

Evidence of Human Exposure to Tamdy Virus, Northwest China

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We report the isolation of Tamdy virus from *Hyalomma asiaticum* ticks in northwest China and serologic evidence of human Tamdy virus infection in the same region. These findings highlight the need to further investigate a potential causal relationship between Tamdy virus and febrile illnesses of unknown etiology in that region.

The species *Tamdy orthonairovirus* (genus *Orthonairovirus*, family *Nairoviridae*) includes 5 viruses: Tamdy virus (TAMV), Burana virus (BURV), Tǎchéng tick virus 1 (TcTV-1), Huángpí tick virus 1 (HpTV-1), and Wēnzhōu tick virus (WzTV) (1). TAMV and BURV were initially isolated from ticks in countries in central Asia (2–4), but little is known about their medical and veterinary importance. TcTV-1, HpTV-1, and WzTV were putative viruses identified by virome sequencing from ticks in China (5); however, their virologic properties and pathogenesis potential remain unclear. One study (6) reported TcTV-1 isolated from a febrile patient in northwest China, providing evidence of the potential public health threat from these viruses. We report TAMV isolated from ticks in northwest China and demonstrate serologic evidence of infection in humans.

The Study

During April and May of 2016 and 2017, we collected *Hyalomma asiaticum* ticks (n = 4,123) from Xinjiang in northwest China and divided the ticks into 55

groups according to the sampling location (n = 50–100 ticks/group) (Figure 1; Appendix Table 1, <https://wwwnc.cdc.gov/EID/article/27/12/20-3532-App1.pdf>). We isolated the virus from homogenates of each tick group in suckling mice. After the first inoculation, we observed symptoms in mice including loss of balance, limb paralysis, tremors, and articulo mortis from 4 (36.37%) of 11 pooled samples from Yuli County, 1 (14.29%) of 7 from the city of Karamay, 3 (60%) of 5 from Luntai County, and 17 (53.13%) of 32 from the city of Wujiaqu (Appendix Table 1). We performed a second inoculation using brain samples from diseased mice from Luntai and Wujiaqu Counties, in which >50% of the mice experienced illness onset after first inoculation. Similar symptoms were reproducibly observed in 1 group from Luntai and 4 groups from Wujiaqu (Appendix Table 1).

Subsequently, we prepared 3 RNA pools of diseased mouse brains and obtained a total of 196,946,814 reads by RNA sequencing. We found TAMV contigs in all 3 pools (Appendix Table 2), confirming findings using real-time reverse transcription PCR (rRT-PCR) (data not shown). We used homogenates of 2 TAMV RNA-positive brain samples from the A-M6 pool (Appendix Table 2) to isolate viruses in Vero E6 cells. As indicated by immunofluorescence assays (IFA) (Appendix), we observed increasing TAMV infection from first to fourth passages in Vero E6 cells, suggesting successful isolation (Appendix Figure 1).

Negative-stain electron microscopy revealed an enveloped spherical viral morphology with a diameter of ~90–110 nm (Figure 2, panel A). We observed viral particles in cytoplasm and vesicles of infected cells (Figure 2, panel B). Although this screening was not exhaustive, IFAs showing varied susceptibility of different cells lines indicate that TAMV seems to have a broad host range, including humans, monkeys, sheep, dogs, and mice (Appendix Figure 2).

