See discussions, stats, and author profiles for this publication at: https://www.researchgate.net/publication/313069140

Capuchin monkey biogeography: Understanding Sapajus Pleistocene range expansion and the curre....

Article *in* Journal of Biogeography · January 2017 DOI: 10.1111/jbi.12945

citatio 3	NS READS 420	S
15 aut	t hors , including:	
6	Marcela Lima Museu Paraense Emilio Goeldi - MPEG 28 PUBLICATIONS 69 CITATIONS SEE PROFILE	Janet C Buckner Iowa State University 8 PUBLICATIONS 55 CITATIONS SEE PROFILE
?	Alexandre Aleixo Museu Paraense Emilio Goeldi - MPEG 245 PUBLICATIONS 2,330 CITATIONS SEE PROFILE	Jean P Boubli University of Salford 90 PUBLICATIONS 897 CITATIONS SEE PROFILE

Some of the authors of this publication are also working on these related projects:



Fundación Proyecto Primates View project



Evolutionary heritage influences Amazon tree ecology Museo de Historia Natural View project

All content following this page was uploaded by Helder Lima Queiroz on 13 December 2017.

The user has requested enhancement of the downloaded file.

ORIGINAL ARTICLE

Capuchin monkey biogeography: understanding *Sapajus* Pleistocene range expansion and the current sympatry between *Cebus* and *Sapajus*

Marcela G. M. Lima^{1,2}* (D), Janet C. Buckner³, José de Sousa e Silva-Júnior², Alexandre Aleixo², Amely B. Martins^{4,5}, Jean P. Boubli⁶, Andrés Link⁷, Izeni P. Farias⁸, Maria Nazareth da Silva⁹, Fabio Röhe^{9,10}, Helder Queiroz¹¹, Kenneth L. Chiou¹², Anthony Di Fiore⁴, Michael E. Alfaro^{1,3} and Jessica W. Lynch Alfaro^{1,13}

¹Institute for Society and Genetics, University of California, Los Angeles, CA, USA, ²Curso de Pós-Graduação em Zoologia, Universidade Federal do Pará/Museu Paraense Emílio Goeldi, Belém, PA, Brazil, ³Department of Ecology and Evolutionary Biology, University of California, Los Angeles, CA, USA, ⁴Department of Anthropology, University of Texas at Austin, Austin, TX, USA, ⁵Centro Nacional de Pesquisa e Conservação de Primatas Brasileiros, ICMBio, MMA, João Pessoa, Brazil, ⁶School of Environment and Life Sciences, University of Salford, Salford, UK, ⁷Department of Biological Sciences and School of Management, Universidad de Los Andes, Bogotá, Colombia, ⁸Universidade Federal do Amazonas UFAM, Manaus, AM, Brazil, ⁹Instituto Nacional de Pesquisas da Amazonia (INPA), Manaus, AM, Brazil, ¹⁰Wildlife Conservation Society, Manaus, Brazil, ¹¹Instituto de Desenvolvimento Sustentável Mamirauá, Tefé, AM, Brazil, ¹²Department of Anthropology, Washington University, St. Louis, MO, USA, ¹³Department of Anthropology, UCLA, Los Angeles, CA, USA

*Correspondence: Marcela G. M. Lima, Departamento de Mastozoologia, Museu Paraense Emílio Goeldi, Av. Perimetral, 1901 – Terra Firme, CEP 66040-170 Belém, PA, Brazil.

E-mail: marcela_gml@yahoo.com.br

ABSTRACT

Aim Our aim was to examine gracile capuchin (*Cebus*) and robust capuchin monkey (*Sapajus*) diversification, with a focus on recent *Sapajus* expansion within Amazonia. We wanted to reconstruct the biogeographical history of the clade using statistical methods that model lineages' occupation of different regions over time in order to evaluate recently proposed 'Out of Amazonia' and 'Reinvasion of Amazonia' hypotheses as alternative explanations for the extensive geographical overlap between reciprocally monophyletic gracile (*Cebus*) and robust (*Sapajus*) capuchin monkeys.

Location Central and South America.

Methods We reconstructed a time-calibrated molecular phylogeny for capuchins under Bayesian inference from three mitochondrial genes. We then categorized 12 capuchin clades across four Neotropical centres of endemism and reconstructed the biogeographical history of the capuchin radiation using six models implemented in 'BioGeoBEARS'. We performed a phylogeographical analysis for a robust capuchin clade that spans the Atlantic Forest, Cerrado, Caatinga and Amazonia.

Results We find support for a late Miocene vicariant *Cebus-Sapajus* divergence and a Pleistocene *Sapajus* invasion of Amazonia from the Atlantic Forest. Our new analyses confirm *Sapajus* diversified first in the Atlantic Forest, with subsequent range expansion into widespread sympatry with *Cebus* in Amazonia, as well as multiple expansions into drier savanna-like habitats. We do not find mitochondrial molecular congruence with morphological species distinctions for *Sapajus flavius*, *S. cay*, *S. macrocephalus*, *S. libidinosus* and *S. apella*; instead, these five morphological types together form a single widespread clade (Bayesian posterior probability = 1) with geographical substructure and shared ancestry during the Pleistocene.

Main conclusions Our results support vicariance dividing ancestral capuchin populations in Amazonia versus the Atlantic Forest, and a Pleistocene 'Amazonian invasion' by *Sapajus* to explain the present-day sympatry of *Cebus* and *Sapajus*.

Keywords

Amazonia, Atlantic Forest, BEAST phylogeography, BioGeoBEARS, Caatinga, *Cebus*, Cerrado, range expansion, *Sapajus*

INTRODUCTION

Neotropical primates are the major diurnal mammalian radiation in the Neotropics, and include over 140 species found across Amazonia, the Andes, the Atlantic Forest, and even habitats such as the Llanos, Caatinga, Cerrado and Central Grasslands that are too hot and dry to support many types of mammals. Neotropical primate diversity is highest within the tropical rain forest habitat in Amazonia. Major questions include whether Neotropical primates diversified first in Amazonia, and how and when they spread throughout the various biomes they now inhabit.

The wide distribution of capuchin monkeys, with species ranging from Central America to northern Argentina, makes them a potential model system for addressing questions of Neotropical primate biogeography. Capuchins comprise two major forms, the more lightly built gracile capuchin, *Cebus*, and the robust capuchin, *Sapajus*. Although recent studies have confirmed the genetic distinctiveness of these two lineages (Lynch Alfaro *et al.*, 2012a; Martins *et al.*, 2014), key questions remain regarding their present-day distribution, including the extensive sympatry between Amazonian *Cebus* and *Sapajus* species.

