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Article

Characterization of Sylvatic Mosquito Viromes in the Cerrado Biome of Minas Gerais, Brazil: Discovery of New Viruses and Implications for Arbovirus Transmission

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Abstract: Studies on animal virome have mainly concentrated on chordates and medically significant invertebrates, often overlooking sylvatic mosquitoes, constituting the major part of mosquito species diversity. Despite their potential role in arbovirus transmission, the viromes of sylvatic mosquitoes remain largely unexplored. These mosquitoes may also harbor insect-specific viruses (ISVs), affecting arboviral transmission dynamics. The Cerrado biome, known for rapid deforestation and its status as a biodiversity hotspot, offers an ideal setting for investigating mosquito viromes due to potential zoonotic spillover risks from land use changes. This study aimed to characterize the viromes of sylvatic mosquitoes collected from various locations within Minas Gerais state, Brazil. Total RNA was extracted from mosquito pools of *Psorophora albipes*, *Sabethes albiprivus*, *Sa. chloropterus*, *Psorophora ferox*, and *Coquillettidia venezuelensis* species, followed by high-throughput sequencing (HTS). Bioinformatic analysis included quality control, contig assembly, and viral detection. Sequencing data analysis revealed 11 near-complete viral genomes (new viruses are indicated with asterisks) across seven viral families and one unassigned genus. These included: *Xinmoviridae* (Ferox mosquito mononega-like virus* and Albipes mosquito Gordis-like virus*), *Phasmaviridae* (*Sabethes albiprivus* phasmavirus*), *Lispiviridae* (Pedras lispivirus isolate MG), *Iflaviridae* (*Sabethes albiprivus* iflavivirus*), *Virgaviridae* (Buriti virga-like virus isolate MG and *Sabethes albiprivus* virgavirus 1*), *Flaviviridae* (*Psorophora ferox* flavivirus*), *Mesoniviridae* (Alphamesonivirus cavallyense isolate MG), and the genus *Negevirus* (Biggie virus isolate MG virus and *Coquillettidia venezuelensis* negevirus*). Phylogenetic analyses revealed evolutionary relationships with varying degrees of similarity to known sequences in distinct host species. In some cases, new viruses and potential new genera were discovered, emphasizing the importance of studying sylvatic mosquito virome to understand viral diversity and evolution in ecologically sensitive regions. Moreover, the presence of ISVs and potential novel arboviruses underscores the need for ongoing surveillance and control strategies to mitigate the risk of emerging infectious diseases.

Keywords: sylvatic mosquitoes; RNA virome; brazilian cerrado; novel viral species; arbovirus transmission; biodiversity hotspot

1. Introduction

Studies on animal virome have predominantly focused on chordates and medically significant invertebrates such as ticks and mosquitoes [1]. Within mosquitoes, urban species like *Aedes aegypti*, *Ae. albopictus* and *Culex quinquefasciatus*, known for their role in arbovirus transmission, have been the primary subjects of virome investigations [2,3]. However, despite an estimated global mosquito species count exceeding 3,700 [4], sylvatic mosquitoes remain underexplored regarding their viromes. This gap is significant given the critical role of certain species, such as *Sabethes spp* in Yellow Fever Virus transmission [5] and *Coquillettidia spp.* in Eastern Equine Encephalitis virus [6] and Oropouche virus transmission [7,8].

In addition to arboviruses, mosquitoes may harbor insect-specific viruses (ISVs) which, although incapable of infecting vertebrates, offer valuable perspectives into viral evolution due to their close phylogenetic relationship with arboviruses. Some ISVs have shown the ability to modulate arboviral transmission from vectors to mammalian hosts, suggesting their potential role in developing innovative strategies for controlling arboviral outbreaks [9,10].

In parallel, high-throughput sequencing (HTS) has emerged as a critical tool for discovering new viruses, including ISVs, and determining virome in mosquitoes [11–13]. HTS enables the comprehensive and unbiased analysis of viral genetic material present in mosquito samples, facilitating the identification of both known and novel viruses [14]. This technology is particularly valuable for exploring the viromes of sylvatic mosquito species, which are often understudied compared to their urban counterparts [15]. By offering intricate perspectives into the diversity and composition of mosquito viromes, HTS enhances our understanding of viral ecology and evolution, supports the development of more effective vector control and surveillance strategies, and can be utilized to discover new viruses in the largely unexplored Brazilian biomes.

The Cerrado, a biodiversity hotspot in Brazil [16], faces alarming rates of deforestation, driven in part by expanding monoculture activities [17]. Given that anthropogenic alterations in land use are major contributors to zoonotic spillover events [18], understanding the viromes associated with arthropod vectors in this biome is crucial for enhancing genomic surveillance of emerging arboviruses. Therefore, the present study aims to deepen our understanding of RNA viromes within sylvatic mosquitoes inhabiting the Cerrado biome.

2. Materials and Methods

2.1. Sample Collection

Sylvatic mosquitoes were collected across five municipalities within the Minas Gerais state (Figure 1). Additionally, a mixed pool comprising diverse mosquito species was included in the study, including *Sabethes spp* and *Ps. ferox* (Table 1).

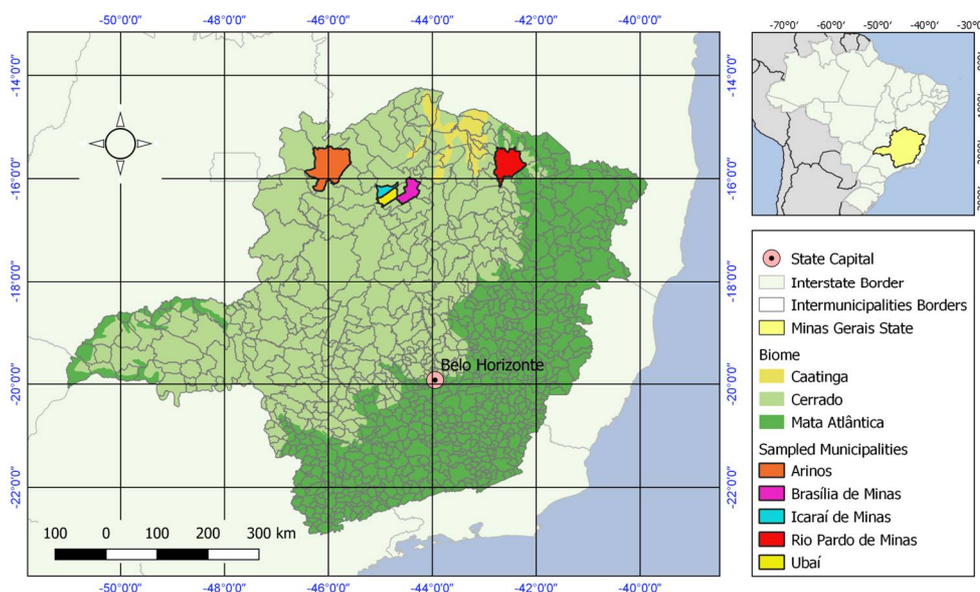


Figure 1. Map of Brazil and Minas Gerais state depicting the municipalities from which mosquito samples were collected and associated biomes.

Table 1. Description of Species, Number of Mosquitoes, and Municipality in Minas Gerais state where mosquito samples were collected.

| Species per pool | Mosquitos per pool | Municipality |
|---------------------------------------------------------------------------------------------------------------------------------------|--------------------|--------------------------------------------|
| <i>Psorophora (Janthinosoma) albipes</i> (Theobald, 1907) ¹ | 20 | Brasília de Minas |
| <i>Sabethes (Sabethes) albiprivus</i> (Theobald, 1903) ² | 200 | Icarai de Minas and Ubaí |
| Mixed pool (<i>Sa. albiprivus</i> , <i>Sa. (Sabethoides) chloropterus</i> (von Humboldt, 1819) ³ , and <i>Ps. ferox</i>) | 55 | Icarai de Minas, Arinos, Brasília de Minas |
| <i>Psorophora (Janthinosoma) ferox</i> (von Humboldt, 1819) ³ | 100 | Rio Pardo de Minas |
| <i>Coquillettidia (Rhynchoaenia) venezuelensis</i> (Theobald, 1912) ⁴ | 100 | Arinos |

¹ Theobald, F.V. 1907. A monograph of the Culicidae or mosquitoes. Volume 4. British Museum (Natural History), London. xix+ 639 pp. (1907.02.23). ² Theobald, F.V. 1903. A monograph of the Culicidae or mosquitoes. Volume 3. British Museum (Natural History), London. xvii + 359 pp. (1903.07.25). ³ von Humboldt, F.H.A. (1819). Voyage aux regions equinoxiales du Nouveau Continent, fait en 1799, 1800, 1801, 1802, 1803 et 1804 par Al. de Humboldt et A. Bonpland; Vol. 2, 722 pp. In Humboldt, F.H.A., Ed., Voyage de MM Alexandre de Humboldt et Aime Bonpland. Pt. I: Relation historique. Paris. ⁴ Note sur le Culicides. Bulletin du museum national d'histoire naturelle 18 (12): 59-61. 29 Février 1912.

Adult mosquitoes were captured employing the protected human attraction method [19] and the Shannon (1939) [20] trap, with assistance from entomological nets and oral aspirators [21]. Following capture, the mosquitoes were sorted by genera and cryopreserved in liquid nitrogen (-196°C) before being transported to the laboratory. Taxonomic classification was conducted on a chilled table at -20°C utilizing a stereoscopic microscope and dichotomous keys [22,23].

2.2. RNA Extraction and High-Throughput Sequencing

Total RNA was extracted from mosquito pools of varying sizes (20 to 200 individuals - Table 1) using the QIAmp Viral RNA (Qiagen) kit. The extracted RNA was quantified using a Quantifluor® RNA system (Promega) kit, following the manufacturer's instructions. For samples containing at least 1 µg of RNA, 0.1 v/v of 3M sodium acetate and 2 v/v of absolute ethanol were added. These samples were then dispatched to Macrogen (South Korea), where Illumina TruSeq stranded Total RNA library + Ribo-zero Gold libraries were generated for NovaSeq 6000, 100bp paired-end sequencing, resulting in 20M to 30M reads per library.

2.3. Bioinformatic Quality Control, Assembly of Contigs, and Viral Detection

We conducted quality control analysis of the raw reads using fastQC [24], Trim-galore for adapter removal v0.6.10 [25] and Trimmomatic v 0.39 [26] to trim 10 bp from each end of the reads. The assembly of trimmed reads into contigs was performed using Spades v3.15.5, with the -rnviral flag [27]. Viral contigs were identified through local diamond blastx searches [28] of RNA-dependent RNA-polymerases compiled in RdRP-scan [29]. To identify M and S segments of *Phasmaviridae*, local diamond blastx searches were conducted using Refseq sequences deposited in Genbank [30] (assessed on 08/07/2023). Viral contigs were verified with BLASTx searches against the non-redundant protein sequences in NCBI, and viral completeness was assessed using CheckV [31]. Viral genomes were annotated utilizing Geneious v. 11.1.5 (Biomatters) and ORFfinder (<https://www.ncbi.nlm.nih.gov/orffinder/>). The obtained viral sequences were deposited in Genbank with accession numbers PP946236 - PP946246.

