

Novel Orthonairovirus Isolated from Ticks near China–North Korea Border

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We isolated a new orthonairovirus from *Dermacentor silvarum* ticks near the China–North Korea border. Phylogenetic analysis showed 71.9%–73.0% nucleic acid identity to the recently discovered Songling orthonairovirus, which causes febrile illness in humans. We recommend enhanced surveillance for infection by this new virus among humans and livestock.

Viruses of the genus *Orthonairovirus*, family *Nairoviridae*, include the consequential tick-transmitted pathogens Crimean–Congo hemorrhagic fever virus and Nairobi sheep disease virus, as well as other poorly characterized viruses that have been found in ticks and mammals. *Orthonairovirus* virions are spherical in shape (80–120-nm diameter) with 3 single-stranded RNA segments 17.1–22.8 kilobases in length and a membrane envelope (1–5). We performed surveillance in areas endemic for tick-borne encephalitis (6) and identified a novel orthonairovirus from *Dermacentor silvarum* ticks collected in 2021 in Jilin Province, China, near the China–North Korea border.

The Study

On April 17, 2021, we dragged corduroy to collect ticks from a forest region in Antu (118°46'E, 43°15'N), a district of the city of Yanbian in eastern Jilin Province, China, near the border with North Korea. We identified captured ticks according to morphologic keys and stored them at 4°C with wet cotton. We

collected 264 ticks of 3 species—29 *Ixodes persulcatus*, 193 *Dermacentor silvarum*, and 12 *Haemaphysalis concinna*—and 30 larvae of unidentified species.

We homogenized ticks using a QIAGEN TissueLyser (QIAGEN, <https://www.qiagen.com>) and inoculated supernatants onto a monolayer of African green monkey kidney (Vero) E6 cells. After 3 successive passages, we observed cells for cytopathic effects. The inoculate from *Dermacentor silvarum* ticks, designated as YB_tick_2021_24, caused cytopathic effects in Vero E6 cells 96 h after inoculation (Figure 1, panels A, B). We collected cells showing cytopathic effects, then fixed and embedded them in epoxy resin. We cut ultrathin (80 nm) sections from the resin block, stained them with citrate lead and uranyl acetate, and observed them under a transmission electron microscope. We observed enveloped virus particles ≈100 nm in diameter that shared morphologic features with *Bunyavirales* viruses (Figure 1, panel C).

We extracted viral RNA from infected culture supernatants using a QIAGEN QIAamp Viral RNA Mini Kit, synthesized cDNA, prepared DNA libraries using an Illumina Nextera XT Kit (Illumina, <https://www.illumina.com>), and performed 150 bp paired-end sequencing using the Illumina MiniSeq System. We filtered reads on the basis of their length and mean quality values. We prepared contigs by de novo assembly and subjected them to BLASTx alignment (<https://blast.ncbi.nlm.nih.gov/Blast.cgi>) at E value <10⁻⁴ against the nonredundant protein and viral proteome databases of the National Center for Biotechnology Information. We used Bowtie 2 (<https://bowtie-bio.sourceforge.net/bowtie2/index.shtml>) to remap the clean reads to the generated virus-related contigs (7). We used rapid

