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SPECIALTY SECTION

This article was submitted to
Virology,
a section of the journal
Frontiers in Microbiology

RECEIVED 25 July 2022

ACCEPTED 18 August 2022

PUBLISHED 08 September 2022

CITATION

Cosentino MAC, D'arc M, Moreira FRR,
Cavalcante LTF, Mouta R, Coimbra A,
Schiffler FB, Miranda TS, Medeiros G,
Dias CA, Souza AR, Tavares MCH, Tanuri A,
Soares MA and Santos AFA (2022)
Discovery of two novel *Torque Teno* viruses
in *Callithrix penicillata* provides insights on
Anelloviridae diversification dynamics.
Front. Microbiol. 13:1002963.
doi: 10.3389/fmicb.2022.1002963

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Miranda, Medeiros, Dias, Souza, Tavares,
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Discovery of two novel *Torque Teno* viruses in *Callithrix penicillata* provides insights on *Anelloviridae* diversification dynamics

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The development of high-throughput sequencing (HTS) technologies and metagenomics protocols deeply impacted the discovery of viral diversity. Moreover, the characterization of novel viruses in the Neotropical primates (NP) is central for the comprehension of viral evolution dynamics in those hosts, due to their evolutionary proximity to Old World primates, including humans. In the present work, novel anelloviruses were detected and characterized through HTS protocols in the NP *Callithrix penicillata*, the common black-tufted marmoset. *De novo* assembly of generated sequences was carried out, and a total of 15 contigs were identified with complete *Anelloviridae* ORF1 gene, two of them including a flanking GC-rich region, confirming the presence of two whole novel genomes of ~3kb. The identified viruses were monophyletic within the *Epsilontorquevirus* genus, a lineage harboring previously reported anelloviruses infecting hosts from the Cebidae family. The genetic divergence found in the new viruses characterized two novel species, named *Epsilontorquevirus callithrichensis I* and *II*. The phylogenetic pattern inferred for the *Epsilontorquevirus* genus was consistent with the topology of their host species tree, echoing a virus-host diversification model observed in other viral groups. This study expands the host span of *Anelloviridae* and provides insights into their diversification dynamics, highlighting the importance of sampling animal viral genomes to obtain a clearer depiction of their long-term evolutionary processes.

KEYWORDS

metagenomic, marmoset, co-divergence, ssDNA viruses, virus evolution

Introduction

The development of high-throughput sequencing (HTS) technologies and metagenomics protocols deeply impacted the discovery of viral diversity (Greninger, 2018; Pérez-Losada et al., 2020). Once a matter of contention, the establishment of novel viral species and higher taxonomic groups based solely on sequencing data became commonplace. Studies describing multiple previously unknown viral groups are increasingly common (Datta, 2015; Shi et al., 2016; Atoni et al., 2019; Zhang et al., 2019), as are studies expanding the diversity of well-established viral families (Smeele et al., 2018; de Sabato et al., 2019; Sawaswong et al., 2019; dos Santos et al., 2020). While these efforts have broad implications on the comprehension of virus diversity, ecology, evolution, and epidemiology, large fractions of the viruses infecting non-model organisms remain to be discovered (Geoghegan and Holmes, 2017). In this regard, Neotropical primates (NP; infraorder Platyrrhini) stand out as a group with great ecological importance, whose viral diversity has not been fully explored. Only 18% (4/22) and 3.4% (6/178) of NP genera and species, respectively, have been previously assessed through viral metagenomics (Ng et al., 2015; Troncoso et al., 2015; Silvestre et al., 2016; Muniz et al., 2018; D'arc et al., 2020; dos Santos et al., 2020).

Currently, NP comprehends 178 species and 22 genera that occupy a myriad of ecological niches in Central and South Americas (IUCN/SSC Primate Specialist Group, 2021). As it is the sister group to Old World primates (OWP; infraorder Catarrhini), including humans, the comprehension of their evolution, immunology, and susceptibility to infectious diseases is central to the characterization of the scope of etiological agents that may impact global health (Devaux et al., 2019). In this sense, previous investigations have shown that multiple viral families might be found in the virome of NP: novel papillomaviruses were found infecting *Callithrix penicillata* (D'arc et al., 2020) and *Alouatta guariba* (Silvestre et al., 2016), anellovirus and genomovirus in *Sapajus nigritus* (dos Santos et al., 2020), smacovirus in *Alouatta caraya* (Ng et al., 2015), and foamy virus in *Brachyteles arachnoides* (Muniz et al., 2018) and *Sapajus xanthosternos* (Troncoso et al., 2015).

The *Anelloviridae* family is one of the most frequently found in metagenomic surveys of NP. This family is composed of non-enveloped virions with circular single-stranded negative sense DNA (ssDNA), whose genomes range from 1.6 to 3.9 kb. Three major ORFs are found in their genomes—named ORF1, ORF2, and ORF3—with putative additional ORFs among isolates. Overlay between ORFs is common and their estimated sizes differ widely among isolates. ORF1 is believed to encode the putative capsid and replication-associated protein, while ORF2 may encode a protein with phosphatase activity or a peptide capable of suppressing NF- κ B pathways (Varsani et al., 2021). Currently, the group is composed of 30 classified genera by the International Committee of Virus Taxonomy (Nishizawa et al., 1997). The *Torque Teno Virus* was first identified in the 1990s in a post-transfusion hepatitis case of unknown etiology in humans (Bendinelli et al., 2001) and was posteriorly discovered in healthy human populations with

prevalence ranging from 10 to 96% (Hino and Miyata, 2007; Freer et al., 2018). No pathology has been confirmed by its infection, although an association with pulmonary and liver diseases has been suggested (Ninomiya et al., 2009). Furthermore, the *Anelloviridae* family is proposed to be a common part of the human virome, with major importance to the immune system maturation (Hrazdilová et al., 2016). A high prevalence in healthy populations and the lack of clinical signs have also been shown in free-living and captive populations of chimpanzees and gorillas (Inami et al., 2000; Okamoto et al., 2000; de Souza et al., 2018). Although new anelloviruses species have been discovered in NP—such as in *Saguinus oedipus*, *Aotus trivirgatus* (Ng et al., 2011), and *Sapajus nigritus* (dos Santos et al., 2020)—an extensive knowledge gap for anelloviruses infecting NP still remains.

