsupials. List of the haplotype code and frequency (n), voucher specimen representing the haplotype, and GenBank accession numbers for *mt-Cytb* sequences of didelphid marsupials used in this study, and map localities were haplotypes were sampled; map numbers are listed in gazetteer. Only haplotypes from Central Brazil have been named.

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8 Gazetteer. List of the decimal geographic coordinates (Longitude / Latitude) of 83
9 sampling localities. Localities are listed by country and Brazilian localities are
10 separate by state (abbreviations in parentheses).

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Figure S1. Bayesian inference tree with the estimates of the time to most recent common ancestor of didelphid species. Blue bars represent the 95 % confidence intervals (95 % CI) of the time to most recent common ancestor derived from BEAST analyses. Details of divergence date estimates are given in Table 1. Geological time scale adapted from Cohen et al. (2013).

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Figure S2. Posterior distributions of the migration rates (m) and of the scaled divergence time (t) between populations of the Araguaia river banks. For *D. marsupialis* three pairwise combinations are shown: eastern and western river banks, eastern river bank and islands, and western river bank and islands.

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Figure S3. Posterior distributions of the effective population size in both populations in eastern and western Araguaia river bank, as well as the ancestral population (*q*EAR, *q*WAR, and *qA*, respectively).

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