Two competing hypotheses have been posited to explain current capuchin distributions (Lynch Alfaro *et al.*, 2012a). The 'Out of Amazonia' (OOA) hypothesis places the ancestral range for all capuchins in Amazonia, an initial diversification of both crown robust and crown gracile capuchins within Amazonia, and subsequent independent and recent invasions of non-Amazonian regions by both gracile and robust capuchins (Lynch Alfaro *et al.*, 2012a). The 'Reinvasion of Amazonia' (ROA) hypothesis proposes that gracile capuchins, *Cebus*, diverged from robust capuchins, *Sapajus*, early in capuchin evolutionary history; crown *Cebus* originated in Amazonia and crown *Sapajus* in the Atlantic Forest; and the presence of *Sapajus* in Amazonia today reflects a recent invasion from the Atlantic Forest into *Cebus*-occupied territory (Lynch Alfaro *et al.*, 2012a).

Phylogeographical analyses using pure dispersal models strongly support the 'ROA' hypothesis (Lynch Alfaro et al., 2012a), and as such have transformed the interpretation of capuchin taxonomy, conservation biology, morphology and ecology (see Lynch Alfaro et al., 2012b, 2014, 2015b; Matthews, 2012; Rylands et al., 2013; Martins et al., 2014; Oliveira et al., 2014; Boubli et al., 2015; Schneider & Sampaio, 2015; Wright et al., 2015). However, limitations of these studies have left key predictions of the OOA hypothesis untested. For example, Lynch Alfaro et al. (2012a) and Nascimento et al. (2015) provided limited sampling of eastern Amazonia and Cerrado/ Caatinga localities, which have the potential to strongly corroborate the OOA scenarios. In addition, capuchin monkey biogeography has yet to be interpreted in light of models that integrate processes beyond pure dispersal (Ronquist, 1997; Ree et al., 2005; Ree & Smith, 2008; Landis et al., 2013).

Here, we assemble an expanded sampling of capuchin monkeys across their range and conduct new biogeographical

analyses using recently developed statistical approaches. Our overall goal is to test hypotheses explaining capuchin biogeography (Boubli et al., 2012; Lynch Alfaro et al., 2012a; Nascimento et al., 2015; Ruiz-García et al., 2016). We thus assemble a new cytochrome b + D-loop + cytochrome oxidase I (COI) matrix with three times the samples and more comprehensive spatial and taxonomic sampling compared to Lynch Alfaro et al. (2012a). We produce a detailed phylogeny for capuchin monkeys, with all major lineages represented, to confirm the monophyly of Cebus and Sapajus and to recover major clades within the capuchin radiation. We compare alternative biogeographical scenarios using a flexible framework for inferring spatial histories that allows for a number of different mechanisms causing range expansion or contraction. We assess whether the employment of a more complete genetic dataset and statistical methods with expanded models increase support for previous conclusions about capuchin biogeographical evolution or support new scenarios altogether. Our sampling allows us to address two new biogeographical hypotheses for this clade. The first hypothesis postulates vicariance between Amazonia and the Atlantic Forest. Divergence between Sapajus and Cebus is caused by isolation of ancestral capuchin populations in the Atlantic Forest versus Amazonian habitats, respectively, with subsequent expansion of Sapajus into sympatry with Amazonian Cebus ('ROA' hypothesis). The second, opposing, hypothesis postulates vicariance within Amazonia due to the formation of the Amazon River. Capuchins arise in Amazonia, and subsequently Cebus is isolated north of the Amazon River, with Sapajus isolated south of the Amazon River and expanding into the Atlantic Forest. Sympatry occurs with subsequent crossing of Amazon River in both directions ('OOA' hypothesis).

MATERIALS AND METHODS

DNA extraction, amplification, sequencing and alignment

We sequenced three mitochondrial genes, cytochrome b (Cyt b, 1133 bp), D-loop (1244 bp) and COI (673 bp), for 108 capuchin samples from poorly known or previously unsampled localities and species (see Table S1 in Appendix S1 in Supporting Information). For Cyt b and D-loop we used primers and PCR protocols from Lynch Alfaro et al. (2012a) and Boubli et al. (2012). We adapted COI primers and protocols from Ward et al. (2005) (see Appendix S2). We combined the 108 new samples with 40 samples analysed previously by Lynch Alfaro et al. (2012a) and 13 new samples with reliable species identification extracted from Gen-Bank, totalling 161 capuchin samples. Note that samples from Ruiz-García et al. (2016) were not available for use in this study. We used only sequences over 500-bp long to improve clade support (Nascimento et al., 2015). We used the most recent comprehensive taxonomic review for capuchins (Rylands et al., 2013), in relation to specimen morphology and collection locality, to assign species names to samples (locality data in Fig. 1 and see Table S1 in Appendix S1). Although the limitations of mitochondrial DNA studies are well understood, *Cyt b*, D-loop and COI have performed well in capturing phylogeny and biogeographical history in a wide range of species (Tobe *et al.*, 2010; Zhang *et al.*, 2011; Robins *et al.*, 2014). In addition, mtDNA sequence can be reliably generated from museum 'crusties' and skins using now standard protocols (e.g. Lynch Alfaro *et al.*, 2012a, 2015a; Mercês *et al.*, 2015).

Phylogenetic reconstruction

We used MRBAYES 3.2.6 (Ronquist et al., 2012) to reconstruct the phylogenetic relationships among capuchins, using Saimiri as an outgroup. We partitioned Cyt b and COI by codon position for the analysis and ran the Markov chain Monte Carlo (MCMC) for 50 million generations. We used 'sumt' to calculate the Bayesian consensus tree. We then used BEAST 1.8.1 (Drummond et al., 2012) to reconstruct a time tree appropriate for our biogeographical analyses by pruning the alignment to a single exemplar taxon for each recovered well-supported clade. We considered branch lengths and recovered polytomies in addition to posterior probabilities at nodes when distinguishing clades. This resulted in a new, pruned alignment with 13 tips, representing 12 capuchin clades and 1 outgroup. We specified a HKY+G model of nucleotide substitution and partitioned by codon position, using Cyt b only to avoid branch length bias due to missing data in the other genes. We used Saimiri as an outgroup and the fossil Neosaimiri (Kay, 2015) to calibrate the split at 12.5 Ma (lognormal prior, offset = 12.5, mean = 0, SD = 1) between capuchins and squirrel monkeys for the time tree. We ran the MCMC analysis for 100 million generations, sampling states every 10,000 generations with a Yule prior on the diversification rate, and an uncorrelated, lognormal relaxed molecular clock.

Biogeographical modelling

Biogeographical scenarios

In addition to testing for evidence for 'OOA' versus 'ROA' hypotheses (see Introduction), we tested for reciprocal monophyly for *Sapajus* in Amazonia (*S. macrocephalus* + *S. apella*); the Cerrado + Caatinga + Central Grasslands (*S. cay* + *S. libidinosus*); and the Atlantic Forest (*Sapajus flavius* + *S. xanthosternos* + *S. robustus* + *S. nigritus*). Lack of reciprocal monophyly across habitats would suggest multiple independent invasions and convergent adaptations to habitat types by independent *Sapajus* lineages.