2.4. Phylogenetic Analyses

Global multiple alignments were created for each viral family and protein marker using Mafft v7.520 [32]. The resulting alignments were subsequently trimmed by employing TrimAI v1.4.15 [33] with the -automated1 flag. The amino acid substitution models for each alignment were determined using modeltest-NG v0.1.7 [34]. Maximum Likelihood trees were then constructed utilizing RAxML-NG v1.2.0 [35] with 1000 bootstraps. All trees, otherwise stated, were mid-point rooted.

3. Results

Each sequencing library comprised approximately 20.5 to 30.4 million reads. Post-assembly, each library produced around 2,014 to 37,238 contigs (Table 2). The higher contig count in the mixed mosquito pool is attributed to its diverse species composition, compared to the single-species composition of the other insect pools.

Table 2. Viral diversity in insect pools included read numbers, contigs, viral families, genome type, and viruses found in this study including new viruses.

| Insect Pool | Read number (Millions) | Contig amount | Viral contigs | Viral families | Genome Type | Viruses found in this study (* new viruses ¹) |
|---------------------------------------------------------------------------------------|------------------------|---------------|---------------|----------------------|---------------------|-----------------------------------------------------------|
| <i>Ps. albipes</i> | 20.5 | 3,882 | 1 | <i>Xinmoviridae</i> | ssRNA- | Albipes mosquito Gordis-like virus* |
| | | | | <i>Phasmaviridae</i> | ssRNA- ² | Sabethes albiprivus phasmavirus* |
| <i>Sa. albiprivus</i> | 25.1 | 13,548 | 4 | <i>Lispividae</i> | ssRNA- | Pedras lispivirus isolate MG |
| | | | | <i>Iflaviridae</i> | ssRNA+ | Sabethes albiprivus iflavivirus* |
| | | | | <i>Virgaviridae</i> | ssRNA+ | Sabethes albiprivus virgavirus 1* |
| Mixed (<i>Sa. albiprivus</i> , <i>Sa. chloropterus</i> , and <i>Ps.ferox</i>) | 26.2 | 37,238 | 1 | <i>Virgaviridae</i> | ssRNA+ | Buriti virga-like virus isolate MG |
| <i>Ps. ferox</i> | 30.4 | 2,859 | 2 | <i>Xinmoviridae</i> | ssRNA- | Ferox mosquito mononega-like virus* |
| | | | | <i>Flaviviridae</i> | ssRNA+ | Psorophora ferox flavivirus* |
| <i>Cq. venezuelensis</i> | 25.6 | 2,014 | 3 | <i>Mesoniviridae</i> | ssRNA+ | Alphamesonivirus cavallyense isolate MG |

Biggie virus isolate MG
Coquillettidia velezuensis
negevirus*

¹ New viruses are indicated with asterisks. ² Segmented.

The HTS data analysis uncovered 11 near-complete viral genomes spanning seven distinct viral families and one genus of an unassigned family. Among these were the ssRNA- families *Xinmoviridae*, *Phasmaviridae*, and *Lispiviridae*, and the ssRNA+ families *Iflaviridae*, *Virgaviridae*, *Flaviviridae*, *Mesoniviridae* along with the genus *Negevirus* (Table 2). Of the 11 viral genomes described below, 7 are new, while the 4 previously known genomes were identified in new host species.

3.1. *Xinmoviridae*

We identified two consensus putative viral sequences, measuring 13 kb and 12 kb, in *Ps. ferox* and *Ps. albipes*, respectively. Their encoded RdRps exhibited 59.02% and 51.10% similarity with their closest matches in Genbank (Guadeloupe mosquito mononega-like virus, MN053735, and Gordis virus, MW435014) within the *Xinmoviridae* family (Figure 2A and 2B). According to ICTV criteria [36], RdRp sequences showing less than 60% similarity within this family may indicate the presence of novel genera. Both of these present genes for a putative nucleoprotein, an envelope glycoprotein, and an RdRp. Despite these viruses being discovered [37,38] in similar hosts (same Culicidae family, but different genera), their positions within the RdRp tree are notably distinct (Figure 2C). Hence, we tentatively named these viruses Ferox mosquito mononega-like virus and Albipes mosquito Gordis-like virus, respectively.

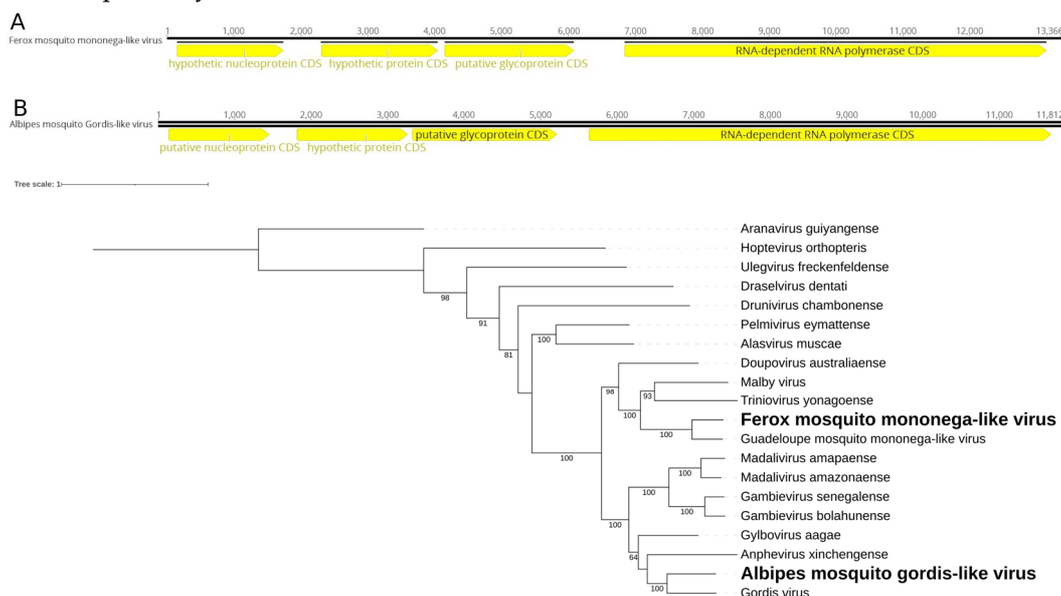


Figure 2. Ferox mosquito mononega-like virus and Albipes mosquito Gordis-like virus genomes discovered in *Ps. ferox* (A) and *Ps. albipes* (B), respectively. C. Phylogenetic analysis of the Ferox mosquito mononega-like virus and Albipes mosquito Gordis-like virus discovered in *Ps. ferox* and *Ps. albipes* hosts, respectively. ML tree was constructed using the RdRp amino acid sequences from the best hits obtained via BLASTx for each xinmovirus identified in this study against the NCBI nr-database. The tree includes currently recognized species in the family by ICTV, with Aranavirus as an outgroup.

3.2. *Phasmaviridae*

Our investigation revealed three distinct putative viral segments - small (S), medium (M), and large (L) segments - in the *Sa. albiprivus* pool, with lengths of 6.5 kb, 2.1 kb, and 1.7 kb, coding for its RdRp, envelope glycoprotein and capsid protein, respectively (Figure 3A). Comparison with Genbank sequences identified an RdRp sequence showing 45.50% similarity to Miglotas virus (QRW41774.1) [38] within the *Phasmaviridae* family. As this RdRp shares less than 95% identity, as

per ICTV criteria, we suggest the classification of a novel virus species, tentatively named *Sabethes albiprivus phasmavirus*. Phylogenetic analysis supports the distinction of this new virus, forming a separate monophyletic branch (Figure 3B).

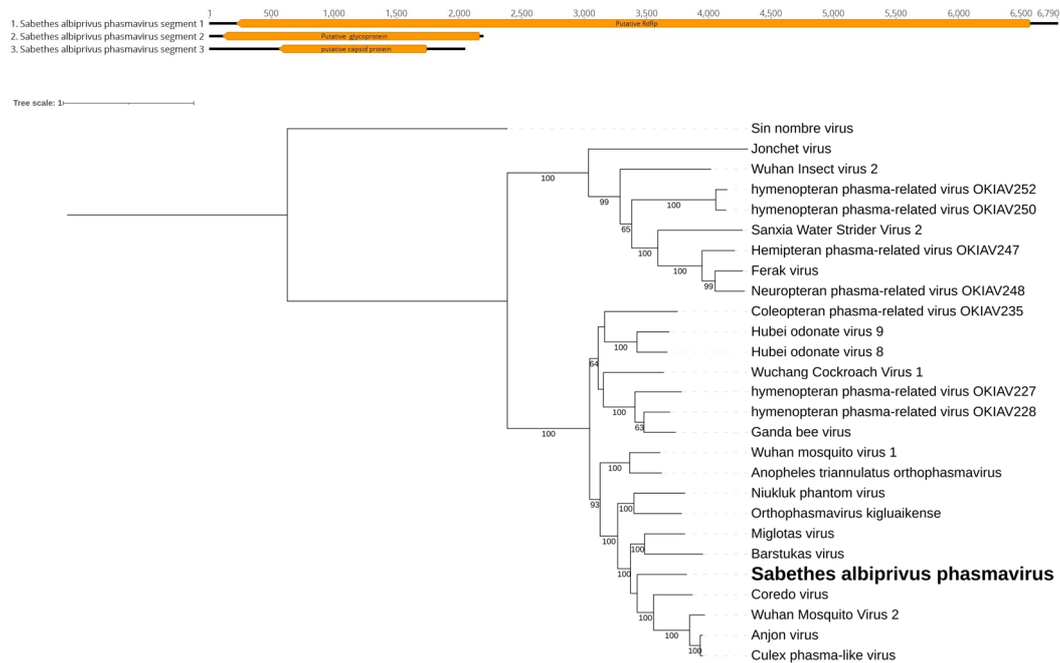
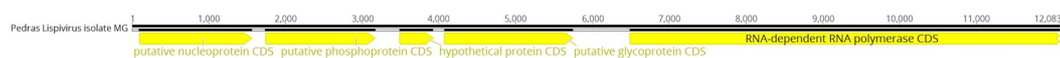


Figure 3. A. *Sabethes albiprivus phasmavirus* genome exhibits L, M, and S segments identified within *Sa. albiprivus*. **B.** Phylogenetic analysis of the *Sabethes albiprivus phasmavirus* discovered in *Sa. albiprivus* host. ML tree was constructed using the RdRp amino acid sequences of the best hits from BLASTx against the NCBI nr-database for the virus in the current study, with Sin nombre virus as an outgroup.

3.3. *Lispiviridae*

We identified a consensus putative viral genome of 12 kb for the *Sa. albiprivus* pool, containing five open reading frames (ORF) (Figure 4A). Its genes include a nucleoprotein, a phosphoprotein, an envelope glycoprotein, and an RdRp. Notably, its RdRp sequence shares more than 85% similarity with that of Pedras lispivirus [39], indicating classification within the same species within the *Lispiviridae* family, according to ICTV criteria [40]. Hence, we designate this virus as Pedras lispivirus isolate MG. In their study [39], previously identified this virus in *Sa. quasicyaneus*, inhabiting a transitional zone between the Cerrado and Amazon biomes. Despite geographical separation, the hosts of this virus share a close evolutionary relationship (same subgenus - *Sabethes Sabethes*) (Figure 4B), suggesting potential occurrence in other species within this mosquito genus, a hypothesis warranting further investigation.



