

Phylogenetic analysis of Bunyamwera and Ngari viruses (family Bunyaviridae, genus *Orthobunyavirus*) isolated in Kenya

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Summary

Orthobunyaviruses, tri-segmented, negative-sense RNA viruses, have long been associated with mild to severe human disease in Africa, but not haemorrhagic fever. However, during a Rift Valley fever outbreak in East Africa in 1997–1998, Ngari virus was isolated from two patients and antibody detected in several others with haemorrhagic fever. The isolates were used to identify Ngari virus as a natural *Orthobunyavirus* reassortant. Despite their potential to reassort and cause severe human disease, characterization of orthobunyaviruses is hampered by paucity of genetic sequences. Our objective was to obtain complete gene sequences of two Bunyamwera virus and three Ngari virus isolates from recent surveys in Kenya and to determine their phylogenetic positioning within the Bunyamwera serogroup. Newly sequenced Kenyan Bunyamwera virus isolates clustered closest to a Bunyamwera virus isolate from the same locality and a Central African Republic isolate indicating that similar strains may be circulating regionally. Recent Kenyan Ngari isolates were closest to the Ngari isolates associated with the 1997–1998 haemorrhagic fever outbreak. We observed a temporal/geographical relationship among Ngari isolates in all three gene segments suggesting a geographical/temporal association with genetic diversity. These sequences in addition to earlier sequences can be used for future analyses of this neglected but potentially deadly group of viruses.

Key words: Arboviruses, bunyaviruses, viral haemorrhagic fever, virus infection, zoonoses.

Introduction

The Bunyaviridae family is divided into five genera; *Orthobunyavirus*, *Phlebovirus*, *Nairovirus*, *Hantavirus* and *Tospovirus* [1]. The largest genus, *Orthobunyavirus*, is composed of over 150 viruses that infect humans and

are transmitted by mosquitoes, midges and ticks from reservoir animals like rodents and livestock. Members of *Orthobunyavirus* are tri-segmented, negative-sense RNA viruses responsible for mild to severe human and animal diseases. The L (large) segment encodes a large protein that consists of the RNA-dependent RNA polymerase activity for replication and transcription of genomic RNA segments. The M (medium) segment encodes a precursor polypeptide which yields the virion surface glycoproteins Gn and Gc and a non-structural protein NSm, and the S (small) segment

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encodes the nucleocapsid (NC) and a non-structural protein (NSs) in overlapping reading frames [2]. Bunyamwera virus is the prototype virus of the *Orthobunyavirus* genus as well as the Bunyaviridae family of arboviruses. Bunyamwera virus is associated with febrile illness, headache, arthralgia, rash and infrequent central nervous system involvement [3]. Viruses of the *Orthobunyavirus* genus were generally not associated with haemorrhagic symptoms until Ngari virus was implicated in haemorrhagic fever outbreaks in Kenya and Somalia, and retrospectively in Sudan [4–6]. Ngari virus was determined to be a reassortment between two segmented viruses (Bunyamwera and Batai) co-circulating within the same environment. Subsequent analysis of isolated sequences from outbreak samples showed that the L and S segment sequences closely matched those of Bunyamwera virus while the M segment was identical to that of Batai virus [5].

Surveillance activities during Rift Valley fever outbreaks have demonstrated co-circulation of arboviruses including Bunyamwera virus [7]. Additionally, ongoing inter-epidemic surveillance activities have indicated continued intense transmission of Bunyamwera virus [8]. Co-circulation of viruses within the same serogroup is likely to provide opportunities for genetic reassortments. Efforts to bring these infections under control and predict their emergence will not be successful unless there is a full understanding of how these viruses are maintained and transmitted within the environment. However, characterization of these emergent arboviral species has been hampered by paucity of genetic sequences making it impossible to accurately estimate their evolutionary trend and public health burden. We have recently isolated Bunyamwera and Ngari viruses from surveillance exercises in Kenya and identified them based on short diagnostic sequences [8, 9]. Our main objective was to provide complete coding sequence of some of these isolates as well as to investigate their phylogenetic positioning within the Bunyamwera serogroup, based on partial and complete genome sequences of similar virus strains and other selected orthobunyaviruses.

Methods

Bunyamwera and Ngari virus isolates (Table 1) obtained from previous surveillance exercises in northern Kenya were inoculated into flasks containing confluent Vero cells and observed daily for cytopathic effects (CPE). Viruses were harvested when more than 75% of cells showed CPE. The supernatants were aliquoted into cryotubes and stored at -70 °C, until use.

