Mitogenomic Phylogeny of *Callithrix* with Special Focus on Human Transferred Taxa

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Abstract

Callithrix marmosets are a relatively young primate radiation, whose phylogeny is not yet fully resolved. These primates are naturally para- and allopatric, but three species with highly invasive potential have been introduced into the southeastern Brazilian Atlantic Forest by the pet trade. There, these species hybridize with each other and endangered, native congeners. We aimed here to reconstruct a robust Callithrix phylogeny and divergence time estimates, and identify the biogeographic origins of autochthonous and allochthonous Callithrix mitogenome lineages. We sequenced 49 mitogenomes from four species (C. aurita, C. geoffroyi, C. jacchus, C. penicillata) and anthropogenic hybrids (C. aurita x Callithrix sp., C. penicillata x C. jacchus, Callithrix sp. x Callithrix sp., C. penicillata x C. geoffroyi) via Sanger and whole genome sequencing. We combined these data with previously published Callithrix mitogenomes to analyze five Callithrix species in total.

Results

We report the complete sequence and organization of the C. aurita mitogenome. Phylogenetic analyses showed that C. aurita was the first to diverge within Callithrix 3.54 million years ago (Ma), while C. jacchus and C. penicillata lineages diverged most recently 0.5 Ma as sister clades. MtDNA clades of C. aurita, C. geoffroyi, and C. penicillata show intraspecific geographic structure, but C. penicillata clades appear polyphyletic. Hybrids, which were identified by phenotype, possessed mainly C. penicillata or C. jacchus mtDNA haplotypes. The biogeographic origins of mtDNA haplotypes from hybrid and allochthonous Callithrix were broadly distributed across natural Callithrix ranges. Our phylogenetic results also evidence introgression of C. jacchus mtDNA into C. aurita.

Conclusion

Our robust Callithrix mitogenome phylogeny shows C. aurita lineages as basal and C. jacchus lineages among the most recent within Callithrix. We provide the first evidence that parental mtDNA lineages of anthropogenic hybrid and allochthonous marmosets are broadly distributed inside and outside of the Atlantic Forest. We also show evidence of cryptic hybridization between allochthonous Callithrix and autochthonous C. aurita. Our results encouragingly show that further development of genomic resources will allow to more clearly elucidate Callithrix evolutionary relationships and understand the dynamics of Callithrix anthropogenic introductions into the Brazilian Atlantic Forest.

Keywords

mtDNA, marmoset, divergence, Atlantic Forest, Brazil, Callitrichidae, primate
Background

*Callithrix* species represent a relatively young radiation, and divergence among lineages within the genus is estimated to be between approximately 0.7 and 2.5 million years ago (Ma) [1, 2, 3]. Two major subgroups occur within the genus, the *aurita* group (*C. aurita/C. flaviceps*) and the *jacchus* group (*C. geoffroyi/C. kuhlii/C. jacchus/C. penicillata*), but the phylogeny of these lineages is not yet fully resolved. *Callithrix* species are naturally para- and allopatric across the Brazilian Atlantic Forest, Cerrado, and Caatinga biomes (Figure 1) [4,5], and natural hybridization occurs between some species [6]. However, *C. geoffroyi*, *C. jacchus*, and *C. penicillata* have high invasive potential [7, 8] and have spread widely outside of their native ranges due to the legal and illegal pet trades. These species have established several allochthonous populations in the southeastern Brazilian Atlantic Forest [6, 9, 10] and hybridize with other allochthonous and autochthonous congeners [6, 9, 11, 10], including endangered *C. aurita* and *C. flaviceps* [12, 13]. Yet, determining evolutionary relationships between autochthonous, allochthonous, and hybrid *Callithrix* populations across Brazil is complicated by the unresolved *Callithrix* phylogeny.

In general, mitochondrial DNA (mtDNA) can be utilized for an initial look into evolutionary relationships among taxa [e.g., 14, 15] as well as track dispersal and gene flow patterns of allochthonous species [16]. MtDNA sequence data can also provide initial genetic insight in the direction of introgression (if sex-biased) when two species hybridize due to incongruences between phenotypes and haplotypes [e.g., 14, 15]. The effective population size of mtDNA is one quarter of that of nuclear DNA from a diploid, bisexual population, which allows mtDNA lineages to coalesce relatively more quickly [17]. MtDNA is also considered a relatively fast mutating genetic marker [18]. As a result, lineage sorting and
reciprocal monophyly are expected to occur faster in mtDNA than nuclear DNA, which can provide insight into shallow evolutionary relationships expected for young radiations.

One major challenge in applying genetic and genomic methods in *Callithrix* studies is an overall lack of genomic resources and sample material for most *Callithrix* species. Studies of *Callithrix* species have utilized mtDNA markers that generally resulted in polytomies and/or poorly supported branching patterns, as well as polyphyly for *C. penicillata* and *C. kuhlii* [19,20,21,22,23]. Also, the few available genetic studies of allochthonous and hybrid *Callithrix* within the Atlantic Forest, all conducted within Rio de Janeiro state, used portions of mtDNA or the Y-chromosome that could not fully resolve the evolutionary relationships of *Callithrix* lineages [e.g., 11,23]. Nonetheless, [24] obtained a well-resolved phylogeny for the *jacchus* group using complete mitogenomes, but they only sampled one individual/species with unknown provenances.