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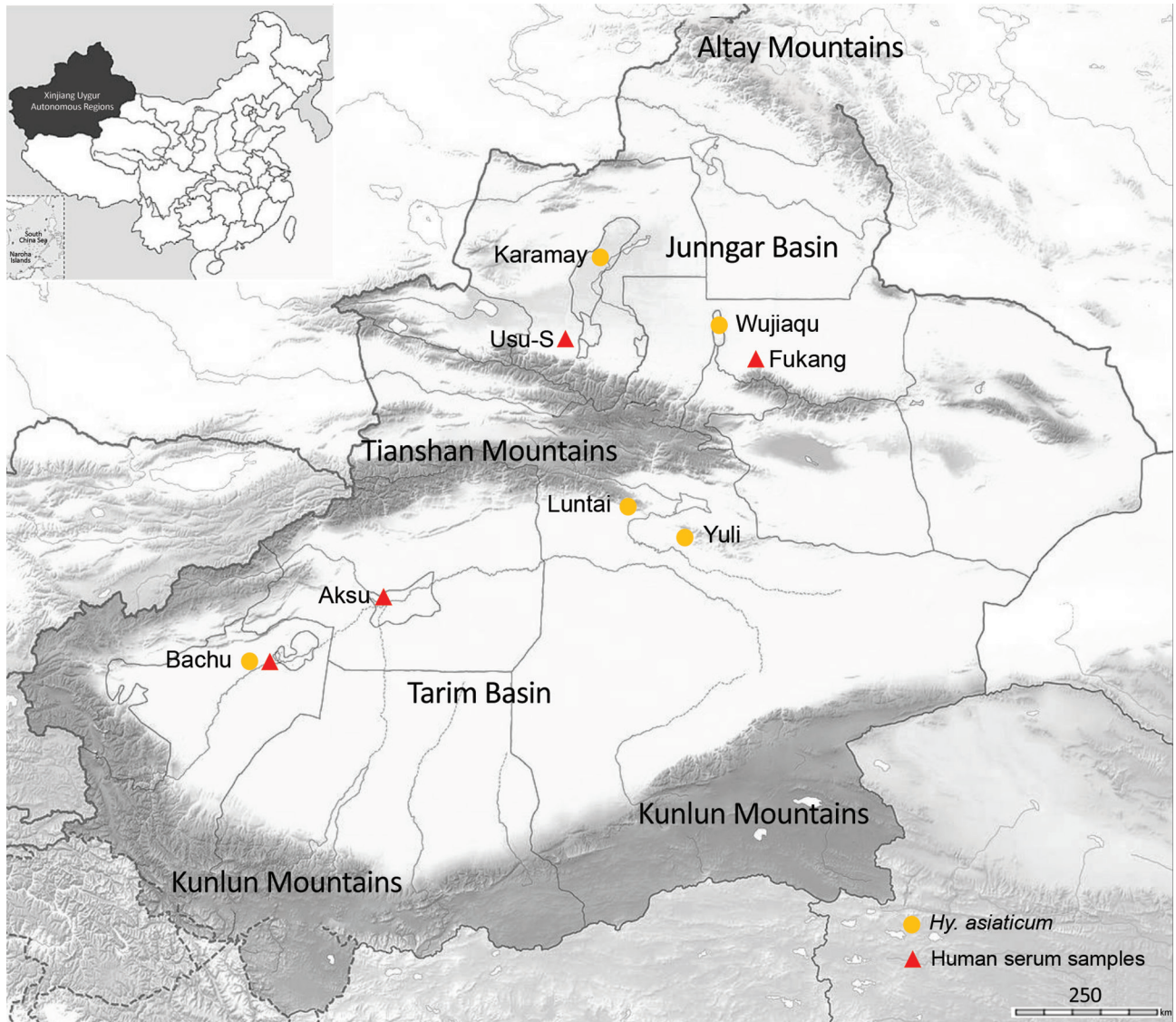


Figure 1. Collection locations for *Hyalomma asiaticum* ticks and human serum samples used in study of human exposure to Tamdy virus, Xinjiang, China. Usu-S, southern area of Usu City.

The 2 TAMV isolates shared very high sequence similarities (99.96% for large, 100% for medium, and 99.95% for small segments). We named the 2 TAMV strains YL16082 and YL16083, including an abbreviation (YL) for the geographic location (Yuli County) where the original tick samples were collected. TAMV genome sequences (YL16082) shared 37%–59% identity with other members of the *T. orthonairovirus* species and 34%–49% identity with Crimean-Congo hemorrhagic fever virus (CCHFV), another *Orthonairovirus* species. Protein sequences shared 44%–62% identity with other members of the *T. orthonairovirus* species and 33%–40% with CCHFV (Table 1). Phylogenetic trees based on nucleotide

sequences of RNA-dependent RNA polymerase, glycoprotein, and nucleoprotein (NP) genes all confirmed the close taxonomic relationships with currently known TAMV strains and other members of the species *T. orthonairovirus* (Appendix Figure 3).