For RNA extraction, the MagNA Pure LC RNA isolation kit I (Roche Applied Science, USA) was used. cDNA was synthesized using the Transcriptor first-strand cDNA synthesis kit (Roche Applied Science) with random hexamers followed by PCR using Phusion high-fidelity PCR kit (Finnzyme Ltd, Finland) and appropriate primers. Overlapping primers (Supplementary Table S1) for each segment were either designed based on sequences of Bunyamwera, Batai and Ngari viruses available in GenBank or obtained from previous publications [10, 11]. Amplified DNA fragments were visualized by electrophoresis on a 1.5% agarose gel. Amplified DNA fragments were purified and prepared for sequencing using ExoSAP-IT PCR clean-up kit (USB Corp., USA) according to manufacturer's instructions and stored at -20 °C.

Sequencing was performed using sets of overlapping primers for the S, M and L segments as designed previously (Supplementary Table S1) using the Big Dye v. 3.1 kit (Applied Biosystems, USA) and run on the 3500XL Genetic Analyzer (Applied Biosystems). The sequences obtained were cleaned and edited using Bioedit software (USA) for both reads from the forward and reverse primers. Sequences obtained were subjected to Basic Local Alignment Search Tool (BLAST) searches in NCBI GenBank (<http://www.ncbi.nlm.nih.gov/blast/Blast>) to identify similar sequences. The clean sequences of each segment were aligned against a selection of corresponding segment sequences of Bunyamwera serogroup viruses, including Bunyamwera, Batai and Ngari viruses, using the multiple alignment fast Fourier transform (MAFFT) sequence alignment program [12]. Phylogenetic trees were constructed using maximum likelihood algorithm and analysed with 1000 replicates for bootstrap testing in Molecular Evolutionary Genetics Analysis (MEGA) v. 5.20 software [13]. La Crosse virus of the California serogroup was used as the out-group for phylogenetic analyses of all three segments of Bunyamwera and Ngari viruses. Nucleotide and amino acid similarities and genetic distances between the Kenyan isolates and selected isolates from diverse regions were computed in MEGA v. 5.20 using the p-distance method [13].

Results and Discussion

The complete S, M and L segment coding regions (704, 4302 and 6717 nt, respectively) were sequenced for all five isolates. For this analysis, we excluded the 3' and 5' non-coding regions. The gene sequences

Table 1. *Bunyamwera and Ngari virus isolates from Kenya sequenced in the study*

Virus	Code	Strain	Site of isolation	Year	Isolation source
Bunyamwera	BUNV_11232	BUNV_GSA/S4/11232_WT	Garissa	2009	<i>Aedes mcintoshi</i>
	BUNV_12060	BUNV_MGD/S1/12060_WT	Magadi	2009	<i>Anopheles funestus</i>
Ngari	NRIV_19801	NRIV_TND/S1/19801_WT	Tana-delta	2011	<i>Anopheles funestus</i>
	NRIV_5170	NRIV_GSA/TS7/5170_WT	Garissa	2009	<i>Amblyomma gemma</i>
	NRIV_5242	NRIV_ISL/TS2/5242_WT	Isiolo	2009	<i>Rhipicephalus pulchellus</i>

determined in this study are deposited in GenBank under the following accession numbers (S segment, M segment, L segment):

KM507344, KM507340 and KM507338 (BUNV_MGD_S1_12060_WT);

KM507345, KM507339 and KM507337 (BUNV_GSA_S4_11232_WT);

KM507343, KM514679 and KM507335 (NRIV_TND_S1_19801_WT);

KM507341, KM514677 and KM507336 (NRIV_GSA_TS7_5170_WT);

KM507342, KM514678 and KM507334 (NRIV_ISL_TS2_5242_WT).

Phylogenetic analyses reveal that within all three genomic segments, Bunyamwera virus isolates sequenced in the current study clustered together with an isolate in GenBank from the same locality (BUNV_AMH001130) with strong bootstrap support. The newly sequenced Kenyan Bunyamwera virus isolates possess high nucleotide sequence identities of over 99.0% (S), 99.4% (M) and 99.4% (L) compared to the previous Kenyan isolate (BUNV_AMH001130). Similarly, the newly sequenced Kenyan Bunyamwera virus isolates possess high percentage amino acid sequence identities of over 99.5% (N ORF), 99.0% (NSs ORF), 99.5% (M polyprotein ORF) and 98.8% (L protein ORF) compared to the previous Kenyan isolate. These results indicate that similar Bunyamwera virus strains may be circulating within Kenya and also regionally considering the close association with the 1994 Bunyamwera virus isolate from the Central African Republic (BUNV_ArB29051).

