Sinu virus, a novel and divergent orthomyxovirus related to members of the genus *Thogotovirus* isolated from mosquitoes in Colombia

María Angélica Contreras-Gutiérreza,b, Marcio R.T. Nunesc, Hilda Guzmand, Sandra Uribeb, Juan Carlos Gallego Gómez, Juan David Suaza Vascoa,b, Jedson F. Cardosoa, Vsevolod L. Popovd, Steven G. Widenf, Thomas G. Woodf, Nikos Vasilakisd,⁎, Robert B. Teshd,⁎

a Programa de Estudio y Control de Enfermedades Tropicales – PECT – SIUSde de Investigación Universitaria – Universidad de Antioquia, Medellín, Colombia
b Grupo de Investigación en Sistemática Molecular-GSM, Facultad de Ciencias, Universidad Nacional de Colombia, sede Medellín, Medellín, Colombia
c Center for Technological Innovation, Evandro Chagas Institute, Ministry of Health, Ananindeua, Para, Brazil
d Department of Pathology, Institute for Human Infections and Immunity, University of Texas Medical Branch, Galveston, TX, USA
e Grupo de Investigación en Medicina Molecular y Traslacional– Universidad de Antioquia, Medellín, Colombia
f Department of Biochemistry and Molecular Biology, University of Texas Medical Branch, Galveston, TX, USA

ARTICLE INFO

Keywords:
Sinu virus
Orthomyxoviridae
Thogotovirus genus
Insect-specific viruses

ABSTRACT

The genome and structural organization of a novel insect-specific orthomyxovirus, designated Sinu virus, is described. Sinu virus (SINV) was isolated in cultures of C6/36 cells from a pool of mosquitoes collected in northwestern Colombia. The virus has six negative-sense ssRNA segments. Genetic analysis of each segment demonstrated the presence of six distinct ORFs encoding the following genes: PB2 (Segment 1), PB1 (Segment 2), PA protein (Segment 3), envelope GP gene (Segment 4), the NP (Segment 5), and M-like gene (Segment 6). Phylogenetically, SINV appears to be most closely related to viruses in the genus *Thogotovirus*.

1. Introduction

Arthropod-borne viruses (arboviruses) are viruses infecting vertebrates that are biologically transmitted by hematophagous arthropods in an alternative vertebrate-arthropod cycle (Bolling et al., 2015). Arboviruses replicate in both their vertebrate and arthropod hosts; and some are capable of causing major diseases outbreaks, such as dengue, yellow fever, Japanese encephalitis, Rift Valley fever and bluetongue, affecting both human and domestic animal populations.

During the past two decades with the advent of sequencing and metagenomics, a second group of viruses has been discovered in hematophagous insects that are now generally referred to as “insect-specific viruses” (ISVs) (Bolling et al., 2015; Blitvich and Firth, 2015; Junglen and Drosten, 2013). Although many of the ISVs are genetically related to some of the important vertebrate pathogenic arboviruses, they differ biologically from the true arboviruses by their inability to infect vertebrates or to replicate in vertebrate cells (Bolling et al., 2015). They are host-restricted to replication in invertebrate cells.

To date, most known ISVs have been associated with blood-feeding Diptera (mosquitoes, sandflies and midges) and relatively few have been described in Acari or other types of hematophagous arthropods (Bolling et al., 2015; Blitvich and Firth, 2015).

The purpose of this report is to describe and characterize a novel ISV orthomyxovirus, designated Sinu virus, that was isolated in C6/36 mosquito cells from a pool of adult mosquitoes collected in Cordoba Department on the Caribbean Coast of Colombia in 2013. SINV appears to be most closely related phylogenetically to the tick-borne viruses in the genus *Thogotovirus*.

2. Materials and methods

2.1. Study area

The study site was located in the municipality of San Bernardo del Viento (9° 21’ 30.97” N, 75° 58’ 37.28”W), Cordoba department, in northern Colombia on the Caribbean Coast. The area encompasses about 321 Km2 (Fig. 1) and is classified as lowland dry tropical forest with high mean annual temperature (29 °C) and relative humidity of 80%. The region has two contrasting seasons: a dry season from December to April and a rainy season from May until November (Climate-Data.org, 2016 http://en.climate-data.org/location/50073/).