Historical biogeography

To test alternative hypotheses of capuchin origins and radiation we used the R-package 'BioGeoBEARS', a recently developed flexible maximum likelihood framework for inferring spatial histories (Matzke, 2013, 2014; see Buckner *et al.*, 2015 and Lynch Alfaro *et al.*, 2015a for its recent application to primate taxa). 'BioGeoBEARS' software allows for a statistical comparison of a suite of models that include alternative biogeographical processes related to cladogenesis and that allow for species distributions across multiple regions (Matzke, 2014).

Our Bayesian consensus tree was pruned to include one representative tip for each 'species' recovered in the MRBAYES analysis. We lumped S. flavius, S. cay, S. libidinosus, S. apella and S. macrocephalus (Sapajus clade 4, see below) as one species. We also combined C. unicolor + C. albifrons, C. olivaceus + C. kaapori and C. versicolor + C. cesarae. This resulted in a new, pruned alignment with 12 tips. We coded presence/absence in four major geographical regions for each tip based on known species distribution using Rylands *et al.* (2013): Atlantic Forest (AF), Cerrado/Caatinga/Central Grasslands (CC), Amazonia (AM) and Central America/Andes (CA) (Fig. 1). We used the

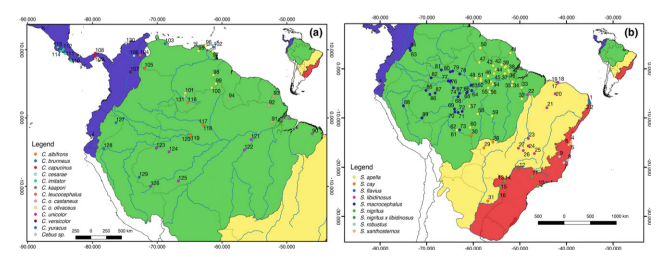


Figure 1 Maps of capuchin monkey sample provenance: (a) *Cebus*, (b) *Sapajus*. Map color: Blue indicates Central America and Andean region (CA); Green indicates Amazonia (AM); Yellow indicates Cerrado, Caatinga and Southern Grasslands (CC) and Red indicates Atlantic Forest region (AF).

pruned, time-calibrated tree with regional coding in 'BioGeo-BEARS' (Matzke, 2013, 2014). Our candidate pool consisted of six models: DEC, DEC-J, DIVA, DIVA-J, BAYAREA-LIKE, BAYAREALIKE-J and we used model weights calculated from corrected Akaike information criterion (AICc) to identify the best model or set of models for our data (Matzke, 2014). The 'BioGeoBEARS'script is available in Appendix S2.

Phylogeographical reconstruction of widespread Sapajus

To examine phylogeographical patterns for a widespread robust capuchin group recovered in the MRBAYES tree, we performed both symmetric and asymmetric discrete states phylogeographical analyses in BEAST under a constant size coalescent prior. These analyses allowed us to take into consideration the uncertainty in the relationships by simultaneously estimating the topology and the distribution history. To compare the asymmetric and symmetric analyses for their fit to our data, we performed marginal likelihood estimation (MLE) using path sampling (PS) and stepping stone (SS) sampling (Baele et al., 2012, 2013). We used the default priors for the MLE. We categorized the locality of each tip for the widespread Sapajus clade in the Bayesian consensus tree as occurring in one biogeographical 'region' (Fig. 2). We divided Amazonia into four regions using the Amazon River and two major tributaries, Negro and Madeira, as described by Wallace (1852) and later Cracraft (1985): Guianas (GU), Negro (NE), Inambari (IN), Rôndonia (RO). We also separated the drier open ecosystems Central Grasslands, Cerrado and Caatinga (CC), from the Atlantic Forest (AF). These regions were modified from Jameson Kiesling et al. (2015) and Cracraft (1985). As mtDNA is inherited maternally, it tracks the movement of female lineages over time. Capuchin monkeys are female philopatric with male-biased dispersal (Lynch Alfaro et al., 2014), so tracing female lineage was a conservative test for the amount of movement over time in this taxon.

Within Amazonia, we tested for isolation by river barriers for widespread *Sapajus*. If gene flow within Amazonian capuchin populations was restricted to major interfluves, we expected a pattern of closely related subclades within, not across, four major Amazonian regions delineated by the Amazon River and its major tributaries. Evidence for the river barrier hypothesis was found for the sister group to capuchins, the Amazonian squirrel monkeys (Lynch Alfaro *et al.*, 2015a), as well as for marmosets and tamarins (Buckner *et al.*, 2015) uakaris, widow titi monkeys, and white-fronted capuchins (Boubli *et al.*, 2015). Robust capuchin lineages with clades spanning multiple interfluves would provide evidence against rivers as significant isolating mechanisms.

RESULTS

Capuchin phylogenetics

We find strong support for reciprocally monophyletic clades of gracile and robust capuchins in the phylogenetic reconstruction from MRBAYES (Fig. 3a). Our species time tree from BEAST (see Figure S1 in Appendix S3) estimates the mean divergence time between *Sapajus* and *Cebus* at 5.8 Ma [95% higher posterior density (HPD) = 2.24–10.94] (late Miocene).

For robust capuchins we recover strong support for the clades S. xanthosternos [posterior probability (pp) = 1] and S. nigritus (pp = 1), but little evidence for reciprocally monophyletic 'species' clades for the rest of Sapajus. Two S. robustus samples form a weakly supported clade (pp = 0.74). All other robust capuchins form one large widely distributed clade ('Sapajus CLADE 4,' pp = 1), comprising S. apella, S. macrocephalus, S. cay, S. libidinosus and S. flavius (Fig. 3b). Subclades within this group are not always congruent with morphological hypotheses about robust capuchin species, a result concordant with the mitochondrial results from Ruiz-García et al. (2012, 2016). Most strikingly we find S. apella sensu stricto to be extensively paraphyletic with respect to other Sapajus species. Within the widespread clade, we recover several geographical subclades: (1) S. flavius (pp = 0.95); (2) S. libidinosus together with eastern S. apella (pp = 0.80); (3) S. cay from Paraguay and Rondônia/Mato Grosso plus S. apella from Guaporé and Apuí (pp = 0.81); (4) S. cay from Chapada dos Guimarães and S. apella from Alta Floresta, both in Mato Grosso, Brazil (pp = 1); (5) S. apella/macrocephalus (pp = 1) from north of the Amazon and Solimões rivers, as well as from regions just south of the Amazon River near the Xingu and Tapajos rivers, and just south of Manaus, together reaching all four Amazonian quadrants, as far northwest as Colombia; (6) S. apella/macro*cephalus* (pp = 0.95) from Brazil and Peru. Minimum clade ranges for each of the six subclades of the widespread Sapajus clade are geographically coherent, although several do not match the current morphological hypotheses about robust capuchin taxonomy (Fig. 3c).