Similarly for Ngari virus, the newly sequenced Kenyan isolates, regardless of the genomic segment, clustered with other Ngari virus isolates (Fig. 1a–c). The newly sequenced Kenyan Ngari virus isolates were closer to Ngari virus isolates associated with the 1997–1998 haemorrhagic fever outbreak in East Africa (NRIV_9800535 and NRIV_9800521) with

strong bootstrap support particularly in analyses of the M and L segments (Fig. 1b, c). This observation suggests that the current Kenyan Ngari virus isolates may have been introduced into Kenya from the Kenya–Somali border. The newly sequenced Kenyan Ngari virus isolates also possess high nucleotide sequence identities of over 98.5% (S), 97.0% (M) and 97.3% (L) compared to other Ngari virus strains regardless of year and place of isolation. Similarly, the newly sequenced Kenyan Ngari virus strains possess high amino acid sequence identities of over 99.1% (N ORF), 100% (NSs ORF), 98.4% (M polyprotein ORF) and 99.1% (L protein) compared to other Ngari virus isolates. The complete sequence data for the three additional isolates of Ngari virus support previous findings that this virus is a reassortant [4–6].

Of additional interest, we observed temporal/geographical clustering of Ngari isolates in all three phylogenetic trees showing that isolates from the same region or outbreak year generally group together within their own strongly supported independent groupings (Fig. 1a–c). Our findings suggest a geographical/temporal association with genetic diversity between strains of Ngari virus. Similar observations have been made for Batai virus in which correlation between geographical and genetic diversity has been suggested [10, 14]. Batai virus isolates from Europe, Asia and Africa generally cluster independently within strongly supported groups with country-specific viruses clustering closest [14]. However, a more in-depth analysis including identification of signature motifs that might be representative of a given geographical/temporal clade of these viruses is restricted by paucity of genetic sequences in GenBank. Furthermore, there is need for more studies on pathogenesis and distribution of these viruses. While public health efforts have focused on well-characterized viruses such as Rift Valley Fever, West Nile, chikungunya and dengue viruses, the emergence of orthobunyaviruses such as Ngari and Schmollenberg viruses as human and veterinary pathogens, respectively, emphasize the need for in-depth characterization and determination of their true public health impact.