To build upon the above previous *Callithrix* studies, we have conducted the largest to-date geographical sampling of *Callithrix* mitogenomes across Brazil (Figure 1) with the following aims: (1) improve resolution of phylogenetic relationships and divergence times estimates between *Callithrix* mtDNA haplotypes; (2) determine which *Callithrix* mtDNA lineages are autochthonous across *Callithrix* ranges; and (3) identify allochthonous *Callithrix* mtDNA lineages in the southeastern Atlantic Forest and their possible biogeographic origins. We sequenced, for the first time, the complete mitogenome of *C. aurita*, and in total obtained 49 new mitogenome sequences from four species (*C. aurita*, *C. geoffroyi*, *C. jacchus*, *C. penicillata*), and four hybrid types (*C. aurita x Callithrix* sp., *C. penicillata x C. jacchus*, *Callithrix* sp. x *Callithrix* sp., *C. penicillata x C. geoffroyi*) for these analyses.
Table 1. Number of *Callithrix* specimens newly sampled by species and hybrid phenotype. Provenance abbreviations are: CRC-Callitrichid Research Center, NEPRC- New England Primate Research Center (no longer in operation), CPRJ-Centro de Primatologia do Rio de Janeiro, and CEMAFUNA-Centro de Conservação e Manejo de Fauna da Caatinga. Letters in parentheses next to numerical values listed in the “N” column for hybrid marmosets correspond to likely maternal species of each hybrid based on phylogenetic analyses presented in Figures 2, and S1-S3. Maternal species abbreviations are- A: *C. aurita*, G: *C. geoffroyi*, J: *C. jacchus*; and P: *C. penicillata*.

| Phenotype | Provenance | Approximate Geographic Coordinates | N |
|-----------|------------|-----------------------------------|---|
| *C. aurita* | Guiricema, Minas Gerais, Brazil | -21.0081075, -42.7231066 | 2 |
| *C. aurita* | Guarulhos Municipal Zoo, Guarulhos, São Paulo, Brazil (wild marmoset originally from Mogi das Cruzes, São Paulo) | -23.5391655, -46.1960017 | 1 |
| *C. aurita* | Guarulhos Municipal Zoo, Guarulhos, São Paulo, Brazil | -23.4425682, -46.5535750 | 2 |
| *C. aurita* | Guarulhos Municipal Zoo, Guarulhos, São Paulo, Brazil (wild marmoset originally from São José dos Campos, São Paulo) | -23.1701146, -45.8938482 | 1 |
| *C. aurita* | CPRJ, Guapimirim, Rio de Janeiro, Brazil (wild marmosets originally from Natividade, Rio de Janeiro) | -21.0605955, -41.9771042 | 2 |
| *C. aurita* | CPRJ, Guapimirim, Rio de Janeiro, Brazil | -22.4885811, -42.9136052 | 1 |
| *C. geoffroyi* | Callitrichid Research Center, Omaha, Nebraska, US | -41.2549350, -95.9768850 | 1 |
| *C. geoffroyi* | CPRJ, Guapimirim, Rio de Janeiro, Brazil | -22.4885811, -42.9136052 | 1 |
| *C. geoffroyi* | Berilo, Minas Gerais, Brazil | -16.9307333, -42.4849902 | 1 |
| *C. geoffroyi* | Serra, Espírito Santo, Brazil | -20.1903636, -40.2374422 | 3 |
| *C. jacchus* | Guarulhos Municipal Zoo, Guarulhos, São Paulo, Brazil | -23.4425682, -46.5535750 | 2 |
| *C. jacchus* | NEPRC, Southborough, Massachusetts, US | 42.2996480, -71.5358183 | 1 |
| *C. jacchus* | CEMAFUNA, Petrolina, Pernambuco, Brazil | -9.3272051, -40.5441172 | 4 |
Using Illumina whole genome sequencing (WGS) and Sanger sequencing approaches, we sequenced complete mitogenomes from 49 *Callithrix* (Figure 1, Table 1, and Table S1). We combined these new mitogenomes with previously published primate mitogenome sequences for downstream analyses (listed in Table S1). The length of the resulting sequence alignment after combining all of these mitogenomes was 17,132 bases. Sampled individuals that possessed the same mtDNA haplotypes are listed in Table S2. The organization of the *C. aurita* mitogenome was consistent with previously published *Callithrix* mitogenomes from [24]. This mitogenome includes 12 protein-coding genes, two rRNAs, and 14 tRNAs on the heavy strand and one protein-coding gene and eight tRNAs on the light strand, as well as the

| Species | Location | Coordinates | Haplotypes |
|---------|----------|-------------|------------|
| *C. jacchus* | CPRJ, Guapimirim, Rio de Janeiro, Brazil | -22.4885811, -42.9136052 | 1 |
| *C. penicillata* | Brasília, Federal District, Brazil | -15.8014526, -47.9234345 | 2 |
| *C. penicillata* | Lavras, Minas Gerais, Brazil | -21.2292301, -44.9988343 | 2 |
| *C. penicillata* | Belo Horizonte, Minas Gerais, Brazil | -19.8703102, -43.9696589 | 2 |
| *C. penicillata* | CEMAFUANA, Petrolina, Pernambuco, Brazil | -9.3272051, -40.5441172 | 2 |
| *C. jacchus* x *C. penicillata* | CPRJ, Guapimirim, Rio de Janeiro, Brazil | -22.4885811, -42.9136052 | 1 (P) |
| *C. penicillata* x *C. Geoffroyi* | Viçosa, Minas Gerais, Brazil | -20.7639782, -42.8990971 | 11 (all P) |
| *C. aurita* x Callithrix sp. | Guarulhos Municipal Zoo, Guarulhos, São Paulo, Brazil | -23.4425682, -46.5535750 | 3 (J,G,P) |
| *C. aurita* x Callithrix sp. | Guarulhos Municipal Zoo, Guarulhos, São Paulo, Brazil (wild marmoset originally from Maripora, São Paulo) | -23.3412818, -46.5754474 | 1 (J) |
| *C. aurita* x Callithrix sp. | CPRJ, Guapimirim, Rio de Janeiro, Brazil | -22.4885811, -42.9136052 | 1 (A) |
| *C. Geoffroyi* x Callithrix sp. | Santa Teresa, Espirito Santo, Brazil | -19.9394185, -40.6009700 | 1 (G) |
| **Total** | | | 49 |
control region (Table S3). The length of the C. aurita mitogenome presented in Table S3 was 16,471 bases.