To investigate potential human infection by TAMV in northwest China, we conducted a seroprevalence study using archived serum samples from 725 healthy persons (collected in 2005 from Fukang City, 2014 from Aksu City, and 2017 from Usu City) and 87 febrile patients (collected in 2007 from Bachu County, which has a history of CCHFV prevalence) (Appendix). Of the 87 febrile patients, 21 (24.14%) were TAMV IgG positive and 17 (19.54%)

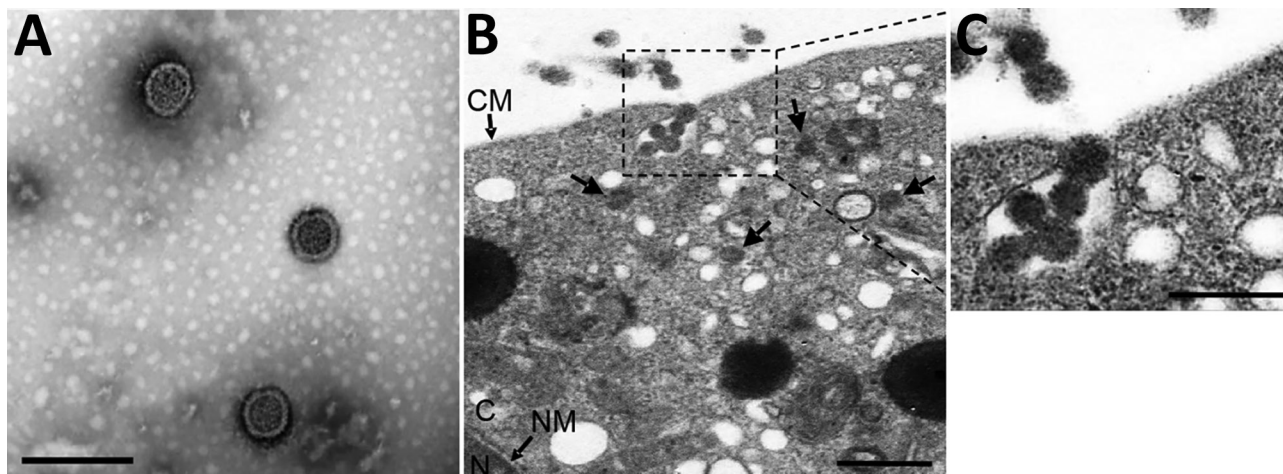


Figure 2. Visualization and subcellular localization of Tamdy virus (TAMV) virions by electron microscopy. A) Negative-staining image of purified TAMV virions. Scale bar indicates 200 nm. B) Image of Vero E6 cells infected with TAMV; arrows indicate TAMV virions in the cytoplasm. Scale bar indicates 500 nm. C) The enlarged image of interest from B. scale bar indicates 200 nm. CM, cell membrane; C, cytoplasm; NM, nuclear membrane; N, nucleus

IgM positive (Table 2; Appendix Table 4), whereas only 1 (0.13%) of the 725 healthy participants we tested IgG positive (data not shown). Neutralization (titers 16–64) was demonstrated in serum samples from 6 febrile patients (6.9%) (Table 2). Moreover, of the 24 tick groups from the same locations as the febrile patients, 10 groups (41.76%, 6 identified in sheep and 4 in fields) tested positive for TAMV RNA by rRT-PCR (data not shown). Partial sequences of large segments from these positive groups clustered together with TAMV strains (Appendix Figure 4). These results showed serologic evidence of human exposure to TAMV and evidence of TAMV presence in *Hy. asiaticum* ticks in northwest China as early as 2007, which warranted more in-depth investigation to establish the potential causal relationship between TAMV and febrile illnesses of unknown etiology in regions where TAMV is present.

Finally, because another *T. orthonairovirus*, TcTV-1, had been identified in a febrile patient in northwest China (6), we thought it important to determine the potential serologic cross-reactivity between these 2 viruses. However, TAMV and TcTV-1 shared limited protein sequence identity (49%–60%) (Table

1), suggesting limited cross-reactivity, if any. This result was confirmed by conducting serologic testing using recombinant NP proteins from the 2 viruses. As shown by both IFA and Western blot analyses, TAMV NP antibodies had no cross-reaction with TcTV-1 NP (Appendix Figure 5, panels A, B). In addition, human serum samples that were positive for TAMV IgM or IgG, or both, showed reactivity with TAMV NP, but not with TcTV-1 NP (Appendix Figure 5, panel C).