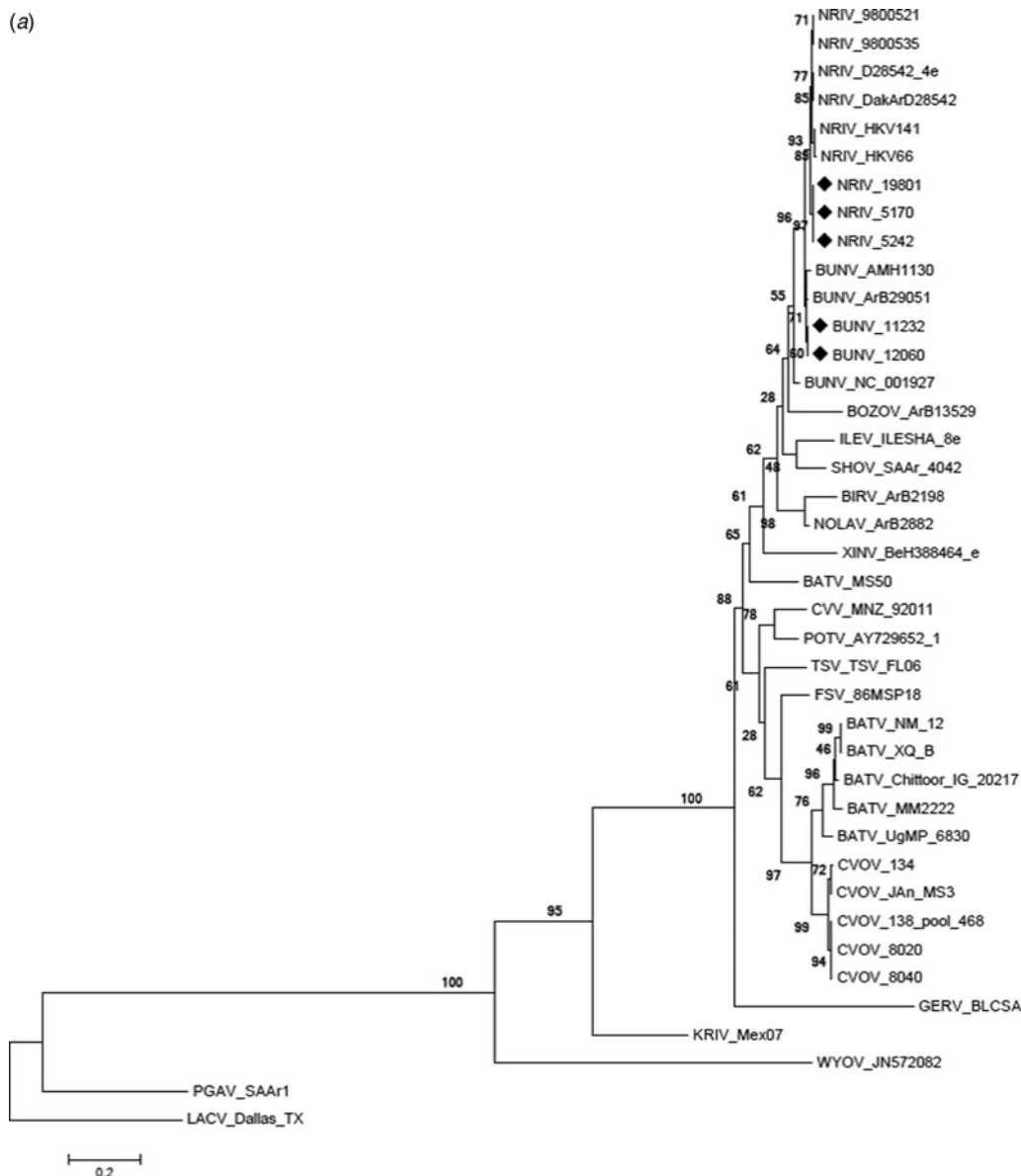


Fig. 1. Phylogenies of the S, M and L segments of the genus *Orthobunyavirus*. According to a limited diversity of data that are available in GenBank, the maximum likelihood method was used to derive phylogenies from (a) the N gene nucleotide ORF (S segment, 702 nt), (b) M polyprotein ORF (M segment, 4302 nt) and (c) L polymerase protein ORF (L segment, 6717 nt) and ORFs of selected orthobunyaviruses. Phylogenies were rooted using the La Crosse virus sequence. Bootstrap values that were determined by 1000 replicates are shown. Scale bars represent the number of nucleotide substitutions per site. Sequences determined as a part of this study are indicated by a rectangle. All *Orthobunyavirus* abbreviations are used according to the International Committee on Taxonomy of Viruses (ICTV, 2005). BATV, Batai virus; BIRV, Biraó virus; BOZOV, Bozo virus; BUNV, Bunyamwera virus; CVV, Cache Valley virus; CVOV, Calovo virus; FSV, Fort Sherman virus; GERV, Germiston virus; ILEV, Ilesha virus; KIRV, Kairi virus; LACV, La Crosse virus; NDOV, Nyando virus; NOLAV, Nola virus; NRIV, Ngari virus; PGAV, Pongola virus; POTV, Potosi virus; SHOV, Shokwe virus; TSV, Tensaw virus; WYOV, Wyomyia virus; XINV, Xingu virus. *S segment accession numbers*: BATV_Chittoor_IG_20217 (JX846598), BATV_MM2222 (JX846595), BATV_MS50 (JX846604), BATV_NM_12 (KJ187040), BATV_UgMP_6830 (JX846601), BATV_XQ_B (KJ398936), BIRV_ArB2198 (AM711131), BOZOV_ArB13529 (AM711132), BUNV_ArB29051 (AM709778), BUNV_NC001927, BUNV_AMH1130 (JF961342), CVOV_134 (KJ542624), CVOV_138_pool_468 (KC608157), CVOV_8020 (KJ542630), CVOV_8040 (KJ542633), CVOV_JAn_MS3 (KJ542627), CVV_MNZ_92011 (KC436108), FSV_86MSP18 (EU564829), GERV_BLCSA (M19420), ILEV_ILESHA_8e (KC608151), KRIV_Mex07 (EU879063), LACV_Dallas_TX_2009 (GU591164), NOLAV_ArB2882 (AM711134), NRIV_9800521 (JX857325), NRIV_9800535 (JX857328), NRIV_D28542_4e (KC608154), NRIV_DakArD28542 (JX857316), NRIV_HKV141 (JX857322), NRIV_HKV66 (JX857319), NRIV_Adrar (KJ716848), PGAV_SAAr1 (EU564828), POTV_AY729652_1 (AY729652), SHOV_SAAr_4042 (EU564831), TSV_TSV_FL06 (FJ943507), XINV_BeH388464_e (EU564830).