Most of the territory is composed of plateau areas, with approximately...
34.2 km of beaches, which are used for ecotourism (Acosta, 2013; Rojas et al., 2010). The natural vegetation varies from intervened dry forests to flooded lands, including mangrove swamps and estuaries with halophytic vegetation (CVS, 2004; Serrano, 2004).

2.2. Mosquito collections

Mosquitoes from San Bernardo were collected, using CDC-light traps, during arbovirus field studies between 2012 and 2013. After collection, mosquitoes were killed by freezing and transported in liquid nitrogen to the University of Antioquia in Medellín, where they were stored at −70 °C. The mosquitoes pools were subsequently transported on dry ice to the University of Texas Medical Branch (UTMB) in Galveston for virus assay, following all national and Institutional regulations.

2.3. Processing mosquitoes for virus isolation

At UTMB, the mosquitoes were separated into pools of approximately 50 insects that were homogenized in plastic tubes, containing 1.0 mL of phosphate-buffered saline, pH 7.4, with 10% fetal bovine serum, 1% penicillin-streptomycin-amphotericin stock (Sigma), and a 3 mm stainless steel ball, using a TissueLyser (Quaigen). After centrifugation at 10,000 rpm for 10 mins, 100 μL of each mosquito homogenate was inoculated into a single well of a 24-multiwell tissue culture plate (Corning), containing a monolayer of Aedes albopictus C6/36 cells. After absorption for 2 h, the inoculum was aspirated and each flask was washed three times with 5 mL of maintenance medium. After the final wash, 500 μL of the medium was removed as a day 0 sample and was frozen at −80 °C for subsequent testing. Cultures were incubated at the temperatures noted above. Each day thereafter, for seven consecutive days, all of the medium was removed by aspiration for testing and 5 mL of fresh medium was added. Total RNA was subsequently extracted from all samples (day 0–7), using Trizol (Ambion) reagent and RNeasy Mini Kit (QIAGEN), following instructions of the manufacturer. The viral RNA extracted was assessed by reverse transcription (RT)-PCR assay using SINV-specific primers targeting the PB1 gene (primer sequences and PCR conditions available upon request).

2.4. Evaluation of CoB 38d (SINUV) growth in mosquito and vertebrate cells

In order to determine the host range of SINUV, a C6/36 cell stock of CoB 38d was inoculated into 12.5 cm² flask cultures of the following cell lines, obtained from the American Type Culture Collection (ATCC), Manassas, VA: Aedes albopictus clone C6/36 (CRL-1660); baby hamster kidney BHK-21 (CCL-10); African green monkey kidney Vero E6 (CRL-1586) and chicken embryonic fibroblast cells (CRL-12203). Cells were grown in 5 mL of culture medium, recommended in the ATCC specification sheets. C6/36 cells were incubated at 28 °C; Vero, baby hamster and chick embryonic cells were maintained at 37 °C. When a confluent monolayer of cells was present in the flasks, 200 μL of stock virus (CoB 38d) was inoculated into each flask. After incubation for 2 h, the inoculum was aspirated and each flask was washed three times with 5 mL of maintenance medium. After the final wash, 500 μL of the medium was removed as a day 0 sample and was frozen at −80 °C for subsequent testing. Cultures were incubated at the temperatures noted above. Each day thereafter, for seven consecutive days, all of the medium was removed by aspiration for testing and 5 mL of fresh medium was added. Total RNA was subsequently extracted from all samples (day 0–7), using Trizol (Ambion) reagent and RNeasy Mini Kit (QIAGEN), following instructions of the manufacturer. The viral RNA extracted was assessed by reverse transcription (RT)-PCR assay using SINUV-specific primers targeting the PB1 gene (primer sequences and PCR conditions available upon request).