Within Cebus, we recover eight strongly supported clades: Cebus versicolor forms a clade (pp = 1) with C. cesarae within the Magdalena river valley in the Colombian Andes, as part of a larger clade including C. capucinus in Panama and Colombia (pp = 1), and C. imitator in Central America (pp = 1). In the eastern Andes, C. brunneus (pp = 1) is sister to C. leucocephalus (pp = 1). Within Amazonia, there is strong support for a C. unicolor + C. albifrons clade (pp = 1). C. yuracus forms a clade with one C. unicolor sample (pp = 1) in western Brazil and Ecuador, and C. o. olivaceus + C. o. castaneus + Cebus kaapori form a strongly supported clade (pp = 1). Our C. kaapori sample is from the holotype, collected by Queiroz (1992), from the type locality Chega-Tudo, in Carutapera Municipality, Maranhão State, Brazil. Here, we provide a correction to the original Queiroz (1992) for the coordinates of that locality, 02°20' S 46°05' W (see Table S1 in Appendix S1).

Reconstruction of capuchin biogeographical history

Model weights based on AICc favoured the DEC model (Table 1, Fig. 4). Under this scenario the ancestral capuchin

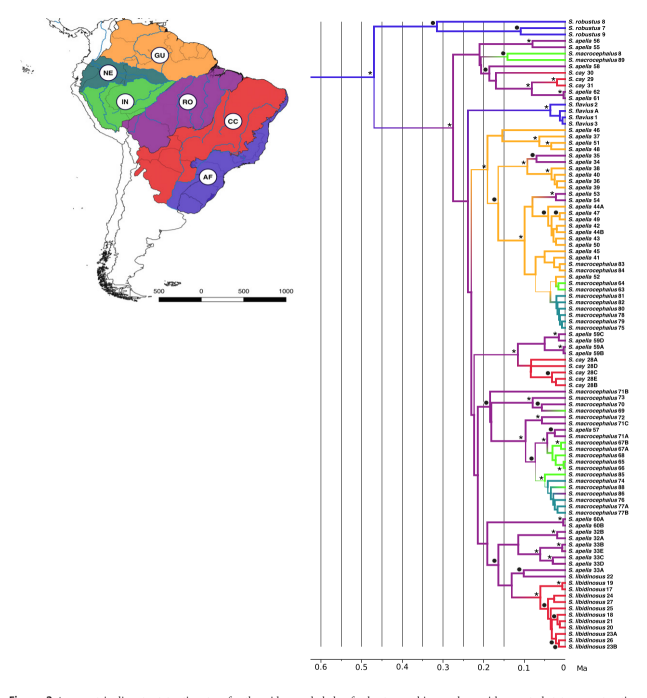


Figure 2 Asymmetric discrete states time tree for the widespread clade of robust capuchin monkeys with ancestral state reconstruction for regions of ancestral nodes and map delineating centres of endemism used for the BEAST phylogeography analyses: GU: Guianas; NE: Negro; RO: Rondonia; IN: Inambari; CC: Caatinga, Cerrado and Central Grasslands; AF: Atlantic Forest. Branches are colour-coded to regions from inset map. Posterior probabilities are shown as asterisks for above 0.95 and as circles for between 0.70 and 0.95.

population is inferred to have had a wide range that included both Amazonia and the Atlantic Forest. The divergence between *Sapajus* and *Cebus* is a vicariance event, where *Sapajus* is isolated in the Atlantic Forest, while the lineage leading to gracile *Cebus* is isolated in Amazonia. Crown *Sapajus* is recovered with a distribution restricted to the Atlantic Forest, with subsequent rapid expansion to the Caatinga/Cerrado/Central Grasslands and Amazonia regions at *c*. 500 ka. This means that *Cebus*, established in Amazonia at *c*. 2.4 Ma, diversified for *c*. 2 Myr before the appearance of any *Sapajus* clade in that region.

For crown *Cebus*, a single expansion event to the Central America/Andes region occurred at *c*. 2.2 Ma, with subsequent range contractions to just Central America/Andes for

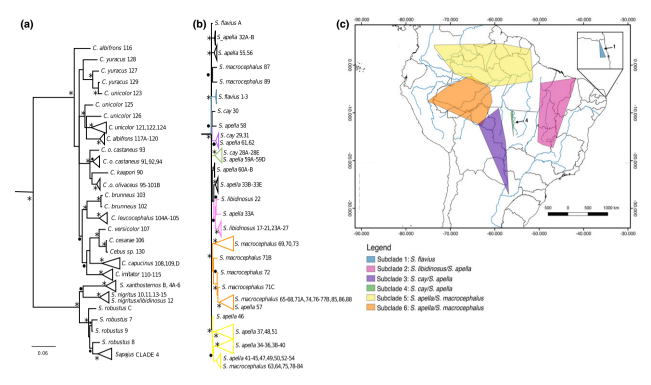


Figure 3 Bayesian consensus tree for capuchin phylogeny: (a) *Cebus* and *Sapajus* phylogeny and (b) details of *Sapajus* CLADE 4; (c) map with minimum convex polygons to show geographical distribution of major subclades within the widespread *Sapajus* CLADE 4. Subclades are presented as minimum convex polygons that connect sample localities. Branches are colour coded to regions from polygons map. Posterior probabilities are shown as asterisks for above 0.95 and as circles for between 0.70 and 0.95. Sample species ID and number correspond to Table S1 in Appendix S1.

the lineage leading to *C. capucinus, C. imitator* and *C. versicolor*, and to Amazonia only for both the lineage leading to *C. brunneus* (note that the 'Amazonia' region extends to the northern Venezuelan coast; see Fig. 4), and for the lineage leading to *C. unicolor*.

Reconstruction of widespread Sapajus radiation

We do not find any statistical difference in support for the symmetric versus the asymmetric analyses (K \approx 0). Both the phylogeographical reconstructions detail the explosive

Table 1 Comparison between models estimated in 'BioGeoBEARS'. For each model implemented in the analysis: values of log-likelihood (LnL), numbers of parameters and Akaike's information criteria (AIC, Δ AIC and AICc weight) are shown.

Model	LnL	No. parameters	AICc	ΔAICc	AICc weight
DEC	-17.29	2	39.92	0	0.68
DIVALIKE	-18.9	2	43.14	3.22	0.14
BAYAREALIKE	-21.41	3	51.81	11.9	0
DEC+J	-17.19	3	43.38	3.46	0.12
DIVALIKE+J	-18.61	3	46.22	6.3	0.03
BAYAREALIKE+J	-18.61	3	46.22	6.3	0.03

Pleistocene range expansion throughout Amazonia, Cerrado, Caatinga and southern grasslands for Sapajus (Fig. 2 and see Figure S2 in Appendix S3). In the asymmetric reconstruction, robust capuchins enter Amazonia by a single expansion from AF to RO at around 300-500 ka, with a second unique expansion across the Amazon River to GU at 200 ka, and at least two independent expansions across the Madeira River from RO to IN. There is also evidence for expansion from GU to IN, and IN to GU, as well as back-crossing from both IN and GU to RO. No strong geographical pattern demarcates rivers as firm barriers separating the four Amazonian regions; instead, there has been significant and recurrent recent movement by Sapajus across the Amazon, Solimões, Madeira and Negro rivers. Sapajus also spread from RO three times into the Cerrado/Caatinga (CC) region, all within the last 200 ka.