Phylogenetic Trees and Divergence Times of Callithrix Mitochondrial Clades

Maximum-likelihood (ML) and Bayesian inference produced well-supported phylogenetic trees that show mostly congruent phylogenetic relationships between the aurita and jacchus groups (Figure 2, Figures S1-S3). The main difference in the topology of the ML and Bayesian trees was in grouping patterns of some haplotypes within the C. jacchus clade described below. A number of nodes in the ML tree possessed 100% bootstrap support but most had bootstrap scores of >70% (Figure S1). Most nodes in the Bayesian trees had posterior probabilities of 1 (Figure 2, Figures S2-S3). Major node names and divergence times within and outside the Callithrix clade are shown in Figure 2, Figure S3, Table 2, and Table S4.

Callithrix diverged from Cebuella approximately 6.83 Ma (Figure 2 node E) and the initial split within Callithrix, separating C. aurita and the jacchus group, occurred approximately 3.54 Ma (Figure 2 node D) (Table 2). Thus, C. aurita formed the Callithrix basal clade, and C. geoffroyi formed the most basal clade within the jacchus group by arising 1.18 Ma (node C). Callithrix penicillata haplotypes grouped into three polyphyletic clades that corresponded to three different biome regions, an Atlantic Forest-Cerrado transition area, Cerrado, and Caatinga. The first of these C. penicillata clades to diverge after C. geoffroyi was the Atlantic Forest-Cerrado transition clade at 0.92 Ma. Afterward, the C. penicillata Cerrado clade appeared at 0.87 Ma, followed by the C. kuhlii clade at 0.82 Ma (Figure 2 node
Table 2. Divergence times in million years (Ma) for *Callithrix* species and select nodes

(MRCA = Most recent common ancestor; values in brackets = 95% highest posterior density).

Node names follow major node destinations shown in Figure 2 in capital letters.

| Node | Taxa Diverging at Node | Age (Ma)       |
|------|------------------------|----------------|
| A    | *C. jacchus* - *C. penicillata* (Caatinga clade) | 0.51 [0.35-0.69] |
| B    | *C. kuhlii* - (*C. jacchus* + *C. penicillata* (Caatinga clade)) | 0.82 [0.59-1.09] |
| C    | *C. geoffroyi* - (*C. kuhlii* + (*C. jacchus* + *C. penicillata*)) | 1.18 [0.87-1.58] |
| D    | *C. aurita* - (*C. geoffroyi* + (*C. kuhlii* + (*C. jacchus* + *C. penicillata*))) | 3.54 [2.37-4.88] |
| E    | *Callithrix* - *Cebuella/Mico* | 6.83 [4.86-9.39] |

B). The *C. penicillata* Caatinga clade and the *C. jacchus* clades represent the two youngest clades within the phylogeny, splitting about 0.51 Ma (Figure 2 node A). As the *C. jacchus* clade showed some of the shallowest branch tips among *Callithrix* haplotypes and poor phylogenetic resolution, a ParsimonySplits network was constructed for haplotypes within this clade (Figure 3).

**Ancestral Origins and Biogeography of Callithrix Mitogenomes**

The ancestral origins of *Callithrix* phylogenetic mitogenome clades and subclades based on BMM biogeographic analysis were largely concordant with the assigned Brazilian states and regions of origin of sampled mitogenomic haplotypes (Figure 4 and Table S5). BMM analyses resulted in > 70% posterior probability of an ancestral origin for Node 93, which represented the basal node of the *C. aurita* clade, in Rio de Janeiro state. Within the *C. aurita* clade, node 92 showed >97% posterior probability of an ancestral origin of Rio de Janeiro state for two haplotypes sampled within this region from *C. aurita*-phenotype individuals and a *C. aurita x Callithrix* sp. hybrid. On the other hand, BMM analysis for
nodes 89-91, which represent the other *C. aurita* subclade, assigned posterior probabilities between 44%-65% for an origin of the Minas Gerais state portion of the natural *C. aurita* range. These haplotypes were obtained from *C. aurita*-phenotype individuals sampled in Minas Gerais, São Paulo, and Rio de Janeiro states, as well as a *C. aurita* x *Callithrix* sp. hybrid from São Paulo state.

Node 87 in Figure 4 represents the basal node of the *C. geoffroyi* clade, and BMM analyses calculated a collective posterior probability of over 75% of this clade originating within the natural range of *C. geoffroyi*. With a BMM posterior probability of 91.93% that node 85 originated in southeastern Espírito Santo state, biogeographic analysis accurately reflected the sampling origin of haplotypes BJT70 and BJT169. These haplotypes come from *C. geoffroyi*-phenotype individuals, as well as one *Callithrix* sp. x *Callithrix* sp. hybrid. For the other *C. geoffroyi* subclade, BMM analyses posterior probabilities support an ancestral origin of associated haplotypes within the natural distribution of *C. geoffroyi*.

