

Characterisation of divergent flavivirus NS3 and NS5 protein sequences detected in *Rhipicephalus microplus* ticks from Brazil

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Transcripts similar to those that encode the nonstructural (NS) proteins NS3 and NS5 from flaviviruses were found in a salivary gland (SG) complementary DNA (cDNA) library from the cattle tick Rhipicephalus microplus. Tick extracts were cultured with cells to enable the isolation of viruses capable of replicating in cultured invertebrate and vertebrate cells. Deep sequencing of the viral RNA isolated from culture supernatants provided the complete coding sequences for the NS3 and NS5 proteins and their molecular characterisation confirmed similarity with the NS3 and NS5 sequences from other flaviviruses. Despite this similarity, phylogenetic analyses revealed that this potentially novel virus may be a highly divergent member of the genus Flavivirus. Interestingly, we detected the divergent NS3 and NS5 sequences in ticks collected from several dairy farms widely distributed throughout three regions of Brazil. This is the first report of flavivirus-like transcripts in R. microplus ticks. This novel virus is a potential arbovirus because it replicated in arthropod and mammalian cells; furthermore, it was detected in a cDNA library from tick SGs and therefore may be present in tick saliva. It is important to determine whether and by what means this potential virus is transmissible and to monitor the virus as a potential emerging tick-borne zoonotic pathogen.

Key words: *Rhipicephalus microplus* - tick - NS3 - NS5 - flavivirus

Ticks transmit a greater variety of infectious agents to humans and other animal species than does any other blood-feeding arthropod (Jongejan & Uilenberg 2004). The pathogens transmitted by ticks include bacteria, protozoa and viruses, with members of the Flaviviridae family among the most common tick-borne viruses. The Flaviviridae family is composed of three genera: Pestivirus, which contains the species responsible for zoonotic infections (ICTV 1995), Hepacivirus, which contains the human hepatitis C virus and (tentatively) the GB viruses (Ferron et al. 2005) and Flavivirus, which contains over 70 species, most of which are arthropod-borne (arboviruses) (Thiel et al. 2005). The tick-borne viral diseases caused by viruses of the Flavivirus genus are mainly

infections of the central nervous system characterised by severe encephalitis in humans. Important and complex environmental interactions involving animal reservoirs of ticks and tick-borne viruses may influence the incidence of these viral diseases (Norman et al. 1999, Laurenson et al. 2003, Cope et al. 2004). In addition to the tick-borne group, the Flavivirus genus contains a mosquito-borne group and an arthropod group with no known vector (NKV) that infects vertebrate hosts, but does not have an identified arthropod vector (Kuno et al. 1998). The Tamana bat virus (TABV) and the cell fusing agent virus are classified as tentative species in the Flavivirus genus (Thiel et al. 2005) because they are considered highly divergent from the flaviviruses.

While analysing a salivary gland (SG) transcriptome from females of the cattle tick *Rhipicephalus microplus*, we identified transcripts with significant similarity to various viral genes. A few of these transcripts were similar to the nonstructural (NS)3 and NS5 genes of viruses in the Flavivirus genus. The Flavivirus NS proteins NS3 and NS5 perform the enzymatic activities necessary for RNA capping and genome replication (Bollati et al. 2010). Although virus-like particles have been found in the SGs of *R. microplus* (Megaw 1978), there have been no reports of Flaviviridae viruses in this tick species.

To ascertain whether the transcripts found in the *R. microplus* SG transcriptome were indeed from an RNA virus, we performed a molecular assay [polymerase chain reaction (PCR)] using DNA and RNA [reverse

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transcribed into complementary DNA (cDNA)] from a different tick sample. The results confirmed that the transcripts were not a product of viral integration into the tick genome. The virus was then isolated from tick extracts in cell culture. The virus samples from the culture supernatants were PCR-positive for the same transcripts detected in the SGs of *R. microplus*, similar to the NS3 and NS5 sequences, suggesting that a potentially novel virus had been isolated. We refer to this possible novel virus as the Mogiana tick virus (MGTV) because of the region where it was isolated.

Viral RNA from the culture supernatant of MGTV-infected Vero cells was deep sequenced and we obtained the complete sequences of NS3 and NS5 from the novel virus. We conducted a comparative molecular analysis of these sequences with those from other Flaviviridae viruses and the results revealed that these sequences were highly divergent from those of other members of the Flavivirus genus. In addition, we detected these divergent NS3 and NS5 sequences in field samples of ticks, which suggests a wide geographical distribution of the virus in Brazil. These findings highlight the importance of studying the presence of arboviruses in *R. microplus* and the associations of these viruses with vectors and vertebrate hosts. These results also increase awareness of possible emerging zoonoses and/or tick-borne viral diseases in bovines.

MATERIALS AND METHODS

cDNA library and bioinformatics analysis - Adult female *R. microplus* ticks were collected from naturally infested cattle and the SGs were immediately dissected from 30 ticks. An *R. microplus* SG library was used and its construction has been described in detail by Maruyama et al. (2010). The bioinformatics analysis of the transcriptome data was performed as previously described (Ribeiro et al. 2006), with some modifications. The alignments were performed using the CLUSTALW program (Thompson et al. 1994) and BioEdit (Hall 1999) sequence alignment editing software. The phylogenetic associations were determined using the neighbour joining (NJ) or maximum likelihood (ML) methods (MEGA 4.0) (Tamura et al. 2007) and the node support of each clade was evaluated using a bootstrap analysis (1,000 replicates). The hydropathy profiles of the proteins were obtained using the web-based tool ProtScale (web.expasy.org/protscale/) from the ExPASy Bioinformatics Resource Portal (Artimo et al. 2012) with the Kyte and Doolittle (1982) scale option and a window size of nine amino acids (aa). The hydropathy plot were constructed based on the alignment of NS3 and NS5 transcripts found in *R. microplus* with NS3 and NS5 transcripts from dengue virus type 2 (DENV-2). Because the transcripts had different lengths the score was zero for gaps. The codon usage adaptation index (CAI) and base composition values were obtained using tools provided by the CAIcal server (genomes.urv.es/CAIcal/) (Puigbò et al. 2008a). The codon usage tables for Flavivirus spp, *R. microplus*, *Bos taurus* and *Homo sapiens* were obtained from the Codon Usage Database (kazusa.or.jp/codon/), which is compiled from the GenBank DNA sequence database

(Kazusa DNA Research Institute). The NS3 and NS5 protein sequences of Flaviviridae viruses used in this work were obtained from National Center for Biotechnology Information (NCBI) RefSeq collection (Supplementary data, Table SI).

Viral isolation - The virus was isolated from ticks collected from Holstein bulls on a farm in Ribeirão Preto, state of São Paulo (SP), Brazil (21°13'09.28''S 48°31'34.91''W). Pools of 20 ticks were crushed in liquid nitrogen, resuspended in 1 mL of sterile phosphate-buffered saline (PBS) (pH 7.0) containing 10% foetal bovine serum (Invitrogen, San Diego, CA, USA), penicillin (500 IU/mL) and streptomycin (500 mg/mL) (Invitrogen) and centrifuged at 2,500 g for 5 min. The supernatant was collected and stored at -70°C until use. Prior to the inoculation of the culture medium, the supernatant was filtered in a 0.22-µm filter.

Viral isolation was performed using both arthropod and mammalian cells. African green monkey kidney (Vero) and baby hamster kidney (BHK)-21 cells were grown at 37°C under 5% CO₂ in minimum essential medium (MEM) (Invitrogen) supplemented with 10% inactivated foetal calf serum (FCS) (Invitrogen), 1% L-glutamine (Cultilab, Campinas, SP, Brazil) and 1% antibiotic-antimycotic (Invitrogen). *Boophilus microplus* cattle tick (BME 26) cells were obtained from Dr Ulrike Munderloh in the Department of Entomology at the University of Minnesota. *Aedes albopictus* mosquito (C6/36) cells were grown at 30°C in Leibovitz's medium (L-15) (Invitrogen) supplemented with 10% FCS and 1% antibiotic-antimycotic.