The addition of a distribution partition to our BEAST analysis provides support for the monophyly of *Sapajus robustus* (0.93) and its position as the sister group to a widespread Amazonian-Grasslands-Atlantic Forest clade (pp = 1.0). *Sapajus robustus* is restricted to the Atlantic Forest, making it clear that the explosive radiation of robust capuchin monkeys began with a recent migration into Amazonia and grasslands.

A significant difference between our asymmetric and symmetric BEAST analyses is related to *S. flavius*. In the

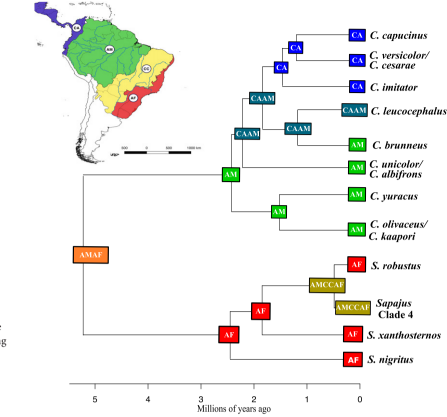


Figure 4 'BioGeoBEARS' ancestral range reconstruction of capuchin monkeys using the DEC model. CA = Central America/ Andes; AM = Amazonia; CC = Caatinga, Cerrado and Central Grasslands; AF = Atlantic Forest. Inset map for four regions of analysis.

symmetric BEAST phylogeography analysis, *S. flavius* is recovered as the sister group to *Sapajus* CLADE 4 (see Figure S2 in Appendix S3). On the other hand, *S. flavius* is embedded within the widespread clade in the asymmetric analysis (Fig. 2), suggesting an expansion back into the Atlantic Forest from the Amazonia/Grasslands regions.

DISCUSSION

Our expanded sample set with comprehensive representation of capuchin species provides further support for the monophyly of robust and gracile capuchins with a split at 5.8 Ma. A late Miocene Sapajus-Cebus split was previously supported using combined nuclear and mitochondrial data in Perelman et al. (2011) (6.0 Ma, 95% confidence interval 3.13-9.35 Ma) and Springer et al. (2012) (5.7 Ma average, 3.55-8.33 Ma composite 95% min-max). However, these time trees did not include all capuchin species; here, we add C. kaapori, C. versicolor and S. flavius. One exception to monophyly of Cebus and Sapajus has been recently reported in Ruiz-García et al. (2016), where some Cebus albifrons sensu lato individuals from the Colombian Eastern Llanos and Ecuadorian Amazonia displayed Sapajus mitochondrial haplotypes, suggesting the possibility of localized past or present introgression across these two genera. Unfortunately, neither sequences nor morphological data are publicly available for any of these individuals at this time.

Our biogeographical analyses reconstruct a widespread capuchin ancestral distribution across forested areas of South America, from Amazonia to the Atlantic Forest, supporting Amazonia versus Atlantic Forest vicariant origin for *Cebus* and *Sapajus*. The distribution of *Sapajus* exclusively in the Atlantic Forest until recently, and *Cebus* exclusively in the Amazonia and Central America/Andes regions from early on, suggests a cladogenetic event (vicariance) led to their initial diversification in isolation.

Throughout capuchin evolution, the Brazilian Shield (within AF, CC, RO Regions) and Guiana Shield (within GU) have been constant highland regions undergoing few physical changes in configuration but perhaps experiencing changes in floral assemblages related to climate (Aleixo & Rossetti, 2007). The divergence between Sapajus and Cebus, dated consistently at c. 6 Ma, coincides with floral assembly of the Cerrado (4-8 Ma), as indicated by the onset of diversification of many Cerrado plant lineages (Simon et al., 2009). The coincidence in timing of the divergence between robust and gracile capuchins and of the Cerrado assemblage suggests that the development of the Cerrado may have acted as a vicariant agent, especially given the reconstructed distributions of ancestral Sapajus (Atlantic Forest) and Cebus (Amazonia). The only extant capuchins distributed in CC belong to our widespread Sapajus CLADE 4 - which invaded those biomes only in the last 500 kyr, and have recent cranial and post-cranial adaptations to drier habitats (Wright et al., 2015).

Combining information from our biogeographical and phylogeographical analyses, we find strong evidence that all of the Atlantic Forest robust capuchin species are monophyletic clades, with the exception perhaps of *S. flavius*, which was found as part of the widespread *Sapajus* 'clade 4' in our MRBAYES phylogeny. Major Amazonian rivers do not appear to be strong barriers for dispersal or gene flow for *Sapajus*, with up to 12 examples of dispersal across significant Amazonian rivers demonstrated through our BEAST analysis by just the sampling included in this study.

Gracile capuchin biogeography and phylogenetics

In our analysis, Cebus shows strong geographical divisions, suggesting a widespread ancestor diversified in Amazonia and was split by range expansion and subsequent range contraction in the Andes and Central America. The Amazonian group (C. yuracus, C. unicolor, C. albifrons, C. olivaceus and C. kaapori) is about the same age as the CA group (C. capucinus, C. imitator, C. versicolor, C. cesarae, C. leucocephalus and C. brunneus). The recently discovered C. kaapori was initially considered to be a subspecies of C. olivaceus (Harada & Ferrari, 1996), but other morphological taxonomists have argued that it is a unique species (Groves, 2001, 2005; Silva-Júnior, 2001), with some morphological characteristics more similar to C. albifrons sensu lato (Masterson, 1995). In our MRBAYES analysis, C. kaapori is recovered within the strongly supported C. olivaceus clade. A close relationship between C. kaapori and C. olivaceus makes sense, as C. o. castaneus is the gracile capuchin geographically most proximate to C. kaapori. Cebus versicolor is recovered as sister to C. cesarae. This is concordant with geographical distribution as both species are found in isolated pockets of the Magdalena Valley in Colombia. MtDNA COII analysis of Cebus albifrons sensu lato by Ruiz-García et al. (2010) found C. cesarae and C. pleei as sister to C. versicolor.

Robust capuchin biogeography and phylogenetics

Our phylogeographical analysis suggests robust capuchins were able to colonize an array of divergent habitat types all within a very short time period, in the last 1 Myr. Robust capuchins show evidence of multiple moves within and between all regions, except that they are absent from the Central American and Andean region. This may be explained because their arrival at the foot of the eastern Andes was so recent that the mountain range was already fully formed and impassable (Lynch Alfaro *et al.*, 2012a).