Conclusions

TAMV was initially found in ticks in countries in central Asia, including Kazakhstan, Kyrgyzstan, Uzbekistan, and Turkmenistan (3,4). Its infection status in humans and livestock animals was not well characterized. Our data, together with reports of TAMV isolated from ticks in Xinjiang, China (7), and identified in Turkey (8), shows that the geographic distribution of TAMV is much wider than originally recognized. In addition, we provide strong serologic evidence of human exposure in TAMV-affected regions.

Findings of a potential role of TcTV-1 in causing human febrile disease (6) suggest that >1 virus in this species group may have the potential to cause

Table 1. Sequence identity of TAMV isolate YL16082 from China compared with other members of the species *Tamdy orthonairovirus* and Crimean-Congo hemorrhagic fever virus*

Virus	Nucleotide identity, %			Amino acid identity, %		
	L segment	M segment	S segment	RdRp	G	NP
Wēnzhōu tick virus	59	50	42	62	51	44
Tāchéng tick virus 1	57	46	44	60	51	49
Huángpí tick virus 1	55	45	42	58	47	46
Burana virus†	59	47	37	62	50	44
Crimean-Congo hemorrhagic fever virus	49	34	39	40	33	36

*G, glycoprotein; L, large; M, medium; NP, nucleoprotein; RdRp, RNA-dependent RNA polymerase; S, small; TAMV, Tamdy virus.

†Partial sequences were available for analyses.

Table 2 Seroprevalence of TAMV among 87 febrile patients with samples collected in 2007 from Bachu County, Xinjiang, China*

Patient characteristics	Serum samples, no.	TAMV IgG positive, %	TAMV IgM positive, %	Neutralization activity, %	Neutralization titers
Sex					
M	20	3 (15.0)	7 (35.0)	1 (5.0)	64
F	29	6 (20.7)	5 (17.2)	3 (10.3)	16, 32, 32
NA	38	12 (31.6)	5 (13.6)	2 (5.3)	16, 16
Age, y†					
<18	6	2 (33.3)	1 (16.7)	0	
18–30	20	1 (5.0)	8 (40.0)	1 (5.0)	16
31–45	7	2 (28.6)	1 (14.3)	0	
46–60	11	2 (18.2)	0	1 (9.1)	32
>60	5	2 (40.0)	2 (40.0)	2 (40.0)	32, 64
Total	87	21 (24.1)	17 (19.5)	6 (6.9)	16–64

*NA, not available; TAMV, Tamdy virus.

†Age stratification based on 49 patients with recorded age information

diseases in humans. However, our data did suggest the possibility of such a relationship because the TAMV-positive ratio was much higher among febrile patients than healthy persons in the study from the same region. In addition, at least 2 febrile patients had both TAMV IgM and IgG at the time of sampling, during or not long after acute illness; 1 of them had neutralization to TAMV (Appendix Table 3).

Among study limitations, the nature of using archived samples limited our ability to provide direct evidence of a causal relationship between TAMV and human febrile illnesses. Also, it is possible that the high TAMV antibody-positive ratio might have resulted not from the recent cases but from a small outbreak of human TAMV infection in northwest China in 2007.

In summary, our study strongly suggests the potential of TAMV as a human pathogen and supports an urgent need to conduct more in-depth epidemiologic and pathogenesis investigations into this group of viruses in China, central Asia, and beyond. While the world's attention is currently on coronavirus disease and batborne viruses, our study highlights the need to pay attention at the same time to emerging zoonoses of tick origin to prevent future outbreaks.

Acknowledgments

We thank the core facility and technical support groups of Wuhan Institute of Virology for providing technical support for electron microscopy analysis, animal experiments, and experimental activities in a Biosafety Level 3 laboratory.

This work was mainly supported by the Science and Technology Basic Work Program (2013FY113500) from the Ministry of Science and Technology of China, the Intergovernmental Special Program of State Key Research and Development Plan from the Ministry of Science and Technology of China (2016YFE0113500), the National

R&D Infrastructure and Facility Development Program of China, “Fundamental Science Data Sharing Platform” (Y706061YZ1), Science Foundation of China (81690369, 81760365), and the Science Research Key Project of Xinjiang Education Department (XJEDU2019I002). Work in L.-F.W.'s group is supported in part by NRF grants NRF2016NRF-NSFC002-013 and NRF2018NRF-NSFC003SB-002, CD-PHRG grant CDPHRG/0006/2014, NMRC grant ZRRF16006, and MINDEF grant DIRP2015-9016102060.