While the diversity and host span of *Anelloviridae* is constantly being updated (Okamoto et al., 2000; Noppornpanth et al., 2001; Pujol et al., 2005; Hon et al., 2008; Holmes, 2010; Ng et al., 2011; Sawaswong et al., 2019; dos Santos et al., 2020; Weber et al., 2020; Kraberger et al., 2021), little is known about the factors driving their diversification dynamics. Specifically, whether the generation of novel viral species is driven by zoonotic spillovers to new host species or co-divergence with long-term host lineages has not been formally assessed. *Anelloviridae* phylogenetics topography suggests a co-divergence within some host lineages, as seen in felids (Noppornpanth et al., 2001) and rodents (Wei et al., 2019; Lam et al., 2020). This hypothesis has been addressed by phylogenetic methods for other groups in large scale meta-analysis (de Souza et al., 2018) as well as in studies focusing on specific viral families (McGeoch et al., 2006; Buck et al., 2016; Silvestre et al., 2016; Smeele et al., 2018; Bando et al., 2021), revealing that the mechanisms that generate viral diversity are often determined by viruses genomic architecture and variability. As the discovery of novel viruses allows a broader evaluation of these mechanisms, metagenomic surveys on non-model organisms have allowed the clarification of the evolutionary dynamics of diverse viral families (Niewiadomska and Gifford, 2013; Kohl et al., 2015; Troncoso et al., 2015; Muniz et al., 2018).

In this study, we characterized the virome of captive *Callithrix penicillata*, the black-tufted marmoset, by HTS and discovered two novel species of anelloviruses, expanding the scope of primate hosts infected by these viruses. Phylogenetic analysis of the novel sequences suggests anellovirus evolution might be driven by the within-species diversity followed by co-divergence model, as suggested for other viral groups (Buchfink et al., 2014; Nurk et al., 2017; Chen et al., 2018; D'arc et al., 2020). Overall, our results shed light on previously unexplored anellovirus diversity and address a hypothesis regarding their diversification dynamics.

Materials and methods

Sample collection

The data analyzed herein was generated on a larger virome project using samples from NP housed at the *Centro de*

TABLE 1 General information of all samples from *Callithrix penicillata* housed at Centro de Primatologia of Universidade de Brasília (CP/UnB).

Species	CP/UnB name	Lab ID	Library Pool ID	Sample type	Gender	Sexual maturity	Enclosure
<i>Callithrix penicillata</i>	Pamonha	SWA1	P1	Anal Swab	F	Adult	1
<i>Callithrix penicillata</i>	Novela	SWA2	P1	Anal Swab	F	Adult	2
<i>Callithrix penicillata</i>	Miuxa	SWA3	P1	Anal Swab	F	Adult	3
<i>Callithrix penicillata</i>	Miúda	SWA4	P1	Anal Swab	F	Adult	3
<i>Callithrix penicillata</i>	Grelina	SWA5	P1	Anal Swab	F	Adult	4
<i>Callithrix penicillata</i>	Rubi	SWA6	P2	Anal Swab	F	Adult	6
<i>Callithrix penicillata</i>	Gonia	SWA7	P2	Anal Swab	F	Adult	8
<i>Callithrix penicillata</i>	Valina	SWA8	P2	Anal Swab	F	Adult	9
<i>Callithrix penicillata</i>	Gora	SWA9	P2	Anal Swab	F	Adult	10
<i>Callithrix penicillata</i>	Miuka	SWA10	P2	Anal Swab	F	Adult	10
<i>Callithrix penicillata</i>	Gontijo	SWA45	P7	Anal Swab	M	Adult	5
<i>Callithrix penicillata</i>	Valente	SWA46	P7	Anal Swab	M	Adult	7
<i>Callithrix penicillata</i>	Supimpa	SWA47	P7	Anal Swab	M	Adult	8
<i>Callithrix penicillata</i>	Miro	SWA48	P7	Anal Swab	M	Adult	9
<i>Callithrix penicillata</i>	Gonero	SWA51	P7	Anal Swab	M	Adult	14
<i>Callithrix penicillata</i>	Birrento	SWA53	P7	Anal Swab	M	Adult	16

Primatologia of Universidade de Brasília (CP/UnB; Brasília—Brazil). General information on the viral diversity characterized can be found in the first publication of the larger project that the present study is part of D'arc et al. (2020). For this study, the data analyzed were sequenced from anal swab samples from 16 black-tufted marmosets (*C. penicillata*), immersed in phosphate-buffered saline (PBS) and stored at -20°C (Table 1). Samples were collected following the national guidelines and provisions of National Council for Animal Experimentation Control (CONCEA), which included animal welfare standard operating procedures. The procedure was conducted under permission of the Brazilian Institute of Environment and Renewable Natural Resources (IBAMA) under license number 1/53/1999/000006-2 and the project was approved by Ethics Committee on the Use of Animals of Universidade Federal do Rio de Janeiro (reference number 037/14).

