Complete mitochondrial genome of the greater round-eared bat, *Tonatia bidens* (Chiroptera: Phyllostomidae) from Brazil and phylogenetic relationships within the family

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**ABSTRACT**

The greater round-eared bat, *Tonatia bidens*, is a locally rare species belonging to the highly diverse family Phyllostomidae. In this study, the complete mitogenome of *T. bidens* was sequenced using optimized protocols of DNA extraction from fixed cells originally prepared for cytogenetic studies. Here we present the complete mitogenome and place our results in a phylogenetic context with other data generated for the family Phyllostomidae. The circular genome had 16,717 bp in size, comprising 37 genes and GC content of 42.24%. Furthermore, the phylogenetic tree indicated a well-supported relationship between the representatives of *Tonatia* into the subfamily Phyllostominae.

Round-eared bats of the genus *Tonatia* (Phyllostomidae, Phyllostominae) are widely distributed throughout Latin America and represented by three extant species: *Tonatia bidens* Spix 1823, *T. bakeri* and *T. maresi* Williams, Willig & Reid 1995, the last two recently elevated from subspecies of *T. saurophila* to species rank (Basantes et al. 2020). Despite their widespread distribution, little information encompassing aspects of their biology is known, and most authors consider the representatives of *Tonatia* as rare species due to few occurrences in nature, generally in small groups, especially *T. bidens* (Willig 1985; Estéberard and Bergallo 2004). *Tonatia bidens* is a medium-sized bat, endemic to South America found in Argentina, Paraguay, Bolivia, and Brazil. Scarce incidence with fragmented and punctual sampling data justifies its categorization as Data Deficient (Barquez and Diaz 2016). Therefore, by describing the mitogenome of *T. bidens*, we provide new insights into extranuclear genome evolution as well as initial steps into the genetic characterization for this species. Herein, one male specimen of *T. bidens* was collected in Parque Estadual Pedra da Boca (6° 27’ 32.2” S, 35° 40’ 45.4” W), Araruna, Paraíba, Brazil, during field expeditions in 2004, as authorized by IBAMA (Licence no. 12264-1 to Santos, N). The voucher specimen was deposited in the mammalian collection of Departamento de Sistemática e Ecologia at Universidade Federal da Paraíba, João Pessoa, Brazil (www.cccn.ufpb.br/museubiologia/) under the voucher number UFPB5719.

Total genomic DNA was extracted from fixed cytological suspension (stored in methanol-acetic acid, 3:1, v/v) and deposited in the Laboratório de Genética e Citogenética Animal e Humana (LGCAH/UFPE; Voucher ID: M829). Fixed cells in suspension were washed twice with 1 × PBS (Life Technologies, pH 7.4) for 20 min before DNA extraction using the DNeasy Blood & Tissue kit (Qiagen), following the manufacturer’s instructions for blood samples. Paired-end libraries were built with Nextera DNA Flex Library Prep kit (Illumina) and sequenced using a high-output v2 kit (300 cycles) on an Illumina NextSeq 500 platform. The mitogenome was assembled using NovoPlasty 3.6 (Dierkskssen et al. 2017) and annotation was performed with MITOS2 (Bernt et al. 2013), with minor manual corrections in Geneious Prime 2020.2 (Biomatters).

The mitogenome of *T. bidens* (GenBank accession MZ391834) comprised 16,717 bp, with two ribosomal RNA (rrnL and rrnS) genes, 13 protein-coding genes (PCGs) and 22 transfer RNA (tRNA). The nucleotide composition was 32.3% A, 29.1% C, 13.1% G, and 25.4% T, with 42.24% of GC content. The average size and gene arrangement were identical to those of other bat species (Meganathan et al. 2012; Botero-Castro et al. 2013, 2018). In addition, two rRNA and most of tRNA and PCGs were encoded in the H-strand, whereas NAD6 was in the L-strand, besides eight tRNAs (trnA, trnC, trnE, trnN, trnP, trnQ, trnS2[nga] and trnY). *Tonatia bidens* had 10 ORFs starting with ATG, two starting with ATA (NAD5 and NAD2), and one starting with ATT (NAD3).
We investigated the relationships among *T. bidens* and other 20 phyllostomid species using available mitogenomes recovered from GenBank, and assigning the sequenced mitogenomes of representatives of the families Mormoopidae, Mystacinidae, and Noctilionidae as outgroups. The phylogenetic analyses were based on all PCGs amino acid (AA) sequences aligned with MAFFT 7.3 (Katoh et al. 2002). Maximum likelihood (ML) trees were obtained with RAxML v8.2 (Stamatakis 2014) using the substitution model PROTGAMMA and rapid bootstrapping (BS) with 1,000 replicates (Figure 1).

The phylogenetic analysis recovered a monophyletic Phyllostomidae clade with strong support (BS = 99) and included mitogenomes from representatives of all eleven subfamilies from the classification proposed by Baker et al. (2003, 2016; Figure 1). Our analysis resulted mostly in well-supported clades (BS > 70). In addition, in contrast with the Baker et al. (2003, 2016) phylogenies, incongruences were observed in the positions of the subfamilies Microcycterinae relative to Macrotritinae, Lonchophyllinae in alternative branching relative to other nectarivorous bats, and Lonchorhininae appeared as a lineage that diverged after Phyllostominae, most of these relations resulted in low-supported clades (Figure 1), although a similar mitogenome AA-tree was observed by Botero-Castro et al. (2018). Within the subfamily Phyllostominae (BS = 74), *T. bidens* was recovered as sister to *T. maresi* (BS = 100), and the genus *Tonatia* was more closely related to *Lophostoma* than to *Chrotoperus* and *Vampyrum* (Figure 1), as already observed in other phylogenetic approaches (Baker et al. 2003; Rojas et al. 2016; Basantes et al. 2020). The present study serves as encouragement to other cytogenetic laboratories to take advantage of cytological material for new purposes, especially in rare species, as provided here useful knowledge of genomic data in this species which will be crucial for the understanding biodiversity and future studies in the taxonomy and molecular systematics of phyllostomid bats.

**Ethics statement**

All procedures performed in this study involving animals followed guidelines established by Animal Care and Use guidelines of ICMBio (Instituto Chico Mendes de Conservação da Biodiversidade), and using the collecting permit number (12264-1) by the Instituto Brasileiro do Meio Ambiente e dos Recursos Naturais Renováveis (IBAMA).

**Author contribution statement**

JCF, SV, and CGS-C contributed to the conception and design of the study, analysis, and interpretation of the data. JCF wrote the first version of the manuscript. SV, GO, NS and CGS-C critically reviewed the article regarding its intellectual content. NS collected biological samples. All authors read, discussed, and approved the final version and all authors agree to be accountable for all aspects of the work.

**Disclosure statement**

The authors report no conflicts of interest.
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Data availability statement

The mitogenome sequence data that support the finding of this study are openly available in GenBank of NCBI at [https://www.ncbi.nlm.nih.gov], under the accession no. MZ391834. The associated BioProject, SRA, and Bio-Sample numbers are PRJNA785884, SRR17118535, and SAMN23588143, respectively.

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