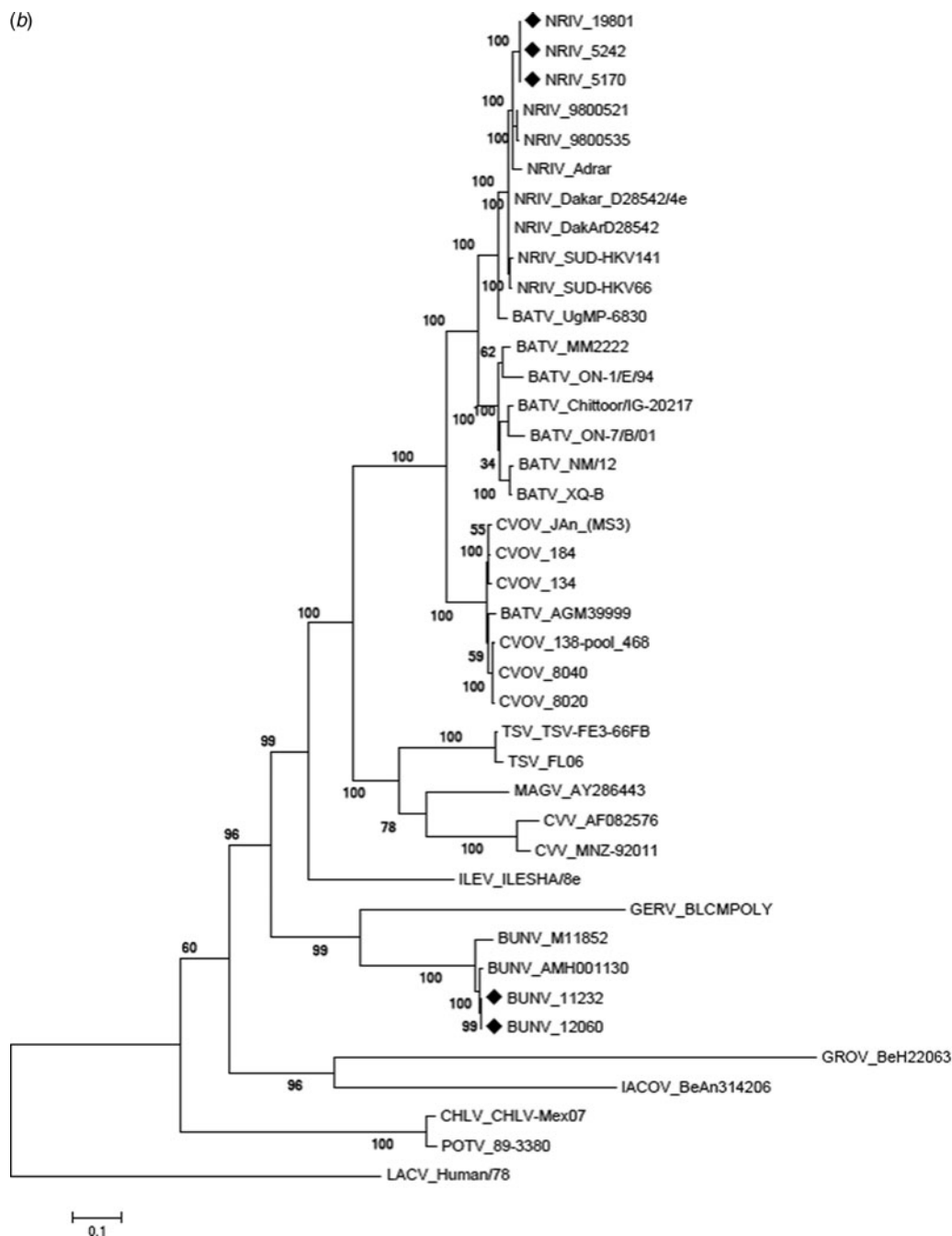


Fig. 1b. *M* segment accession numbers: BATV_ON_7_B_01 (AB257765), BATV_ON_1_E_94 (AB257764), BATV_NM_12 (KJ187039), BATV_XQ+B (KJ398937), BATV_UgMP_6830 (JX846602), BATV_Chittoor_IG_20217 (JX846599), BATV_MM2222 (JX846596), BATV_Italy_2009 (KC168047), BUNV_M11852 (M11852), BUNV_AMH001130 (JF961341), CHLV_CHLV_Mex07 (JN808310), CVV_MNZ_92011 (KC436107), CVV_AF082576 (AF082576), CVV_807270 (AF186243), CVOV_138_pool_468 (KC608156), CVOV_JAnMS3 (KJ542628), CVOV_134 (KJ542625), CVOV_184 (DQ334335), GROV_BeH22063 (AY380581), GERV_BLCMPOLY (M21951), IACOV_BeAn314206 (JN572066), ILEV_ILESHA_8e (KC608150), LACV_Human_78 (AF528166), MAGV_AY286443 (AY286443), NRIV_Dakar_D28542_4e (KC608153), NRIV_9800521 (JX857326), NRIV_9800535 (AY593725), NRIV_DakArD28542 (JX857317), NRIV_HKV141 (JX857323), NRIV_HKV66 (JX857320), POTV_89_3380 (EU004189), TSV_FE3_66FB (FJ943508), TSV_FL06 (FJ943506), NRIV_Adrar (KJ716849).