High-throughput sequencing

The samples were shipped to the *Laboratório de Diversidade e Doenças Virais* of Universidade Federal do Rio de Janeiro (LDDV/UFRJ; Rio de Janeiro—Brazil) and then the homogenized PBS aliquots were pooled into three arbitrarily sampled groups (arbitrary named P1, P2, and P7) to maximize sequencing efficiency with 4–6 samples each (Table 1). Pooled samples were filtered using Millex-HV 0.45 μm SLHV0135L (Merck, German) and sacrose gradient by ultracentrifugation at 35,000g, 4°C for 90 min was conducted to separate the enriched viral bottom fraction. Supernatants were discarded and the remaining 0.2 ml were digested with DNases (Promega and Epicentre, United States), RNase A (Ambion, United States), and Benzonase (Sigma, United States). The QIAamp MinElute Virus Spin Kit (QIAGEN, Germany) was used for viral nucleic acid extraction

(DNA + RNA), following the manufacturer's protocol with slight modifications to improve the recovery of total nucleic acids in accordance to the CDC protocol methodology (Ondov et al., 2011): (i) the "Carrier RNA" was not used in the AL Buffer; (ii) the protease was resuspended with AVE Buffer, instead of Protease Resuspension Buffer; (iii) the washing step with AW1 was not performed; and (iv) the final elution was done in 20 μl of ultra-pure water. The material was then subjected to a complementary DNA (cDNA) synthesis using the Superscript III First Strand Synthesis Supermix Kit (Thermo Fisher Scientific, United States). The second strand cDNA synthesis was performed using Klenow fragment 3'-5' exo (New England Biolabs Inc., United States), following the manufacturer's instructions. Sequencing libraries were constructed using mixed DNA (DNA + cDNA) with Nextera XT DNA Library Preparation Kit (Illumina, United States), following the standard protocol. In a MiSeq V2 500-cycle cartridge (Illumina), 4 pM of each library were loaded and massive sequencing was conducted on the Illumina MiSeq platform of the Department of Genetics of UFRJ.

Bioinformatic analysis

Sequencing data were processed with a custom pipeline. Raw sequencing reads were filtered with Fastp v.0.20.1 (Kearse et al., 2012), removing short reads (<50 bp) and low-quality (Phred score < 30) bases. Reads for each sample were then *de novo* assembled with the Meta-spades software v.3.15.3 (Peter et al., 2000). Diamond v.2.0.14 (Edgar, 2004) was used to taxonomically assign both reads and their respective contigs, with an *e*-value cut-off of 10^{-5} and parameters "--more-sensitive" and "--max-target 1." A similarity search was performed against the complete non-redundant protein (nr) database from NCBI (as of July 27, 2021). Krona v.2.7.1 (Capella-Gutiérrez et al., 2009) was

used to generate interactive visualization plots, allowing the fast identification of viral families of interest. To avoid false positive results due to sequencing index-hopping (incorrect assignment of a viral read to a given sample), we considered that samples presenting reads for a viral family with a read count of less than 1% than the highest count among all libraries sequenced in the same MiSeq run to be product of contamination and hence excluded from further analyses.

Given that *Anelloviridae* was the major viral family identified across the samples (see the section “High-throughput sequencing and taxonomic assignments” in Results), we were prompted to further explore the structure and diversity of the novel viral genomes. The genome sequences were annotated by the Transfer annotations tool in Geneious Prime v.2021.2.2 (Minh et al., 2020) with a 40% similarity threshold, using as template TTV tamarin (NCBI accession number NC_014085.1) from *S. oedipus*, as well as identifying ORFs through the Find Orf tool in Geneious. Complete genomes were circularized also in Geneious by identifying the GC-rich region in *Anelloviridae*.

Phylogenetic analysis of novel *Anelloviridae* genomes

To contextualize the novel genomes within the *Anelloviridae* diversity, all annotated contigs were searched with the getorf/EMBOSS v.6.6.0.0 (Le et al., 2012) for the presence of ORF1 fragments with at least 1,500 nucleotides. These sequences were aligned to a dataset previously assembled containing all species from the viral family (Nishizawa et al., 1997), composing a comprehensive dataset of 618 ORF1 sequences. These sequences were translated and aligned with MUSCLE v.3.8.425 (Kalyaanamoorthy et al., 2017). The alignment was trimmed with TrimAl v.1.4 (Guindon et al., 2010) with the “-gappypout” option. For quality control, a complete recombination scan was performed with RDP4 (Rodriguez et al., 1990). A maximum likelihood tree was then inferred with IQ-Tree v.2.1.4 (Yang, 1994) under the LG+F+G4 model (R Core Team. R: A, 2018), suggested by ModelFinder (Paradis and Ape, 2019), and with the implementation of the SH-like approximate likelihood ratio test (SH-aLRT; Suchard et al., 2018). A complete list of used sequences, as well as the described alignment and tree are available in [Supplementary File S1](#).

Molecular clock analysis of TTVs from NP

A separate nucleotide dataset comprising all TTVs from NP was assembled and a maximum likelihood tree was inferred under the GTR+F+I+G4 model (Perelman et al., 2011; Baele et al., 2012). The tree and alignment were used for identity analysis with the R software (Baele et al., 2013), using the APE package v.5.5 (Griffiths and Tavare, 1994). As the phylogenetic pattern observed at the genus level was consistent with the ancient within-species

diversity model, we were prompted to infer a time-scaled phylogenetic tree for *Episilotorquevirus* genus, which comprehends TTVs from capuchin (*S. nigrurus*), tamarin (*S. oedipus*), and the new marmoset (*C. penicillata*) viruses. This analysis was performed with BEAST v1.10.4 (Drummond et al., 2005). Following Perelman et al. (2011) and (Gernhard, 2008), estimated host divergence dates were used to calibrate internal nodes of the viral tree. A calibration time point between TTV capuchin and other *Episilotorquevirus* was set by a normal distribution of mean 19.95 million years ago (mya; 95% CI = 15.66–24.03 mya), while the calibration time point between TTVs tamarin and marmoset was set by a normal distribution of mean 14.89 mya (95% CI = 11.04–18.48 mya). Marginal likelihoods for eight different combinations of tree priors and clock models have been evaluated with path-sampling and stepping-stone sampling algorithms (Drummond et al., 2006; Ferreira and Suchard, 2008). The evaluated tree priors were: speciation yule and coalescent constant, exponential and skyline (Hino and Prasetyo, 2009; Rambaut et al., 2018; Li et al., 2019). Both the strict and uncorrelated relaxed clock models have been assessed (Bédarida et al., 2017; Deb et al., 2021). For each model combination, a Markov chain Monte Carlo run with 50 million generations was executed, sampling every 5,000 steps. Mixing (effective sample size > 200) for all models parameters was verified on Tracer v.1.7.0 (Arze et al., 2021), and a maximum clade credibility tree was inferred with TreeAnnotator v1.10.4 (Arze et al., 2021) for the best fit model. All BEAST xmls, logs and maximum clade credibility are available in [Supplementary File S2](#).