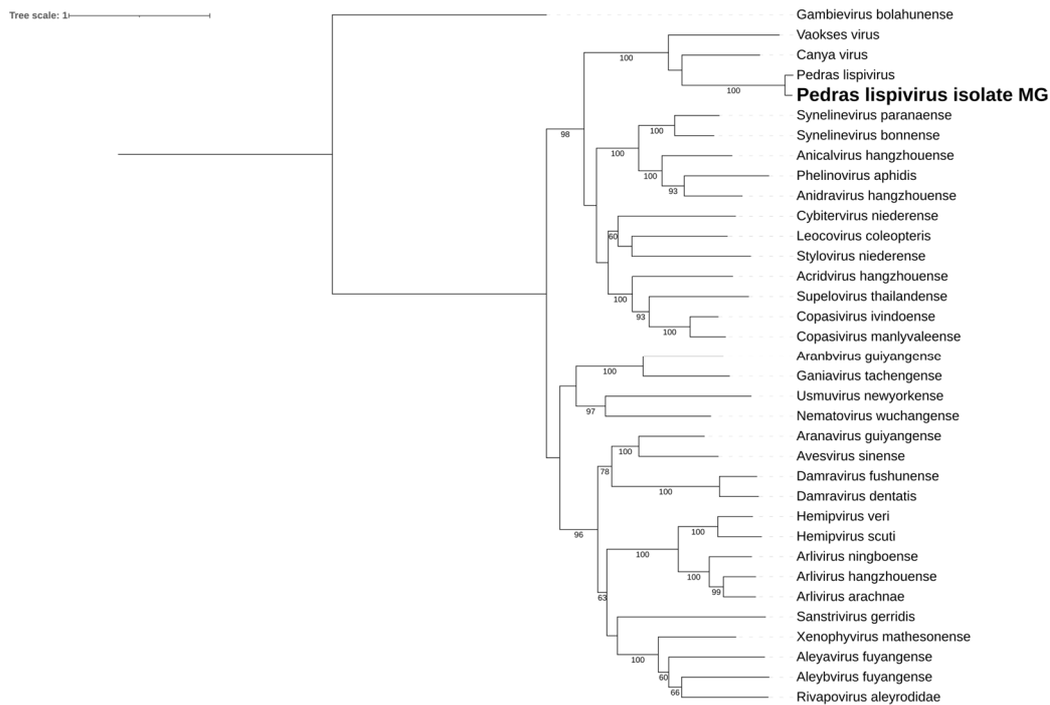
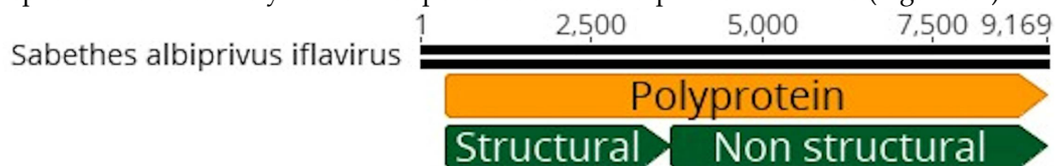


Figure 4. A. Pedras lispivirus isolate MG genome identified within *Sa. albiprivus*. **B.** Phylogenetic analysis of the Pedras lispivirus isolate MG discovered in the *Sa. albiprivus* host. ML tree was constructed with the RdRp aminoacid sequences of the best hits found in the present study against the NCBI nr-database, currently recognized species in the family by ICTV, and Gambievirus as an outgroup.

3.4. Iflaviridae

We identified a consensus putative viral genome of approximately 9 kb for the *Sa. albiprivus* pool (Figure 5A). This genome encodes a single polyprotein comprising both structural and non-structural components, showing homology to viruses from the *Iflaviridae* family. The structural component, responsible for coding the capsid proteins, has its closely related sequence in Genbank with 62% coverage and 71.65% similarity to Aedes Iflavi-like virus 1 (QQD36915.1), a virus isolated from *Ae. aegypti* in the Brazilian Amazon [41]. According to the criteria set by the ICTV [42], the structural component's similarity is lower than 90%, and its occurrence in *Sa. albiprivus*, suggests that this is a novel species. We tentatively name this species *Sabethes albiprivus iflavivirus* (Figure 5B).



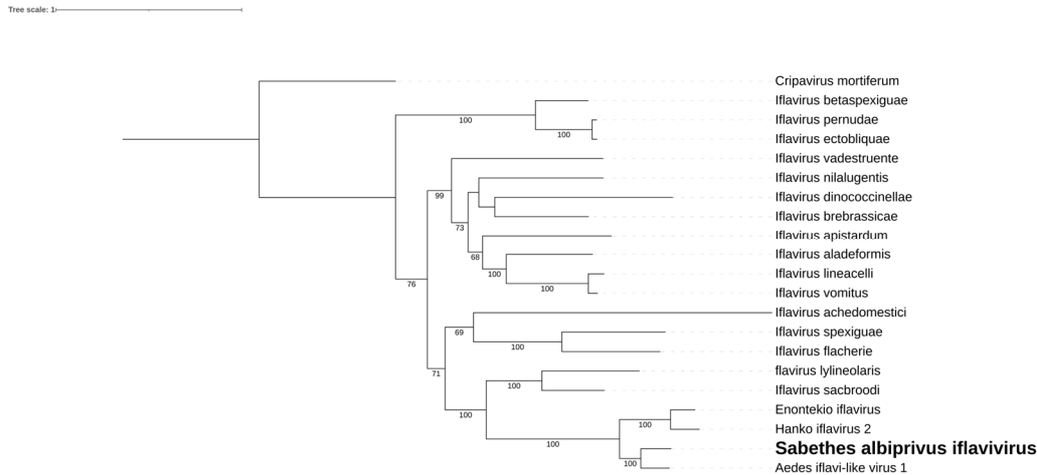


Figure 5. A. Sabethes albiprivus iflavivirus genome identified within *Sa. albiprivus*. **B.** Phylogenetic analysis of the Sabethes albiprivus iflavivirus discovered in the *Sa. albiprivus* host. ML tree was constructed using the polyprotein sequences of the best BLASTx hits found in the present study against the NCBI nr-database, currently recognized species in the family by ICTV, and Cripavirus as an outgroup.

3.5. Virgaviridae

We identified two putative viral consensus genomes of approximately 8.8 kb and 9.1 kb in the mixed pool (*Sa. albiprivus*, *Sa. chloropterus*, and *Ps.ferox*) and *Sa. albiprivus* pool, respectively (Figure 6A), showing homology to viruses from the *Virgaviridae* family, each with a polyprotein coding gene and a putative capsid coding gene, albeit in differing synthesis. We tentatively name these viruses, respectively, Buriti virga-like virus isolate MG and Sabethes albiprivus virga virus 1. In the phylogenetic analysis, Buriti virga-like virus isolate MG clusters with Buriti virga-like virus found in *Sa. chloropterus* [39] (Figure 6B). Considering the ICTV criteria for taxa classification within this family (https://ictv.global/report_9th/RNApos/Virgaviridae), metadata such as the mode of transmission is required. Therefore, we are currently unable to propose a taxon for Sabethes albiprivus virgavirus 1.

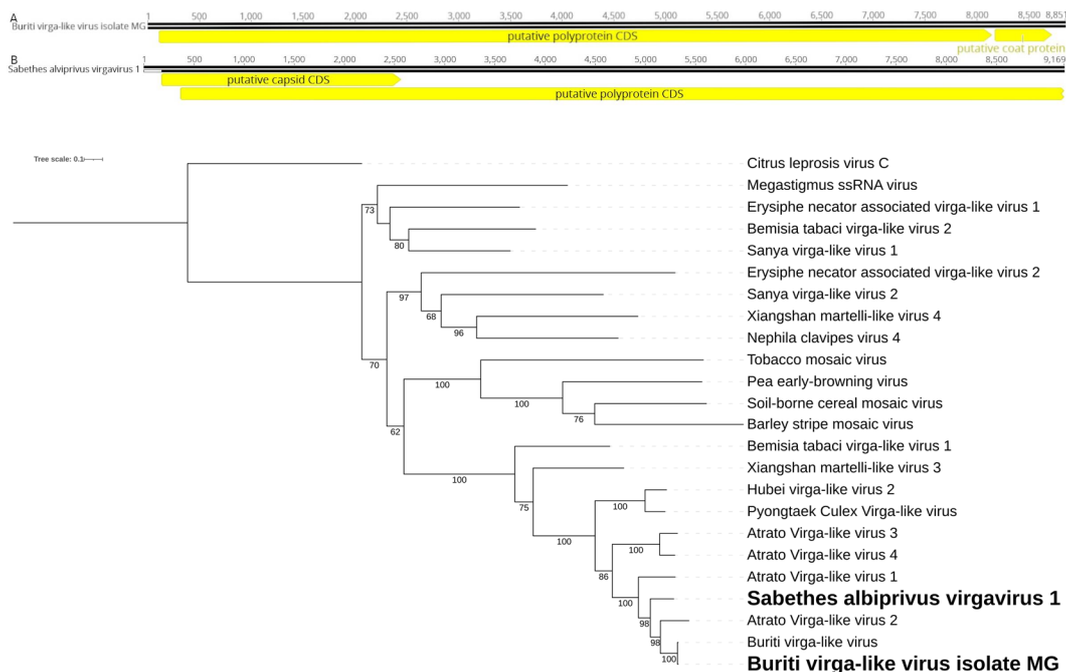


Figure 6. A. Buriti virga-like virus isolate MG (A) and Sabethes albiprivus virgavirus 1 (B) genomes identified within *Sa. albiprivus*. **B.** Phylogenetic analysis of the Buriti virga-like virus isolate MG and Sabethes albiprivus virgavirus 1 discovered in *Sa. albiprivus* host. ML tree was constructed using the polyprotein aminoacid sequences of the best hits in BLASTx against the NCBI nr-database for the viruses in the current study, with Citrus leprosis virus as an outgroup.

3.6. Flaviviridae

In the *Ps. ferox* pool, we identified a consensus putative viral genome of approximately 11 kb (Figure 7A) showing homology to viruses from the *Flaviviridae* family. The genome includes the ORFs characteristic of flaviviruses, encoding the polyprotein that is cleaved into structural (C, prM, E) and non-structural (NS1, NS2A, NS2B, NS3, NS4A, NS4B, NS5) proteins. NS5 encoded RdRp that presented 55.73% identity with its most closely related sequence in GenBank, *Mansonia flavivirus* (LC567153) [43]. We have tentatively named this virus *Psorophora ferox flavivirus*. Considering the ICTV criteria for the family (https://ictv.global/report_9th/RNApos/Flaviviridae), classification requires metadata beyond sequence information, including antigenicity and ecological characteristics, which are beyond the scope of this study. Therefore, we lack sufficient data to propose a new taxon for *Psorophora ferox flavivirus*. Given its placement within the phylogenetic tree, it is most likely an ISV (Figure 7B).