For the three *C. penicillata* clades, BMM analysis showed high posterior probabilities for each clade’s corresponding geographic area as also being each respective clade’s ancestral region. Nodes 79-81 (Figure 4), which represent the *C. penicillata* Atlantic Forest-Cerrado transition clade, each possessed > 98% posterior probabilities of originating in the Atlantic Forest-Cerrado transition zone of Minas Gerais. This clade contained several haplotypes from *C. penicillata*-phenotype individuals sampled in this transition zone, as well as a hybrid sampled in São Paulo state. BMM posterior probability for the central Brazil Cerrado being the ancestral region for node 77 (Figure 4), which encompassed the *C. penicillata* Cerrado clade, was 98.16%. The *C. penicillata* Cerrado clade included haplotypes from *C. penicillata*-phenotype individuals sampled in Brasília. Finally, nodes 69-73 (Figure 4), representing the *C. penicillata* Caatinga clade, possessed BMM posterior probability support between
96.14%-99.60% for the Caatinga of Bahia state as the ancestral region of this clade. The clade contained haplotypes from *C. penicillata*-phenotype animals sampled at CEMAFUNA. Two haplotypes, representing eleven *C. penicillata* × *C. geoffroyi* hybrids sampled in Viçosa as well as a *C. jacchus* × *C. penicillata* hybrid, clustered within the *C. penicillata* Caatinga group.

BMM biogeographic analysis of the *C. jacchus* clade calculated high posterior probability (>99%) that haplotypes associated with nodes 60-64 originated in Ceará and/or Pernambuco states, regions whose dominant biome is the Caatinga. These haplotypes were obtained from marmosets with *C. jacchus* phenotypes sampled at CEMAFUNA and the Guarulhos Zoo, as well as three *C. aurita* phenotype individuals sampled within São Paulo. For nodes 65-68, BMM analyses calculated posterior probabilities of the associated haplotypes originating first from Pernambuco, and then from Ceará and/or Pernambuco. In particular, nodes 66 and 67 had respective posterior probabilities of 49.68% and 76.88% of originating in Pernambuco state. Haplotypes associated with these nodes came from a *C. aurita* × *Callithrix* sp. hybrid sampled in São Paulo state and a CEMAFUNA *C. jacchus-*phenotype individual.

**Genetic Distance between Callithrix Phylogenetic Clades**

Pairwise genetic distances between the above established phylogenetic clades are shown in Table 3 as measures of $D_{xy}$. The *C. aurita* clade was the most genetically distant from all other *Callithrix* clades, with $D_{xy}=0.055-0.056$. The smallest genetic distance can be observed between *C. jacchus* and the *C. penicillata* Caatinga clade at $D_{xy}=0.009$. The remaining pairwise genetic distances varied between $D_{xy}=0.013-0.015$, but the *C. geoffroyi* clade was the most distant relative to all other *jacchus* group clades.
Table 3. Array of pairwise $D_{xy}$ genetic distances between *Callithrix* phylogenetic clades.

|                  | C. jacchus |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
|------------------|------------|--|---|---|---|---|---|---|---|---|---|---|---|---|---|---|
| **C. penicillata** (Caatinga clade) | 0.009 | C. penicillata (Caatinga clade) |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| **C. kuhlii**     | 0.014 | 0.013 | C. kuhlii  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| **C. penicillata** (Cerrado clade) | 0.016 | 0.015 | 0.014 | C. penicillata (Cerrado clade) |  |  |  |  |  |  |  |  |  |  |  |  |
| **C. penicillata** (Atlantic Forest/ Cerrado Transitional clade) | 0.015 | 0.016 | 0.014 | 0.015 | C. penicillata (Atlantic Forest/ Cerrado Transitional clade) |  |  |  |  |  |  |  |  |  |  |  |
| **C. geoffroyi**  | 0.018 | 0.017 | 0.017 | 0.018 | 0.017 | C. geoffroyi |  |  |  |  |  |  |  |  |  |  |
| **C. aurita**     | 0.056 | 0.056 | 0.055 | 0.056 | 0.055 | 0.056 |  |  |  |  |  |  |  |  |  |  |

**Discussion**

*Callithrix Mitochondrial Phylogenetic Relationships and Divergence Times*

Our ML and Bayesian phylogenies were generally well supported and corroborated *Callithrix* divergence patterns from previous nuclear and mtDNA studies [3,24,25,26]. In ours and these previous phylogenies, the *C. aurita* clade was the most basal within the genus, the *C. geoffroyi* clade was most basal within the *jacchus* group, and *C. penicillata* and *C. jacchus* was the most recently diverged sister clade. Finally, our mtDNA analysis also showed that *C. penicillata* mitochondrial clades are polyphyletic, similar to the results obtained by [21] and [23]. The latter two studies also showed that *C. kuhlii* mitochondrial clades are polyphyletic.

Given the recent divergence times of *Callithrix* species, *Callithrix* polyphyly may be explained by incomplete lineage sorting when ancestral polymorphisms at a given locus are not fixed before population divergence [27]. Another possibility to explain *Callithrix* polyphyly may be due to past hybridization between these species and other *Callithrix* taxa or
perhaps recent migrations of *C. kuhlii* outside of its native range. However, we do not believe it is likely for any of these cases. For these alternative scenarios, we would expect to find at least some instances of allochthonous *C. kuhlii* mtDNA lineages, which to our knowledge have not yet been reported. Additionally, we did not observe any discordance involving *C. kuhlii* genotypes/phenotypes with that of any other *Callithrix* species, nor did mitogenome haplotypes from any hybrids sampled in this or previous studies group with *C. kuhlii* phylogenetic clades. Finally, locations where we sampled *Callithrix* species within native ranges were far removed from any natural hybridization zones with *C. kuhlii*, so natural secondary contact between *C. kuhlii* and other *Callithrix* species was unlikely at our sampling locations. Thus, incomplete lineage sorting is the most parsimonious explanation for *Callithrix* polyphyly observed in this and previous *Callithrix* studies.