Results

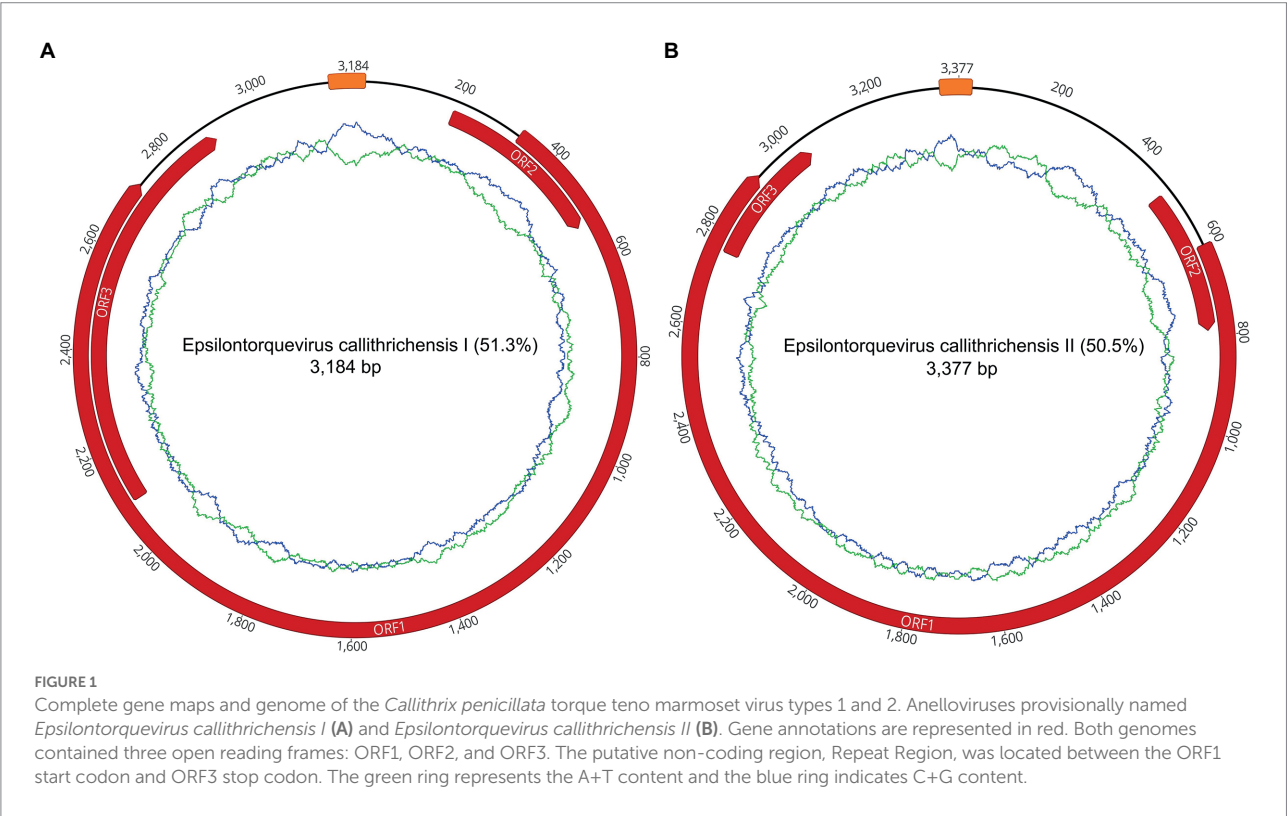
High-throughput sequencing and taxonomic assignments

The HTS protocol generated a total of 8,352,114 reads, harboring a total of 1,487,358 raw reads for P1 library, 3,319,936 for P2, and 3,544,820 for P7. After reads were filtered, a total of 363,161 (24.4%), 696,397 (21%), and 544,058 (15.3%) reads were obtained for libraries P1, P2, and P7, respectively. Reads were taxonomically assigned through similarity search against the complete nr database from NCBI. Viral reads corresponded to approximately 0.142% of the total generated data (1,109 viral reads in P1, 9,200 in P2, and 1,547 in P7). The meta-spades software *de novo* assembled more than 54,000 contigs, 15,519 contigs assembled in P1, 17,500 in P2, and 21,671 in P7. In the assembled contigs, 312 (0.57% of total) were taxonomically assigned as viruses, from which 78 (0.5%), 186 (1%), and 48 (0.2%) belonged to the sequenced libraries P1, P2, and P7, respectively.

Several viral families have been identified among the evaluated libraries. A complete table with viral taxonomic assignments can be found in [Supplementary Table S1](#). Among the vertebrate viral families identified, *Anelloviridae* stands out as the one presenting the vast majority of viral reads and contigs (3,816 reads and 164

TABLE 2 Summary of the number of total reads and contigs identified as Anelloviridae and number of ORF1 regions and complete genomes obtained.

Library	Filtered reads	Anellovirus reads	Contigs	Anellovirus contigs	ORF1	Genomes
P1	363,161	216	15,519	27	1	0
P2	696,397	3,514	17,500	129	14	2
P7	544,058	86	21,671	8	0	0



contigs in total), being hence the main group explored throughout this study. *Anelloviridae* characterization.

The *Anelloviridae* family was identified in all evaluated pools, including 216 assigned reads for library P1, 3,514 for P2, and 86 for P7. All of them had a read count greater than the contamination cut-off previously established ($n=35$), supporting the *bona fide* presence of the viral family. A total of 27, 129, and eight assembled contigs were identified as *Anelloviridae* (Table 2) for P1, P2, and P7, respectively.