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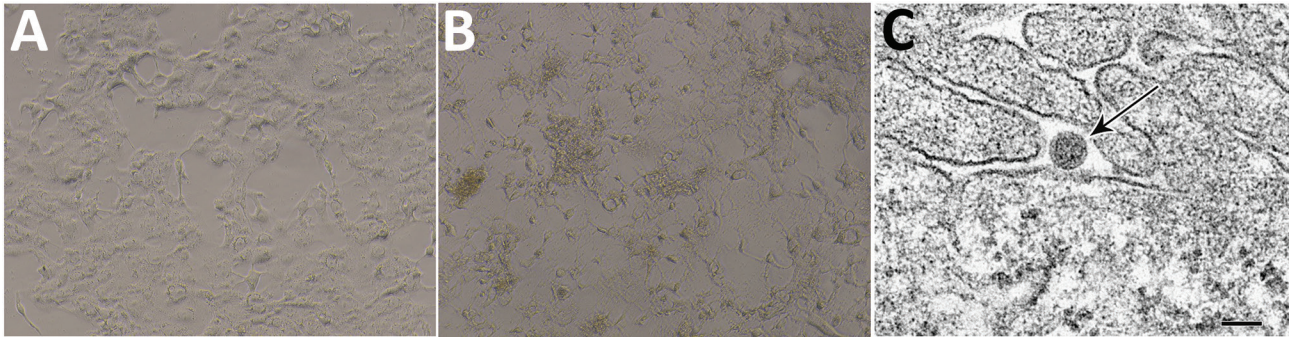


Figure 1. Discovery and characterization of novel orthonairovirus Antu virus isolate YB_tick_2021_24 from *Dermacentor silvarum* ticks in China. A) Vero E6 cells without YB_tick_2021_24 infection. Original magnification $\times 10$. B) YB_tick_2021_24-infected Vero E6 cells showing cytopathic effects visible by light microscopy. Original magnification $\times 10$. C) Ultrathin section electron micrograph of an isolated particle (black arrow) on a cell surface. Scale bar = 100 nM

amplification of cDNA ends (RACE) PCR and Sanger sequencing to confirm the terminal sequences of virus genomes, and deposited the new genome in GenBank (accession nos. OQ207701–3). We identified open read frames (ORFs) using ORF finder (<https://www.ncbi.nlm.nih.gov/orffinder>) and

calculated sequence similarities using BLAST.

Our procedure generated 40,826,350 reads (6.1 Gbp), which produced 266 virus-related contigs. Three contigs, the 1,516 bp small (S), 3,936 bp medium (M), and 12,133 bp large (L) segments, were annotated to Songling virus (SLV), a previously reported