The *Callithrix* divergence time estimates from our study, being between approximately 0.5 and 6.8 Ma, are within the range of previously published estimates [6,25,26]. These time estimates place the divergence of *Callithrix* species into the Pleistocene. In this epoch, climatic oscillations that may have promoted para- and allopatric speciation in South America, including that of *Callithrix* species, through repeated contractions and expansions of forested refuge [28,29].

**Biogeography Origins of Autochthonous and Allochthonous Callithrix Mitogenomes**

Previous phylogenetic and biogeographic analyses support a *Callithrix* origin in the southeastern Brazilian Atlantic Forest and then a northward expansion [3,4,24,25,26]. Our biogeographic analysis of *Callithrix* mitogenome lineages reflected similar biogeographic patterns for *Callithrix* species, as well as a natural geographic separation of major phylogenetic clades. When considering sampled mitogenome lineages of known provenance,
the biogeographic origins of our reconstructed phylogenetic clades was strongly influenced by the geographic origin of our samples across natural *Callithrix* ranges. For example, within the *C. geoffroyi* clade, the Minas Gerais state lineage formed a separate clade from the Espírito Santo state lineages. *Callithrix penicillata* possesses the largest natural geographic distribution of all *Callithrix* species [5] and biogeographic origins of paraphyletic clades were defined by where samples were collected within the Cerrado, Atlantic Forest, and Caatinga biomes. Although most sampled *C. jacchus* clade haplotypes did not possess known provenance and showed shallow tips, biogeographic analyses showed strong evidence that these haplotypes likely originated from within the Caatinga biome. Further evidence that *C. jacchus* mtDNA lineages tend to group geographically under denser sampling was shown by [23] with a geographically broader sampling of *C. jacchus* mtDNA D-loop sequences.

Allochthonous *Callithrix* species began appearing in portions of the southeastern Brazilian Atlantic Forest within approximately the last 20-30 years [30,31, pers. obs. C. Igayara]. Our biogeographic analyses can be used to infer the probable origins of parental populations of these allochthonous *Callithrix* species and anthropogenic hybrid *Callithrix* found in the southeastern Brazilian Atlantic Forest. Overall, biogeographic patterns show that the parental populations of these *Callithrix* likely possess multiple geographic origins from within and outside Atlantic Forest. For example, our biogeographic results show that mtDNA haplotypes of three *C. aurita* x *Callithrix* sp. hybrids we sampled in São Paulo state respectively originated from northeastern Minas Gerais or Espírito Santo states, the Atlantic Forest-Cerrado Transitional region, and the Caatinga. On the other hand, the likely provenance of a haplotype of *C. aurita* x *Callithrix* sp. individual was Guapimirim, Rio de Janeiro state. The three haplotypes from *C. penicillata* x *C. geoffroyi* hybrids we sampled in
Minas Gerais state and a *C. jacchus* x *C. penicillata* hybrid sampled from Rio de Janeiro state likely originated from the Caatinga.

**Implications of Biological Invasions for Callithrix Genetic Integrity, Hybridization, and Conservation**

Several species of non-native fauna and flora have been introduced to the Brazilian Atlantic Forest [32] - one of the most anthropogenically disturbed, yet highly biodiverse biomes on Earth [33,34]. Deliberate and accidental relocations of species beyond their natural geographic ranges by humans may lead to the establishment of non-autochthonous populations and biological invasions within new geographic localities. Such introductions alter the ecological relationships among taxa, and in cases of closely related species, gene flow may occur due to hybridization [35,36,37].

Indeed, our analyses show, for the first time, evidence of introgression of genetic material from allochthonous *Callithrix* species into the genetic background of an endangered, autochthonous *Callithrix* species. Two mtDNA haplotypes that grouped within the *C. jacchus* clade were associated with three individuals with pure *C. aurita* phenotypes sampled within São Paulo state. Two of these individuals were sampled in two different regions of São Paulo state in municipalities (Mogi Das Cruzes and São José dos Campos) that lie 60 km apart. These data not only show the first genetic evidence for cryptic hybridization within the *aurita* group marmosets, but also suggest two independent occurrences of a *C. jacchus* female mating with a *C. aurita* male that led to genetic introgression. Under scenarios of biological invasions, theoretical and empirical data show that hybridization between allochthonous species and endangered, native species creates extinction risk for the latter [38]. Our initial mtDNA data strongly prompt for the development of diagnostic genetic markers to detect the
actual extent of allochthonous *Callithrix* genetic introgression in *C. aurita* populations, particularly within São Paulo state.

Contemporary anthropogenic hybrid *Callithrix* and allochthonous *Callithrix* species are normally found in urban or peri-urban areas of southeastern Brazil, due to releases of exotic pet marmosets into such locales [6,9,31,10], where they may encounter autochthonous *Callithrix* species. Indeed, cases exist of native *C. aurita* and *C. flaviceps* meeting up and interbreeding with hybrid and allochthonous *Callithrix* at urban fringes [6,9,10,11,39,40]. Such interactions likely facilitate gene flow from invasive *C. jacchus* and *C. penicillata* into marmoset populations in southeastern Brazil, with consequences that may include outbreeding depression, admixture, hybrid swamping, or introgressive replacement [41,42,43,44].