2.5. Transmission electron microscopy

For ultrastructural analysis, infected C6/36 cells were fixed for at least 1hr in a mixture of 2.5% formaldehyde prepared from paraformaldehyde powder and 0.1% glutaraldehyde in 0.05 M cacodylate buffer (pH 7.3), to which 0.03% picric acid and 0.03% CaCl2 were added. The monolayers were washed in 0.1 M cacodylate buffer, and cells were scraped off and processed further as a pellet. The pellets were postfixed in 1% OsO4 in 0.1 M cacodylate buffer (pH 7.3) for 1 h, washed with distilled water, and en bloc stained with 2% aqueous uranyl acetate for 20 min at 60 °C. The pellets were dehydrated in ethanol, processed through propylene oxide, and embedded in Poly/Bed 812 (Polysciences, Warrington, PA). Ultrathin sections were cut on a Leica EM UC7 ultramicrotome (Leica Microsystems, Buffalo Grove, IL), stained with lead citrate, and examined in a Philips 201 transmis-
2.6. Titration of SINUV in mosquito cells

A SINUV stock, prepared in the C6/36 cells, was titrated in 24-well culture plates with monolayers of C6/36 cells. Serial 10-fold dilutions of the virus stock were prepared in phosphate buffered saline, pH 7.4 (PBS) with 10% fetal bovine serum (FBS). Then 100 µL of each dilution was inoculated into four microplate wells. After absorption at 28 °C for 2 h, 1.5 mL of maintenance medium was added to each well. Maintenance medium was minimal essential medium (Gibco), supplemented with 2% fetal bovine serum, 2% MEM non-essential amino acid solution (Sigma), 1% NaHCO₃ solution (1.5%) and 1% of a l-glutamine-penicillin-streptomycin 100X solution (Sigma). Microplate cultures were incubated at 28 °C in a 5% CO₂ atmosphere. Six days post-infection, mosquito cells from each well were spotted onto Cel-Line 12 well glass slides (Thermo Fisher Scientific) for examination by indirect fluorescent antibody technique (IFAT), as described before (Xiao et al., 2001). A standard IFAT was done, using a mouse hyperimmune polyclonal antibody prepared against Thogoto virus, obtained from the World Reference Center for Emerging Viruses and Arboviruses (WRCEVA). The tissue culture infectious dose (TCID₅₀) per mL was calculated, using the Reed–Muench method (Reed and Muench, 1938).

2.7. RNA extraction, construction of RNA library, sequence assembly and analysis

Viral RNA was extracted from culture fluid of C6/36 cells infected with isolate CoB 38d (SINUV) using Qiamp RNA mini kit (Qiagen) as previously described by (Vasilakis et al., 2014). The genome sequence of CoB 38d was obtained using parallel sequencing and an Illumina TruSeq RNA v2 Kit, following the manufacturer's protocol. Briefly, a highly efficient protocol was made including the rRNA removal (Ribo-Zero technology) followed by a rapid, ligation-free cDNA synthesis procedure for preparing directional RNA-sequencing libraries (ScriptSeq v2 technology) (Pease and Sookman, 2012). The data generated were assembled using a de novo strategy implemented in the Mira Software (Chevreux et al., 2004). The resulting contigs were visually inspected and sequences were annotated using the Geneious v. 9 and deposited in the GenBank database (http://www.ncbi.nlm.nih.gov/genbank/) (Table 1).

2.8. Genome characterization

The recovered genome for isolate CoB 38d was characterized, based on its genetic traits such as genome size, terminal regions, genome organization, potential Open Reading Frames (ORFs), encoded proteins, potential glycosylation sites, cysteine residues, and conserved motifs, using a set of applications available in the Geneious software v. 9 (Kearse et al., 2012) and the open source NetGlyce v.1.0 server (http://www.cbs.dtu.dk/services/NetNGlyc/).

2.9. Genetic variability

Nucleotide and deduced amino acid sequences of orthomyxoviruses closely related to the CoB 38d strain were obtained from the GenBank database (Tables 1 and 2). Regions referred to the ORFs were translated into amino acid sequences, using the Genious software v.9 to determine the best amino acid substitution model by using the ProtTest v3.4 software (Darriba et al., 2011). Alignments were visually checked and identity calculations were performed, using Geneious software v.9, as well as by the Boxplot analysis implemented in the R package v.1.14.4 (Maechler et al., 2016) and T test as previously described (Williamson et al., 1989).