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Appendix

Collection of ticks in Xinjiang, China

In April and May of 2016 and 2017, we collected a total of 4,123 *Hyalomma asiaticum* ticks from wild fields locating in Karamay City, Yuli County, Luntai County, and Wujiaqu City in Xinjiang using methods described elsewhere (1) (Figure 1). We first identified tick species by morphologic characteristics under microscope and confirmed identifications using sequencing of the rRNA internal transcribed spacer 2 gene, as described elsewhere (2). We divided the ticks into 55 groups (n = 50–100 ticks/group) by sampling locations. We prepared homogenates of the 55 tick groups and used them to inoculate mice as described elsewhere (3) (Appendix Table 1). In addition, 24 groups of *Hy. asiaticum* ticks (17 groups each from different individual sheep and 7 from wild fields) were collected in 2007 from Bachu, the same origin as archived samples from the febrile patients were collected (Figure 1).

Virus Isolation by Mouse Inoculation and Molecular Characterization

We isolated viruses by mouse inoculation as described elsewhere (3) in an animal biosafety level 2 laboratory according to the animal welfare and related regulations specified in the Directory of Pathogenic Microorganisms Transmitted among Humans issued by the Chinese Ministry of Health (<https://www.nhc.gov.cn/wjw/gfxwj/201304/64601962954745c1929e814462d0746c.shtml>). We harvested the brains of diseased mice and stored them in glycerol at -80°C .

We identified the viruses in the mouse brains using RNA-seq. We generated clarified homogenates from the brain of 1 diseased mouse randomly selected from each group as described elsewhere (3). We purified total RNA from 250 μL of clarified brain homogenates using TRIzol (Invitrogen, <https://www.thermofisher.com>) according to manufacture instructions.

We prepared 3 RNA pools, each containing a mixture of 5 µg total RNA from the mouse brain samples (Appendix Table 1). We reverse transcribed the rest of the RNA from each mouse brain into complementary DNA (cDNA) and stored it at –80°C until used for further investigation. RNA-seq library preparation was performed according to the protocol provided by Illumina (4). We used HiSeq 3000 platform (Illumina, <https://www.illumina.com>) for 150 bp paired-end sequencing of the libraries, and analyzed sequencing data to identify viral-related sequences as described elsewhere (5).

For rRT-PCR analysis of TAMV-specific viral RNA, we purified total RNA from homogenates from mouse brains (100 µL), tick groups (100 µL), or cell culture supernatants (300 µL) using TRIzol (Invitrogen), and further transcribed into cDNA using M-MLV reverse transcription (TaKaRa Biotechnology, <https://www.takarabio.com>) according to manufacturer instructions. We performed PCR with primers TAMV-L-F: 5'-CTCAATCATCATCCTCGTCTCAT-3' and TAMV-L-R: 5'-GGACAATAGTTCGCTTAGGCTCT-3' to amplify a partial (423 nt) fragment of the L segment in a 50 µL reaction volume containing 1–3 µL cDNA templates.

Characterization of TAMV in Cell Culture Using Electron Microscopy

We isolated viruses by cell culture as described elsewhere (2) in a biosafety level 3 laboratory according to the Directory of Pathogenic Microorganisms Transmitted among Humans. We inoculated Vero E6 cells with 200 µL of clarified brain homogenates for 2 h at 37°C. We removed supernatants and replaced them with fresh GIBCO Dulbecco's Modified Eagle Medium (ThermoFisher) containing GIBCO 2% fetal calf serum (ThermoFisher) and continued incubation for 5–7 days at 37°C. We monitored infection susceptibility and virus replication using immunofluorescence assays (IFA) as described elsewhere (2) using the polyclonal α-TAMV-NP serum produced in-house (see below).

To test susceptibility of cell lines derived from different hosts to TAMV, U87MG, SW13, 293, Vero E6, MODK, MDBK, PK15, DH82, and BHK21 cells were incubated with 500 µL of clarified supernatants from fourth passage (P4) in Vero E6 cells. Virus infection was confirmed by IFA. For morphologic studies, we purified TAMV from supernatants and visualized it using

negative-staining electron microscopy as described (1). We examined subcellular localization of viral particles in Vero E6 cells by transmission electron microscopy as described elsewhere (1).