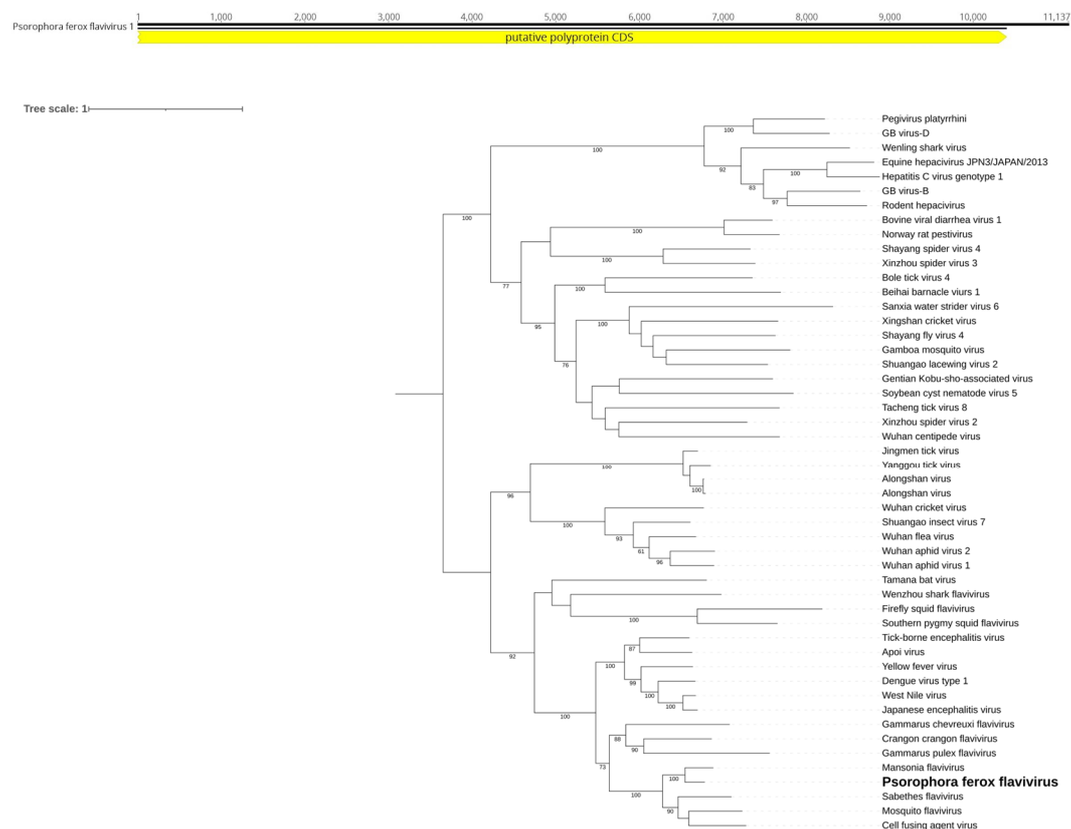


Figure 7. A. Flavivirus genome identified within *Ps. ferox*. **B.** Phylogenetic analysis of the *Psorophora ferox flavivirus* discovered in the *Ps. ferox* host. ML tree was constructed using the polyprotein sequences of the best hits from BLASTx found in the present study against the NCBI nr-database.

3.7. Mesoniviridae

We identified a consensus putative viral genome consisting of 20,198 bases in *Cq. venezuelensis* (Figure 8A), showing homology to viruses within the *Mesoniviridae* family. Its gene synteny is the same as observed in the species, including genes encoding for two polyproteins for its non-structural proteins and smaller structural coding genes near its 3' end. The encoded putative RdRp has over

99% similarity to that of *Alphamesonivirus cavallyense* (AXL48235.1) found in *Culex pipiens* [44]. Notably, *Cq. venezuelensis* and *Cx. pipiens* differ significantly in terms of phylogeny, behavior, and geographic distribution. These species belong to different tribes, Mansonini and Culicini, respectively. *Cq. venezuelensis* exhibits wild habits and is restricted to the Americas [45], whereas *Cx. pipiens* is urban and invasive across multiple continents [46]. In the phylogenetic analysis, this virus found in *Cq. venezuelensis* formed a monophyletic branch (Figure 8B). We tentatively name this *Alphamesonivirus cavallyense* isolate MG.

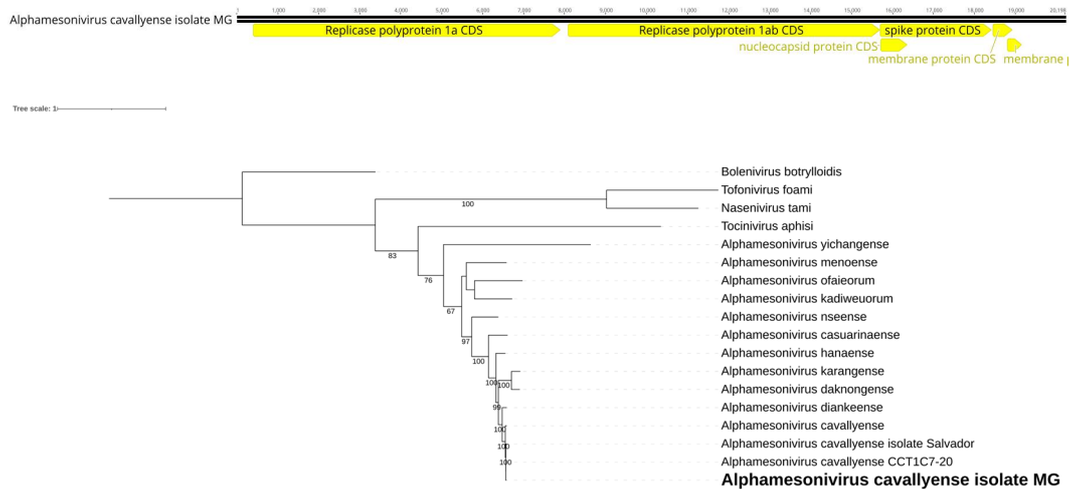
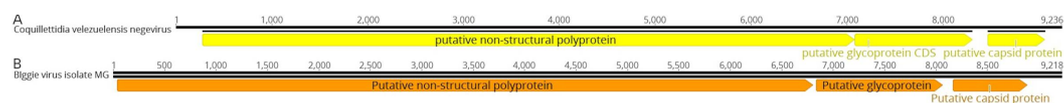


Figure 8. A. *Alphamesonivirus cavallyense* isolate MG genome identifies within *Cq. venezuelensis*. **B.** Phylogenetic analysis of the *Alphamesonivirus cavallyense* isolate MG discovered in *Cq. venezuelensis* host. ML tree was constructed using the amino acid sequences of the ORF1a replicase from representatives of the four currently recognized genera within the family, along with the two best BLASTx hits from the nr-database and *Bolenivirus* as an outgroup.

3.8. Negevirus

We identified two putative consensus viral genomes of 9.2 Kb in *Cq. velezuelensis* (Figure 9A and 9B), showing homology to viruses within the Negevirus genus. Both of these present a gene synteny typical of the genus, including a non-structural polyprotein, an envelope glycoprotein, and a capsid protein. One of these genomes encodes a replicase with 74% coverage and over 99% similarity to its closest sequence in Genbank, Biggie virus Mos11 (KX924639.1) found in *Culex pipiens* collected in the USA [47]. We have tentatively named this Biggie virus isolate MG. As previously mentioned, these hosts differ significantly, which may suggest considerable plasticity of this virus. The second genome encodes a replicase with 73% coverage and 66% similarity to its closely related sequence, Aqua Salud Negevirus (MT197494.1) detected in *Culex declarator* in Panama [48], also a quite different host. We tentatively named this virus *Coquillettidia velezuelensis* negevirus (Figure 9C).



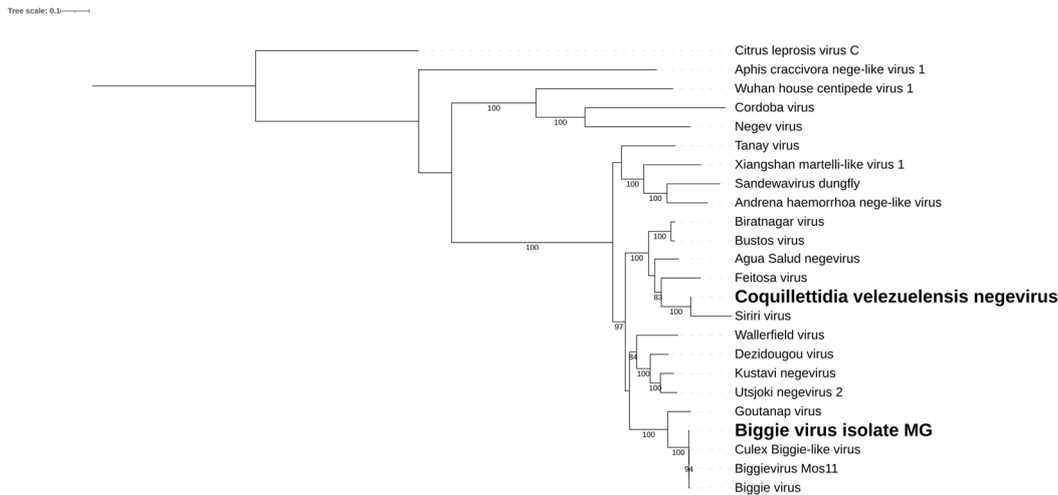


Figure 9. Negevirus genomes identified within *Cq. velezuelensis*. A) *Coquillettidia velezuelensis* negevirus. B) Biggie virus isolate MG. C. Phylogenetic analysis of the negevirus discovered in *Cq. velezuelensis* host. ML tree was constructed using the non-structural polyprotein amino acid sequences of the best hits from BLASTx against the NCBI nr-database for the virus in the current study, with *Citrus leprosis virus* as an outgroup.

4. Discussion

In this study, we identified seven novel viruses and four previously known genomes identified in new host species, all of which are underrepresented in virological research. The discovery of new ISVs and potential arboviruses is particularly significant as it broadens our understanding of viral diversity within mosquito populations and reveals the potential ecological roles these viruses may play. Given the capacity of ISVs to interact with or impede arbovirus transmission [49], our findings underscore the necessity of further exploring these interactions. Moreover, HTS once again demonstrates its remarkable ability to discover new viruses, even in pools of mosquitoes from different species, thereby enhancing our understanding of comprehensive virome surveillance in regions undergoing rapid environmental changes, such as the Cerrado Brazilian biome. Compared to the work of Silva et al. (2023) [13], the number of full RdRp sequences detected was similar to our study. However, the viruses' characterization by authors was partial. Interestingly, only one mosquito species, *Cq. venezuelensis*, was the same host searched, but the viruses identified were different. This highlights the diversity of sylvatic mosquitoes and underscores the need for further research. Below we discuss in detail the viruses detected and characterized here.