A 30-µL aliquot of each tick pool filtrate was diluted in 310 µL of PBS (pH 7.0) with 2% antibiotic-antimycotic (Invitrogen). This solution was inoculated into paired wells of six-well plates (Corning, Corning, NY, USA) containing semiconfluent monolayers of C6/36, BME 26, Vero and BHK cells. The filtrate was allowed to adsorb for 60 min at 37°C (Vero and BHK cells) or 30°C (C6/36 and BME 26). To prevent the detection of residual, nonreplicating viruses from the inoculum after the adsorption of the virus from the tick extracts to the cell monolayers, the plates were thoroughly washed with sterile saline to remove the inoculum before the maintenance medium was added. The cells were incubated for seven days at 37°C in a 5% CO₂ atmosphere (Vero and BHK) or at 30°C (C6/36 and BME 26) with maintenance medium (for Vero and BHK cells: MEM with 2% FCS, 1% L-glutamine and 2% antibiotic-antimycotic; for C6/36 and BME 26 cells: L-15 supplemented with 2% FCS and 2% antibiotic-antimycotic). The plates were observed daily for any cytopathic effects (CPE) and infection was confirmed by the molecular detection of the virus. RNA was extracted from the cell culture supernatants of the first and second passages using a QIAamp Viral RNA kit (Qiagen, Valencia, CA, USA) and reverse-transcription (RT)-PCR was performed for molecular detection using primers for the NS3 and NS5 coding sequences of MGTV. Additionally, the first passage virus was replicated in Vero cells. Cells were infected as described above and the supernatant was collected 1 h, 3 h and 6 h after infection and on days one,

three, five, seven and nine after infection. The viral load (VL) was determined by amplifying the NS5 region herein described. A KAPA SYBR® FAST Universal Kit was used the NS5 region through real-time PCR, using the KAPA SYBR (KAPA Biosystems, Woburn, MA, USA) under the amplification conditions recommended by the manufacturer. The reactions were performed in a 7500 Fast Real-Time PCR System (Applied Biosystems, Foster City, CA, USA). A standard curve was generated by cloning the 401 region using the InsTAclone PCR Cloning Kit (Thermo Scientific) and the final value was expressed as MGTV RNA copies/mL calculated as copies/mL.

Deep sequencing of viral RNA - Purification of viruses by sucrose cushion - Cultures of Vero cells (2×10^6) were infected with tick pools as described. The culture supernatants were collected six days post-infection. The cellular debris was removed by centrifugation at 10,000 g for 30 min at 4°C. The supernatants were layered onto a 20% sucrose cushion prepared in TNM buffer [10 mM Tris HCl (pH 7.5), 5 mM MgCl₂ and 150 mM NaCl]. The solution was centrifuged in an AH-629 Thermo Scientific rotor at 27,000 rpm for 9 h at 4°C. The viral pellet was resuspended in 0.3 mL of cold PBS and used for RNA extraction.

RNA preparation - A 0.15-mL aliquot of purified virus was treated with 100 units of RNase I and four units of DNase I at 37°C for 1 h to remove nucleic acids that were not protected by viral capsids. After incubation, 2 µL of dithiothreitol and 2 µL of RNase inhibitor were added and RNA was extracted using a QIAamp viral RNA mini kit (Qiagen). The viral RNA was sequenced by an external company.

Deep sequencing using the Illumina MiSeq platform - Sequence data were generated by the High-Throughput Sequencing and Genotyping Unit at the University of Illinois in Urbana-Champaign. The deep sequencing was performed according to the manufacturer's (Illumina, San Diego, CA) instructions. Briefly, after a sample quality control analysis, the RNA was converted into cDNA using random hexamers and was then nebulised, adapted and quantitated. The paired-end reads [150 nucleotides (nt) in length] were sequenced in one lane of a MiSeq instrument for 2 x 150 cycles and were further analysed using Casava 1.8.2. A total of 11,307,270 Illumina paired-end reads were obtained.

Bioinformatics analysis of the short reads - The workflow for the bioinformatic analysis consisted of rounds of mapping steps to reference genomes to filter out of unwanted sequences. The TopHat2 (Kim et al. 2013) and Bowtie2 (Langmead & Salzberg 2012) aligners were used with the default options to remove the following: sequences that were identical to host cell sequences [Rhesus monkey *Macaca mulatta* genome assembly MMUL_1, downloaded from Ensembl (ensembl.org/index.html)], then to sequences from bacterial contaminants [*Mycoplasma hyorhinitis* SK6 genome assembly GCA_000313635.1, downloaded from EnsemblBacteria (bacteria.ensembl.org/index.html)] and ultimately to unwanted viral sequences [artificial reference

genome built from DNA viruses, retrovirus genomes and sequences from endogenous viruses in primates, all downloaded from RefSeq-NCBI (ncbi.nlm.nih.gov/refseq/)]. After the successive purging steps, 6,512,646 paired-end reads were used as input for *de novo* assembly with the program SOAPdenovo-Trans (Luo et al. 2012). Variable k-mer values ranging from k = 13 to k = 31 (in steps of 2) and from k = 33 to k = 73 (in steps of 4) and the parameters -d 3 -D 5 -L 150 -u -e 3 for every selected k-mer size were employed. All the resulting assembled sequences larger than 150 nt were combined into a FASTA file and submitted as input to an assembly pipeline consisting of BLASTN (Altschul et al. 1997) and CAP3 (Huang & Madan 1999) iterations, as previously described (Karim et al. 2011). The assembled contigs were BLASTed against the NCBI nonredundant (NR) protein database Swiss Prot (O'Donovan et al. 2002), the Conserved Domains Database (CDD) (Marchler-Bauer et al. 2002), a customised protein database containing only viral proteins from the NR database and a second customised database containing only Flavivirus proteins from RefSeq-NCBI. The data from the bioinformatics analysis are presented in Supplementary data (Table SII), a hyperlinked Excel spreadsheet.

Collection of the ticks - Ticks were collected in March 2006 from 19 livestock farms located in different regions of Brazil (Supplementary data, Table SIII) in a partnership with Vallee SA [Montes Claros, state Minas Gerais (MG), Brazil]. All the samples were placed in RNAlater (Invitrogen) and stored at -70°C for later nucleic acid extraction.

DNA and total RNA extraction - Pools of ticks (organised according to locality and life stage) (Supplementary data, Table SIII) were crushed and resuspended in PBS for simultaneous DNA and RNA isolation. DNA was extracted from an aliquot of the homogenate using the QIAamp DNA kit (Qiagen). The remaining homogenate was mixed with Trizol (Invitrogen) and total RNA was isolated from it using the SV Total RNA Isolation Kit (Promega, Madison, WI, USA). cDNA was synthesised from the total RNA from the tick samples with the ImProm-II™ Reverse Transcription System kit (Promega). All the procedures were performed according to the manufacturers' instructions.

Molecular detection - PCR was performed to detect the divergent NS3 and NS5 fragments in genomic DNA (gDNA) and cDNA. The contig sequences were used as templates for primer design. Previously described actin primers (de la Fuente et al. 2008) were used as reaction controls. The reactions were performed using the primers listed in Supplementary data (Table SIV).