Cells, Viruses, and Antibodies

We purchased cell lines derived from human (U87MG [HTB-14], SW13 [CCL-105], and 293 [CRL-1573]), monkey (Vero E6 [CRL-1586]), sheep (MDOK [CRL-1633]), cattle (MDBK [CCL-22]), swine (PK15 [CCL-33]), dog (DH82 [CRL-10389]), and mouse (BHK-21 [CCL-10]) cells from American Type Culture Collection. The 2 isolates of TAMV are deposited in the National Virus Resource Center (IVCAS 6.7499 for strain YL16082; IVCAS 6.7500 for strain YL16083). We performed IFAs, western blots, and neutralization assays using the YL16082 isolate. We used mouse anti-hemagglutinin tag polyclonal antibody (α -HA) for IFAs as the primary antibody (ABclonal Technology, <https://abclonal.com>). We used β -actin antibody (α - β -actin) (Sangon Biotech, <http://www.life-biotech.com>) as the control for western blots. We used fluorescein isothiocyanate-conjugated sheep anti-human IgG antibody, rabbit anti-human IgM μ chain (Alexa Fluor 488, ThermoFisher), goat anti-mouse IgG H&L (Alexa Fluor 488, ThermoFisher), and goat anti-mouse IgG H&L (horseradish peroxidase) as secondary antibodies for IFAs and western blots (Abcam, <https://www.abcam.com>).

In-house Production of α -TAMV-NP in Mice

We amplified the 1,452 bp open reading frame coding region of TAMV NP from strain YL16082 from TAMV-positive mouse brain by PCR using 2 \times Rapid Taq Master Mix (Vazyme Biotech, <https://www.vazymebiotech.com>) according to manufacture instructions. We cloned PCR products into the plasmid pET-32a to generate the expression plasmid pET-32a-TAMV-NP and confirmed the insert by sequencing. We conducted protein expression, purification, and rabbit immunization as described elsewhere (1).

Serologic Investigation of Human Infection by TAMV

Using serum samples archived in a repository, we investigated TAMV seroprevalence in Xinjiang among 725 healthy persons: 465 from a southern area of Usu City in 2017 (1), 80 from Fukang City in 2005, and 180 from Aksu City in 2014, and in June 2007, among 87 febrile

patients from 1 hospital in the downtown area and 2 clinics in the village with the largest population in Bachu County. The 87 serum samples tested negative for Crimean-Congo hemorrhagic fever virus RNA and antibodies (data not shown). We recorded sex and age except from 38 febrile patients whose records lacked personal information. We tested for IgG and IgM with IFA as described elsewhere (1) using TAMV infected cells as antigen and α -TAMV-NP as the positive control antibody. We performed microneutralization tests using the TAMV strain YL16082 as described elsewhere (1).

Cross-Reactivity between TAMV Antiserum and TcTV-1 Proteins.

The protein sequence comparison between the 2 viruses revealed a sequence identity of \approx 48%–60% (Table 1). We cloned full-length open reading frames of both TAMV (1,452nt) and TcTV-1 NP (1,473 nt) proteins in pCAGGS-P7 vector, generating the TAMV-NP expression plasmid and the TcTV-1-NP expression plasmid fused with HA tag at C-terminus (TcTV-1-NP-HA). We transfected Vero E6 cells with TAMV-NP expression plasmid, TcTV-1-NP-HA expression plasmid, and the control vector pCAGGS-P7. At 48 h after transfection, we conducted IFA using plasmid-transfected cells as described elsewhere (3) with α -TAMV-NP, α -HA, and IFA-positive serum samples from febrile patients. For western blot analyses, we also harvested plasmid transfected cells and blotted them with α -TAMV-NP, α -HA, and α - β -actin.

Sequences and Bioinformatics

We deposited the complete genome sequences of 2 TAMV strains in GenBank under the accession numbers from MT815989 to MT815994. The datasets of sequence reads from 3 pools of mouse brains are publicly available in the NCBI Sequence Read Archive (<http://www.ncbi.nlm.nih.gov>) under the ID PRJNA649646.