2.10. Phylogenetic analysis and signal

Phylogenetic trees of the amino acid alignments were created, using the neighbor-joining method and maximum likelihood phylogenetic reconstructions with the RAxML v 8.1.21 hybrid version (Randomized Axelerated Maximum Likelihood) (Pfeiffer and Stamatakis, 2010). A bootstrap analysis was performed using 1,000 replicates (reliability value of 95%) and trees were visualized in the FigTree graphic viewer (Rambaut, 2008). In addition, Bayesian phylogenetic analysis was performed, using Beast v1.8 (Drummond et al., 2012), under the HKY+Γ+I model, estimating the evolutionary rate using both a strict and an uncorrelated log-normal relaxed clock model.

The phylogenetic signal of each RNA segment was assessed, using the maximum likelihood method (Guindon and Gascuel, 2003). The initial step in this phylogenetic exploration involved the alignment of genomes. This was followed by the reconstruction the phylogenetic signal determined for each 200 nt window, using the bootscanning sliding window approach implemented in the Simplot software. The phylogenetic signal and number of resolved trees for each of the three quartets were estimated in percentage values (Alcantara et al., 2009; Escoto-Delgadillo et al., 2008).

2.11. Reassortment analyses of CoB 38d and other orthomyxoviruses

Because viruses with segmented genomes have the potential for reassortment (Gomberg and Raming, 1986), nucleotide sequences for each RNA segment of the CoB 38d isolate were used to investigate potential genome reassortment events with other members of the Thogotovirus and Quaranjavirus genera, using the Simplot V. 3.5.1 software (Loé et al., 1999). Nucleotide sequence similarities are represented in percentage values. For each segment, values ranging from 80% to 100% indicate reassortment.

2.12. Molecular modeling and electrostatics maps of the CoB 38d GP protein

A BLASTP (Altschul et al., 1990) search against the PDB (Berman et al., 2000) database was used to identify suitable templates. Sequence alignments between target and template amino acid sequences were performed with PROMALS3D (Pei et al., 2008). Structural models were constructed, using Modeller v916 (Webb and Sali, 2014). A total of 100 models were constructed for each target. The best model, based on DOPE score (Shen and Sali, 2006), was selected for further evaluation with PROSA (Wiederstein and Sippl, 2007) and Ramachandran plots generated with PROCHECK (Laskowski et al., 1996). Pymol (http://www.pymol.org) was used to inspect the 3D structure models, calculate structural alignments and to generate electrostatic maps with APBS plugin (Baker et al., 2001). Geneious v9 was used to inspect sequence alignments.

3. Results

3.1. Virus Isolation and in vitro characterization

Virus strain CoB 38d was isolated from a single pool of 50 adult mosquitoes (species not identified) collected in June 2013 in northern Colombia. Based on the locality where the mosquito pool was collected, the virus has been designated Sinu virus (SINUV).

The original homogenate of mosquito pool CoB 38d produced CPE in a culture of C6/36 on the fifth day after inoculation. After passage a second time in C6/36 cells, a filtrate of the culture medium was inoculated into flask cultures of Vero and BHK cells, which were inoculated at 37 °C for 14 days. No CPE was observed. A portion (about 10–12 µL) of the second C6/36 culture material was also inoculated intracranially into a litter of newborn mice. No illness or death occurred in the pups after 14 days. Animal work was done at UTMB.
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under an approved IACUC protocol (ID# 9505045).