Accessions - All the sequences obtained from the ticks and culture cells have been deposited in GenBank (NCBI). The accessions include the following: (i) the partial sequences of isolates from tick pools (extract and culture) are JQ289026-JQ289041, as detected in ticks, (ii) the Illumina reads are the short reads archive SRA 055953, (iii) the complete sequences of NS3 and NS5 are JX390985 (protein_id: "AGL39759") and JX390986

(protein_id: “AGL39760”), respectively, and (iv) the partial sequences from the field samples of ticks and the cDNA library are HS586608-HS586670.

RESULTS AND DISCUSSION

Similarities to the NS NS3 and NS5 proteins in the tick cDNA library - The cDNA library from the SGs of female *R. microplus* generated 1152 expressed sequence tags (ESTs) clustered into 533 contigs (Maruyama et al. 2010), which were compared to the NCBI NR protein database using BLASTX. The annotation of the contigs revealed that five contigs aligned with NS flaviviral proteins (Table I) spanning seven ESTs. NS3 and NS5 proteins from three different Flavivirus species (TABV, Apoi virus and Kamiti River virus) represented the best matches in the NR protein database, as indicated by low e-values (except for contig 319, with an e-value of 0.002).

To evaluate the presence of this putative Flavivirus member in a different tick sample and to confirm that our sequences were not derived from a viral insertion into the tick genome, as has been reported for the mosquito *Ae. albopictus* (Crochu et al. 2004), we performed molecular detection assays in different samples of *R. microplus* ticks. These samples were collected from two Brazilian farms in Ribeirão Preto and Araguari (MG); these cities are located in the Mogiana macroregion (between SP and MG). We designed primer pairs for , sequences 317, 401 and 2743, which produced amplicons of 258 bp, 281 bp and 248 bp, respectively. A protocol for extracting gDNA and total RNA from the same tick sample was developed to facilitate viral detection through PCR with gDNA or cDNA as a template. The 317, 401 and 2743 primers amplified one conserved motif from the nucleotidases domain of NS3 (motif VI: QRRGRVGR) (Wu et al. 2005), one conserved motif from the methyltransferase domain of NS5 (motif IV: DTLLFDGGE) (Egloff et al. 2002) and one conserved motif from the RNA-de-

pendent RNA polymerase domain (RdRp) of NS5 (motif B: SG_VVTYALNLT) (Poch et al. 1989), respectively. PCR using each set of primers (317, 401, 2743 and actin as a positive control) was used to amplify the gDNA and cDNA samples. The agarose gel electrophoresis analysis revealed that no products were obtained using the RNase-treated gDNA as a template for the detection of fragments of the 317 (NS3), 401 (NS5) and 2743 (NS5) sequences, except in the positive control reactions using actin primers. The PCR-negative results for the gDNA template indicated that these sequences were not derived from the tick genome. In contrast, when cDNA (RNA) was used as a template, all nine tick samples were PCR-positive for the 317 primer set detection, whereas seven tick samples were PCR-positive for the 401 and 2743 primer set detection (Supplementary data, Fig. S1). These results indicated that the viral-like transcripts observed in the tick cDNA library were genuinely derived from RNA (cDNA) and were likely from an RNA virus.

Viral isolation in cultured cells and deep sequencing of RNA in viral capsids - To confirm the presence of the presumed Flavivirus found in the tick samples, viral isolation was performed in mammalian and arthropod cells. Five pools of ticks (4 from females and 1 from larvae) from the Ribeirão Preto farm, all of which were PCR-positive for the 317, 401 and 2743 sequences, were used to inoculate the cell cultures. No CPE was observed in the mosquito (C6/36), tick (BME 26) or hamster (BHK) cell lines; a CPE was observed only in the Vero cells (data not shown). Upon primary isolation, the CPE was minimal and developed slowly (for 5 days after the inoculation), resulting in the rounding and shrinkage of the cells, which became refractile and detached from the plate surface. After the second passage, the CPE disappeared; therefore, we could not titre it with a plaque-forming assay. The CPE observed during the initial passage alone

TABLE I
Contigs of *Rhipicephalus microplus* ticks salivary glands transcriptome that presented matches with nonstructural (NS) proteins NS3 and NS5 of flaviviruses

Contigs ^a	Number of ESTs	Best mach to NR database		
		BLASTX	e-value	Virus
317	2 (gi 317017979; gi 317017980)	Putative NS protein NS3 (gi 27735332)	5e-009	TABV
319	1 (gi 317017985)	Putative NS protein NS3 (gi 27735332)	0.002	TABV
400	2 (gi 317017981; gi 317017982)	NS protein NS5 (gi 27697405)	3e-006	APOIV
401	1 (gi 317017983)	NS5 protein (gi 37695589)	2e-005	KRV
2,743	1 (gi 317017984)	Putative NS protein NS5 (gi 27735336)	1e-010	TABV

^a: these five contigs are part of a transcriptome analysis that generated more than 3,000 contigs; APOIV: Apoi virus; KRV: Kamiti virus; TABV: Tamana bat virus.

might have been due to toxicity in the inoculum; however, no CPE was observed in the C6/36, BME 26 or BHK cells inoculated under the same conditions.

The molecular detection of the virus was performed using supernatants from the four cell lines and primer sets 317, 401 and 2743. Each supernatant was inoculated with a different pool of ticks. The cell cultures were inoculated with the tick extracts, the cell monolayers were thoroughly washed after 1 h of incubation to remove any residue from the inoculum and maintenance medium was added. Seven days later, the culture supernatants were subjected to PCR. Sequences 317, 401 and 2,743 were successfully amplified from all the cell cultures inoculated with female tick pools, but not from the cell cultures inoculated with unfed larvae (UL) pools (Supplementary data, Fig. S2 displays the results from primer set 401). The culture supernatants from the second passage of the four cell lines remained PCR-positive for primer set 401 (Supplementary data, Fig. S3 shows the results from the Vero cells). Thus, the positive PCR results suggested that the virus replicated in the mammalian and arthropod cell lines. The viral strains isolated from the four tick pools were sequenced using primer sets 317 and 401 and the results confirmed that these strains represented the same fragments previously identified in the *R. microplus* SG cDNA library (data not shown).

The VL from the first passage was further evaluated in Vero cells and the viral titre was highest on the seventh day, at 1.7×10^4 copies/mL (data not shown). This result differs from findings from other flaviviruses, which exhibit the highest VL between the third and fifth days after infection (Bonaldo et al. 2007, Orlinger et al. 2011). The low observed VL may have been responsible for the absence of a CPE in the second passage. Because this was a field isolate, it may have required more passages to increase the titre.

To obtain longer sequences than those of tick cDNA library contigs 307, 401 and 2,743, we first performed PCR using the cDNA produced from viral RNA isolated from the tick extracts and culture supernatants described. A previously described set of universal Flavivirus primers was employed (Gaunt & Gould 2005, Maher-Sturgess et al. 2008). None of the amplifications with the universal primers produced amplicons (data not shown), indicating that the genome of the isolated virus may be highly divergent from those of classical flaviviruses. We then purified the virus from an infected Vero cell supernatant (2nd passage). Following RNase and DNase treatment to degrade the nucleic acids that were not protected by viral capsids, the viral RNA was extracted and the molecular detection of the divergent NS5 sequence was confirmed (Supplementary data, Fig. S3). This viral RNA was used for deep sequencing on an Illumina platform. The MiSeq run generated 11,898,134 paired-end reads, which were purged of reads with sequence identity to the cellular host bacterial contamination (*M. hyorhinis*), DNA viruses, retroviruses or endogenous viruses of primates. The purged paired-end read files were then assembled using the SOAPdenovo-Trans program. An automatic annotation was performed using a customised bioinformatics workflow on the BLASTX

protein database. The RNA-seq dataset, which included over 6,500 contigs, was plotted in a hyperlinked Excel spreadsheet (Supplementary data, Table SII). We searched this dataset for significant BLAST results against the Flavivirus protein database that were identical to the tick cDNA library contigs (317, 401 and 2,743) described in Table I.