Consistent with this recent explosive radiation, we find little support for the internal topology of a morphologically diverse and geographically expansive clade of robust capuchins. Morphological species contained in this group include *S. libidinosus*, *S. cay*, *S. flavius*, *S. macrocephalus* and *S. apella*. While there is significant individual variation within *Sapajus* populations, these five morphological species each display distinct morphological patterns with clear geographical correspondence (Silva-Júnior, 2001), possibly as a consequence of repeated founder effects. While the recent divergence times among all the Amazonian and open region grasslands capuchins (all under 400 ka) suggest there probably has not been time enough for speciation among these morphological types, our data do point to geographical clades that deserve further study for their morphological and behavioural population characteristics. For example, morphological differences between S. macrocephalus north and south of the Amazon River have already been described (Rylands et al., 2013); our analysis here provides new hypotheses for geographical boundaries between major population centres within Amazonian Sapajus, and suggests the need for new morphological and morphometric analyses for undersampled regions. We suggest nuclear genomic methods will be necessary to provide a complementary perspective to help resolve the relationships within this clade.

Rapid Pleistocene expansion throughout Amazonia has been recovered for the sister taxon to capuchin monkeys, the squirrel monkeys (Chiou et al., 2011; Lynch Alfaro et al., 2015a). Sapajus presence today in western Amazonia is explained in the BEAST phylogeography analysis as resulting from two distinct pathways: one was expanding from RO west across the Madeira River, into IN, and then spreading north into NE. The second was a move first from RO across the Amazon River into GU, then expansion west across the Negro River into NE, and from there south to IN. The exact distribution of S. macrocephalus in western Amazonia is contested (Groves, 2001, 2005; Silva-Júnior, 2001). Rylands et al. (2013) depict S. macrocephalus morphological diversity with Brazilian, Peruvian, Bolivian and Colombian forms; one explanation for this variation congruent with our results is that western Amazonia has been colonized multiple times from different Sapajus seed populations from different parts of Amazonia. In contrast, most of Sapajus diversity in eastern Amazonia can be explained by a single founder event from AF into RO, and another single founder event from RO into GU, with some later expansions back to RO from GU.

Sapajus cay has been the subject of controversy within robust capuchin taxonomy, and two major authorities (Groves, 2001, 2005; Silva-Júnior, 2001) disagree significantly about its distribution, in part due to differences in the localities of samples available in each of their studies. Sapajus cay (sensu Rylands et al., 2013) is recovered here as paraphyletic, with evidence for at least two distinct Sapajus populations moving independently into gallery forests in more open habitat types. Sapajus cay samples from Paraguay, Rondônia and Mato Grosso cluster with S. apella from Rondônia; while S. cay from northern central Mato Grosso forms a clade with nearby S. apella from Alta Floresta. These two populations have apparently converged morphologically to a lighter pelage phenotype in the drier, open regions. Sapajus libidinosus (sensu Rylands et al., 2013) marks a third expansion into CC from Amazonian Sapajus, with a resultant lighter pelage (Fig. 2). All expansions into drier habitats appear to have occurred in the last 200–100 kyr, similar to findings for other Neotropical primates (Lynch Alfaro *et al.*, 2015b).

We expected to recover S. flavius as sister to S. xanthosternos because of geographical proximity or sister to S. libidinosus based on a gradation of morphology between the two morphotypes (Silva, 2010); or possibly sister to all other Sapajus species, based on its unusual morphology (Lynch Alfaro et al., 2012a). In our BEAST phylogeography and MRBAYES analyses, S. flavius is recovered as part of the widespread Sapajus CLADE 4 (Figs 2 & 3b). However, in the symmetric BEAST phylogeography analysis, S. flavius is recovered as the sister group to Sapajus CLADE 4 (see Figure S2 in Appendix S3), suggesting all four Atlantic Forest Sapajus taxa may have diverged from each other first prior to Sapajus expansion into other regions. Given the difference in topology and the range of support values among analyses, the phylogenetic placement of S. flavius remains uncertain, although close phylogenetic affinity to Sapajus CLADE 4 is clear.

Sympatry between robust and gracile capuchins

Overall, our data and analyses support the 'ROA' hypothesis (Lynch Alfaro *et al.*, 2012a) by *Sapajus* as indicated by the ancestral reconstructions of the crown *Cebus* and crown *Sapajus* clades. The two capuchin genera diversify in isolation, with *Sapajus* species only later invading a *Cebus*-occupied Amazonia at *c*. 500 ka. The BEAST phylogeography analysis shows *Sapajus* invasion of Amazonia occurring first in Rondonia (south-western Amazon) and quickly spreading to all four quadrants of Amazonia. The recovered timing of entry into sympatry for gracile and robust capuchins indicate an event in Amazonia < 1 Ma, as a result of *Sapajus* counter-invasion into a region already occupied by *Cebus*.

ACKNOWLEDGEMENTS

Thanks to Robert Whittaker for inviting us to contribute this paper, and to Alexandre Antonelli, Marcelo Tejedor and anonymous referees for their excellent suggestions that have improved this article substantially. Special thanks to L.E. Olson and K. Fales, University of Alaska Museum. Support to M.G.M.L.'s PhD research was provided by a CNPq PhD fellowship (142141/2012-7) and CNPq SWE fellowship (201172/2014-3). Funding for this research was provided in part by NSF BCS 0833375 to J.W.L.A. Some of the molecular analyses and field expeditions were funded by CNPq/ FAPEAM SISBIOTA Program (563348/2010-0) and CAPES (AUX no. 3261/2013) to I.P.F. and CNPq (grants 'INCT em Biodiversidade e Uso da Terra da Amazônia' 574008/2008-0; and 310880/2012-2) and NSF-FAPESP 2012/50260-6 (grant 1241066 - Dimensions US-BIOTA-São Paulo: Assembly and evolution of the Amazonian biota and its environment: an integrated approach) to A.A. This is article number 24 of the Amazonian Mammals Research Group.