4.1. *Xinmoviridae*

Infections with anpheviruses (family *Xinmoviridae*; order *Mononegavirales*) seem to be relatively prevalent, evidenced by their presence in laboratory colonies and wild populations of *Aedes albopictus* from various geographic locations over several years [50,51]. Notably, *Wolbachia* bacteria enhance *Aedes* anphevirus replication, while this virus marginally inhibits DENV replication in mosquito cell lines [50]. Anpheviruses have also been identified in populations of *Culex* spp. [52] and anophevirus-like species have been found in *Amazonian anophelines*, including *Anopheles marajoara* and *An. darlingi* [53]. Here, we described two putative new *Xinmoviridae* species in two *Psorophora* sp. mosquitoes. The widespread distribution of most *Psorophora* species in the Americas [23] poses a potential threat to public health due to their capacity to transmit several arboviruses such as Yellow Fever Virus, Rocio, West Nile Virus, Eastern Equine Encephalitis and Ilheus Virus [54–59]. The capability of arboviruses to infect *Psorophora* species suggests a potential for disease transmission. The discovery of Ferox mosquito mononega-like virus and Albipes mosquito Gordis-like virus in *Ps. ferox* and *Ps. albipes*, respectively, presents new avenues for future studies aiming to understand the interactions between ISVs and other arboviruses in the context of pathogen transmission. The

identification of these novel viruses characterized by their distinct positions within the RdRp phylogenetic tree and less than 60% similarity to known viruses, hints at the potential presence of new genera within the *Xinmoviridae* family, emphasizing the necessity for further research into the diversity and ecological roles of ISVs in mosquito populations.

4.2. *Phasmaviridae*

Phasmaviruses remain relatively enigmatic, with our understanding largely shaped by HTS endeavors. In our investigation, we introduce novel viral entities identified within *Sa. albiprivus*, tentatively classified as *Sabethes albiprivus* phasmavirus. These findings contribute to the expanding spectrum of phasmaviral diversity. Across arthropod hosts spanning Diptera, Hymenoptera, and Coleoptera orders, phasmaviruses have been sporadically detected [60]. Presently, the ICTV acknowledges seven genera within the *Phasmaviridae* family [61]. Nevertheless, our grasp of the physicochemical attributes of these entities remains rudimentary, with detailed characterizations limited to merely two of these genera [61]. The dearth of comprehensive *in vitro* analyses underscores the pressing need for elucidating the fundamental properties of phasmaviruses, including their constituent subunits. Additionally, exploring the socio-virological dynamics associated with these viruses represents a fertile ground for future inquiry, promising perspectives into their ecological roles and potential impacts on host populations. Thus, bridging these knowledge gaps stands as imperative for unraveling the enigmatic realm of phasmaviral biology.

4.3. *Lispiviridae*

Members of the *Lispiviridae* family have been detected in various hosts across different regions worldwide, including arthropods such as hemipterans, odonatanans, hymenopterans, and orthopterans [60,62–66]. Infection by viruses representing the *Lispiviridae* family appears rare in mosquitoes. In this study, we identified a lispivirus (Pedras lispivirus isolate MG) in the host species *Sa. albiprivus*, previously identified in *Sa. quasicyaneus* [39], suggesting a close relationship between virus species and their hosts within the genus *Sabethes*, subgenus *Sabethes* (*Sabethes*). Members of the genus *Sabethes* are significant vectors of the wild yellow fever virus in the Americas [67]. Notably, *Sa. albiprivus* has already been found naturally infected by the amaryllic virus [5,68]. Future studies are warranted to investigate the occurrence of this virus in other species within this mosquito genus to enhance our understanding of its epidemiology and potential impact on public health.

4.4. *Iflaviridae*

Iflaviruses have been described in insects across several orders, including Lepidoptera, Hymenoptera, and Hemiptera, as well as in bee parasitic mites [69]. In bees, iflaviruses are major pathogens and are primarily associated with wing deformation [70,71]. The iflavivirus found in *Sa. albiprivus* (named *Sabethes albiprivus* iflavivirus) in this study shares similarities with *Aedes* Iflavi-like virus 1, isolated from *Ae. aegypti* in the Brazilian Amazon [41]. Despite the phylogenetic distinction and geographical / behavioral separation between the host mosquitoes *Ae. aegypti* and *Sa. albiprivus*, they harbor similar Iflaviruses. This discovery suggests a potential broader host range and adaptability of iflaviruses across different mosquito species and regions. Further investigations are necessary to comprehend the ecological and epidemiological implications of these viruses, particularly regarding their impact on mosquito populations and their potential role in disease transmission.

4.5. *Virgaviridae*

The *Virgaviridae* family has emerged prominently across diverse mosquito virome studies, underscoring its prevalence and potential significance in mosquito ecology [72,73]. In our investigation, we introduce two novel viral entities, tentatively designated as Buriti virga-like virus isolate MG and *Sabethes albiprivus* virgavirus 1, expanding the repertoire of known *Virgaviridae* members. Traditionally associated with plant hosts, the detection of virgavirus in mosquito viromes

raises intriguing questions regarding their ecological roles and transmission dynamics within insect populations. While it is conceivable that certain virgavirus species have adapted to exploit mosquito hosts for replication, an alternative explanation implicates dietary factors in their presence within mosquito viromes. The multifaceted nature of mosquito viromes encompasses a broad spectrum of viruses, including ISVs, arboviruses, and even plant viruses [74]. The phylogenetic positioning of Buriti virga-like virus isolate MG alongside its counterpart in *Sa. chloropterus* suggests a shared evolutionary ancestry, indicative of potential host-specific adaptations. Conversely, the distinct monophyletic branch formed by Sabethes albiprivus virgavirus 1 underscores its unique evolutionary trajectory within the *Virgaviridae* family. Further exploration into the ecological interactions between virgaviruses and their mosquito hosts promises valuable perspectives into their adaptive strategies and ecological impacts within vector populations.

4.6. *Flaviviridae*

The *Flaviviridae* family harbors a notable array of ISVs with intriguing implications for arboviral dynamics. Key among these are ISVs like Nhumirim virus [75], Cell fusion agent virus [76], and *Culex flavivirus* [77], which have been documented to exhibit inhibitory effects on their arboviral counterparts, albeit with variable outcomes across studies. While the majority of these findings stem from cell-line investigations, insights gleaned from mosquito-based studies are also noteworthy. In our study, we introduce *Psorophora ferox flavivirus* as a novel member of the *Flaviviridae* family. This discovery adds to the growing catalog of ISVs within the flavivirus lineage and underscores the diverse virome composition within mosquito populations. The inhibitory mechanisms attributed to ISV flaviviruses are postulated to stem from their close genetic relatedness to arboviruses within the family (reviewed in Carvalho and Long, 2021) [78]. However, the nuanced nature of these interactions is underscored by the variability in outcomes observed across different experimental settings. Despite the discordance among studies, the intricate interplay between flaviviruses holds promise for arboviral control strategies and enhances our comprehension of vector competence across mosquito species. Continued exploration of these viral dynamics, both in laboratory settings and within natural vector populations, is crucial for elucidating the full spectrum of ISV effects on arboviral transmission and for informing targeted interventions aimed at mitigating vector-borne disease burdens.

4.7. *Mesoniviridae*

Alphamesonivirus stands out as a ubiquitous presence across diverse mosquito species (over 34) and geographic regions, encompassing *Aedes spp.*, *Culex spp.*, *Anopheles spp.*, *Armigeres subalbatus*, and *Cq. xanthogaster* [3]. Our study adds a significant dimension to this narrative by documenting the presence of Alphamesonivirus cavallyense isolate MG for the first time in *Cq. venezuelensis*, a notable inclusion given the species' unique ecological and geographical context. The robust adaptability of Alphamesonivirus across different host species and environments underscores its remarkable success in host colonization, warranting comprehensive investigation into the factors driving its epidemiological dynamics. Notably, the Yichang virus, a close relative within the *Mesoniviridae* family, has demonstrated both horizontal and vertical transmission among *Ae. albopictus* and *Cx. quinquefasciatus*, alongside the intriguing ability to hinder flavivirus replication in mosquito cell lines [79]. However, elucidating similar relationships between other mesonivirids and arboviruses remains an open avenue for research. The discovery of Alphamesonivirus cavallyense isolate MG in *Cq. venezuelensis*, a species with distinct phylogenetic, behavioral, and ecological attributes compared to its typical hosts like *Cx. pipiens*, offers a unique lens into the dynamics of mesonivirus-host interactions. By forming a distinct monophyletic branch within the phylogenetic tree, this novel isolate underscores the genetic diversity and evolutionary adaptability of mesoniviruses across divergent mosquito taxa. Further exploration into the ecological drivers shaping the distribution and dynamics of mesoniviruses, alongside their potential implications for arboviral transmission dynamics, remains imperative. Unraveling the intricate interplay between mesonivirids and arboviruses holds promise for advancing our understanding of vector-borne disease ecology and informing targeted interventions aimed at mitigating mosquito-borne disease burdens.

4.8. Negevirus

Several viruses in this genus have been first isolated from mosquito cell lines exposed to filtered mosquito and phlebotomine specimens, particularly using C6/36 cell lines [80]. However, recent reports of new negeviruses have expanded to agricultural pests, including Hemiptera [81,82]. The detection of mosquito-associated negeviruses, using either cell cultures or HTS, has occurred across the globe [3,80,83], indicating wide host ranges and genetic variability. Their closest relatives include plant arboviruses, such as kitavirids [82,83]. Furthermore, diverse viruses in this genus also negatively modulate the multiplication of flaviviruses and alphaviruses in co-cultures in mosquito cell lines, possibly through superinfection exclusion [84,85]. This modulator capacity may be explored in the future for arbovirus control, which could help explain the lack of vectorial capacity of diverse mosquito genera in transmitting arboviruses. The negeviruses from the current study (named Biggie virus isolate MG and *Coquillettidia vlezuelensis* negevirus) represent a promising avenue for further exploration, particularly in isolating viruses from sylvatic mosquitoes using mosquito cell lines for further characterization.

Our study shares several limitations common to RNA virome studies. First, host attribution of the identified viruses is because these viruses might replicate in other organisms, such as plants consumed by mosquitos. This limitation is exacerbated by using whole-body mosquitoes for pools, rather than dissected parts such as salivary glands. To better determine the hosts of the viruses identified in our study, complementary approaches are necessary, such as virus culturing in mosquito cell lines and small RNA sequencing. The latter can provide sequences derived from the RNAi machinery of the cell, indicating active replication. Second, although none of the virus genomes we identified possess genes similar to host genes, we cannot exclude the possibility that these viruses might be endogenous viral elements (EVEs) [9]. Third, associating different segments into a single genome for multipartite viral genomes is very challenging without isolated viruses. We were able to associate the phasmavirus segments into a single genome because they were the only segments found for these viral taxa in their pool. However, this is only sometimes the case for different samples. Lastly, we used sequence-similarity approaches to identify RdRps in contigs. While RdRps are hallmarks enzymes of RNA viruses, this strategy cannot identify highly divergent RdRp sequences, leaving them in the realm of viral dark matter. Small RNA sequencing may aid in properly characterizing viral dark matter by associating sequence cleavage patterns with otherwise unrelated contigs to form complete viral genomes.