Invasive *C. jacchus* and *C. penicillata* represent a potential risk for genetic extinction of the other two *jacchus* group species, *C. geoffroyi* and *C. kuhlii*, and [10] recently showed that *C. penicillata* is encroaching on the range of *C. geoffroyi*. We sampled one *Callithrix* sp. x *Callithrix* sp. hybrid in Santa Teresa, Espírito Santo state, a city that straddles the native ranges of *C. flaviceps* and *C. geoffroyi*. Although this hybrid possessed a *C. geoffroyi* clade mitogenome lineage, the individual had a phenotype that strongly suggested some level of *C. penicillata* or *C. jacchus* ancestry- a white “star” on the forehead [30]. Anthropogenic hybridization of *jacchus* group species generally results in the formation of hybrid swarms, admixed populations that lost parental phenotypes and genotypes [6,30,45]. Should large numbers of exotic *C. jacchus* or *C. penicillata* ever invade native ranges of *C. kuhlii* or *C. geoffroyi*, the latter two species may be threatened with genetic swamping by the former two species, a process through which parental lineages are replaced by hybrids that have admixed
genetic ancestry [38]. As *C. kuhlii* is considered vulnerable [46], biological invasions by other marmosets present potential conservation risks for this species.

Brazil already possesses several legal instruments for the conservation and protection of wildlife [discussed in 47]. These instruments include national species plans that legally lay out action and aims for the protection of specific groups of endangered species. A national species plan already exists that includes *C. aurita* and *C. flaviceps*, the National Action Plan for the Conservation of Atlantic Forest Primates and Collared Sloth [PAN PPMA, 47], and this plan may eventually need to include *C. kuhlii*. The PAN PPMA considers hybridization as a major threat to the survival of the *aurita* group marmoset species. Thus, the expanded perspective on marmoset hybridization provided by this work should be considered within the context of Brazilian legal instruments that protect endangered marmosets. Such an evaluation is important for incorporating new biological information about marmoset hybridization, as it may call for adopting new legal measures or modifying existing ones to further protect endangered Brazilian primates.

**Conclusions**

We provide a robust *Callithrix* phylogeny based on the largest to-date geographical sampling of *Callithrix* mitogenomes across Brazil, showing that the *aurita* group is basal to the *jacchus* group. Our divergence time estimates show these two groups diverged approximately 3.54 Ma, and within the *jacchus* group, *C. jacchus* diverged most recently from the *C. penicillata* Caatinga clade approximately 0.51 Ma. With future sampling of *C. flaviceps*, full mitogenomes can likely be utilized to fully resolve the *Callithrix* phylogeny. Nonetheless, we used our current well-supported phylogenies and biogeographic analyses to elucidate, for the first time, evolutionary relationships among autochthonous, allochthonous,
and anthropogenic hybrid marmosets across Brazil. We show that parental populations of allochthonous and anthropogenic hybrid marmosets within the southeastern Brazilian Atlantic Forest incorporate local populations and populations broadly distributed outside of the regions. We also show, for the first time, evidence of allochthonous *Callithrix* species genetic introgression into the genetic background of endangered, autochthonous *C. aurita*. At this time, further determination is needed of the ancestry of *Callithrix* anthropogenic hybrids in southeastern Brazil as well as the fitness and viability of these hybrids. Such data will help determine to what extent anthropogenic hybrids and allochthonous *Callithrix* species threaten the genetic integrity, or ability of a population to preserve its genotypes over generations [48], of autochthonous Atlantic Forest *Callithrix* species.

**Methods**

**Samples**

In 2011, skin samples were collected from two *C. penicillata* individuals that were captured in Brasília, Federal District. Between 2010 and 2016, skin tissue was collected from: (1) wild marmosets in Minas Gerais and Espírito Santo states as well as the Brazilian Federal District; (2) captive-born, wild-caught, and confiscated marmosets housed at the Guarulhos Municipal Zoo, Guarulhos, São Paulo, CEMAFAUNA (Centro de Manejo de Fauna da Caatinga), Petrolina, Pernambuco, and Centro de Primatologia do Rio de Janeiro (CPRJ), Guapimirim, Rio de Janeiro; (3) a wild group from Natividade, Rio de Janeiro that was caught and housed at CPRJ; (4) a captive-born *C. geoffroyi* sample donated by the Callitrichid Research Center (CRC), Omaha, Nebraska, US; (5) a captive born *C. jacchus* donated by the New England Primate Research Center (NEPRC, now closed), Southborough, Massachusetts, US. Sampling consisted of a total of 49 *Callithrix* individuals as described in Table 1, Table
S1, and Figure 1. Table S1 also lists information on utilized sequences that were published elsewhere. Marmoset capture and sampling methodology has been described elsewhere [23]. All individuals were allowed to recover after sample collection, and wild marmosets were released at their point of capture. Specimens were classified phenotypically as pure *C. aurita*, *C. geoffroyi*, *C. jacchus* and *C. penicillata* or hybrid (*C. aurita* x *Callithrix* sp., *C. jacchus* x *C. penicillata*, and *C. penicillata* x *C. geoffroyi*) based on published descriptions [2,11,23,30].

**Laboratory Protocols**

DNA from skin samples was extracted using a standard proteinase K/phenol/chloroform protocol [49]. Buffers used for extraction, precipitation and elution of DNA from blood and skin tissue are listed elsewhere [24]. DNA from the Callitrichid Research Center samples was extracted at Arizona State University (ASU). DNA from Brasilia individuals was extracted at Northern State Fluminense University, Rio de Janeiro State, Brazil, and then exported to ASU (CITES permit #11BR007015/DF). DNA from all other individuals was extracted at the Federal University of Viçosa (UFV), Viçosa, Minas Gerais, Brazil.