To further contextualize the novel sequences within the *Anelloviridae* diversity, all contigs were searched for the presence of ORF1 fragments with at least 1,500 nucleotides. We found a total of one contig with an ORF1-like in P1 and 14 contigs in P2. No contig with ORF1-like was identified in P7. Among those 15 contigs, two of them possessed a flanking GC-rich region and all other anellovirus ORFs could be completely annotated, confirming the presence of two whole novel genomes of ~3 kb in P2 library (Figure 1; Tables 2, 3).

The two novel genomes sequenced were submitted to ICTV and were provisionally named as *Epsilontorquevirus callithrichensis*

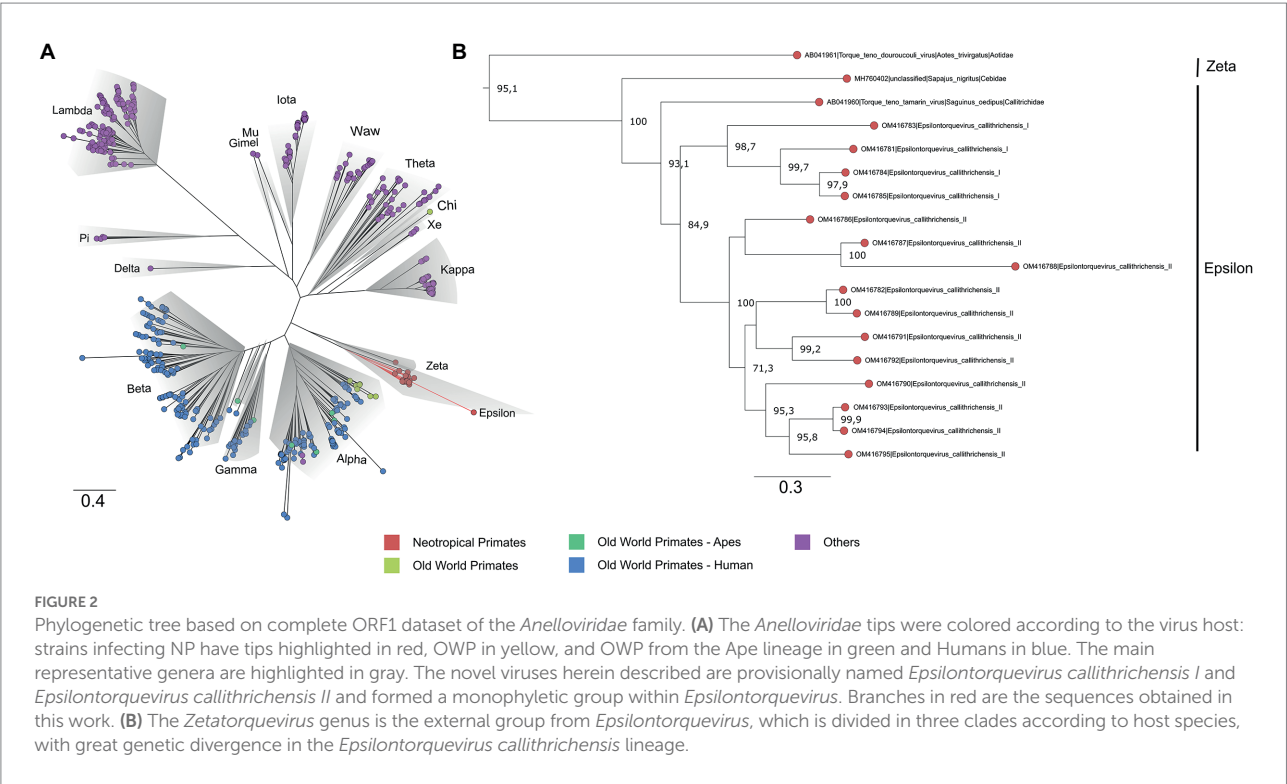
(*E. callithrichensis*) types I and II. The *E. callithrichensis* I is 3,184bp long with a guanine-cytosine (GC) content of 51.3%, while *E. callithrichensis* II is 3,337 bp long with a GC content of 50.5%. Both genomes contain the three major ORFs found in *Anelloviridae*. The complete ORF1 gene, a fundamental gene to species determination in the family, had an estimated length of 2,403 and 2,313 bp for the *E. callithrichensis* I and II, respectively, and shared less than 53.49% identity to previously known *Epsilontorquevirus*. Particularly, the ORF3 gene from *E. callithrichensis* I had an estimated length of 801 bp, superior to the estimated ORF3 in *E. callithrichensis* II with 282 bp and to the TTV tamarin (Accession Number NC_014085.1) ORF3, with 497 bp.

A viable ORF4 was not found in any *E. callithrichensis*, but the Transfer annotations tool in Geneious software found regions with moderate similarity (52.89%, nucleotide positions 2,189--2,896) to TTV capuchin ORF4 in *E. callithrichensis* I. The same region similarity was also found in *E. callithrichensis* II (49.26%, nucleotide positions 2,356--3,040), as well as regions with moderate similarity to TTV tamarin ORF4 (48.15%, nucleotide

TABLE 3 Genome characteristics of the two novel TTV marmosets characterized.

Type	Genome length	%GC	Nucleotide position and size (amino acid)				Most closely related Anellovirus	% ORF1 Seq ID
			ORF1	ORF2	ORF3	Repeat Region		
<i>Epsilontorquevirus callithrichensis I</i>	3.184bp	51.3	327–2.729 (801)	196–534 (113)	2.095–2.895 (267)	3.137–21 (69) (Rambaut et al., 2018) ^a	TTV Tamarin	53.49 ^b
<i>Epsilontorquevirus callithrichensis II</i>	3.377bp	50.5	615–2.927 (771)	484–789 (102)	2.758–3.039 (94)	3.339–26 (65) (Drummond et al., 2005) ^a	TTV Tamarin	51.7 ^b

^aNucleotide size of Repeat region.
^bResults are based on ORF1 pairwise identity in Clustal Omega. TTV Tamarin, Torque teno tamarin virus (Accession number NC_014085).