5. Conclusions

This study expands our understanding of the RNA virome of sylvatic mosquitoes from the Brazilian Cerrado biome, uncovering eleven near-complete viral genomes in seven distinct viral families and one genus of an unassigned family, including *Ferox* mosquito mononega-like virus* and *Albipes* mosquito Gordis-like virus* (*Xinmoviridae*), *Sabethes albiprivus* phasmavirus* (*Phasmaviridae*), *Pedras* lispivirus isolate MG (*Lispiviridae*), *Sabethes albiprivus* iflavivirus* (*Iflaviridae*), *Buriti virga*-like virus isolate MG and *Sabethes albiprivus virga* virus 1* (*Virgaviridae*), *Psorophora ferox* flavivirus* (*Flaviviridae*), *Alphamesonivirus cavallyense* isolate MG (*Mesoniviridae*), and *Biggie* virus isolate MG and *Coquillettidia vlezuelensis* negevirus* (*Negevirus*), with seven being new and four previously known genomes identified in new host species. The discovery of two xinmovirids in *Ps. ferox* and *Ps. albipes* presents new avenues for future studies aimed at understanding the interactions between ISVs and other arboviruses in the context of pathogen transmission. Additionally, HTS once again demonstrates its remarkable capacity to discover new viruses, even in pools of mosquitoes from different species, thereby enhancing our understanding of comprehensive virome surveillance in regions undergoing rapid environmental changes, as such the Brazilian Cerrado biome. Moreover, the presence of ISVs and potential novel arboviruses underscores the need for ongoing surveillance and control strategies to mitigate the risk of emerging infectious diseases. Further research is needed to elucidate the ecological roles of these viruses, their potential impacts on vector competence, and their implications for human and animal health in the

context of emerging infectious diseases. Continuing to explore the viromic landscape of sylvatic mosquitoes can better prepare us for and mitigate the threats posed by future viral outbreaks.

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Data Availability Statement: The authors declare that all data supporting the findings of this study are available within the paper. The viruses' genome sequences have been deposited in GenBank under the accession numbers PP946236 to PP946246. HTS raw data has been deposited in the SRA platform under BioProject accession number PRJNA1126168, with BioSample accession numbers SAMN41926027 to SAMN41926031 and SRA (SRR) accession numbers SRR29474085 to SRR29474089.

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References

1. Harvey, E.; Holmes, E.C. Diversity and evolution of the animal virome. *Nat Rev Microbiol.* **2022**, *20*(6), 321-334. doi: 10.1038/s41579-021-00665-x.
2. Atoni, E.; Zhao, L.; Karungu, S.; Obanda, V.; Agwanda, B.; Xia, H.; Yuan, Z. The discovery and global distribution of novel mosquito-associated viruses in the last decade (2007-2017). *Rev Med Virol.* **2019**, *29*(6), e2079. doi: 10.1002/rmv.2079.
3. Moonen, J.P.; Schinkel, M.; van der Most, T.; Miesen, P.; van Rij, R.P. Composition and global distribution of the mosquito virome - A comprehensive database of insect-specific viruses. *One Health* **2023**, *16*:100490. doi: 10.1016/j.onehlt.2023.100490.
4. Harbach, R.E. Mosquito Taxonomic Inventory. Available online: <https://mosquito-taxonomic-inventory.myspecies.info/> (accessed on 18 January 2024). 2013.
5. de Oliveira, C.H.; Andrade, M.S.; Campos, F.S.; da C. Cardoso, J.; Gonçalves-Dos-Santos, M.E.; Oliveira, R.S.; Aquino-Teixeira, S.M.; Campos, A.A.; Almeida, M.A.; Simonini-Teixeira, D.; da P. Sevá, A.; Temponi, A.O.D.; Magalhães, F.M.; da Silva Menezes, A.S.; Lo-pes, B.T.; Almeida, H.P.; Pedroso, A.L.; Gonçalves, G.P.; Chaves, D.C.C.; de Menezes, G.G.; Bernal-Valle, S.; Müller, N.F.; Janssen, L.; Dos Santos, E.; Mares-Guia, M.A.; Albuquerque, G.R.; Romano, A.P.; Franco, A.C.; Ribeiro, B.M.; Roehe, P.M.; Lourenço-de-Oliveira, R.; de Abreu, F.V.S. Yellow Fever Virus Maintained by Sabethes Mosquitoes during the Dry Season in Cerrado, a Semiarid Region of Brazil, in 2021. *Viruses* **2023**, *15*(3), 757. doi: 10.3390/v15030757.
6. Sherwood, J.A.; Stehman, S.V.; Howard, J.J.; Oliver, J. Cases of Eastern equine encephalitis in humans associated with *Aedes canadensis*, *Coquillettidia perturbans* and *Culiseta melanura* mosquitoes with the virus in New York State from 1971 to 2012 by analysis of aggregated published data. *Epidemiol Infect.* **2020**, *148*, e72. doi: 10.1017/S0950268820000308.
7. Anderson, C.R.; Spence, L.; Downs, W.G.; Aitken, T.H. Oropouche virus: a new human disease agent from Trinidad, West Indies. *Am J Trop Med Hyg.* **1961**, *10*, 574-8. doi: 10.4269/ajtmh.1961.
8. Romero-Alvarez, D.; Escobar, L.E. Oropouche fever, an emergent disease from the Americas. *Microbes Infect.* **2018**, *20*(3), 135-146. doi: 10.1016/j.micinf.2017.11.013.
9. de Almeida, J.P.; Aguiar, E.R.; Armache, J.N.; Olmo, R.P.; Marques, J.T. The virome of vector mosquitoes. *Curr Opin Virol.* **2021**, *49*, 7-12.

10. Olmo, R.P.; Todjro, Y.M.H.; Aguiar, E.R.G.R.; de Almeida, J.P.P.; Ferreira, F.V.; Armache, J.N., et al. Mosquito vector competence for dengue is modulated by insect-specific viruses. *Nat Microbiol.* **2023**, *8*(1), 135–49.
11. Maia, L.J.; Oliveira, C.H.; Silva, A.B.; Souza, P.A.A.; Müller, N.F.D.; Cardoso, J.D.C.; Ribeiro, B.M.; Abreu, F.V.S.; Campos, F.S. Arbovirus surveillance in mosquitoes: Historical methods, emerging technologies, and challenges ahead. *Exp Biol Med.* **2023**, *248*(22), 2072–2082. doi: 10.1177/15353702231209415.
12. Pérez-Losada, M.; Arenas, M.; Galán, J.C.; Bracho, M.A.; Hillung, J.; García-González, N.; González-Candelas, F. High-throughput sequencing (HTS) for the analysis of viral populations. *Infect Genet Evol.* **2020**, *80*, 104208. doi: 10.1016/j.meegid.2020.104208.
13. Silva, A.F.; Machado, L.C.; Silva, L.M.I.; Dezordi, F.Z.; Wallau, G.L. RNA virome of sylvatic mosquitoes from northeast Brazil reveals a divergent and diverse insect-specific viral community. *bioRxiv* **2023**, 2023.06.27.546706.
14. Moutailler, S.; Yousfi, L.; Mousson, L.; Devillers, E.; Vazeille, M.; Vega-Rúa, A.; Perrin, Y.; Jourdain, F.; Chandre, F.; Cannet, A.; Chantilly, S.; Restrepo, J.; Guidez, A.; Dusfour, I.; Vieira Santos de Abreu, F.; Pereira Dos Santos, T.; Jiolle, D.; Visser, T.M.; Koenraadt, C.J.M.; Wongsokarijo, M.; Diallo, M.; Diallo, D.; Gaye, A.; Boyer, S.; Duong, V.; Piorkowski, G.; Paupy, C.; Lourenco de Oliveira, R.; de Lamballerie, X.; Failloux, A.B. A New High-Throughput Tool to Screen Mosquito-Borne Viruses in Zika Virus Endemic/Epidemic Areas. *Viruses* **2019**, *11*(10), 904. doi: 10.3390/v11100904.
15. Gómez, M.; Martínez, D.; Páez-Triana, L.; Luna, N.; Ramírez, A.; Medina, J.; Cruz-Saavedra, L.; Hernández, C.; Castañeda, S.; Bohórquez Melo, R.; Suarez, L.A.; Palma-Cuero, M.; Murcia, L.M.; González Páez, L.; Estrada Bustos, L.; Medina, M.A.; Ariza Campo, K.; Padilla, H.D.; Zamora Flórez, A.; De Las Salas, J.L.; Muñoz, M.; Ramírez, J.D. Influence of dengue virus serotypes on the abundance of *Aedes aegypti* insect-specific viruses (ISVs). *J Virol.* **2024**, *98*(1), e0150723. doi: 10.1128/jvi.01507-23.
16. Myers, N.; Mittermeier, R.A.; Mittermeier, C.G.; da Fonseca, G.A.; Kent, J. Biodiversity hotspots for conservation priorities. *Nature* **2000**, *403*(6772), 853–8. doi: 10.1038/35002501.
17. Da Silva, J.M.C.; Bates, J.M. Biogeographic Patterns and Conservation in the South American Cerrado: A Tropical Savanna Hotspot: The Cerrado, which includes both forest and savanna habitats, is the second largest South American biome, and among the most threatened on the continent. *BioScience* **2002**, *52*, 225–234. doi: 10.1641/0006-3568(2002)052[0225:BPACIT]2.0.CO;2.
18. Plowright, R.K.; Reaser, J.K.; Locke, H.; Woodley, S.J.; Patz, J.A.; Becker, D.J.; Oppler, G.; Hudson, P.J.; Tabor, G.M. Land use-induced spillover: a call to action to safeguard environmental, animal, and human health. *Lancet Planet Health* **2021**, *5*(4), e237–e245. doi: 10.1016/S2542-5196(21)00031-0.
19. Brazil. 2019. Guia para o Planejamento das Ações de Captura de Anofelinos pela Técnica de Atração por Humano Protegido (TAHP) e Acompanhamento dos Riscos à Saúde do Profissional Capturador. Ministério da Saúde, Secretaria de Vigilância em Saúde, Departamento de Imunização e Doenças Transmissíveis. Brasília Ministério da Saúde, Brasil, 27 p. Available online: <https://www.gov.br/saude/pt-br/centrais-de-conteudo/publicacoes/svsa/malaria/vigilancia-entomologica-e-controle-vetorial-da-malaria/guia-para-o-planejamento-das-acoes-de-captura-de-anofelinos-pela-tecnica-de-atracao-por-humano-protegido-tahp-e-acompanhamento-dos-riscos-a-saude-do-profissional-capturador.pdf>. (accessed on 17 may 2024).
20. Shannon, R.C. Methods for collecting and feeding mosquitoes in jungle yellow fever studies. *Am. J. Trop. Med. Hyg.* **1939**, *19*, 131–140.
21. Abreu, F.V.S.; Ribeiro, I.P.; Ferreira-de-Brito, A.; Santos, A.A.C.D.; Miranda, R.M.; Bonelly, I.S.; Neves, M.S.A.S.; Bersot, M.I.; Santos, T.P.D.; Gomes, M.Q.; et al. *Haemagogus leucocelaenus* and *Haemagogus janthinomys* are the primary vectors in the major yellow fever outbreak in Brazil, 2016–2018. *Emerg. Microbes Infect.* **2019**, *8*, 218–223.
22. Consoli, R.A.G.B.; Oliveira, R.L. Principais mosquitos de importância sanitária no Brasil [online]. Rio de Janeiro: Editora Fiocruz, 1994. 228 p. ISBN 85-85676-03-5. Available online: <https://static.scielo.org/scielobooks/th/pdf/consoli-9788575412909.pdf> (accessed on 17 may 2024).
23. Forattini, O.P. Culicidologia Médica: Identificação, Biologia, Epidemiologia. Editora da Universidade de São Paulo: São Paulo, Brasil, 2002; Volume 2, p. 1924.
24. Andrews, S. 2010. FastQC: A Quality Control Tool for High Throughput Sequence Data [Online]. Available online: <https://www.bioinformatics.babraham.ac.uk/projects/fastqc/>. (accessed on 17 may 2024).
25. Krueger, F.; James, F.; Ewels, P.; Afyounian, E.; Boeckler, B.S. FelixKrueger/TrimGalore: v0.6.7 - 2021. Available online: <https://doi.org/10.5281/zenodo.5127899>. (accessed on 17 may 2024).
26. Bolger, A.M.; Lohse, M.; Usadel, B. Trimmomatic: a flexible trimmer for Illumina sequence data. *Bioinformatics* **2014**, *30*(15), 2114–20. doi: 10.1093/bioinformatics/btu170.
27. Meleshko, D.; Hajirasouliha, I.; Korobeynikov, A. coronaSPAdes: from biosynthetic gene clusters to RNA viral assemblies. *Bioinformatics* **2021**, *38*(1), 1–8. doi: 10.1093/bioinformatics/btab597.
28. Buchfink, B.; Reuter, K.; Drost, H.G. Sensitive protein alignments at tree-of-life scale using DIAMOND. *Nature Methods* **2021**, *18*, 366–368. doi:10.1038/s41592-021-01101-x.