To determine if SINUV replication occurred in vertebrate cells without producing CPE, additional experiments were carried out in C6/36, Vero, BHK and chick embryonic cell cultures to assay for virus replication by RT-PCR. Samples of medium from the four SINUV-inoculated cell lines were collected from day 0 to day 7, as described in the Methods section. After RNA extraction, a partial region of the PBI gene of SINUV was amplified and run on gels with expected band size between 500 and 550 pb. SINUV RNA extracted from culture fluid from the C6/36 cells from days 0–7 post-inoculation (dpi) displayed strong bands on all days (data not shown). In contrast, extracted and amplified viral RNA from the Vero, BHK and chick cells showed decreasing intensity of the RNA bands from day 0 to day 3. On days 4–7, no bands were visible, indicating that SINUV did not replicate in the three vertebrate cell lines (data not shown).

3.2. Virion morphology

In ultrathin sections of infected C6/36 cells spherical virus particles 130–140 nm in diameter were found free in the cytosol. They had an envelope ~7 nm thick covered with a crown of projections ~14 nm long with ~15 nm periodicity (Fig. 2).

3.3. Titration of SINUV in mosquito cells

As CoB 38d virus did not cause illness in mice, it was not possible to produce a specific mouse immune ascitic fluid (MIAF) for use in serologic tests. However, in preliminary indirect fluorescent antibody tests (IFAT), it was observed that MIAFs produced against Thogoto virus, and to a lesser extent to Dhori virus (Sbrana et al., 2007), reacted

Table 2.
Details of the orthomyxovirus described based on NGS, used in this study.

<table>
<thead>
<tr>
<th>Putative Classification</th>
<th>Source Isolation</th>
<th>Putative arthropod host</th>
<th>Country of Isolation</th>
<th>Genbank Accession N° to PB1 sequence</th>
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<td>Culicoides imicola</td>
<td>Culicoides imicola</td>
<td>Senegal</td>
<td>Provided by autor (Temmam et al., 2016)</td>
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Fig. 2. Ultrastructure of CoB 38d virus in C6/36 cells. Enveloped spherical virions 130–140 nm in diameter with multiple spikes at the surface are localized free in the cytosol. Bar=100 nm.

Fig. 3. Indirect fluorescent antibody (IFA) assay. (a) CoB 38d antigen in infected C6/36 cells, as detected by indirect fluorescent antibody technique (IFAT), using a Thogoto virus hyperimmune mouse ascitic fluid (HMAF). (b) C6/36 cell control (uninfected) stained by IFAT, using the same HMAF.
with C6/36 cells infected with CoB 38d (Fig. 3). These observations confirmed the ultrastructural studies, indicating that CoB 38d was an orthomyxovirus, probably related to Thogoto virus. For this reason, a Thogoto virus MIAF, obtained from World Reference Center of Emerging Viruses and Arboviruses, was used in IFATs to determine the growth of CoB 38d in C6/36 cells. The TCID50 of the second passage stock of CoB 38d was calculated as 10^6 TCID50/mL.

3.4. Genetic characterization of SINUV

The complete CoB 38d virus genome consisted of six RNA segments. The genome of the virus was determined to be 10,833 nucleotides (nt) in length, including the 5′ and 3′ UTRs. The length of the segments ranged from 1009 nt (segment 1) to 2376 nt (Segment 6) (Table 3). Highly conserved terminal sequences for CoB 38d were determined to be: 5´ AGTATTAACAAGAG[A/C/G]TTTC and 3´ AAAAA[T/G/C][C/T] [T/C] CTTTGTACTACTACC. Each segment encodes for a structural protein with related functional properties as follows:

3.5. Segments 1 to 3: polymerase proteins (PA, PB1, PB2)

CoB 38d RNA segments 1–3 ranged from 2,376 to 2,000 nt in length, and encode the proteins of a replicative polymerase complex, based on their identity to the influenza virus proteins. The length open reading frame (ORF) of the segment 1 was shown to be 2,298 nucleotides in length, encoding a protein of 765 amino-acid (aa). The largest segment of SINUV shows sequence homology to orthomyxoviral Polymerase basic protein 2 or PB2 gene sequence (Pfam Id PF00604; “Flu_PB2”) (Table 3). ORF of of SINUV segment 2 is 2,133 nt long (710 aa) and corresponds to orthomyxoviral Polymerase basic protein 1 or PB1 sequence (Pfam Id PF00602; “Flu_PB1”) (Table 3).