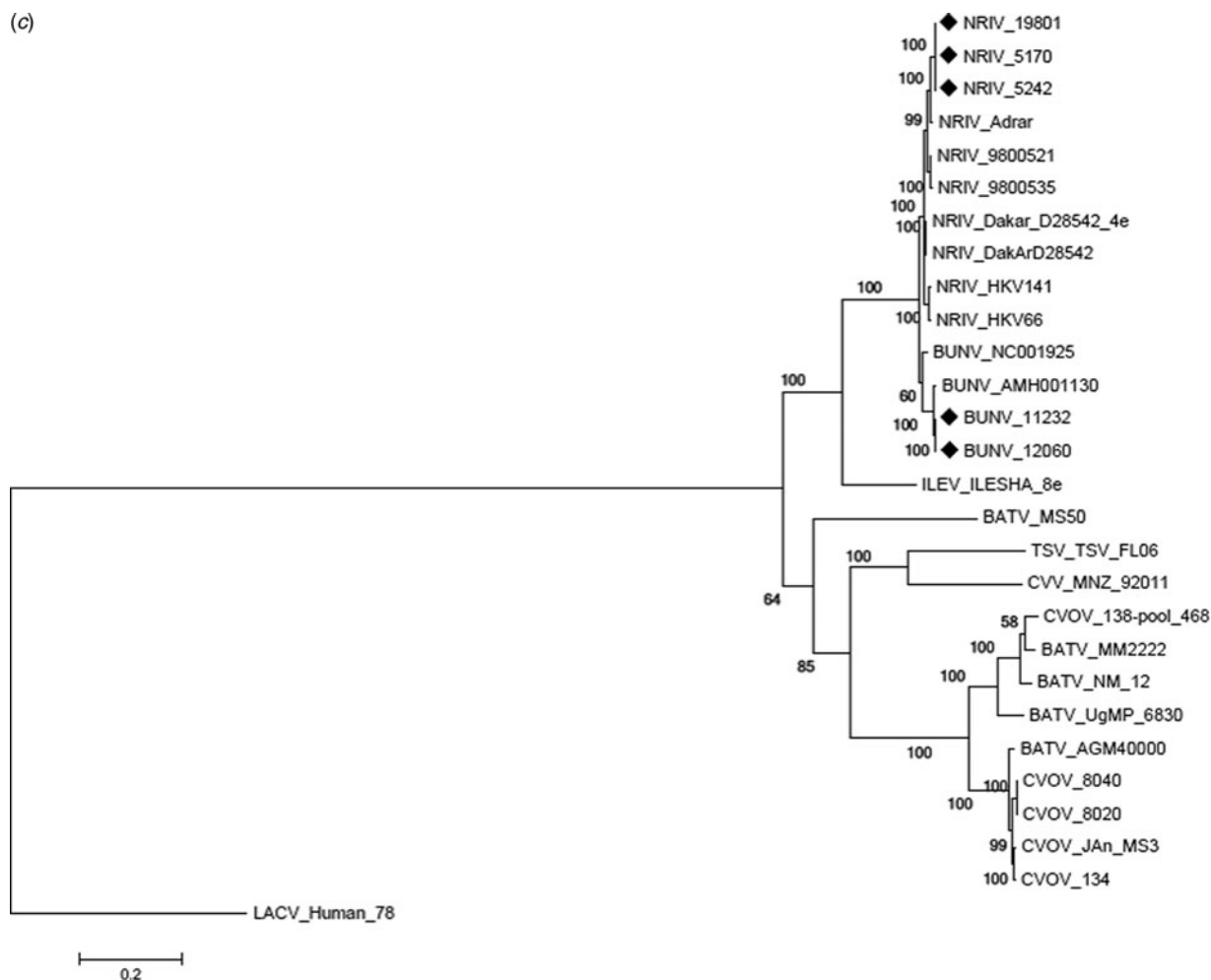


Fig. 1c. *L* segment accession numbers: BATV_NM_12 (KJ187038), BATV_UgMP_6830 (JX846603), CVOV_138-pool_468 (JX846600), BATV_MM2222 (JX846597), BATV_Italy_2009 (KC168048), BUNV_AMH001130 (F961340), BATV_MS50 (JX846606), BUNV_NC001925 (NC001925), CVOV_8040 (KJ542635), CVOV_8020 (KJ542632), CVOV_JAn_MS3 (KJ542629), CVOV_134 (KJ542626), TSV_FL06 (FJ943509), CVV_MNZ_92011 (KC436106), ILEV_ILESHA_8e (KC608149), ILEV_R5964 (KF234075), LACV_Human_78 (AF528165), NRIV_D28542_4e (KC608152), NRIV_9800521 (JX857327), NRIV_9800535 (JX857330), NRIV_DakArD28542 (JX857318), NRIV_HKV141 (JX857324), NRIV_HKV66 (JX857321), NRIV_Adrar (KJ716850).

Additionally, as more sequences are continually generated, a more extensive survey with regard to sample size and geography is necessary to better understand the distribution of Ngari and Bunyamwera viruses.

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Declaration of interest

None.

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Supplementary Material

Supplementary Table S1. Primers used in sequencing of Kenyan Bunyamwera and Ngari virus isolates

Target protein	Primer sequence (5'-3')	Annealing position (5'-3')	Reference
BUNV N protein			
BUNS1	AGTAGTGTACTCCACACTACAACT	1-25	Yandoko et al, 2007 [1]
BUNS9	AGGAATCCACTGAGGCGGTGGAGG	381-358	
BUNS4	CTGGCAACCGGAACAACCCAGTT	318-340	Yandoko et al, 2007 [1]
BUNS5	GAGACAACCTGTCAGTGCAGACTGAA	711-687	
BUNS10	TCAGTCTGCACTGACAGTTGTCTC	688-711	Yandoko et al, 2007 [1]
BUNS2	AGTAGTGTGCTCCACCTAAAACCTTA	961-937	
BUNV M Polyprotein			
Bunya M14C	CGGAATTCAGTAGTGTACTACC*	1-14	Yanase et al, 2006 [2]