We used the coding regions of the L, M, and S segments of viruses from the genus *Orthonairovirus*, family *Nairoviridae*, to construct the maximum likelihood trees. We used sequences of severe fever with thrombocytopenia virus, Rift Valley fever virus, and Hantaan virus as outgroup sequences. We built the maximum likelihood tree of TAMV strains using the 423 nt partial L sequences obtained from PCR products for TAMV-positive tick groups. We

constructed trees using Mega 6.0 (6) and tested them using the bootstrap method with 1,000 replications.

Ethical Statement

Animal inoculation experiments and human serologic investigation were approved by the ethics committee of Wuhan Institute of Virology, Chinese Academy of Sciences under the approval numbers WIVA33201702 and WIVH01201501.

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Appendix Table 1. Summary of tick samples used for mouse inoculation*

Year	Location	Ticks	Tick groups, no.	Mouse groups showing illness at P1	Mouse groups for further inoculation	Mouse groups showing illness at P2
2016	Yuli County	550	11	4	NA	NA
	Karamay City	383	7	1	NA	NA
	Luntai County	250	5	3	1	1
2017	Wujiaqu City	2,940	32	17	6	4
Total		4,123	55	25	7	4

*NA, not applicable; P1, first inoculation; P2, second inoculation.

Appendix Table 2. Summary of RNA pools of mouse brains used for RNA sequencing

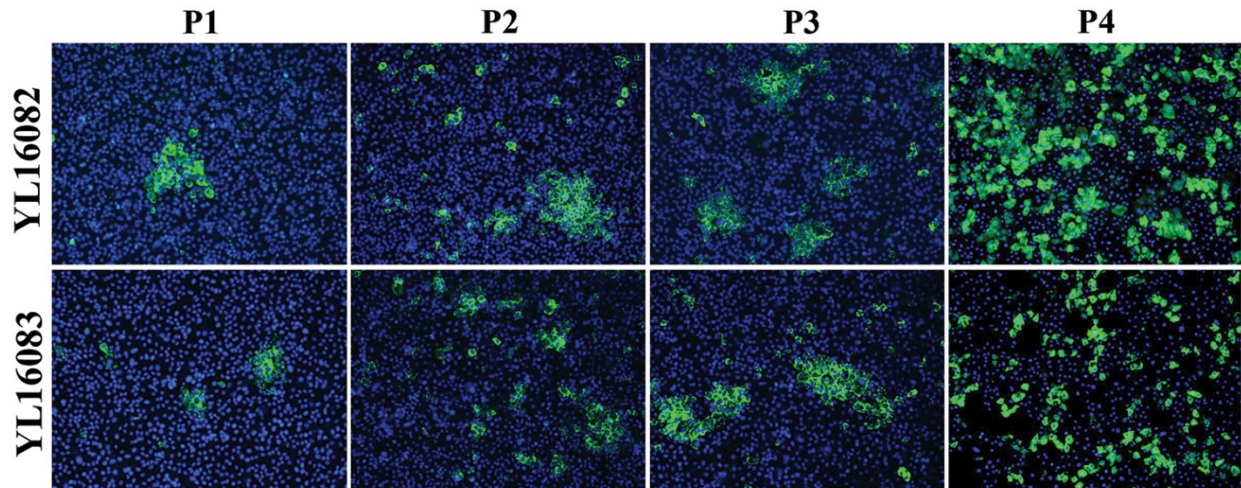
RNA pool	Mouse brains, no.	Location of ticks used to inoculate mice	Generation of inoculated mice	Total reads	TAMV-related reads	nt Sequence identity to TAMV, %
A-M6	4	Yuli County	P1	52,476,380	64,425	94.92–96.96
ML	1	Karamay City	P1	95,862,588	382,724	94.80–96.86
	4	Luntai County	P1 (×3) P2 (×1)			
XJ4	4	Wujiaqu City	P2	48,607,846	69,965	94.75–96.55
Total				196,946,814	517,114	

Appendix Table 3. Personal and clinical characteristics of febrile patients from Bachu County, June 2007*