ORF encoded by segment 3 of SINUV is 1,911 nt long and encodes a 636 aa of Flu-PA protein (Polymerase acidic protein or PA) (Pfam Id PF00603; “Flu_PA”) (Table 3).

3.6. Segment 4: Glycoprotein

ORF of segment 4 is 1,497 nt long and encodes a 498 aa. The putative glycoprotein (GP) of SINUV displayed conservation with the “baculovirus gp64 envelope glycoprotein family” (Pfam Id PF03273; “Baculo_gp64”) (Fig. 4, Table 3).

The GP amino acid sequence of SINUV displayed conserved domains, which were identified as domain I (aa position 36–199), II (aa positions 21–35 and 200–267), III (271–345), IV (400–448) and V (408–427). Fusion peptides were determined as YSYQGGPLDPNTGDIE (L2) and NNNHFA (L1). A total of 16 Cys residues and 7 potential N-Glyc sites were found over the GP amino acid sequence. Supplementary Fig. 1 summarizes the conserved domains, cysteine residues, and potential glycosylation sites for SINUV and other Thogotovirus members.

3.7. Segment 5: Nucleoprotein

ORF encoded by segment 5 is 1,422 nt long and encodes a 473 aa. The open reading frame (ORF) encoded by segment 5 of CoB 38d virus is conserved with respect to flu NP-like protein (Pfam Id: SSF161003; “flu NP-like”) (Table 3).

3.8. Segment 6: M-like THOV protein

The smallest RNA segment of CoB 38d virus genome encodes a ORF with 816 nt long (271aa). This segment encodes a undescribed protein of unknown function by InterProScan. Analysis using the translated nucleotide-protein Blast X program (https://blast.ncbi.nlm.nih.gov/Blast.cgi) showed low percentage of aa similarity (23%) with the M
3.9. Genetic variability

Amino acid and nucleotide alignment analysis of individual segments of SINUV were variable and demonstrated a relative similarity with members of the genus Thogotovirus, family Orthomyxoviridae based on PB1, PA, PB2 and GP sequences (Table 4), as follows:

Segment 1 (PB2) of SINUV shares a 26.1–29.3% amino acid identity (39.8–42.7% nt identity) with Dhori virus (DHOV), Bourbon virus (BRBV) and Thogoto virus (THOV). A much higher percentage of identity was displayed in the PB1 segment, which shares 54.8–55.3% amino acid identity (55.2–56.8% nt identity) with THOV and DHOV. The PA segment shares 31.8–35.5% amino acid identity (36.2–36.4% nt identity) with DHOV, BRBV, Batken virus (BKNV) and THOV (Table 4). The GP of SINUV shares 29.3–29.8% amino acid identity to DHOV, BRBV and BKNV, as well as with Upolu, Aransas Bay and Jos viruses (Table 4). The NP of CoB 38d shares 32.4–43.8% amino acid identity to the NP of Thogotovirus genus (Table 3 and 4). Segment 6 of CoB 38d shares at a low percentage of amino acid identity (18.8% to 20.7.0%) with the putative matrix (M) of THOV (Table 3 and 4).

3.10. Boxplot analysis

Genetic distances found among members of the genus Thogotovirus were significantly smaller than the values obtained between members other orthomyxovirus genera. The SINUV sequence showed 60% mean genetic variability (p values < 0.05, overall p value=0.0003567) within all genera in the family Orthomyxoviridae (Supplementary Fig. 2, supplementary Table 1). Interestingly, genetic distances within the genera Quaranjavirus and Thogovirus showed relative high genetic variability with mean values of 50% and 37% respectively, in comparison with other orthomyxoviruses (Supplementary Fig. 2, supplementary Table 1).

3.10.1. Phylogenetic analysis

Phylogenetic analysis, using the protein sequences for all segments (1–6) of SINUV in comparison to other orthomyxoviruses, displayed a similar topology. Phylogenetic relationships showed that CoB 38d is a new member of the family Orthomyxoviridae, grouping in a monophyletic cluster distinct from other orthomyxoviruses described previously (Fig. 5a–f). Based on the high bootstrap supports (98–100%) in the phylogenetic trees, SINUV clearly clusters with members of the Thogotovirus genus (Fig. 5a–f). Importantly, SINUV consistently placed in a basal position to the Thogotovirus genus clade, suggesting that this virus diverged at an earlier time.