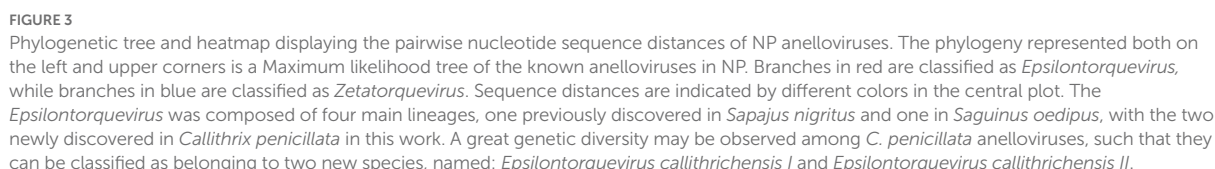
positions: 2,978–3,145). Those hypothetical ORF4 found in *E. callithrichensis* do not possess the typical start codons from other annotated ORFs in *Anelloviridae*.

Evolutionary analysis

To contextualize the novel *Anelloviridae* genome sequences, a maximum likelihood phylogeny was reconstructed with a comprehensive dataset, following Varsani et al. (2021). No recombination signal was detected in the dataset. The obtained tree presented phylogenetic consistency with the previously proposed *Anelloviridae* phylogeny, being all genera inferred as monophyletic with moderate to high support values (SH-aLRT = 67.4–100). The marmoset anelloviruses composed a monophyletic group within *Epsilontorquevirus* (SH-aLRT = 100), a genus comprehending virus species described for the NP. The external group of *Epsilontorquevirus* was the *Zetatorquevirus*

genus, which comprehends the TTV douroucouli sampled in *Aotus trivirgatus* (Ng et al., 2011; Figure 2). This major clade comprised all characterized NP TTVs.

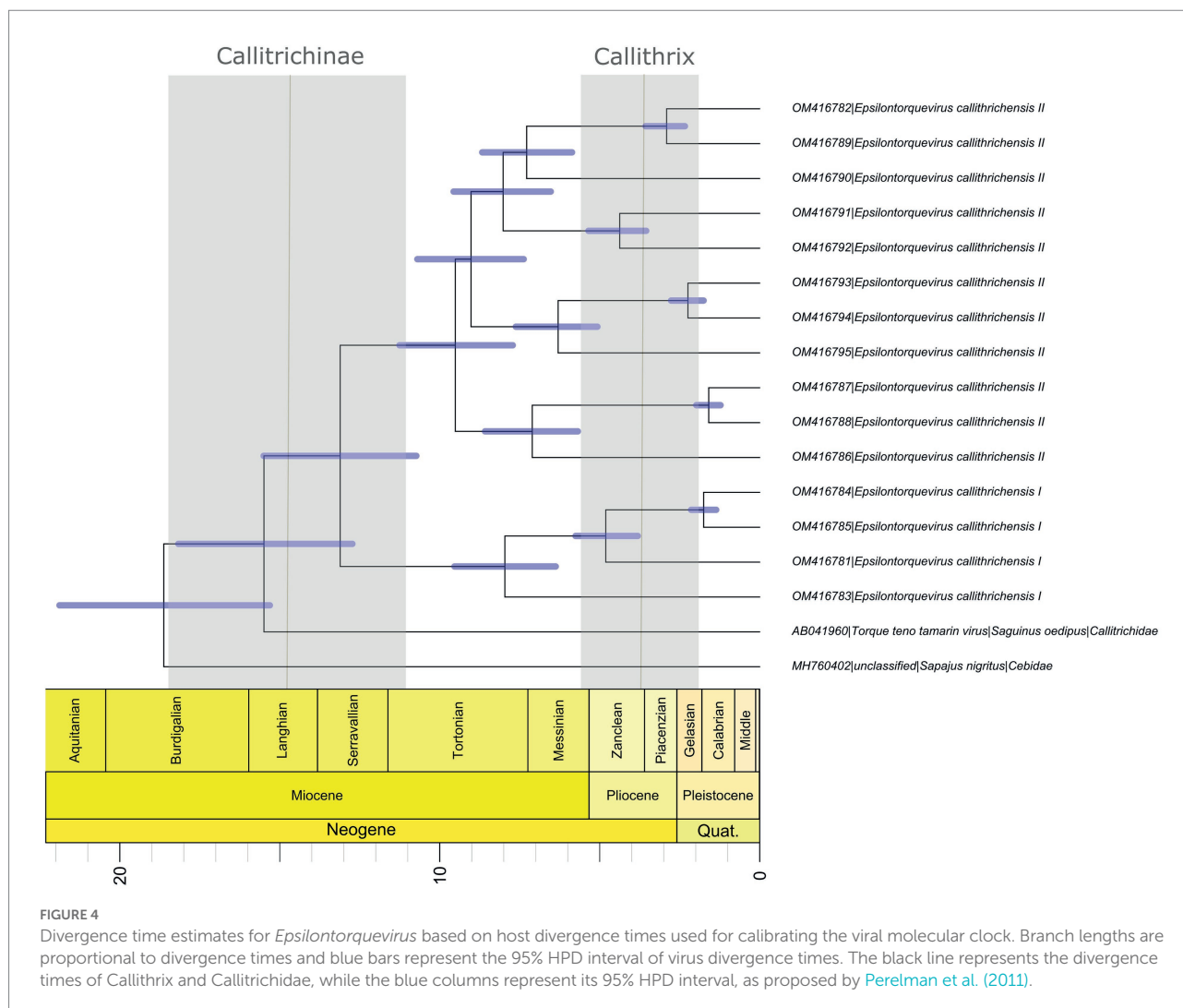
To verify whether the newly identified marmoset viruses belonged to a new species or genus, a separate dataset for NP TTVs was assembled and analyzed. Pairwise nucleotide sequence distances were obtained and visualized through heatmap (Figure 3). The ORF1 sequences from the novel viruses clustered in two monophyletic groups, presenting pairwise identities between 55.89 and 59.35% to the previously described *Epsilontorquevirus* in tamarin. For the TTV capuchin, distances ranged from 53.68 to 56.58%. The ICTV criteria for TTV classification establishes genetic distances thresholds for species (69%; Nishizawa et al., 1997). When compared to the single *Zetatorquevirus* described, the marmoset anelloviruses demonstrated diminished identity, ranging from 42.19 to 44.96%. The first marmoset anellovirus monophyletic lineage, named *E. callithrichensis I*, had a pairwise nucleotide sequence distance