REFERENCES

- Aleixo, A. & Rossetti, D.D.F. (2007) Avian gene trees, landscape evolution, and geology: towards a modern synthesis of Amazonian historical biogeography? *Journal of Ornithol*ogy, 148, 443–453.
- Baele, G., Lemey, P., Bedford, T., Rambaut, A., Suchard, M.A. & Alekseyenko, A.V. (2012) Improving the accuracy of demographic and molecular clock model comparison while accommodating phylogenetic uncertainty. *Molecular Biology and Evolution*, 29, 2157–2167.
- Baele, G., Li, W.L.S., Drummond, A.J., Suchard, M.A. & Lemey, P. (2013) Accurate model selection of relaxed molecular clocks in Bayesian phylogenetics. *Molecular Biol*ogy and Evolution, **30**, 239–243.
- Boubli, J.P., Rylands, A.B., Farias, I.P., Alfaro, M.E. & Lynch Alfaro, J. (2012) *Cebus* phylogenetic relationships: a preliminary reassessment of the diversity of the untufted capuchin monkeys. *American Journal of Primatology*, 74, 381–393.
- Boubli, J.P., Ribas, C., Lynch Alfaro, J.W., Alfaro, M.E., Silva, M.N.F., Pinho, G.M. & Farias, I.P. (2015) Spatial and temporal patterns of diversification on the Amazon: a test of the riverine hypothesis for all diurnal primates of Rio Negro and Rio Branco in Brazil. *Molecular Phylogenetics and Evolution*, 82, 400–412.
- Buckner, J.C., Lynch Alfaro, J., Rylands, A.B. & Alfaro, M.E. (2015) Biogeography of the marmosets and tamarins (Callitrichidae). *Molecular Phylogenetics and Evolution*, 82, 413–425.
- Chiou, K.L., Pozzi, L., Lynch Alfaro, J.W. & Di Fiore, A. (2011) Pleistocene diversification of living squirrel monkeys (*Saimiri* spp.) inferred from complete mitochondrial genome sequences. *Molecular Phylogenetics and Evolution*, 59, 736–745.
- Cracraft, J. (1985) Historical biogeography and patterns of differentiation within the South American avifauna: areas of endemism. *Ornithological Monographs*, **36**, 49–84.
- Drummond, A.J., Suchard, M.A., Xie, D. & Rambaut, A. (2012) Bayesian phylogenetics with BEAUti and the BEAST 1.7. *Molecular Biology and Evolution*, **29**, 1969–1973.
- Groves, C.P. (2001) *Primate taxonomy*. Smithsonian Institution Press, Washington, DC.
- Groves, C.P. (2005) Order Primates. Mammal species of the world: a taxonomic and geographic reference (ed. by D.E. Wilson and D.M. Reeder), pp. 111–184. Johns Hopkins University Press, Baltimore, MD.
- Harada, M.L. & Ferrari, S.F. (1996) Reclassification of Cebus kaapori Queiroz 1992 based on new specimens from eastern Pará, Brazil. Abstracts. XVIth Congress of the International Primatological Society, XIXth Congress of the American Society of Primatologists. pp. 729. Wisconsin Regional Primate Center, University of Wisconsin, Madison, WI, USA.

- Jameson Kiesling, N.M., Yi, S.V., Xu, K., Sperone, F.G. & Wildman, D. (2015) The tempo and mode of New World monkey evolution and biogeography in the context of phylogenomic analysis. *Molecular Phylogenetics and Evolution*, **82**, 386–399.
- Kay, R.F. (2015) Biogeography in deep time what do phylogenetics, geology and paleoclimate tell us about early platyrrhine evolution? *Molecular Phylogenetics and Evolution*, **82**, 358–374.
- Landis, M.J., Matzke, N.J., Moore, B.R. & Huelsenbeck, J.P. (2013) Bayesian analysis of biogeography when the number of areas is large. *Systematic Biology*, 62, 789–804.
- Lynch Alfaro, J.W., Boubli, J.P., Olson, L.E., Di Fiore, A., Wilson, B., Gutierrez-Espeleta, G.A., Chiou, K.L., Schulte, M., Neitzel, S., Ross, V., Schwochow, D., Farias, I., Janson, C. & Alfaro, M.E. (2012a) Explosive Pleistocene range expansion leads to widespread Amazonian sympatry between robust and gracile capuchin monkeys. *Journal of Biogeography*, **39**, 272–288.
- Lynch Alfaro, J.W., Silva-Júnior, J.S. & Rylands, A.R. (2012b) How different are robust and gracile capuchin monkeys? An argument for the use of *Sapajus* and *Cebus*. *American Journal of Primatology*, **74**, 273–286.
- Lynch Alfaro, J.W., Izar, P. & Ferreira, R.G. (2014) Capuchin monkey research priorities and urgent issues. *American Journal of Primatology*, **76**, 705–720.
- Lynch Alfaro, J.W., Boubli, J.P., Paim, F.P., Ribas, C.C., da Silva, M.N.F., Messias, M., Röhe, F., Mercês, M.P., Silva Júnior, J.S., Silva, C.R., Pinho, G.M., Koshkarian, G., Nguyen, M.T.T., Harada, M.L., Rabelo, R.M., Queiroz, H.L., Alfaro, M.E. & Farias, I.P. (2015a) Biogeography of squirrel monkeys (genus *Saimiri*): South-central Amazon origin and rapid pan-Amazonian diversification of a lowland primate. *Molecular Phylogenetics and Evolution*, **82**, 436–454.
- Lynch Alfaro, J.W., Cortés-Ortiz, L., Di Fiore, A. & Boubli, J.P. (2015b) Special issue: Comparative biogeography of Neotropical primates. *Molecular Phylogenetics and Evolution*, 82, 518–529.
- Martins, A.M.G., Jr, Amorim, N., Carneiro, J.C., Affonso, P.R.A.M., Sampaio, I. & Schneider, H. (2014) *Alu* elements and the phylogeny of capuchin (*Cebus* and *Sapajus*) monkeys. *American Journal of Primatology*, **77**, 368–375.
- Masterson, T.J. (1995) Morphological relationships between the Ka'apor capuchin (*Cebus kaapori* Queiroz 1992) and other male *Cebus* crania: a preliminary report. *Neotropical Primates*, **3**, 165–171.
- Matthews, L.J. (2012) Variations in sexual behavior among capuchin monkeys function for conspecific mate recognition; a phylogenetic analysis and a new hypothesis for female proceptivity in tufted capuchins. *American Journal of Primatology*, **74**, 287–298.
- Matzke, N. (2013) 'BioGeoBEARS': biogeography with Bayesian (and likelihood) evolutionary analysis in R Scripts. University of California, Berkeley, CA. R package, version