Mitogenomes were obtained for a subset of the samples (Table S1) following the long-range PCR (LR-PCR) methodology of [24], and sequenced on an ABI 3730 sequencer with the BigDye Cycle Sequencing Kit (Applied Biosystems) by the ASU School of Life Science DNA Core Laboratory. The remainder of mitogenomes was obtained from whole genome sequencing (WGS). Individual WGS sequencing libraries were prepared at UFV and ASU with Illumina Nextera DNA Flex Library Prep Kits (catalog #20018704) following manufacturer’s instructions. Individual libraries were barcoded with Illumina Nextera DNA CD Indexes (catalog # 20018707), and pooled in equimolar amounts and sequenced on an Illumina NextSeq using v2 chemistry for 2 x 150 cycles at the ASU Genomic Core Facilities.
Mitogenome Alignment and Data Analysis

Genetic samples collected since 2015 have been registered in the Brazilian CGen SISGEN database (Supplementary Table S6) and newly sequenced mitogenomes have been deposited in GenBank (Table S1). Trace files of resulting forward and reverse reads from LR-PCR products for each individual sequence were inspected by eye and merged into a single contig for each sampled individual using SEQMAN PRO software from the DNAStar Lasergene Core 10 suite (DNASTAR, Madison, WI). Mitogenomes from WGS data were assembled with NOVOPlasty 2.6.4 [50] (scripts available at https://github.com/Callithrix-omics/Callithrix_mtDNA.git). We downloaded mitogenome sequences of several primate species from GenBank (Supporting Information Table S1). All mitogenomes were aligned in MAAFT (https://www.ebi.ac.uk/Tools/mafft/) with default settings and this MAFFT alignment was confirmed visually in Mesquite 3.5 [51]. Gene, tRNA, rRNA, and control region features within the newly generated marmoset mitogenomes were manually annotated based on the GenBank record of C. kuhlii (Accession number KR817257). To check for the presence of nuclear mitochondrial DNA (numts) in mitochondrial sequence data, we followed the strategy described in [24].

We kept mitogenomes in their entirety, but trimmed part of tRNA-Phe, 12s rRNA and the control region to accommodate the length of all utilized sequences. Mitogenome haplotypes were determined with DnaSP 6.12.03 [52], and haplotypes were used for phylogenetic reconstruction. Individuals that possess identical mtDNA haplotypes are listed in Table S3, and these groups are represented in phylogenetic reconstructions by a single haplotype. We added data from several other New World monkeys (Table S1) and reconstructed phylogenetic trees with ML and Bayesian algorithms using IQ-TREE 2.0.3 [53].
and MrBayes 3.2.6 [54,55], respectively. For the ML phylogeny, we used the optimal substitution model (GTR+F+R4) as calculated with ModelFinder [56,57] in IQ-TREE under the Bayesian Information Criterion (BIC). We performed the ML analysis in IQ-TREE with 10,000 ultrafast bootstrap (BS) replications [58]. In MrBayes, we used the closest available substitution model GTR+G. The Bayesian tree was reconstructed via Markov Chain Monte Carlo (MCMC) runs with 10,000,000 generations and tree and parameter sampling occurring every 100 generations. Upon completion of the two runs, the first 25% of generations were discarded as burn-in. To check convergence of all parameters and the adequacy of the burn-in, we assessed the uncorrected potential scale reduction factor (PSRF) [59] and that all parameter Estimated Sample Size (ESS) values were above 200. We calculated posterior probabilities (PP) and a phylogram with mean branch lengths from the posterior density of trees using MrBayes. Phylogenetic trees were visualized and edited with FigTree 1.4.2 (http://tree.bio.ed.ac.uk/software/figtree/). Pairwise genetic distances between each of the resulting Callithrix mitochondrial clades was measured in DnaSP 6.12.03 as $D_{xy}$, the average number of per site nucleotide substitutions between clades.

The divergence time calculation was performed with the BEAST 2.4.8 package [60] using a relaxed lognormal clock model of lineage variation [61] and by applying a Yule tree prior and the best-fit model of sequence evolution as obtained by ModelFinder. To calibrate the molecular clock, we applied fossil data to constrain the splits between Cebinae and Saimirinae and between Callicebinae and Pitheciinae with hard minimum and soft maximum bounds using a log normal prior following settings and fossils described in detail in [62]. Briefly, for the Cebinae - Saimirinae split, we used an offset of 12.6, mean of 1.287 and standard deviation of 0.8, which translates into a median divergence of 16.2 million years ago (Ma) (95% highest posterior density [HPD]: 13.4-30.0 Ma). For the Callicebinae - Pitheciinae
split, we used an offset of 15.7, mean of 1.016 and standard deviation of 0.8, resulting in a median divergence of 18.5 Ma (95% HPD: 16.3-28.9 Ma). We performed two independent runs each with 50 million generations and tree and parameter sampling setting in every 5000 generations. To assess the adequacy of a 10% burn-in and convergence of all parameters, we inspected the trace of the parameters across generations using Tracer 1.6 [63]. We combined sampling distributions of both replicates with LogCombiner 2.4.8 and summarized trees with a 10% burn-in using TreeAnnotator 2.4.8 (both programs are part of the BEAST package). A ParimonySplits network of a subset of mtDNA haplotypes was made with default settings in SplitsTree4 [64].