Timescale analyses

of their host species tree, echoing the within-species diversity followed by co-divergence model of viral diversification. The TTV capuchin was the first lineage to diverge within *Epsilontorquevirus*, followed by TTV tamarin, which is sister to the clade comprehending all TTVs marmoset. This branching pattern is consistent with the Cebidae molecular phylogeny inferred (Gernhard, 2008). In this sense, we inferred a time-scaled phylogenetic tree for *Episilontorquevirus* genus using host divergence dates to calibrate internal nodes of the viral tree. Among all models tested, the one with the best statistical fit was characterized by the strict clock and the Yule speciation prior (Supplementary File S2, Path and Stone Sampling table).

Overall, our time scale inference found that the most recent common ancestor of the *Episolontorquevirus* genus was estimated around 18.7 mya (95% HPD = 15.31–21.88), in accordance with the prior established for the analysis. In addition, the divergence



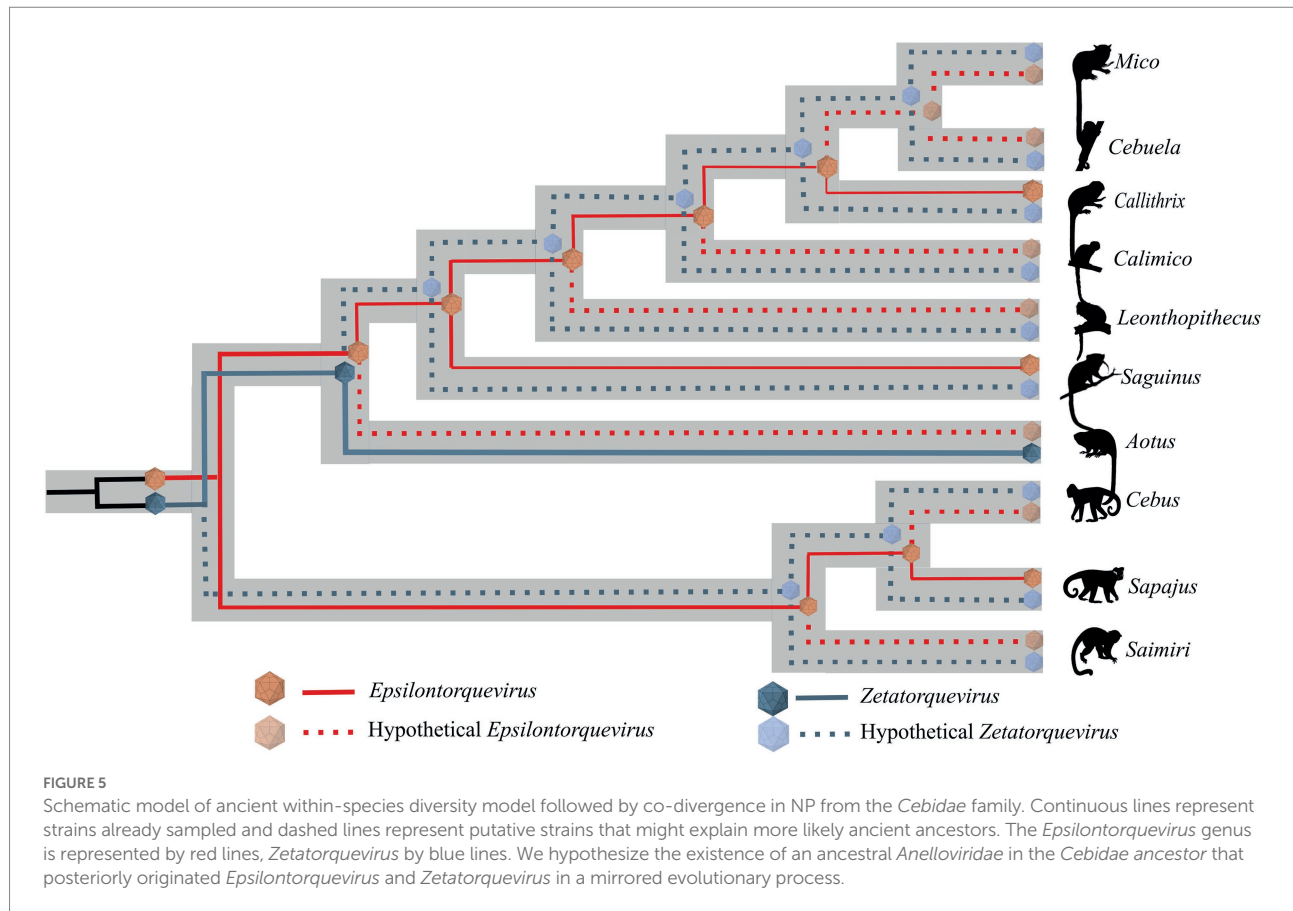
across the TTV tamarin and *E. callithrichensis* was estimated around 15.48 mya (95% HPD=12.73–18.17). The divergence between the *E. callithrichensis I* and *E. callithrichensis II* was estimated around 13.1 mya (95% HPD=10.73–15.50 mya). The ancestor for *E. callithrichensis I* was dated around 7.9 mya (95% HPD=6.37–9.54 mya) and the one for *E. callithrichensis II* around 9.6 mya (95% HPD=7.71–11.2 mya) (Figure 4).