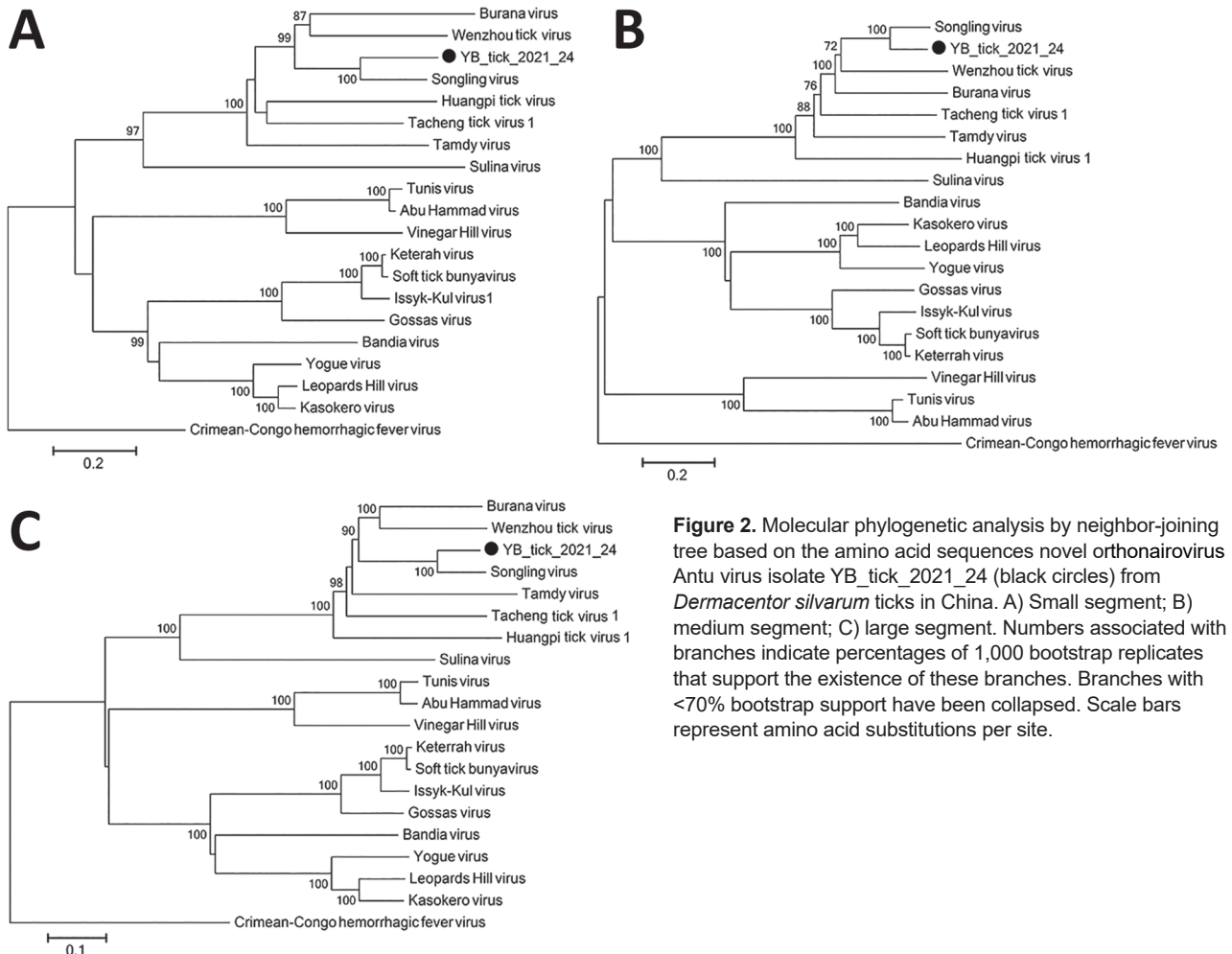


Figure 2. Molecular phylogenetic analysis by neighbor-joining tree based on the amino acid sequences novel orthonairovirus Antu virus isolate YB_tick_2021_24 (black circles) from *Dermacentor silvarum* ticks in China. A) Small segment; B) medium segment; C) large segment. Numbers associated with branches indicate percentages of 1,000 bootstrap replicates that support the existence of these branches. Branches with $<70\%$ bootstrap support have been collapsed. Scale bars represent amino acid substitutions per site.

Table. Homology comparisons of the sequence of novel orthonairovirus Antu viruses from China and other related viruses*

Protein/virus	Antu virus	SLGV	WTV	TTV	BURV	TDYV	HTV	CCHFV
Small								
Antu virus		71.5	52.6	53.7	52.1	48.9	47.4	37.3
SLGV	71.9		55.3	51.2	51.8	49.6	45.9	34.9
WTV	60.3	60.6		50.0	54.1	43.8	46.0	34.0
TTV	59.5	57.9	56.6		46.6	49.1	51.0	33.4
BURV	58.6	59.9	60.5	55.2		44.3	42.4	35.0
TDYV	56.5	58.2	54.0	57.9	53.1		45.0	34.3
HTV	55.5	55.8	55.9	58.7	52.9	55.8		35.2
CCHFV	47.6	46.8	46.8	46.5	47.9	47.7	46.3	
Medium								
Antu virus		79.5	59.9	56.3	58.7	53.8	46.9	25.4
SLGV	72.4		58.2	53.9	57.0	51.2	46.5	24.5
WTV	61.9	61.8		51.8	54.0	50.0	46.5	24.5
TTV	57.2	58	56.4		51.6	51.1	48.1	24.5
BURV	61	61.2	58.9	58		49.4	47.2	24.0
TDYV	56.6	57.5	55.9	55.1	55.4		45.7	24.4
HTV	53.7	53.9	53.4	52.9	54.2	52.1		24.6
CCHFV	41.8	40.7	40.5	40.6	41.3	42.2	40.3	
Large								
Antu virus		84.6	66.5	64.1	66.0	62.2	60.1	39.2
SLGV	73.0		65.7	64.0	65.1	61.4	60.1	38.5
WTV	63.4	63.7		63.5	69.7	61.7	60.0	38.5
TTV	61.9	62.0	62.2		63.2	59.3	60.0	38.7
BURV	63.4	63.3	66.1	61.8		61.5	60.3	38.7
TDYV	60.4	60.5	60.5	59.4	60.8		58.0	39.2
HTV	59.8	60.1	59.5	61.5	60.3	58.2		38.2
CCHFV	48.1	48.3	48.2	48.0	48.8	48.1	48.5	