Despite the use of high-throughput sequencing, the assembled reads did not reveal the full-length genome sequence of the potentially novel virus. Although steps to remove background had been performed, most of the assembled reads were assigned to bacteria and miscellaneous endogenous virus sequences. This contamination, together with the low MGTV VL in the cell cultures, appears to have hindered the sequencing of transcripts expressed by MGTV at a lower frequency. The deep sequencing results revealed that the majority of the reads from our viral RNA sample from cultured MGTV-infected cell supernatants may have been derived from an endogenous primate retrovirus present in Vero cells. The chemical induction of endogenous retroviruses in Vero cells has been reported (Ma et al. 2011). Furthermore, the mobilisation of endogenous retroviruses in mice following infection with an exogenous retrovirus has occurred (Evans et al. 2009). Because we did not investigate whether viral RNA was present in supernatants of uninfected Vero cells (control), we could not determine whether these endogenous viral-like particles were constitutive or induced by MGTV infection.

Only 0.025% of the reads exhibited identity with contig sequences 401 and 317. Although viral RNA sequences from MGTV were underrepresented, we found two RNA-seq contigs, 1,961 and 2,579, with greater than 90% nt identity with the tick cDNA library contigs (Supplementary data, Table SII, highlighted rows). Both contigs exhibited significant similarity to the TABV polyprotein. RNA-seq contigs 1,961 (2,993 nt) and 2,579 (2,721 nt) were similar to the Flavivirus NS5 and NS3 NS proteins, respectively. Several contigs exhibited intermediate similarity with other Flavivirus structural and NS proteins (Supplementary data, Fig. S6). However, because most of these contigs represented short consensus sequences, we were unable to build a proper scaffold genome for MGTV. Therefore, RNA-seq contigs 1,961 and 2,579 were the only ones confirmed as MGTV-derived sequences. Although the complete genomic sequence could not be identified through deep sequencing, the complete NS3 and NS5 sequences from MGTV were obtained. Because NS3 and NS5 are the two largest and most conserved Flavivirus proteins (Chambers et al. 1990), we performed a comparative sequence analysis between these two MGTV proteins and NS3 and NS5 from Flaviviridae viruses to confirm our findings concerning the presence of Flavivirus-like transcripts in *R. microplus* ticks and to investigate the relationship between this presumed flavivirus and other members of the Flaviviridae family.

Molecular characterisation of the NS3 and NS5 protein sequences found in R. microplus ticks - The NS3 and NS5 proteins are the two main components of the flaviviral replication machinery. To obtain the complete NS3 and NS5 protein sequences from MGTV, open

reading frames +1 and -2 of the nt consensus sequences of RNA-seq contigs 1,961 and 2,579, respectively, were translated into aa sequences and aligned with different flaviviral NS3 and NS5 protein sequences [the Flavivirus sequences used are listed in the Supplementary data (Table SI)]. The alignment results were used to obtain the complete protein sequences for NS3 (554 aa, GenBank accession JX390985) and NS5 (866 aa, GenBank accession JX390986) from the potentially novel virus. Both proteins were shorter than known proteins from other flaviviruses, except for NS5 from TABV (831 aa).

Conserved motifs - To analyse sequence conservation in motifs of the MGTV NS3 and NS5 proteins, we performed sequence alignments using flavivirus sequences from one representative virus for each group vector and highlighted the conserved motifs. The N-terminal region of the flavivirus NS3 protein encodes a viral serine protease (Gorbalenya et al. 1989, Falgout et al. 1991), whereas the C-terminal portion encodes a helicase (Lain et al. 1989). The alignment of the flavivirus NS3 protein sequences revealed that residues in important regions for these functions were conserved in NS3 from MGTV (Fig. 1). The catalytic triad of histidine, aspartate

and serine residues required for serine protease activity (Polgar 2005) was conserved in the MGTV sequence at positions 45, 69 and 126, respectively. Together with a glycine residue at position 124 [Fig. 1A, Supplementary data (Fig. S7)], these four residues provide an electrostatic environment for the active site of the enzyme. The flavivirus helicase/NTPase catalyses the unwinding of the RNA strand (Dumont et al. 2006) to facilitate the initiation of viral replication and contains seven conserved motifs (Gorbalenya & Koonin 1993). Compared with the set of motifs described for the yellow fever virus (Wu et al. 2005), all seven motifs were well conserved in the corresponding novel sequence (Fig. 1B). Of note, motifs I and VI were the most strongly conserved. The complete sequence alignment of the NS3 proteins displayed in Fig. 1 can be viewed in the Supplementary data (Fig. S4).

The flaviviral NS5 protein contains a C-terminal (RdRp), whereas the N-terminus possesses the methyltransferase activity implicated in capping the 5'-end of the flaviviral RNA genome (Egloff et al. 2002, Davidson 2009). The alignment of the flavivirus NS5 protein sequences revealed that NS5 from MGTV contained conserved motifs (Fig. 2). The NS5 N-terminus also con-

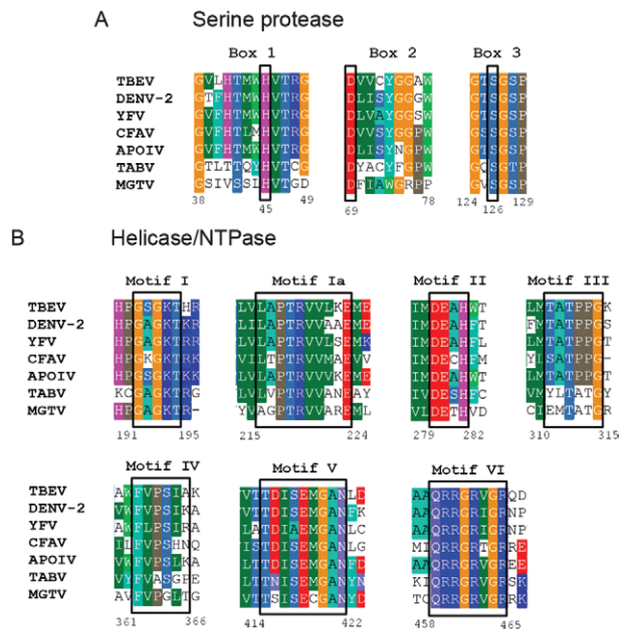


Fig. 1: conserved motifs in nonstructural (NS)3 of Mogiana tick virus (MGTV). Multiple sequence alignment of NS3 from MGTV and representative of tick-borne encephalitis virus (TBEV), mosquito-borne [dengue virus type 2 (DENV-2) and yellow fever virus (YFV)], insect-only [cell fusing agent virus (CFAV)] and not known vector [Apoi virus (APOIV) and Tamana bat virus (TABV)] flavivirus groups. For better visualisation, only blocks of conserved regions along the alignment are displayed. Motifs are delimited accordingly. A: N-terminal regions of NS3 contain a serine protease, amino acids (aa) of the catalytic triad of which are highlighted; B: C-terminus portion of NS3 contain a helicase/nucleotidases (NTPases) where seven conserved motifs (I, Ia and II-VI) are found. Numbers at the bottom of the alignments refer to MGTV sequence. The threshold for shading colours of aa similarity was 50%.