29. Charon, J.; Buchmann, J.P.; Sadiq, S.; Holmes, E.C. RdRp-scan: A bioinformatic resource to identify and annotate divergent RNA viruses in metagenomic sequence data. *Virus Evol.* **2022**, *8*(2), veac082. doi: 10.1093/ve/veac082.
30. National Center for Biotechnology Information (NCBI). Bethesda (MD): National Library of Medicine (US); 1988. Available online: <https://www.ncbi.nlm.nih.gov/>. (accessed on 17 may 2024).
31. Nayfach, S.; Camargo, A.P.; Schulz, F.; Eloie-Fadrosch, E.; Roux, S.; Kyrpides, N.C. CheckV assesses the quality and completeness of metagenome-assembled viral genomes. *Nat Biotechnol.* **2021**, *39*(5), 578-585. doi: 10.1038/s41587-020-00774-7.
32. Katoh, K.; Standley, D.M. MAFFT multiple sequence alignment software version 7: improvements in performance and usability. *Mol Biol Evol.* **2013**, *30*(4), 772-80. doi: 10.1093/molbev/mst010.
33. Capella-Gutiérrez, S.; Silla-Martínez, J.M.; Gabaldón, T. trimAl: a tool for automated alignment trimming in large-scale phylogenetic analyses. *Bioinformatics* **2009**, *25*(15), 1972-3. doi: 10.1093/bioinformatics/btp348.
34. Darriba, D.; Posada, D.; Kozlov, A.M.; Stamatakis, A.; Morel, B.; Flouri, T. ModelTest-NG: A New and Scalable Tool for the Selection of DNA and Protein Evolutionary Models. *Mol Biol Evol.* **2020** *37*(1), 291-294. doi: 10.1093/molbev/msz189.
35. Kozlov, A.M.; Darriba, D.; Flouri, T.; Morel, B.; Stamatakis, A. RAxML-NG: a fast, scalable and user-friendly tool for maximum likelihood phylogenetic inference. *Bioinformatics* **2019**, *35*(21), 4453-4455. doi: 10.1093/bioinformatics/btz305.
36. Sharpe, S.; Paraskevopoulou, S. ICTV Virus Taxonomy Profile: Ximoviridae 2023. *J Gen Virol.* **2023**, *104*(10). doi: 10.1099/jgv.0.001906.
37. Shi, C.; Beller, L.; Deboutte, W.; Yinda, K.C.; Delang, L.; Vega-Rúa, A.; Failloux, A.B.; Matthijnsens, J. Stable distinct core eukaryotic viromes in different mosquito species from Guadeloupe, using single mosquito viral metagenomics. *Microbiome* **2019**, *7*(1), 121. doi: 10.1186/s40168-019-0734-2.
38. Batson, J.; Dudas, G.; Haas-Stapleton, E.; Kistler, A.L.; Li, L.M.; Logan, P.; Ratnasiri, K.; Retallack, H. Single mosquito metatranscriptomics identifies vectors, emerging pathogens and reservoirs in one assay. *Elife* **2021**, *10*, e68353. doi: 10.7554/eLife.68353.
39. Aragão, C.F.; da Silva, S.P.; do Nascimento, B.L.S.; da Silva, F.S.; Nunes Neto, J.P.; Pinheiro, V.C.S.; Cruz, A.C.R. Shotgun Metagenomic Sequencing Reveals Virome Composition of Mosquitoes from a Transition Ecosystem of North-Northeast Brazil. *Genes (Basel)* **2023**, *14*(7), 1443. doi: 10.3390/genes14071443.
40. Li, J.M.; Wang, F.; Ye, G.; Paraskevopoulou, S. ICTV Virus Taxonomy Profile: Lispiviridae 2023. *J Gen Virol.* **2023**, *104*(7). doi: 10.1099/jgv.0.001869.
41. Ribeiro, G.O.; Morais, V.S.; Monteiro, F.J.C.; Ribeiro, E.S.D.; Rego, M.O.D.S.; Souto, R.N.P.; Villanova, F.; Tahmasebi, R.; Hefford, P.M.; Deng, X.; Delwart, E.; Cerdeira Sabino, E.; Fernandes, L.N.; da Costa, A.C.; Leal, É. *Aedes aegypti* from Amazon Basin Harbor High Diversity of Novel Viral Species. *Viruses* **2020**, *12*(8), 866. doi: 10.3390/v12080866.
42. Valles, S.M.; Chen, Y.; Firth, A.E.; Guérin, D.M.A.; Hashimoto, Y.; Herrero, S.; de Miranda, J.R.; Ryabov, E.; Ictv Report Consortium. ICTV Virus Taxonomy Profile: Iflaviridae. *J Gen Virol.* **2017**, *98*(4), 527-528. doi: 10.1099/jgv.0.000757.
43. Orba, Y.; Matsuno, K.; Nakao, R.; Kryukov, K.; Saito, Y.; Kawamori, F.; Loza Vega, A.; Watanabe, T.; Maemura, T.; Sasaki, M.; Hall, W.W.; Hall, R.A.; Pereira, J.A.; Nakagawa, S.; Sawa, H. Diverse mosquito-specific flaviviruses in the Bolivian Amazon basin. *J Gen Virol.* **2021**, *102*(3). doi: 10.1099/jgv.0.001518.
44. Hang, J.; Klein, T.A.; Kim, H.C.; Yang, Y.; Jima, D.D.; Richardson, J.H.; Jarman, R.G. Genome Sequences of Five Arboviruses in Field-Captured Mosquitoes in a Unique Rural Environment of South Korea. *Genome Announc.* **2016**, *4*(1), e01644-15. doi: 10.1128/genomeA.01644-15.
45. Alencar, J.; Pacheco, J.B.; Correa, F.F.; Silva, J. dos S.; Guimarães, A.É. New report on the bionomics of *Coquillettidia venezuelensis* in temporary breeding sites (Diptera: Culicidae). *Rev Soc Bras Med Trop.* **2011**, *44*(2), 247-8. doi: 10.1590/s0037-86822011000200023.
46. Farajollahi, A.; Fonseca, D.M.; Kramer, L.D.; Marm Kilpatrick, A. "Bird biting" mosquitoes and human disease: a review of the role of *Culex pipiens* complex mosquitoes in epidemiology. *Infect Genet Evol.* **2011**, *11*(7), 1577-85. doi: 10.1016/j.meegid.2011.08.013.
47. Sadeghi, M.; Altan, E.; Deng, X.; Barker, C.M.; Fang, Y.; Coffey, L.L.; Delwart, E. Virome of > 12 thousand *Culex* mosquitoes from throughout California. *Journal of Virology* **2018**, *523*, 74-88. doi: 10.1016/j.virol.2018.07.029.
48. Hermanns, K.; Marklewitz, M.; Zirkel, F.; Overheul, G.J.; Page, R.A.; Loaiza, J.R.; Drost, C.; van Rij, R.P.; Junglen, S. Agua Salud alphavirus defines a novel lineage of insect-specific alphaviruses discovered in the New World. *J Gen Virol.* **2020**, *101*(1), 96-104. doi: 10.1099/jgv.0.001344.
49. Patterson, E.I.; Villinger, J.; Muthoni, J.N.; Dobel-Ober, L.; Hughes, G.L. Exploiting insect-specific viruses as a novel strategy to control vector-borne disease. *Curr Opin Insect Sci.* **2020**, *39*, 50-56. doi: 10.1016/j.cois.2020.02.005.