Bunya M619	GACATATGYTGATTGAAGCAAGCATG*	586-576	
BUNM11F	TCAGCACTGGCTGGTTTAAG	481-500	This study
BUNM11R	ACCTGCACCGAAGAGTGATG	1178-1159	
BunM12F	GAGATAGCAGATGTCCTTCAAGC	1092-1114	This study
BunM12R	CAAGCAAGTGCATTCTGTGG	1725-1706	
Bun M3F	TTCCTAATAAATGTGGGTTTTG	1311-1333	This study
Bun M3R	ATGCTGACTGCCTGATAGGG	2071-2052	
Bun M4F	TCCAGGCACTGCTTATGTTT	2003-2022	This study
Bun M4R	TCTTCCCCTGGTATGTGGTC	2671-2652	
Bun M5F	TGTGATTCTGGTCCGCTCTAC	2625-2645	This study
Bun M5R	GCATCCCCATCTGCTAGTTC	3236-3217	
Bun M6F	CAAAGCATGACGAACATTGC	3136-3155	This study
Bun M6R	TCAAGGCTGCAGATTGTGTC	3919-3900	
Bunc7MF	CCAGATTGATGCGAAATGTG	3821-3840	This study
BAT 3' end R	GAATTCAGTAGTGTGCTACC*	4458-4445	
BUNV L Protein			
M13 BunL 1C Yanase	TAAAACGACGGCCAGTAGTGTACTCT*	1-14	Yanase et al, 2006 [2]
BunL605R Yunase	RGTGAARTCNCCATGTGC	614-597	
Bun2LF	GTTGCTGGACAAGTTTGCTG	551-570	This study
Bun2LR	TTGCCAATCTTAACCGCTTC	1219-1200	
Bun3LF	AAAATGATTGCCAGGTCAAC	1152-1171	This study
Bun3LR	TATGAAAAGCCCAGGTGATG	1889-1870	
Bun5LF	AAGAAACAACATGTCAATCTACC	2421-2443	This study
Bun5LR	TAGCGGATTTCTCTTCTGCT	3085-3066	
Bunya L For JV	AATATAATAGACATAATACATTTAGAGT	1617-1645	This study
Bunya L Rev JV	CTCCATTTDGCATRTCTGCA)	3175-3154	
BUNL6F	GCTGAACACGGACGAGATG	3003-3020	This study
BUNL6R	GGCCCAATGTAAGCACAATC	3784-3765	
BUNL7F	GGAGAACCACTATCTGTCTTTGG	3717-3739	This study
BUNL7R	ATTGTGCTGGGTTTTGGATG	4491-4472	
BUNL8F	GCAGTTCATGCAATCTGTCC	4412-4431	This study
BUNL8R	TGTTGCCCTGAATCAATGTG	5163-5144	
BUNL9F	TCACAGCTGCAAACTTCC	4729-4748	This study
BUNL9R	TCAATCTGCTTGCCTTCTCC	5334-5315	
BUNL10F	AAGCGAGCAAATAATAGCAAATG	5256-5274	This study
BUNL10R	TACCTAGGGCCTCTGGATTG	5866-5847	