Phylogenetic signals for each RNA segment (in nucleotides) were assessed and determined in overall percentage values in comparison to other members of the family Orthomyxoviridae (Quaranjavirus, Influenzavirus A, Influenzavirus B, Influenzavirus C, Influenzavirus D, and Isavirus genera) with high bootstrap values for NJ (98%) and ML trees (100%). Values of overall percentages under the three quartets and percentages of resolved trees over each genomic segment are summarized in supplementary material (Table S2, Supplementary Fig. 3).

3.11. Molecular modeling and electrostatic maps of the SINUV GP protein

The stereochemical quality of the generated models presented ~90% of their residues in the most favored regions of the plot. The overall quality of the models was also evaluated using z-scores obtained from ProSA-web server. The calculated z-scores were negative and in the range of −5.02 to −6.84; these values are in agreement with good quality structures and with the z-score found for the template (−5.32) (Table S3, Supplementary Fig. 4). The 3D models were also aligned with the selected template. The domains I, III and V were highly conserved in all proteins, with an average R.M.S.D of 0.67 Å, 2.41 Å and 0.21 Å, respectively. On the other hand, domain II was less conserved, with an average R.M.S.D of 6.13 Å. Notably, some of the beta-sheets on the interface of domain II and III became less structured in envDhori (Fig. 6A). While domain II of envSINU and envTHogoto have two disulphide bridges stabilizing antiparallel beta sheets, envDhori lacks one disulphide bridge, explaining its less ordered structure (Fig. 6B). The predicted structures and electrostatic maps of SINUV, THOV and DHOV glycoproteins are displayed in Fig. 6C and D.

4. Discussion

Based on its genome sequences and sizes, number of segments, and genetic characteristics (ORFs, and 5’ –3’ termini), SINUV is similar to members of the family Orthomyxoviridae. The SINUV genome is characterized by six negative-sense, single-stranded RNA segments with translated functional ORFs. The similar genes, proteins and functions (S1 to S3 related to the polymerase subunits PB1, PB2 and PA proteins; S4 encoding for the glycoprotein; S5 encoding the nucleoprotein, and S6 encoding the M-like), all support the inclusion of SINUV within the family Orthomyxoviridae (Fig. 4, Table 3).

In order to confirm the genetic relationship of SINUV to other orthomyxoviruses, we performed a phylogenetic reconstruction, based on all complete ORFs, that confirms the multiple alignment and box plot approaches. Regardless of the analyzed RNA segment analyzed, the inferred phylogenetic trees depicted SINUV as a separate clade most closely related to members of the Thogotovirus genus (Fig. 5a–f). Importantly, SINUV was consistently placed at a basal phylogenetic relationship to all recognized species of the genus isolated from various

![Fig. 4. Genome structure of the isolate CoB 38d. The six RNA segments are shown in positive-sense orientation (5'–3'). Numbers indicates the nucleotide positions along the segments. Black line represents the start (ATG) and stop (TGA/TAA) codons respectively.](image-url)
Fig. 5. Phylogenetic analysis based on (a) PB2, (b) PB1, (c) PA, (d) GP, (e) NP and (f) M-like proteins based of full length ORFs. Values over each node of the tree correspond to the bootstrap support values for NJ and values into () correspond to mL. Bayesian posterior probability values are placed within parenthesis. Number over the bars represents the number of amino acid substitutions/site. Scale bar is representing the amino acid substitution/site.
arthropods and vertebrates including humans. Combining all results obtained for the current analyses (genetic divergence and phylogeny) we conclude that the unique clade position, as well as the genetic similarity (protein similarity) of less than 35% for all encoded proteins indicate that SINUV represents a unique member within the family Orthomyxoviridae. To this date SINUV is the only known member of this genus to have a host-restricted range. Given that the observed phylogenetic reconstructions do not show a deep node distance to the rest of the genus members, at this time, it is impossible to reconstruct or identify ancestral host switching processes, as well as to infer the direction of the virus spread to new hosts. Understandably the genetic diversity of this genus will be greatly enriched with future virus discovery studies in various arthropods and should provide valuable information on the properties of new divergent viruses, such as host range restriction to insects and deconstruction of their evolutionary ancestry and pathogenic potential.