0.2.1. Available at: http://CRAN.R-project.org/package= BioGeoBEARS (accessed 27 July 2013).

- Matzke, N.J. (2014) Model selection in historical biogeography reveals that founder-event speciation is a crucial process in Island Clades. *Systematic Biology*, **63**, 951– 970.
- Mercês, M.P., Lynch Alfaro, J.W., Ferreira, W.A.S., Harada, M.L. & Silva Júnior, J.S. (2015) Morphology and mitochondrial phylogenetics reveal that the Amazon River separates two eastern squirrel monkey species: *Saimiri sciureus* and *S. collinsi. Molecular Phylogenetics and Evolution*, **82**, 426–435.
- Nascimento, F., Lazar, A., Seuánez, H. & Bonvicino, C. (2015) Reanalysis of the biogeographical hypothesis of range expansion between robust and gracile capuchin monkeys. *Journal of Biogeography*, **42**, 1349–1363.
- Oliveira, S.G., Lynch Alfaro, J.W. & Veiga, L.M. (2014) Activity budget, diet and habitat use in the critically endangered Ka'apor capuchin monkey (*Cebus kaapori*) in Pará State, Brazil: a preliminary comparison to other capuchin monkeys. *American Journal of Primatology*, **76**, 919–931.
- Perelman, P., Johnson, W.E., Roos, C., Seuánez, H.N., Horvath, J.E., Moreira, M.A.M., Kessing, B., Pontius, J., Roelke, M., Rumpler, Y., Schneider, M.P.C., Silva, A., O'Brien, S.J. & Pecon-Slattery, J. (2011) A molecular phylogeny of living primates. *PLoS Genetics*, 7, e1001342.
- Queiroz, H.L. (1992) A new species of capuchin monkey, genus *Cebus* Erxleben 1977 (Cebidae, Primates), from eastern Brazilian Amazonia. *Goeldiana Zoologia*, **15**, 1–3.
- Ree, R.H. & Smith, S.A. (2008) Maximum-likelihood inference of geographic range evolution by dispersal, local extinction, and cladogenesis. *Systematic Biology*, **57**, 4–14.
- Ree, R.H., Moore, B.R., Webb, C.O. & Donoghue, M.J. (2005) A likelihood framework for inferring the evolution of geographic range on phylogenetic trees. *Evolution*, **59**, 2299–2311.
- Robins, J.H., Tintinger, V., Aplin, K.P., Hingston, M., Matisoo-Smith, E., Penny, D. & Lavery, S.D. (2014) Phylogenetic species identification in *Rattus* highlights rapid radiation and morphological similarity of New Guinean species. *PlosOne*, **9**, e98002.
- Ronquist, F. (1997) Dispersal-vicariance analysis: a new approach to the quantification of historical biogeography. *Systematic Biology*, **46**, 195–203.
- Ronquist, F., Teslenko, M., van der Mark, P., Ayres, D.L., Darling, A., Höhna, S., Larget, B., Liu, L., Suchard, M.A. & Huelsenbeck, J.P. (2012) MRBAYES 3.2: efficient Bayesian phylogenetic inference and model choice across a large model space. *Systematic Biology*, 61, 539–542.
- Ruiz-García, M., Castillo, M.I., Vásquez, C., Rodríguez, K., Pinedo, M., Shostell, J. & Leguizamon, N. (2010) Molecular phylogenetics and phylogeography of the white-fronted capuchin (*Cebus albifrons*; Cebidae, Primates) by means of mtCOII gene sequences. *Molecular Phylogenetics and Evolution*, 57, 1049–1061.

- Ruiz-García, M., Castillo, M.I., Lichilín-Ortiz, N. & Pinedo-Castro, M. (2012) Molecular relationships and classification of several tufted capuchin lineages (*Cebus apella*, *Cebus xanthosternos* and *Cebus nigritus*, Cebidae), by means of mitochondrial cytochrome oxidase II gene sequence. Folia Primatologica, 83, 100–125.
- Ruiz-García, M., Castillo, M.I. & Luengas-Villamil, K. (2016) Is it misleading to use *Sapajus* (robust capuchins) as a genus? A review of the evolution of the capuchins and suggestions on their systematics. *Phylogeny, molecular population genetics, evolutionary biology and conservation of the Neotropical primates* (ed. by M. Ruiz-Garcia and J.M. Shostell). pp. 209–268. Nova Science Publisher Inc., New York.
- Rylands, A.B., Mittermeier, R.A., Bezerra, B.M., Paim, F.P. & Queiroz, H.L. (2013) Species accounts of Cebidae. *Hand*book of the mammals of the world: vol. 3. Primates (ed. by R.A. Mittermeier, A.B. Rylands and D.E. Wilson), pp. 390–413. Lynx Edicions, Barcelona.
- Schneider, H. & Sampaio, I. (2015) The systematics and evolution of New World primates – a review. *Molecular Phylogenetics and Evolution*, 82, 348–357.
- Silva, T.C.F. (2010) Estudo da variação na pelagem e da distribuição geográfica em Cebus flavius e Cebus libidinosus do Nordeste do Brasil. MSc Dissertation. Programa de Pós-Graduação em Ciências Biológicas – Zoologia, Universidade Federal da Paraíba, Paraíba, Brazil. 41p.
- Silva-Júnior, J.S. (2001) Especiação nos macacos-prego e caiararas, gênero Cebus Erxleben, 1777 (Primates, Cebidae). PhD Thesis, Universidade Federal do Rio de Janeiro, Rio de Janeiro, Brazil. 377p.
- Simon, M.F., Grether, R., Queiroz, L.P., Skema, C., Pennington, R.T. & Hughes, C.E. (2009) Recent assembly of the Cerrado, a Neotropical plant diversity hotspot, by in situ evolution of adaptations to fire. *Proceedings of the National Academy of Sciences USA*, **106**, 20359–20364.
- Springer, M.S., Meredith, R.W., Gatesy, J., Emerling, C.A., Park, J., Rabosky, D.L., Stadler, T., Steiner, C., Ryder, O.A., Janecka, J.E., Fisher, C.A. & Murphy, W.J. (2012) Macroevolutionary dynamics and historical biogeography of primate diversification inferred from a species supermatrix. *PlosOne*, 7, e49521.
- Tobe, S.S., Kitchener, A.C. & Linacre, A.M.T. (2010) Reconstructing mammalian phylogenies: a detailed comparison

of the Cytochrome *b* and Cytochrome Oxidase Subunit 1 mitochondrial genes. *PlosOne*, **5**, e14156.

- Wallace, A.R. (1852) On the monkeys of the Amazon. Proceedings of the Zoological Society of London, 20, 107–110.
- Ward, R.D., Zemlak, T.S., Innes, B.H., Last, P.R. & Hebert, P.D.N. (2005) DNA barcoding Australia's fish species. *Philosophical Transactions of the Royal Society B: Biological Sciences*, 360, 1847–1857.
- Wright, K.A., Wright, B.W., Ford, S.M., Fragaszy, D., Izar, P., Norconk, M., Masterson, T., Hobbs, D.G., Alfaro, M.E. & Lynch Alfaro, J.W. (2015) The effects of ecology and evolutionary history on robust capuchin morphological diversity. *Molecular Phylogenetics and Evolution*, 82, 455–466.
- Zhang, Y., Zhang, H., Gao, T. & Miao, Z. (2011) Structure of mitochondrial DNA control region and molecular phylogenetic relationship among three founders of genus *Pleuronectes. Biochemical Systematics and Ecology*, **39**, 627–634.

SUPPORTING INFORMATION

Additional Supporting Information may be found in the online version of this article:

Appendix S1 Details of individual and GenBank accession numbers.

Appendix S2 Supplementary details of methods. **Appendix S3** Supporting results.

BIOSKETCH

Marcela G. M. Lima is a biologist interested in biogeography, phylogenetics and population genetics, with a focus on Neotropical vertebrates.

Author contributions: M.G.M.L., J.C.B., M.E.A. and J.W.L.A. designed and ran phylogenetic and biogeographical analyses and developed the article; M.G.M.L., J.S.S.J., A.B.M., A.L., I.P.F., J.P.B., F.R., H.Q., M.N.S., A.D.F. and J.W.L.A. collected and curated sample material; M.G.M.L., A.B.M. and I.P.F. generated new genetic sequences; K.C., M.G.M.L., J.C.B. and J.W.L.A. designed maps and figures; and all authors contributed to the writing of the article.

Editor: Alexandre Antonelli