Novel Viruses Isolated from Mosquitoes in Pantanal, Brazil

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Genomic sequences are described from five novel viruses and divergent strains of Brejeira and Guáico Culex viruses from mosquitoes collected in Pantanal, Brazil, in 2010.

We describe here the genomes of five novel viruses and divergent strains of Brejeira (BRJV) and Guáico Culex (GCXV) viruses isolated from pooled mosquitoes captured with CDC light traps from Pantanal, Mato Grosso do Sul state, in west-central Brazil, in April 2010. Genomic sequences were derived from mosquito cell cultures by Illumina technology (1–3) (Table 1).

Four BRJV isolates from *Culex* (*Culex*) spp. yielded 99% identity among themselves, but 88% to 90% identity with a BRJV isolated in northern Brazil, indicating the detection of a divergent strain of this recently described *Negevirus*-like virus (4).

We report here the first Brazilian detections of GCXV, a recently described multicomponent Jingmenivirus (5). The five genomic segments from *Culex* (*Culex*) spp. pool number 407 aligned with coverages from 78% to 87% and identities from 97% to 100% with published sequences. A second isolate from pool number 313, also *Culex* (*Culex*) spp., was the same strain, based on partial sequences (data not shown). In Brazil, the first detection of a Jingmenivirus was from tick collections in southeast Brazil, and called Mogiana tick virus (6).

Five novel viruses were detected and named to honor indigenous tribes of Mato Grosso do Sul state (Terena, Ofaié, Kadiwéu, Kaiowá, and Guató viruses). Translated nucleotide identities ranged between 32% and 73% to known viruses. The following abbreviations are suggested for the novel viruses: TERV, OFAV, KADV, KAIV, and GUTV.

TERV isolated from *Mansonia* spp. pool aligns with viruses of a newly described genus within *Bunyaviridae* called *Phasmavirus* (7). It contains the standard three-segmented genome with the small (S), medium (M), and large (L) segments of other bunyaviruses.

OFAV derived from the same mosquito pool as TERV. OFAV has a large genome of 19.8 kb and most closely aligns (<66% predicted amino acid sequence identity) with Méno virus, recently described within the family *Mesoniviridae* (8). KADV is also related to Méno virus (<71% identical) and is similarly large

### TABLE 1 Genomic sequences identified from 12 isolates of 7 viruses isolated from mosquitoes in Pantanal, Brazil, in April 2010

| GenBank accession no. | Virus name | Segment | Virus family | Pool no. | Mosquito species (no. of specimens) |
|-----------------------|------------|---------|--------------|----------|-------------------------------------|
| KT966481              | Kaiówa virus |              | Unclassified | 313      | *Culex* (*Culex*) spp. (15)         |
| KT966482              | Brejeira virus |              | Unclassified | 313      | *Culex* (*Culex*) spp. (15)         |
| KT966483              | Brejeira virus |              | Unclassified | 324A     | *Culex* (*Culex*) spp. (27)         |
| KT966484              | Guató virus |              | Unclassified | 324A     | *Culex* (*Culex*) spp. (27)         |
| KT966485              | Brejeira virus |              | Unclassified | 324B     | *Culex* (*Culex*) spp. (23)         |
| KT966486              | Guató virus |              | Unclassified | 324B     | *Culex* (*Culex*) spp. (23)         |
| KT966487              | Brejeira virus |              | Unclassified | 332A     | *Culex* (*Culex*) spp. (26)         |
| KT966490              | Guató virus |              | Unclassified | 332A     | *Culex* (*Culex*) spp. (26)         |
| KT966491              | Ofaié virus |              | *Mesoniviridae*-like | 360      | *Mansonia* spp. (25)                |
| KT966492              | Terena virus | S          | *Bunyaviridae*-like | 360      | *Mansonia* spp. (25)                |
| KT966493              | Terena virus | M          | *Bunyaviridae*-like | 360      | *Mansonia* spp. (25)                |
| KT966494              | Terena virus | L          | *Bunyaviridae*-like | 360      | *Mansonia* spp. (25)                |
| KT966495              | Kadiwéu virus |              | *Mesoniviridae*-like | 407      | *Culex* (*Culex*) spp. (50)         |
| KT966498              | Guáico Culex virus | 1       | Unclassified | 407      | *Culex* (*Culex*) spp. (50)         |
| KT966499              | Guáico Culex virus | 2       | Unclassified | 407      | *Culex* (*Culex*) spp. (50)         |
| KT966500              | Guáico Culex virus | 3       | Unclassified | 407      | *Culex* (*Culex*) spp. (50)         |
| KT966501              | Guáico Culex virus | 4       | Unclassified | 407      | *Culex* (*Culex*) spp. (50)         |
| KP762047              | Guáico Culex virus | 5       | Unclassified | 407      | *Culex* (*Culex*) spp. (50)         |

* The taxonomy, source, and relationship to similar known viruses were determined by nucleotide BLAST (“BLASTx” in GenBank).
(20.4 kb). OFAV and KADV align with each other with 70% nucleotide identity over a query coverage of 68%.

GUTV and KAIV isolated from Culex (Culex) spp. most closely align with a recently described virus from China called Wuchang Cockroach virus 3. The three isolates of GUTV are 99% identical to each other, whereas KAIV and GUTV are 30% divergent at the nucleotide level. These viruses consistently accompanied isolates of BRJV. Whether the accompaniment of these viruses is coincidence, or they derived from coinfected mosquitoes within each pool, is unknown. However, KAIV and GUTV genomes could in fact be independent genomic segments within a multisegmented genome that includes BRJV.

To evaluate the host range of these virus isolates, attempts to generate cytopathic effect (CPE) were made with human, monkey, frog, bat, mosquito, and tick cell cultures. A clear CPE was observed only in mosquito cells. However, the capacity of these novel viruses to cause CPE in vertebrate and invertebrate cells was not fully evaluated.

Pantanál’s biodiversity has recently produced other mosquito-borne viruses, and may be useful for studying virus evolution (9–14).

Accession number(s). The sequences have been deposited in GenBank under the accession numbers listed in Table 1.

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