Segment 6 of SINUV contains a predicted open reading frame of 271 aa which does not share sequence homology with any known protein, using protein databases (e.g. RCSB PDB, ViPR, Uniprot); but it displays a low amino acid identity of 23% with the M protein of THOV. Further studies are necessary to know the role of this segment of SINUV. Differences in the conserved segment termini are also compatible with a significant evolutionary distance of SINUV from other members of the Thogotovirus genus.

Although the current International Committee on Taxonomy of Viruses (2015) (ICTV) (www.ictvonline.org/virustaxonomy.asp) only lists two species within the genus Thogotovirus, there are now actually eight viruses within the genus: Thogoto (THOV), Dhori (DHOV), Batken (BKNV), Upolu (UPOV), Aransas Bay (ABV), Jos (JOSV), Bourbon (BRBV) and Dielmo orthomyxovirus (DOV) (Allison et al., 2015, Bussetti et al., 2012, Lvov et al., 2015, Briese et al., 2014, Lambert et al., 2015, Temmam et al., 2016). The addition of SINUV to the genus will bring the total number of thogotoviruses to nine. Most of the thogotoviruses have been associated with ticks, although Batken virus has been isolated from both ticks and mosquitoes (Lvov et al., 1974), Sinu virus was isolated from mosquitoes and Dielmo orthomyxovirus was found in Culicoides midges (Temmam et al., 2016).

Seven of the nine thogotoviruses (THOV, DHOV, BKNV, UPOV, ABV, JOSV, and BRBV) replicate in vertebrate (Vero or BHK) cells and cause illness and death in newborn mice (CDC, 2016; Lambert et al., 2015). SINUV does not, and only grew in C6/36 cells. The status of DOV is unknown, since it was identified by next-generation sequencing and not isolated. DHOV and THOV have been associated with disease (encephalitis and febrile illness) in humans (CDC, 2016), and THOV has caused abortion in sheep (Davies et al., 1984). BRBV has also been

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**Fig. 6.** Three dimensional structure of the glycoprotein (GP). (a1) DHOV; (a2) isolate CoB 38d and (a3) THOV. (b) Fusion peptide domain in detail for each virus respectively. (c) top and vertical views of the trimmer 3D structures and electrostatic charges (1a,1b,1c,2a,2b,2c) distribution in the structure (red areas = negatively charged; white areas = neutral charge; blue areas = positively charged). (d) Alignment of the fusion peptides of DHOV, THOV and SINUV.
associated with febrile illness and death in humans (Kosoy et al., 2015). The disease potential of the other thogotoviruses, including SINUV, for humans and animals is unknown. With the exception of DOV, all of the other eight thogotoviruses were initially detected by culture (in vitro) in vertebrate or insect cell lines or by inoculation of newborn mice. However, given the tremendous number and diversity of arthropods on Earth and the increasing use of metagenomics to search for novel virus genomes in all life forms (Li et al., 2015, Temmam et al., 2016, Shi et al., 2016), it can be assumed that other new thogotoviruses (and quaranjaviruses) will be discovered in non-hematophagous arthropods and other vertebrates.

Acknowledgements and support

MAC is supported by a Ph.D scholarship from the Colombian Department of Science (Colciencias Convocatoria 567). The fieldwork in Colombia was supported in part by Colciencias grant 111549326198. Laboratory work in the USA was funded in part by NIH grant R24 AI120942. We also acknowledge the help of Sandro Patroca da Silva in doing the Boxplot analysis and Richard Hoyos for mosquito collections.

Appendix A. Supporting information

Supplementary data associated with this article can be found in the online version at doi:10.1016/j.virol.2016.11.014.

References