To reconstruct the biogeographic history of *Callithrix* mitochondrial lineages, we applied the Bayesian Binary Method (BBM) in Reconstruct-Ancestral-States-in-Phylogenies 4.0 (RASP) [65,66]. The ML phylogeny obtained with IQTREE was used for the BBM analysis, which was conducted as two independent runs of 10 chains that ran for 5,000,000 generations and sampled every 100 generations. The fixed Jukes-Cantor+Gamma evolutionary model was implemented for each run. For haplotypes states of origin within a given phylogenetic clade, presence and absence of each associated taxon was determined using a combination of information of known provenance for sampled individuals and recognized *Callithrix* geographical distribution following [5]. For haplotypes obtained from exotic *Callithrix* species and anthropogenic hybrids, we noted where each of these haplotypes clustered among resulting phylogenetic clades, and assigned probable origin for these haplotypes according to the likely natural geographic range associated with each clade.
Abbreviations

ASU - Arizona State University

BBM - Bayesian Binary Method

BIC - Bayesian Information Criterion

BS - Bootstrap

CEMAFAUNA-Centro de Conservação e Manejo de Fauna da Caatinga

CPRJ-Centro de Primatologia do Rio de Janeiro

CRC-Callitrichid Research Center

ESS - Estimated Sample Size

HPD - Highest posterior density

LR-PCR - long-range PCR

mtDNA - mitochondrial DNA

Ma - million years ago

MCMC - Markov Chain Monte Carlo

MRCA - Most recent common ancestor

ML - Maximum likelihood

NEPRC-New England Primate Research Center

PP - Posterior probabilities

PSRF - potential scale reduction factor

UFV - Universidade Federal de Viçosa

WGS - Whole genome sequencing

Declarations

Ethics approval and consent to participate
Tissues were collected under the approval of the ASU Institutional Animal Care and Use Committee Animals (ASU IACUC, protocols #11-1150R, 15-144R), Brazilian Environmental Ministry (SISBIO protocols #47964-2 and #28075-2), and a CPRJ internal review. Biological tissue sampling complied with all institutional, national, and international guidelines.

Consent for publication

Not applicable.

Availability of data and materials

Newly sequenced and annotated *Callithrix* mitogenomes have been deposited in NCBI GenBank. Accession numbers for each sequence are given in Table S1.

Competing interests

The authors declare that they have no competing interests.

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Authors' contributions

JM formulated the idea for the study, collected samples, obtained funding, conducted wet and dry laboratory work, and wrote the original manuscript. RAC provided study guidance, and logistical support. NHAC provided field assistance and logistical support. JAD provided study guidance, and logistical support. CM assisted in phenotypic identification of hybrids and generation of Appendix A. CSI gave access and provided logistical support to collect samples from animals kept at Guarulhos Zoo. SBM provided logistical support and veterinary assistance to collect samples from animals kept at the Rio de Janeiro Primatology Center. AN gave assistance with data processing. PAN gave access and provided logistical support to collect samples from animals kept at CEMAFAUNA. MP provided field assistance and logistical support. LCMP gave access and provided logistical support to collect samples from animals kept at CEMAFAUNA. AP gave access and provided logistical support to collect samples from animals kept at CPRJ. CRRM and DZ were major contributors in writing the manuscript. DLS performed DNA extractions, methodological optimization, and carried out PCR. ACS was a major contributor in writing the manuscript, provided study guidance, and logistical support. CR gave assistance in data processing, carried out divergence analysis, gave significant guidance and input in the development of this study. All authors read and approved the final manuscript.

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Main Figure Titles and Legends

Figure 1. Approximate distribution of Callithrix species in Brazil (2012 IUCN Red List Spatial Data; [http://www.iucnredlist.org/technical-documents/spatial-data](http://www.iucnredlist.org/technical-documents/spatial-data)) and geographic origins of Brazilian samples, as indicated by capital letter symbols. Locations of three biomes where Callithrix occur naturally, the Caatinga, Cerrado, and Atlantic Forest, are also indicated.
Figure 2. Phylogenetic relationships and divergence ages in million years (Ma) among *Callithrix* haplotypes as calculated from complete mitogenomes (complete tree with outgroups is presented in Figure S3). Major nodes are identified by capital letters, and blue bars at nodes indicate 95% highest posterior densities (HPD) of divergence times. Node support is shown for major nodes where either posterior probability was < 1 in the BEAST tree, posterior probability was < 1 in the MRBAYES tree, or bootstrap support < 70% in the ML tree. Haplotype colors at tips correspond to the ‘Species and Hybrid Phenotypes’ legend, and indicate phenotypes associated with each given haplotype.
Figure 3. ParimonySplits network of haplotypes from phylogenetic *C. jacchus* clade.
Haplotype colors at tips follow Figure 1 ‘Species and Hybrid Phenotypes’ legend, and indicate phenotypes associated with each given haplotype.
Figure 4. Ancestral state reconstructions performed by the Bayesian Binary MCMC analysis as implemented in RASP v4.2 using the ML rooted tree. Donut charts at each node represent ancestral host estimations. Each node is internally identified with a number. The posterior
probabilities of ancestral origins of major nodes are shown in Table S5. Localities where species associated with each phylogenetic clade were sampled or known to occur: A-Ceará state; B-Rio Grande do Norte state; C-Paraíba state; D-Alagoas state; E-Piauí/Maranhão/Western Bahia states; F-Pernambuco state; G-Caatinga biome in Bahia state; H-Atlantic Forest biome in Bahia state; I-southern Espírito Santo state; J-northeastern Minas Gerais; K-northern Espírito Santo state; L-southeastern Minas Gerais; M-Rio de Janeiro state; N-São Paulo state; O-Atlantic Forest and Cerrado transitional areas in southern Minas Gerais; P-Cerrado Brazilian Federal District. These localities are color coded in the map inset on the left side.