BUNL11F	CCGGCTATTTAAGATCAATAAGG	5575-5597	This study
BUNL11R	TGCTCATCGCCATTACATTC	6187-6168	
BUNL12F	TCTAGCATTGTCCCCTTTTG	6138-6157	This study
BUNL12R	TTTTGCCACATAGTGCTTTTG	6823-6803	
BUNL12F	TCTAGCATTGTCCCCTTTTG	6138-6157	This study
Bun 3'end LR	<u>GTAAAACGACGGCCAGTAGTGTGCTCC*</u>	6875-6863	
NRIV Polyprotein M segment			
BunyaM14C	CGGAATTCAGTAGTGTACTACC*	1-14	This study
TrialR2	TGACCCGCAATTTGTAAAGG	845-826	
BATM13F	CCAAACCGAGAAGTTGAACC	419-438	This study
BATM13R	AATCCTTCCAGGACATCAGC	1099-1080	
BATM14F	TGCTCATGCTGTGGTCTAGC	798-817	This study
BATM14R	ACCTCCACTTTGCCTGTGAG	1788-1769	
BATAIM3F	CCTGGGGAAGCATTGTGATTACT	1704-1726	Jost et al, 2011 [3]
BATAIM3R	CTAGCCAGCGACTCTTGCCTTCC	2228-2206	
BATAIM4F	GTCGCTGGCTAGTGCTACCTCTGG	2217-2240	Jost et al, 2011 [3]
BATAIM4R	CTGATTATTGTCGGATTTATTGGGAACCT	2726-2698	
BATAIM5F	AAAGGTTCCCAATAAATCCGACAA	2696-2719	Jost et al, 2011 [3]
BATAIM5R	CAAATTCTTCACATCCCCAACGACTA	3220-3195	
BATAIM6F	AGAATTTGGGTGCCTTGCTGTCA	3213-3235	Jost et al, 2011 [3]
BATAIM6R	AGATGTTTGGTCCCCTGTGCTTATTT	4086-4061	
BAT 3'end F	TGTTTCGCAGATAACCATGAAAC	3688-3709	This study
BAT 3'end R	<u>GAATTCAGTAGTGTGCTACC*</u>	4438-4425	
NRIV L segment			
NgariL6F	TTTTGAGAAGGAATAATGAAGG	5821-5842	This study
NgariL6R	CCCCTGCTGCAACTAACTTC	6339-6320	

* Underlined sequence corresponds to virus segment ends.

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