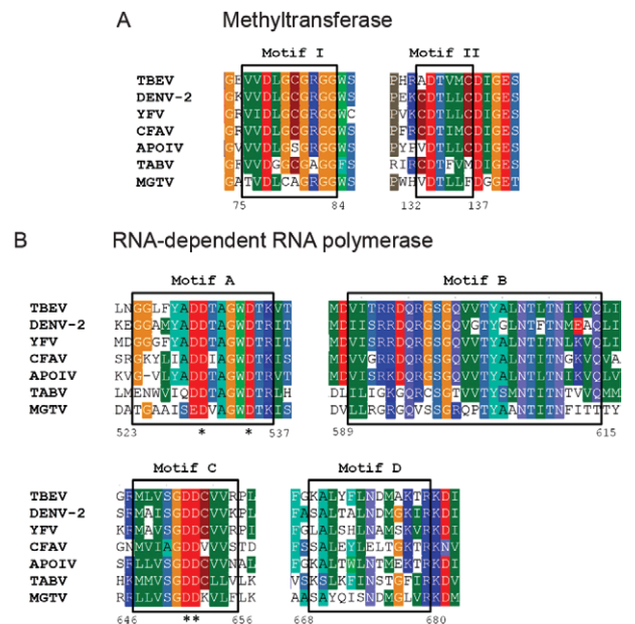


Fig. 2: conserved motifs in nonstructural (NS)5 of Mogiana tick virus (MGTV). Multiple sequence alignment of NS5 from MGTV and representative of tick-borne encephalitis virus (TBEV), mosquito-borne [dengue virus type 2 (DENV-2) and yellow fever virus (YFV)], insect-only [cell fusing agent virus (CFAV)] and no known vector [Apoi virus (APOIV) and Tamana bat virus (TABV)] flavivirus groups. To better visualisation, only blocks of conserved regions along the alignment are displayed. Motifs are accordingly delimited. A: N-terminal region of NS5 presents methyltransferase activity; two conserved motifs are found (I and II); B: C-terminus portion of NS5 contains an RNA-dependent RNA polymerase; four conserved motifs (A-D) are found; asterisk indicates the conserved aspartate residues important for enzyme activity. Numbers at the bottom of the alignments refer to MGTV sequences. The threshold for shading colours of amino acid similarity was 50%.

tains an RNA guanylyltransferase, which uses GTP as a substrate and its activity is stimulated by the NS3 protein (Issur et al. 2009). The N-terminal methyltransferase domain displayed two conserved motifs, I and II, involved in S-adenosyl methionine binding (Koonin 1993). The novel (MGTV) NS5 sequence exhibited approximately 70% identity with residues from motifs I and II of the methyltransferase region (Fig. 2A). In addition to capping the viral genome, NS5 is involved in the synthesis of RNA through the RdRp activity at the C-terminus; thus, it plays a key role in viral replication. The RdRp domain includes four conserved motifs: A, B, C and D (Poch et al. 1989). Of these, the D motif appears to be the least conserved among the analysed flaviviruses; the RdRp from the MGTV NS5 sequence showed conserved residues mainly in motifs A and C (Fig. 2B). Remarkably, the four strictly conserved aspartate residues, D-X₄-D and G-D-D (which are located in motifs A and C, respectively) were also strictly conserved in the MGTV NS5 sequence at positions 530-535 and 650-652, respectively (Fig. 2B, indicated by an asterisk). The complete sequence alignment of the NS5 proteins displayed in Fig. 2 can be viewed in the Supplementary data (Fig. S5).

Phylogenetic analysis - To verify the phylogenetic associations between MGTV and other Flaviviridae members, we performed ML analyses of the NS3 and NS5 sequences. In addition to the Flavivirus genus (including the divergent TABV), members of the Pestivirus and Hepacivirus genera were included in the analyses [the sequences IDs are listed in Supplementary data (Table SI)]. The phylograms of NS3 and NS5 indicated that MGTV is positioned closer to the Flavivirus genus than to the other genera of Flaviviridae (Fig. 3), which was supported by significant bootstrap values. However, MGTV is highly divergent from the other flaviviruses and is distantly related to all Flavivirus species described to date, both vectored and non-vectored, including TABV.

The phylogeny of the Flavivirus genus has been studied extensively (Kuno et al. 1998, Billoir et al. 2000, Gaunt et al. 2001) and different patterns of phylogenetic positions have been observed, depending on the gene analysed. All these studies used the divergent insect-only viruses and did not consider the divergent TABV virus. We included TABV in our analyses due to the BLASTX results, which showed that the best hit was a TABV polyprotein [Table I, Supplementary data (Table SII)]. A more recent comprehensive phylogenetic analysis of NS3, NS5 and entire flaviviruses genomes sequences using the ML method revealed that the NS3 and complete genome trees exhibited the same phylogenetic topology in the flavivirus groups (Cook & Holmes 2006). Because the NS3 tree appears to reflect information from relationships between complete flavivirus genome sequences, the association of MGTV with the Flavivirus genus is also likely reliable (Fig. 3A) despite the high divergence of this novel virus. The complete genomic sequence of MGTV will be critical for clarifying the taxonomic organisation of the Flaviviridae family.

Hydropathy profile - The aa chemical properties of a protein are reflected in its hydrophobicity/hydrophilicity

profile. The MGTV NS3 and NS5 proteins exhibited a hydrophobic content of 33% (Supplementary data, Table SV). Because the flaviviral NS3 and NS5 proteins are hydrophilic (Chambers et al. 1990), we compared the hydrophathy profiles of NS3 and NS5 from MGTV with the profiles of the corresponding proteins from a classical flavivirus, DENV-2. Both NS3 and NS5 from MGTV displayed distributions of hydrophilic residues (Fig. 4) similar to those of DENV-2. Together with the observed similarities in the conserved motifs (Figs 1, 2), this result highlights the physiochemical similarities between the MGTV NS3 and NS5 proteins and those that form the flaviviral replication complex.

Codon usage - RNA viruses evolve quickly; however, particular genomic sites may be more conserved due to purifying selection. The resulting genome reflects these opposing pressures and specific sites may reflect more conservative forces in the form of signatures that may be identified by an analysis of their base composition. We explored the NS3 and NS5 nt sequences from MGTV and from other flaviviruses by analysing their base composition. We used three different approaches [Nc (effective number of codons), overall GC (G + C bases composition) and GC3 (G + C content of third-base codon position) and CAI] that have been applied to analyse codon bias; these methods are explained in the Supplementary data (Fig. S7) together with details of the results of the analy-

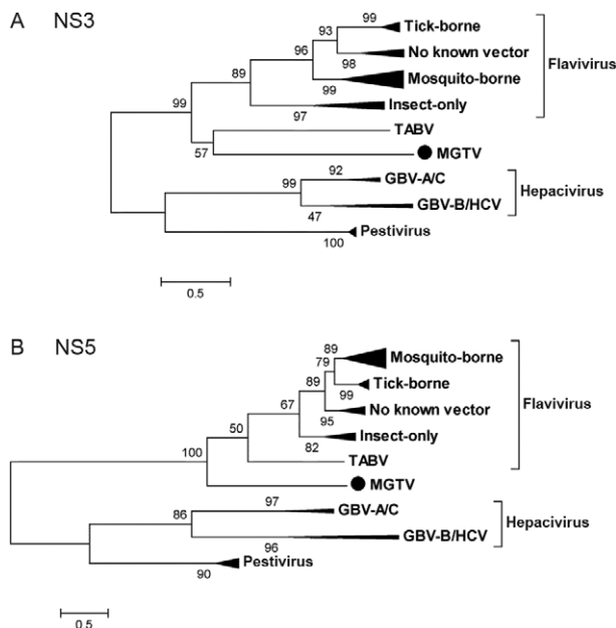


Fig. 3: phylogenetic analysis of nonstructural (NS)3 and NS5 sequences of Mogiana tick virus (MGTV) virus within the Flaviviridae family. Maximum likelihood analyses were performed with 1,000 bootstrap replicates. The evolutionary distances were computed using the JTT matrix-based method. A: NS3 tree; B: NS5 tree. Virus groups were condensed for better visualisation and all viruses used in trees construction are listed in Supplementary data (Table SIII). The bar at the bottom indicates 50% amino acid divergence. GBV: GB virus; HCV: hepatitis C virus; TABV: Tamana bat virus (TABV).

ses. All the codon usage computations were performed using the CAIcal server (Puigbò et al. 2008a). The CAI calculation for NS3 and NS5 from the potentially novel virus indicated similarity with flaviviral codon usage.