*Percentage nucleotide sequence identity presented below and amino acid identity above blank cells. BURV, Burana virus; CCHFV, Crimean-Congo hemorrhagic fever virus; HTV, Huangpi tick virus; SGLV, Songling virus; TDYV, Tamdy virus; TTV, Tacheng tick virus; WTV, Wenzhou tick virus.

orthonairovirus (8). Average sequencing coverages remapped to the 3 contigs were 48× (S), 63× (M), and 234× (L). The final genome lengths confirmed by RACE sequencing were 1,848 bp encoding 488 aa for the S segment, 4,099 bp encoding 1,263 aa for the M segment, and 12,001 bp encoding 3,950 aa for the L segment. We performed multiple alignments using MAFFT version 7 (<https://mafft.cbrc.jp/alignment/server>) (9) and constructed a phylogenetic tree in MEGA7 (<https://www.megasoftware.net>) by using the neighbor-joining method with a bootstrap test for 1,000 replicates (10).

Phylogenetic analysis showed the strain belongs to the genus *Orthonairovirus*, family *Nairoviridae*, and is genetically related to SLV (Figure 2) (4,5,8,11). The terminal nucleotides of the S segment were identical to those of orthonairoviruses (3' segment terminus AGAGUUUCU and 5' segment terminus AGAAACUCU) (5). The termini of the M and L segments were different (Appendix Figure, <https://wwwnc.cdc.gov/EID/article/29/6/23-0056-App1.pdf>). Homology analysis comparing YB_tick_2021_24 with SLV sample YC585 showed 71.9% nucleic acid (na) and 71.5% aa identities for the S segment, 72.4% na and 79.5% aa identities for the M segment, and 73.0% na and 84.6% aa identities for the L segment (Table 1) (8). Those results indicate that the isolate represents a unique

Orthonairovirus species. For purposes of archiving, we designated novel YB_tick_2021_24 as Antu virus and deposited the strain in the National Pathogen Resource Center (accession no. NPRC 2.3.9401).

Conclusion

We identified a novel orthonairovirus, Antu virus, in *Dermacentor silvarum* ticks collected in China near the China–North Korea border. Nucleotide and amino acid sequence homologies, combined with phylogenetic analysis of other orthonairovirus genomes, suggested that Antu virus is a new member of the genus *Orthonairovirus*, genetically related to SLV. Tamdy virus and SLV are orthonairoviruses reportedly able to infect human and livestock (8,12,13). Lacking direct evidence of the ability of Antu virus to infect and cause illness among humans and livestock animals, we recommend enhanced monitoring and surveillance for Antu virus infection among humans and livestock in potentially endemic areas.