Discussion

With the development of HTS protocols to explore the diversity of existing viruses, previous works have identified new anelloviruses species in wildlife—bats, rodents (Niewiadomska and Gifford, 2013), seals (Weber et al., 2020), wild felids (Noppornpanth et al., 2001), canids (Pujol et al., 2005), and non-human primates (Okamoto et al., 2000; Hon et al., 2008; Holmes, 2010; Ng et al., 2011; Sawaswong et al., 2019; dos Santos et al., 2020). In this work, we sequenced and characterized two novel *Anelloviridae* species genomes in NP, increasing the host

range for this family. To the best of our knowledge, this is the first report of this viral family in the black-tufted marmoset (*C. penicillata*; Okamoto et al., 2000; Ng et al., 2011; de Souza et al., 2018), the known diversity in NP is yet remarkably limited. Until the present study, only two studies identified anelloviruses (Ng et al., 2011; dos Santos et al., 2020), though no effort to characterize their evolutionary dynamics has been presented. Hence, this is the first attempt to investigate in depth the evolution of novel NP anellovirus.

The anelloviruses sequenced in this work were unique to the known NP *Anelloviridae* diversity, comprehending two novel species by the ICTV species criterion (Figures 1, 2; Table 3). Both viruses' complete genomes had the expected ORFs found in this viral family: the putative capsid protein and replication-associated protein (ORF1) and the ORF2 protein (75), as well as the putative ORF3. By particularly looking at some regions of the novel genomes, we could identify regions similar to TTV tamarin and TTV capuchin ORF4, though with numerous stop codons and frameshift mutations, implying lack of transcriptional function. This suggests ORF4 is not fundamental for *E. callithrichensis*



infection and shows an instance of genomic structure plasticity in ssDNA viruses. The adoption of models and cell systems to study anellovirus infections *in vitro* shall improve the knowledge regarding the function and structure of its proteins, deeply scarce at the present time.

Phylogenetic analysis of the new genome sequences suggests a topological pattern consistent with a previously proposed model for the diversification of multiple viral families, the within-species diversity followed by co-divergence model (Buchfink et al., 2014; Nurk et al., 2017; Chen et al., 2018; D'arc et al., 2020). Under this model, viral lineages diversify either alongside their host species or through a speciation process within a single host, occasionally caused by niche adaptation to specific cell types or tissues. This process culminates in a viral phylogenetic pattern where multiple clades reflect the host phylogeny, as is the case of primate TTVs (Ng et al., 2011). We explored the consequences of this model to calibrate the molecular clock and obtain, for the first time, a time-scaled phylogenetic tree for NP TTVs (Figure 4). This analysis suggests that the lineages that lead to both species described in this study diverged around 13.1 mya, while the time for the ancestors of *E. callithrichensis* I and II were dated around 7.9 and 9.6 mya, respectively. Both dates are much older than the origins of *Callithrix*, implying that these viral lineages circulated among the ancestors of *Callitrichinae* (around 14.9 mya; Gernhard, 2008). We thus hypothesize that its descendant extant species may harbor TTVs

related to the described viral lineages (Figure 5). A central limitation of the presented analysis lies in the limited sampling of NP TTVs. Increasing the availability of anellovirus genomes from diverse NP species will allow a deeper evaluation of their diversification dynamics and a further verification of our hypothesis.

We observed a high genetic diversity between the *E. callithrichensis* characterized herein (Figure 3), a phenomenon already described in anelloviruses due to their high mutation rate (76–78). The presence of multiple lineages co-infecting a host has already been observed as three distinct *Anelloviridae* genera were described in humans (79). The same pattern is present in OWP, as chimpanzees are infected by *Alpha*, *Beta*, and *Gammatorquevirus* (Inami et al., 2000; Okamoto et al., 2000; Ng et al., 2011; de Souza et al., 2018). Once the NP *Anelloviridae* diversity is further explored, and the diversity of *Epsilontorquevirus* and *Zetatorquevirus* is thoroughly addressed, the pattern of coinfection by multiple TTV strains may be further verified in NP (Figure 5).

While previous observations of co-divergence patterns have been made for other TTVs (Noppornpanth et al., 2001; Ng et al., 2011), no clear evolutionary model has been proposed to assess TTV diversity. Although this study focused on NP TTVs, we argue that the within-species diversity followed by co-divergence model might explain the drivers of diversification for the whole *Anelloviridae* family. Further characterization of TTVs diversity in non-model organisms through metagenomics

might play a key role in unraveling the validity of the proposed model. Finally, this study showcases an instance of the importance of sampling wildlife viruses to obtain a better characterization of the long-term evolutionary processes driving the diversification of viral families.

Data availability statement

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found at: <https://www.ncbi.nlm.nih.gov/genbank/>, OM416781–OM416795; https://github.com/matheus-cosentino/Cosentino_2022_Epsilontorquevirus.

Ethics statement

The animal study was reviewed and approved by Ethics Committee on the Use of Animals of Universidade Federal do Rio de Janeiro (reference number 037/14).

Author contributions

The idea of characterizing anelloviruses originated from MD's work on viral metagenomics. CD, AS, and MT were responsible for sample collection. Sequencing data by MD. Analysis and curation of generated data by MC, MD, FM, LC, RM, AC, FS, TM, and GM. Genome annotations, phylogenetic analyses, molecular dating, and images by MC, MD, and FM. Manuscript ideas and development by all authors. All authors contributed to the article and approved the submitted version.

Funding

This work was supported in part by the Conselho Nacional de Desenvolvimento Científico e Tecnológico/CNPq (grant

313005/2020–6 to ASa) and Fundação de Amparo à Pesquisa do Estado do Rio de Janeiro/FAPERJ (grants E-26/112.647/2012 to MS and E26/202.738/2018, E26/210.122/2018, and E26/211.040/2019 to ASa) for funding this study.

Acknowledgments

We thank to all the workers of the Centro de Primatologia de Brasília, Universidade de Brasília (Distrito Federal, Brazil) for assistance with sample collection and to the fantastic LDDV members, in special to Déa Mota, for the wonderful teamwork. We are also grateful to Carolina Furtado and Nicole Scherer from Instituto Nacional de Câncer José de Alencar (Rio de Janeiro, Brazil) for assistance with HTS and analysis.

Conflict of interest

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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Supplementary material

The Supplementary material for this article can be found online at: <https://www.frontiersin.org/articles/10.3389/fmicb.2022.1002963/full#supplementary-material>

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