Because the CAI value (e-CAI) indicates a codon bias towards a reference set, we used the reference set from the codon usage table of Flavivirus spp to determine the e-CAI of the sequences of MGTV NS3 and NS5 and of the flaviviruses addressed in this study. A normalised e-CAI of ≥ 1.0 indicates that the observed CAI is equal to or greater than the expected value (eCAI); such a result could be interpreted as a codon usage adaptation towards the Flavivirus genus. The normalised CAI_{NS3} and CAI_{NS5} values for MGTV were 1.00247 and 1.00732, respectively, suggesting that both genes indeed exhibit a codon usage adaptation towards the Flavivirus genus (Fig. 5). Interestingly, the e-CAI of both MGTV genes showed greater accordance with the Flavivirus codon usage patterns than some well-described flaviviruses.

The circulation of RNA arboviruses in nature, such as mosquito-borne and tick-borne Flavivirus species, rely on their capacity of replication in a range of hosts (vertebrates and invertebrates). Some experimental studies of the evolution of RNA arboviruses have shown that the host alternation cycles constrain the adaptation and evolution of these viruses towards a single host, corroborating the theory that an adaptation selected by one host involves a fitness trade-off for the other (Weaver et al. 1999, Coffey et al. 2008). Analyses of the dinucleotide compositions and codon usage patterns of Flaviviridae viruses and their hosts have suggested that host-induced pressure shapes the viral codon usage pattern (Lobo et al. 2009): the NKV mosquito-borne and tick-borne groups displayed dinucleotide usage patterns that were more closely related to the vertebrate genomic signature.

We used the normalised CAI data to explore the relationship between the codon usage patterns of MGTV and its potential invertebrate host, the cattle tick *R. microplus*. Furthermore, we extrapolated this analysis to virtual vertebrate (bovine and human) hosts because we had observed the replication of MGTV in mammalian cells (Vero and BHK) and arthropod cells (BME 26 and

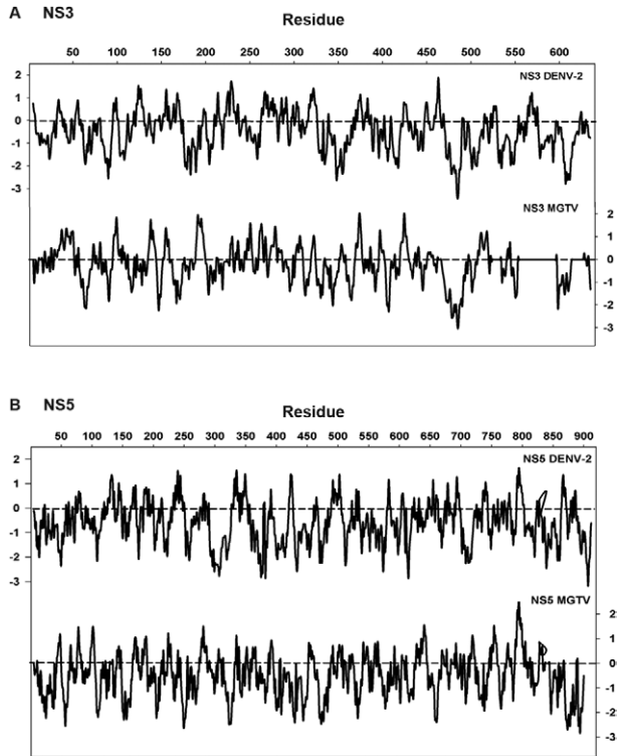


Fig. 4: hydropathy profile. Hydropathy values were determined by Kyte and Doolittle scale using ProScale web-based tool. Plot of non-structural (NS)3 (A) and NS5 (B) of Mogiana tick virus (MGTV). Both are compared with dengue virus type 2 (DENV-2) proteins. Negative scores mean a hydrophilic residue. Dashed line is the upper threshold for hydrophilicity.

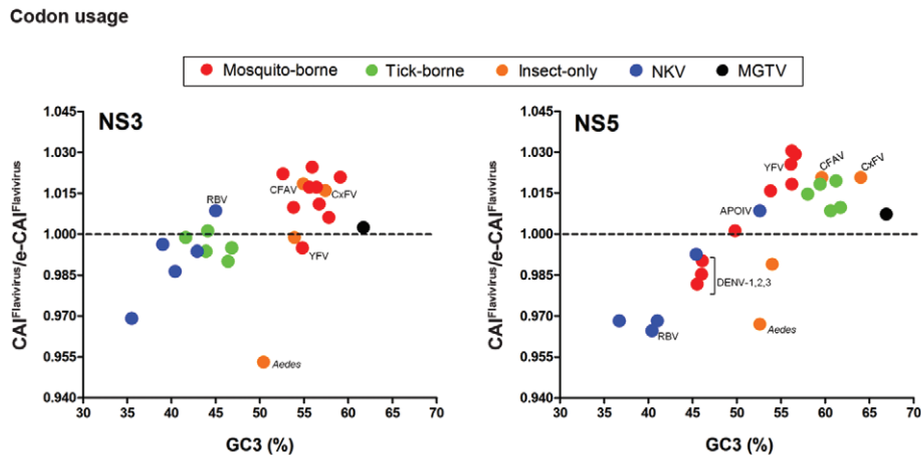


Fig. 5: codon usage pattern in Mogiana tick virus (MGTV) sequences. Codon bias in MGTV sequences were analysed with CAIcal Server. Codon usage adaptation index (CAI) calculation used Flavivirus sp. codon usage table as reference. The values were normalised with the expected CAI value (eCAI) for codon usage pattern in Flavivirus, calculated with input sequences. The dashed line indicates the threshold for which equal or higher values are interpreted as a codon usage adaptation towards Flavivirus codon usage pattern. APOIV: Apoi virus; CFAV: cell fusing agent virus; CxFV: *Culex flavivirus*; DENV: dengue virus; NKV: no known vector; NS: nonstructural; RBV: Rio Bravo virus; YFV: yellow fever virus.

C6/36), which suggested that the virus was a true arbovirus. The CAI and eCAI values of the NS3 and NS5 genes were calculated using the codon usage tables for *R. microplus*, *B. taurus* and *H. sapiens* and normalised CAI values were obtained for each condition (Table II). The three normalised CAI values, referring to the hosts, fluctuated for NS3 and were more stable for NS5. However, all three values were lower than 1.0, indicating a constrained codon usage adaptation towards a single host (neither a tick nor a vertebrate). This finding was theoretically consistent with what it would be expected for arboviruses. In conclusion, the codon usage analyses revealed characteristics of MGTV codon usage that are similar to those of flaviviruses.