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Appendix

	5' terminus	3' terminus
CCHFV-L	TCTCAAAGATATCAATCCCCCGTT	CTCTTCTTAGCTATATCTTTGAGA
CCHFV-M	TCTCAAAGA AATACTTGCGGCACG	GCGTGCCGCCACTATATCTTTGAGA
CCHFV-S	TCTCAAAGAAACACGTGCGGCTTA	CTGTGCGGCAACGGTATCTTTGAGA
HUGV-L	TCTCAAAGATATAAATCCCTACACC	GGTTTAGGGAACACTTTCTTTGAGA
HUGV-M	TCTCAAAGAAAGACCTGCAGCAAC	GTTAGCTGCCACAATATCTTTGAGA
HUGV-S	TCTCAAAGA AAGACGTGCCTATCC	GTTAGCGGCAACAATATCTTTGAGA
ERVEV-L	TCTCAAAGA AAGCAATCCCCCA	TAGGGGGAATACTATCTTTGAGA
ERVEV-M	TCTCAAAGAAAGACTAGCGGCAA	GATGCGGCTCCTATATCTTTGAGA
ERVEV-S	TCTCAAAGAAAGTTGTGCTTACT	GATGCAGCAACACTATCTTTGAGA
TDYV-L	CACACCCA AATACATAGAACCAGG	CCGGTGTATGGTATTGGGTGTG
TDYV-M	CACACCAAGCATTATAAACCAGTT	CAGGTGGATAAGATTTCTGGGTGTG
TDYV-S	CACACCCA AACTTTACACTTTAGG	CCTTGAGCAATTTGCTTTGGGTGTG
TTV-L	TCTCAAAGATATATATCCTGCACAC	GCTGGCACAAAATCGGTGAAATTGT
TTV-M	CTGCAGCACACCAAAAGCCTTTCA	GCCGTGAAAAAGAAAGAAATACA
TTV-S	AACGTGCTGCACACCAATAGCATT	ATCAGTTTACTACTGGTGAAGTT
HTV-L	AAGATATATATCCTGCACACCCAAA	GTGACCAACTCTTCTGGTCTGATA
HTV-M	AAACAGTGTGAAGGCAATGATGAG	AAACAAACAAAAGAGAAAA
HTV-S	CTGCACACCAAAACCTAAAGCAAC	ATATAAGAGATCGGAAGAGCGTCG
WTV-L	CCTAACACCACTTAACATCTGCCAA	TAGGGTTARGGGTGTGCAGGAACA
WTV-M	CCCTACTAAAGGCTAAAGGTTAGCG	TGGTGTAGGTGATTGGTGTGCTG
WTV-S	ATCACCTACATCGAATACCCATCCC	GTAAGTTAAAGGGTGTGCAGCAACA
SGLV-L	TCTCAAAGATATATATCCTGCACA	GTGTGGGAAGGTTGATCCCATGT
SGLV-M	ACATGGGATAGTAACCTGTGCTAG	CCTGTCTCGACCATGCCCCCATGT
SGLV-S	TCTCAAAGAAACACGTGCTGCACAC	AGAAACAAAACAAATTCCCATGT
ATV-L	GATATATATCCTGCACACCCAAAC	AGCTGCTGTGGAGTCTGGTGACCT
ATV-M	TGTCAGCATGAAGGAGGGAACA	AAGTCCCGCTCTGCAACTGCCT
ATV-S	TCTCAAAGAAAAACGTGCTGCACA	GTGCAGCAACAATATCTTTGAGA

Appendix Figure. Terminal sequences of ATV and other orthonairoviruses. Typical terminal sequences of orthonairoviruses are highlighted in yellow and putative terminal sequences in red. L, Large segment;

M, medium segment; S, small segment; CCHFV, Crimean-Congo hemorrhagic fever virus (L, NC_005301; M, NC_005300; S, NC_005302); HUGV, Hughes virus (L, NC_040512; M, NC_040513; S, NC_040514); ERVEV, Erve virus (L, JF911697; M, JF911698; S, JF911699); TDYV, Tamdy virus (L, MK757580; M, MK757581; S, MK757582); TTV, Tacheng tick virus 1 (L, NC_031284; M, NC_031285; S, NC_031286); HTV, Huangpi tick virus 1 (L, NC_031135; M, NC_031136; S, NC_031137); WTV, Wenzhou tick virus (L, NC_031291; M, NC_031288; S, NC_031289); SGLV, Songling virus (L, MT328776; M, MT328775; S, MT328777); ATV, Antu virus.