50. Parry, R.; Asgari, S. Aedes Anphevirus: an Insect-Specific Virus Distributed Worldwide in Aedes aegypti Mosquitoes That Has Complex Interplays with Wolbachia and Dengue Virus Infection in Cells. *Journal of Virology* **2018**, *92*(17), e00224-18. doi: 10.1128/JVI.00224-18.
51. Manni, M.; Zdobnov, E.M. A Novel Anphevirus in Aedes albopictus Mosquitoes Is Distributed Worldwide and Interacts with the Host RNA Interference Pathway. *Viruses* **2020**, *12*(11), 1264. <https://doi.org/10.3390/v12111264>.
52. Shi, M.; Neville, P.; Nicholson, J.; Eden, J.S.; Imrie, A.; Holmes, E.C. High-Resolution Metatranscriptomics Reveals the Ecological Dynamics of Mosquito-Associated RNA Viruses in Western Australia. *Journal of Virology* **2017**, *91*(17), e00680-17. doi: 10.1128/JVI.00680-17.
53. Scarpassa, V.M.; Debat, H.J.; Alencar, R.B.; Saraiva, J.F.; Calvo, E.; Arcà, B.; Ribeiro, J.M.C. An insight into the sialotranscriptome and virome of Amazonian anophelines. *BMC Genomics* **2019**, *8*, 20(1), 166. doi: 10.1186/s12864-019-5545-0.
54. Moreno, E.S.; Rocco, I.M.; Bergo, E.S.; Brasil, R.A.; Siciliano, M.M.; Suzuki, A.; Silveira, V.R.; Bisordi, I.; Souza, R.P. Yellow Fever Working Group. Reemergence of yellow fever: detection of transmission in the State of São Paulo, Brazil, 2008. *Revista da Sociedade Brasileira de Medicina Tropical* **2008**, *44*(3), 290–296. doi: 10.1590/s0037-86822011005000041.
55. Lopes, O.S.; Sacchetta, L.A.; Francy, D.B.; Jakob, W.L.; Calisher, C.H. Emergence of a new arbovirus disease in Brazil III. Isolation of Rocio virus from Psorophora ferox (Humboldt, 1819). *American Journal of Epidemiology* **1981**, *113*(2), 122-125.
56. Kulasekera, V.L.; Kramer, L.; Nasci, R.S.; Mostashari, F.; Cherry, B.; Trock, S.C.; Glaser, C.; Miller, J.R. West Nile virus infection in mosquitoes, birds, horses, and humans, Staten Island, New York, 2000. *Emerging infectious diseases* **2017**, *4*, 722–725. doi: 10.3201/eid0704.010421.
57. Cupp, E.W.; Zhang, D.; Yue, X.; Cupp, M.S.; Guyer, C.; Sprenger, T.R.; Unnasch, T.R. Identification of reptilian and amphibian blood meals from mosquitoes in an eastern equine encephalomyelitis virus focus in central Alabama. *The American Journal of Tropical Medicine and Hygiene*, **2004**, *71*(3), 272–276.
58. Turell, M.; O'Guinn, M.; Jones, J.; Sardelis, M.; Dohm, D.; Watts, D.; Fernandez, R.; Travassos Da Rosa, A.; Guzman, H.; Tesh, R.; Rossi, C.; Ludwig, G.; Mangiafico, J.; Kondig, J.; Wasieloski, L.; Pecor, J.; Zyzak, M.; Schoeler, G. Isolation of viruses from mosquitoes (Diptera: Culicidae) collected in the Amazon Basin region of Peru. *Journal of Medical Entomology* **2005**, *42*(5), 891-8. doi: 10.1093/jmedent/42.5.891.
59. Vasconcelos, P.F. Febre amarela [Yellow Fever]. *Revista da Sociedade Brasileira de Medicina Tropical* **2003**, *36*(2), 275–293. doi: 10.1590/s0037-86822003000200012.
60. Käfer, S.; Paraskevopoulou, S.; Zirkel, F.; Wieseke, N.; Donath, A.; Petersen, M.; Jones, T.C.; Liu, S.; Zhou, X.; Middendorf, M.; Junglen, S.; Misof, B.; Drosten, C. Re-assessing the diversity of negative strand RNA viruses in insects. *PLoS pathogens* **2019**, *15*(12), e1008224. doi: 10.1371/journal.ppat.1008224.
61. ICTV, 2024. Phasmaviridae family. Available on: <https://ictv.global/report/chapter/phasmaviridae/phasmaviridae>. (accessed on 11 June 2024).
62. Viljakainen, L.; Holmberg, I.; Abril, S.; Jurvansuu, J. Viruses of invasive Argentine ants from the European Main supercolony: characterization, interactions and evolution. *The Journal of General Virology*, **2018**, *99*(8), 1129–1140. doi: 10.1099/jgv.0.001104.
63. Lay, C.L.; Shi, M.; Buček, A.; Bourguignon, T.; Lo, N.; Holmes, E.C. Unmapped RNA Virus Diversity in Termites and their Symbionts. *Viruses* **2020**, *12*(10), 1145. doi: 10.3390/v12101145.
64. Huang, H.J.; Ye, Z.X.; Wang, X.; Yan, X.T.; Zhang, Y.; He, Y.J.; Qi, Y.H.; Zhang, X.D.; Zhuo, J.C.; Lu, G.; Lu, J.B.; Mao, Q.Z.; Sun, Z.T.; Yan, F.; Chen, J.P.; Zhang, C.X.; Li, J.M. Diversity and infectivity of the RNA virome among different cryptic species of an agriculturally important insect vector: whitefly Bemisia tabaci. *NPJ Biofilms and Microbiomes* **2021**, *7*(1), 43. doi: 10.1038/s41522-021-00216-5.
65. Wang, F.; Yuan, B.; Xiao, S.; Zhang, J.; Jia, W.; Fang, Q.; Wang, F.; Song, Q.; Ye, G. Diverse RNA Viruses Discovered in Three Parasitoid Wasps of the Rice Weevil Sitophilus oryzae. *mSphere* **2021**, *6*(3), e00331-21. doi: 10.1128/mSphere.00331-21.
66. Ye, Z.X.; Wang, S.M.; Lu, G.; Chen, J.P.; Zhang, C.X.; Xu, L.Y.; Li, J.M. Complete genome sequence of a novel arlivirus from a yellow spotted stink bug (Erthesina fullo (Thunberg, 1783)). *Archives of Virology* **2022**, *167*(4), 1205–1209. doi: 10.1007/s00705-022-05399-6.
67. Hervé, J.P.; Dégallier, N.; Travassos-da-Rosa, A.P.A.; Pinheiro, F.P.; Sá Filho, G.C. Arboviroses: aspectos ecológicos. In: Instituto Evandro Chagas: 50 anos. Belém: *Instituto Evandro Chagas* **1986**, 408-437.
68. Goenaga, S.; Fabbri, C.; Dueñas, J.C.; Gardenal, C.N.; Rossi, G.C.; Calderon, G.; Morales, M.A.; Garcia, J.B.; Enria, D.A.; Levis, S. Isolation of yellow fever virus from mosquitoes in Misiones province, Argentina. *Vector Borne and Zoonotic Diseases* **2012**, *12*(11), 986–993. doi: 10.1089/vbz.2011.0730.
69. van Oers, M.M. Genomics and Biology of Iflaviruses. In K. Johnson, & S. Agari (Eds.), Caister Academic Press. *Insect Virology* **2010**, 231-250.
70. Bowen-Walker, P.L.; Martin, S.J.; Gunn, A. The transmission of deformed wing virus between honeybees (Apis mellifera L.) by the ectoparasitic mite Varroa jacobsoni Oud. *J. Invertebr. Pathol* **1999**, *73*, 101-106.

71. Posada-Florez, F.; Childers, A.K.; Heerman, M.C.; Egekwu, N.I.; Cook, S.C.; Chen, Y.; Evans, J.D.; Ryabov, E.V. Deformed wing virus type A, a major honey bee pathogen, is vectored by the mite *Varroa destructor* in a non-propagative manner. *Sci Rep* **2019**, *9*, 12445. doi: 10.1038/s41598-019-47447-3.
72. Konstantinidis, K.; Dovrolis, N.; Kouvela, A.; Kassela, K.; Rosa Freitas, M.G.; Nearchou, A.; de Courcy Williams, M.; Veletza, S.; Karakasiliotis, I. Defining virus-carrier networks that shape the composition of the mosquito core virome of a local ecosystem. *Virus Evol.* **2022**, *8*(1), veac036.
73. Liu, Q.; Cui, F.; Liu, X.; Fu, Y.; Fang, W.; Kang, X.; Lu, H.; Li, S.; Liu, B.; Guo, W.; Xia, Q.; Kang, L.; Jiang, F. Association of virome dynamics with mosquito species and environmental factors. *Microbiome* **2023**, *11*(1), 101.
74. Bonning, B.C. The Insect Virome: Opportunities and Challenges. *Curr Issues Mol Biol.* **2020**, *34*, 1-12. doi: 10.21775/cimb.034.001.
75. Pauvolid-Corrêa, A.; Solberg, O.; Couto-Lima, D.; Kenney, J.; Serra-Freire, N.; Brault, A.; Nogueira, R.; Langevin, S.; Komar, N. Nhumirim virus, a novel flavivirus isolated from mosquitoes from the Pantanal, Brazil. *Arch Virol.* **2015**, *160*(1), 21-7. doi: 10.1007/s00705-014-2219-8.
76. Stollar, V.; Thomas, V.L. An agent in the *Aedes aegypti* cell line (Peleg) which causes fusion of *Aedes albopictus* cells. *Virology* **1975**, *64*, 367-377. doi: 10.1016/0042-6822(75)90113-0.
77. Hoshino, K.; Isawa, H.; Tsuda, Y.; Yano, K.; Sasaki, T.; Yuda, M.; Takasaki, T.; Kobayashi, M.; Sawabe, K. Genetic characterization of a new insect flavivirus isolated from *Culex pipiens* mosquito in Japan. *Virology* **2007**, *359*(2), 405-14. doi: 10.1016/j.virol.2006.09.039.
78. Carvalho, V.L.; Long, M.T. Insect-Specific Viruses: An overview and their relationship to arboviruses of concern to humans and animals. *Virology* **2021**, *557*, 34-43. doi: 10.1016/j.virol.2021.01.007.
79. Ye, G.; Wang, Y.; Liu, X.; Dong, Q.; Cai, Q.; Yuan, Z.; Xia, H. Transmission competence of a new mesonivirus, Yichang virus, in mosquitoes and its interference with representative flaviviruses. *PLoS Negl Trop Dis.* **2020**, *14*(11), e0008920. doi: 10.1371/journal.pntd.0008920.
80. Vasilakis, N.; Forrester, N.L.; Palacios, G.; Nasar, F.; Savji, N.; Rossi, S.L.; Guzman, H.; Wood, T.G.; Popov, V.; Gorchakov, R.; González, A.V.; Haddow, A.D.; Watts, D.M.; da Rosa, A.P.; Weaver, S.C.; Lipkin, W.I.; Tesh, R.B. Negevirus: a proposed new taxon of insect-specific viruses with wide geographic distribution. *J Virol.* **2013**, *87*(5), 2475-88. doi: 10.1128/JVI.00776-12.
81. Qi, Y.H.; Xu, L.Y.; Zhai, J.; Ye, Z.X.; Lu, G.; Chen, J.P.; Zhang, C.X.; Li, J.M. Complete genome sequence of a novel nege-like virus in aphids (genus *Indomegoura*). *Virol J.* **2021**, *18*(1), 76. doi: 10.1186/s12985-021-01552-w.
82. Quito-Avila, D.F.; Reyes-Proano, E.; Armijos-Capa, G.; Alcalá Briseño, R.I.; Alvarez, R.; Flores, F.F. Analysis of a new negevirus-like sequence from *Bemisia tabaci* unveils a potential new taxon linking nelorpi- and centiviruses. *PLoS One* **2024**, *19*(5), e0303838. doi: 10.1371/journal.pone.0303838.
83. Nunes, M.R.T.; Contreras-Gutierrez, M.A.; Guzman, H.; Martins, L.C.; Barbirato, M.F.; Savit, C.; Balta, V.; Uribe, S.; Vivero, R.; Suaza, J.D.; Oliveira, H.; Nunes Neto, J.P.; Carvalho, V.L.; da Silva, S.P.; Cardoso, J.F.; de Oliveira, R.S.; da Silva Lemos, P.; Wood, T.G.; Widen, S.G.; Vasconcelos, P.F.C.; Fish, D.; Vasilakis, N.; Tesh, R.B. Genetic characterization, molecular epidemiology, and phylogenetic relationships of insect-specific viruses in the taxon Negevirus. *Virology* **2017**, *504*, 152-167. doi: 10.1016/j.virol.2017.01.022.
84. Patterson, E.I.; Kautz, T.F.; Contreras-Gutierrez, M.A.; Guzman, H.; Tesh, R.B.; Hughes, G.L.; Forrester, N.L. Negevirus Reduce Replication of Alphaviruses during Coinfection. *J Virol.* **2021**, *95*(14), e0043321. doi: 10.1128/JVI.00433-21.
85. Carvalho, V.L.; Prakoso, D.; Schwarz, E.R.; Logan, T.D.; Nunes, B.T.D.; Beachboard, S.E.; Long, M.T. Negevirus Piura Suppresses Zika Virus Replication in Mosquito Cells. *Viruses* **2024**, *16*, 350. doi: 10.3390/v16030350.

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