Molecular detection of the NS3 and NS5 sequences from MGTV in field samples of ticks - The cattle tick *R. microplus* is broadly distributed in midwestern, southeastern and southern Brazil (Estrada-Pena 1999), where it causes heavy infestations in dairy cattle and is the main factor limiting animal health and production. We obtained a collection of tick samples from 19 dairy farms located in these three regions and searched for potentially novel viruses in these field samples through the molecular detection of partial sequences of NS3 (using primer set 317) and NS5 (using primer set 401). Different stages of the *R. microplus* life cycle were evaluated and both DNA and RNA from identical tick samples were used as templates for the reactions. Tick actin primers were used for positive control experiments. PCR amplifications using each set of primers (317, 401 and actin) were performed for each of the 81 gDNA and 81 cDNA samples. The agarose gel electrophoresis analysis revealed that there were no products of the PCR amplification of the gDNA templates, except from the reactions using actin primers (data not shown), as previously observed (Supplementary data, Fig. S1).

Interestingly, most of the analysed farms provided tick samples that were PCR-positive for the MGTV NS3 and/or NS5 sequences (Fig. 6). The analysed tick samples were collected from a large area in Brazil that includes the main cattle-producing regions (Fig. 6A). Most of the sampled farms received positive results for at least one tick life stage (Fig. 6B). The MGTV NS3 and NS5 fragments were detected in 42 samples, representing 51.8% of the total tick pools. Notably, the NS3 and/

or NS5 fragments were also detected in most of the UL samples. Because this is an unfed life stage, it is possible that the vertical transmission of this virus occurs in nature. The only tick samples in which the NS3 and NS5 fragments were not detected were from farms located in Presidente Prudente (SP) and Ribas do Rio Pardo, state of Mato Grosso do Sul (MS). Interestingly, although these regions are separated by the Paraná River, they are joined by a highway along which there are several abattoirs that receive carcasses from MS.

The positive samples (Fig. 6B) were sequenced to confirm that they corresponded to the same fragments found in the *R. microplus* SG cDNA library. The aa compositions of these samples were identical to those of contigs 317 and 401 (data not shown). However, we observed synonymous substitutions along the sequences, a common feature of RNA viruses due to the high mutation rates and consequently rapid evolution of RNA genomes (Holland et al. 1982). A high level of genetic diversity has been reported in populations of West Nile virus, a mosquito-borne flavivirus; this diversity confers a fitness benefit to mosquitoes (Jerzak et al. 2005, Fitzpatrick et al. 2010). To examine the diversity among our samples, phylogenetic trees were constructed for sequences 317 (NS3-derived) and 401 (NS5-derived) (Fig. 7). The sequences were distributed along numerous branches, demonstrating high sequence variability (nt substitutions) in the NS3 (Fig. 7A) and NS5 trees (Fig. 7B).

To confirm the origin of the samples, we highlighted several branches composed of samples from the same farm. Branch I of the NS3 tree (Fig. 7A) was formed by samples from farm Q (Santa Vitória, MG, Southeast Region) and branch I of the NS5 tree (Fig. 7B) was formed by samples from farm M [Piracanjuba, state of Goiás (GO), Central-West Region]. In both phylogenetic trees, branch II comprised samples from farm N (Itauçu, GO), whereas branches III and IV of the NS3 tree were formed by samples from farms T (Água Clara, MS) and H (South Region), respectively (Fig. 7A). Samples from farm X (Ribeirão Preto, Southeast Region) were grouped in both phylogenetic trees, on branches V and III of the NS3 and NS5 trees, respectively.

Final remarks - The original viral sequences described herein were detected in the SGs of female *R. microplus* ticks and in UL, suggesting a potential viral

TABLE II

Codon usage pattern observed for nonstructural (NS) genes in Mogiana tick virus and its relation with potential hosts

Gene	Length (bp)	Cattle tick			Bovine			Human		
		CAI _{Rm}	eCAI _{Rm} *	CAI _{Rm} /eCAI _{Rm}	CAI _{Bt}	eCAI _{Bt} *	CAI _{Bt} /eCAI _{Bt}	CAI _{Hs}	eCAI _{Hs} *	CAI _{Hs} /eCAI _{Hs}
NS3	1,662	0.740	0.776	0.95361	0.687	0.745	0.92215	0.729	0.781	0.93342
NS5	2,598	0.757	0.776	0.97551	0.724	0.745	0.97181	0.764	0.781	0.97823

codon usage tables of cattle tick, bovine and human were used as reference set for codon usage adaptation index (CAI) computation (asterisk means $p = 0.05$). eCAI: expected CAI value; *Rm*: *Rhipicephalus microplus*; *Bt*: *Bos taurus*; *Hs*: *Homo sapiens*.

transmission through saliva and the maintenance of a viral reservoir in ticks by vertical transmission. More pools of larvae should be analysed to confirm this possible mechanism of transmission because we could not isolate the virus from the UL, only from the female ticks. The vertical transmission of arboviruses in mosquitoes has been reported for mosquito-borne Flavivirus viruses, such as DENV (Rosen et al. 1983). This phenomenon has also been described in ixodid ticks; the best known case is the transovarial transmission of tick-borne encephalitis virus (Danielova et al. 2002). An interesting survival strategy of arboviruses in nature involves vertical transmission in the arthropod host so that the viruses are maintained in the environment even under adverse conditions. *R. microplus* is a monoxenic tick, i.e., it spends its entire cycle on a single host. Thus, it is essential that the UL are vertically infected to complete the parasite's life cycle, as is the case for Babesia protozoa transmission by *Rhipicephalus* spp (Guglielmone 1995).

Brazil has the largest number of commercial cattle worldwide. Other species of ticks also affect cattle, such as *Amblyomma* spp, which are heteroxenic ticks that

parasitise many different hosts, including humans. In a study concerning the potential transmission of tick-borne pathogens in Brazil, a Flavivirus was isolated from *Amblyomma cajennense* ticks (Figueiredo et al. 1999); however, this virus has not been further characterised. Therefore, increasing awareness about the possibilities of emerging zoonoses is important because other tick species, such as those of the *Amblyomma* genus, can infest cattle and then parasitise humans, occasionally infecting them with a new pathogen.

Our findings suggest that the potential novel virus (MGTV) may be an arbovirus belonging to Flavivirus genus because MGTV was able to infect both vertebrate and invertebrate cells and because viral sequences were found in tick SGs. The molecular characterisation of NS3 and NS5, the largest and most highly conserved flaviviral proteins, showed that MGTV is more closely related to the Flavivirus genus, although it is a highly divergent member of this genus. The complete genome sequence of this novel virus will be necessary to define its correct taxonomic position. Unexpectedly, the results of the deep sequencing of viral RNA purified from cul-

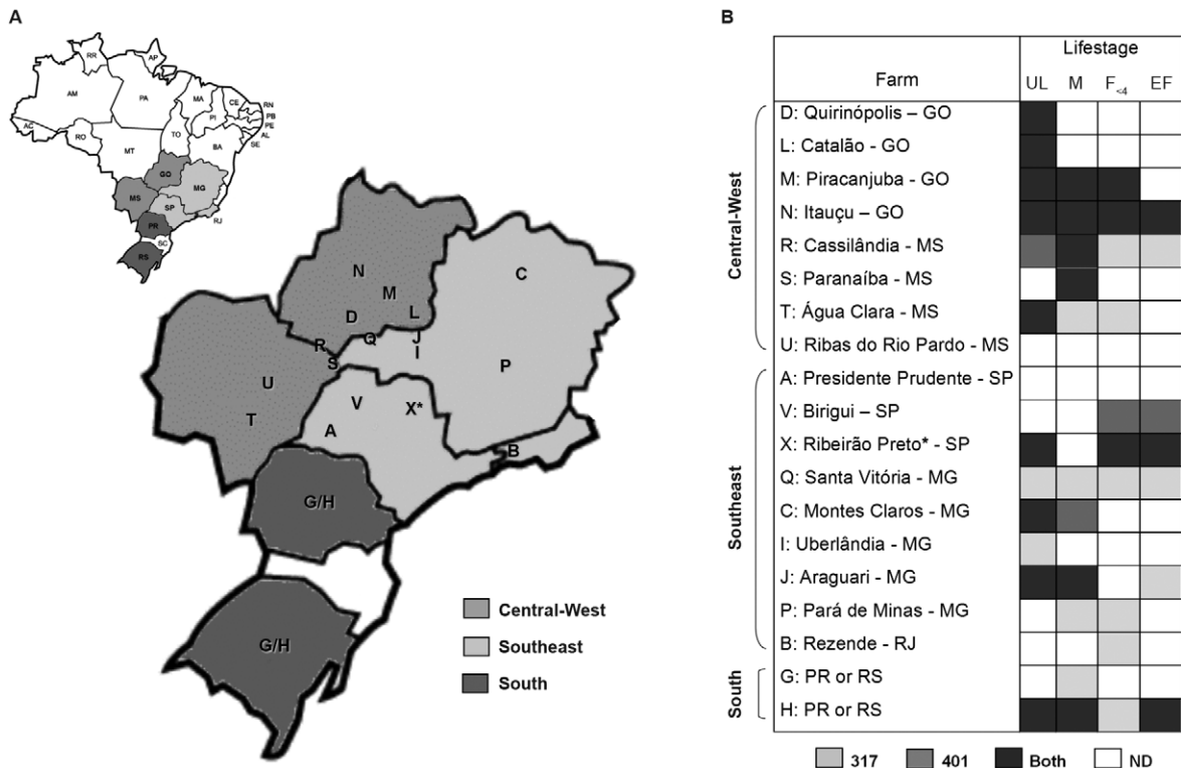


Fig. 6: molecular detection of viral transcripts in total RNA of tick samples collected in farms from seven Brazilian states. Detections were done through reverse-transcription-polymerase chain reaction using primers that target the 317 contig sequence [nonstructural (NS)3 fragment] or primers that target the 401 contig sequence (NS5 fragment). A: tick samples were collected from seven Brazilian states: Mato Grosso do Sul (MS) and Goiás (GO) (Central-West Region), São Paulo (SP), Minas Gerais (MG) and Rio de Janeiro (RJ) (Southeast Region), Paraná (PR) and Rio Grande do Sul (RS) (Southeast Region); B: panel for molecular detection linking farms to results for primers 317 (light grey), 401 (medium grey), both (dark grey) or no detection (ND) using any of the primers. Life stages of ticks: EF: engorged female (F); F₄: F less than 4 mm (before the rapid engorgement phase of feeding); M: male; UL: unfed larvae; *: samples from farm X (Ribeirão Preto, SP) were the only ones that presented positive detection when 2,743 primers were tested. Other Brazilian states: AC: Acre; AL: Alagoas; AP: Amapá; AM: Amazonas; BA: Bahia; CE: Ceará; DF: Distrito Federal; ES: Espírito Santo; MA: Maranhão; MT: Mato Grosso; PA: Pará; PB: Paraíba; PE: Pernambuco; PI: Piauí; RN: Rio Grande do Norte; RO: Rondônia; RR: Roraima; SC: Santa Catarina; SE: Sergipe; TO: Tocantins;

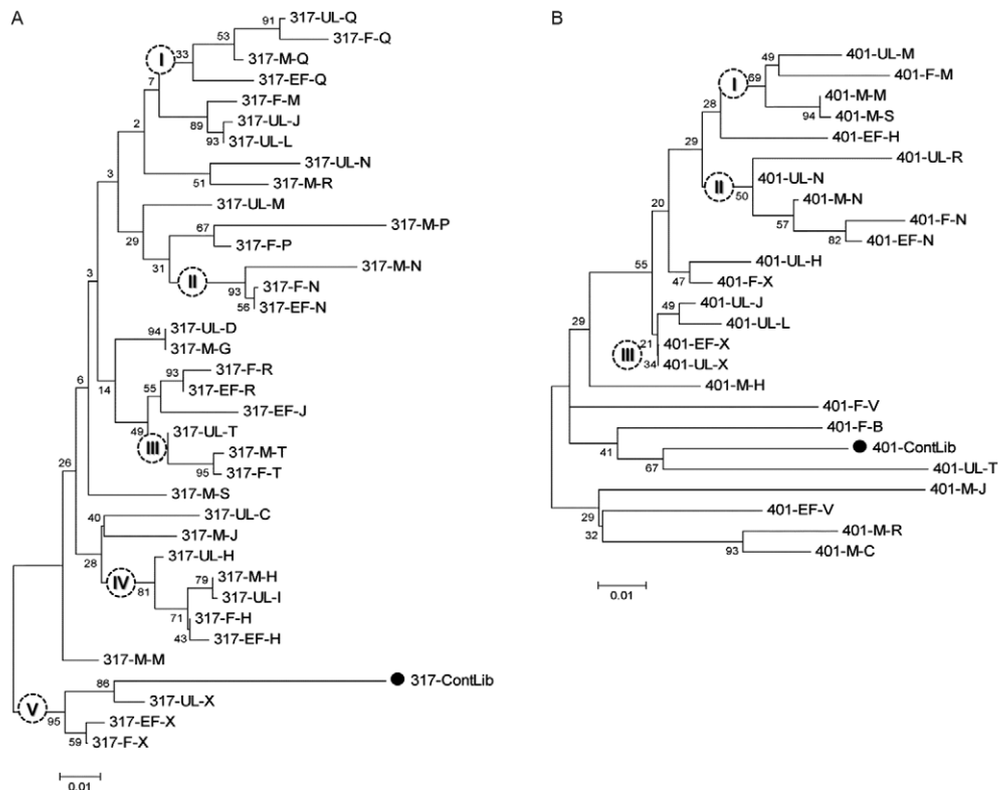


Fig. 7: phylogenetic trees of nonstructural (NS)3/NS5 sequences from positive tick samples. Sequences are named according to detection primers (317 or 401) followed by tick life stage [UL: unfed larvae; EF: engorged female (F) less than 4 mm; M: male] and the farm alphabetic label (see Table I for corresponding location). A: 36 sequences positive for detection using primers 317 (NS3 fragment) plus original contig sequence from the complementary DNA (cDNA) library (named ContLib highlighted in closed circle); B: 21 sequences positive for detection using primers 401 (NS5 fragment) plus original contig sequence from the cDNA library (named ContLib highlighted in closed circle); I-V: branches composed of samples from same farm. Neighbour-joining analysis was performed with 1,000 bootstrap replicates. The bar at the bottom indicates 1% nucleotide substitution.

tured infected cells did not provide the complete genome sequence of MGTV. This failure may have been caused by a technical artefact during the isolation and purification of the virus from the Vero cell culture, during the subsequent processing of the samples to obtain the viral RNA, or during the freezing and thawing prior to sequencing. Alternatively, this difficulty in obtaining the full genome sequence may reflect in inherent characteristics of this novel virus, such as the low VL observed. This virus is being further passaged to increase the VL.

The PCR assays as molecular detection of viral RNA fragments for molecular detection of viral RNA fragments for assigning viral infection has been debated. For example, Telis (2012) criticised a study by Bingham et al. (2012) that described the presence of the eastern equine encephalomyelitis virus in snakes using quantitative RT-PCR. Despite this controversy, our findings should be publicised due to their importance for Public Health authorities, who should be alerted to the possibility of emerging zoonoses and/or tick-borne viral diseases. In addition to the complete genome sequence, the potential transmission cycle and pathogenicity of MGTV have yet to be defined. Further research to characterise MGTV genotypically and phenotypically is important.

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