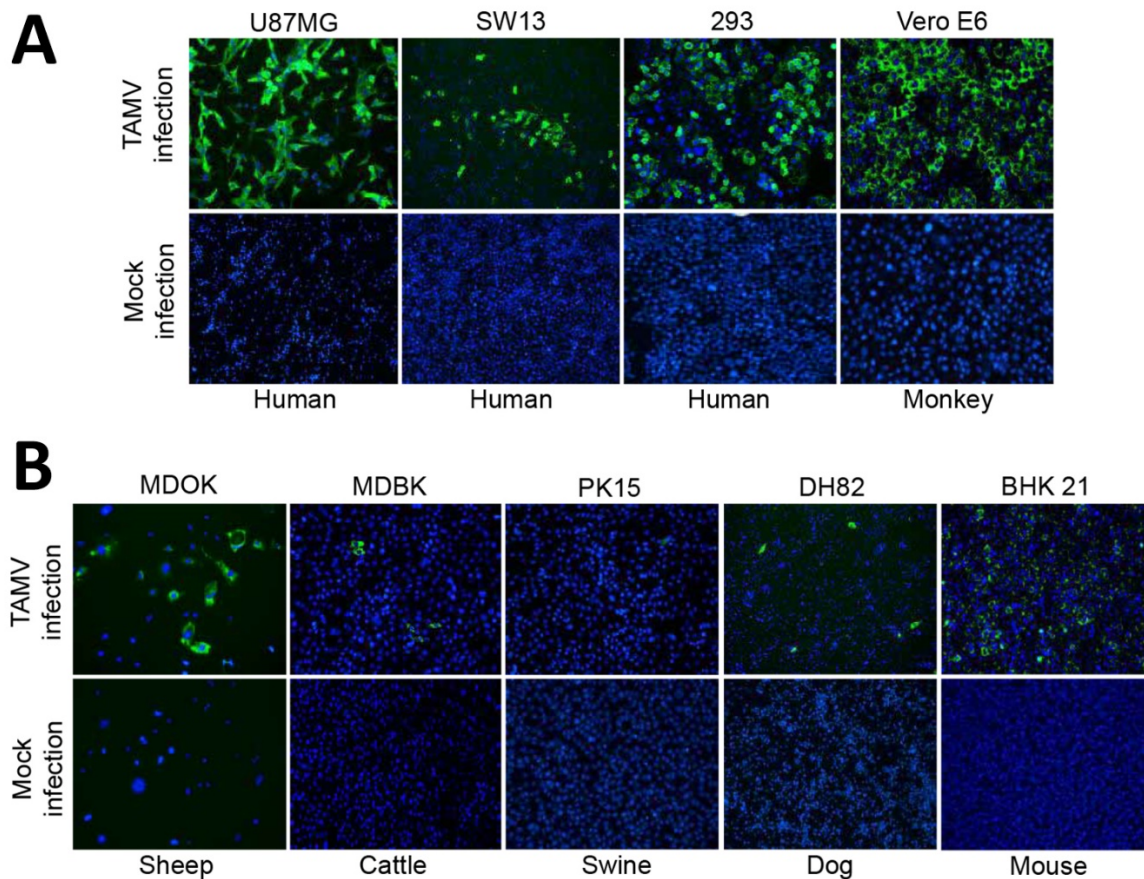
Case	Sex/age, y	Inpatient/outpatient	Clinical diagnosis and parameters	IFA		Neutralization, titer
				TAMV IgM	TAMV IgG	
1	M/30	O	Liver damage	–	–	NA
2	M/45	I	Hyperlipemia	–	–	NA
3	F/50	I	Coronary heart disease; liver damage	–	–	NA
4	F/18	O	Liver damage	–	–	NA
5	M/6	I	Liver damage	+	–	NA
6	F/21	O	Leukopenia (leukocyte: $9.6 \times 10^8/L$); liver damage	+	–	NA
7	F/22	O	NA	+	–	16
8	NA	O	Postpartum infection	+	–	NA
9	F/16	O	NA	–	–	NA
10	NA	O	NA	+	+	NA
11	F/18	O	NA	+	–	NA
12	NA	O	NA	–	+	NA
13	NA	O	NA	–	–	NA
14	F/24	O	NA	–	–	NA
15	M/21	O	NA	+	–	NA
16	F/23	O	NA	–	–	NA
17	F/22	O	NA	–	–	NA
18	M/23	O	NA	+	–	NA
19	M/27	O	NA	+	–	NA
20	M/28	O	NA	+	–	NA
21	M/28	O	NA	+	–	NA
22	NA	O	NA	+	–	NA
23	F/29	I	NA	–	–	NA
24	F/25	I	NA	–	–	NA
25	F/15	O	NA	–	+	NA
26	NA	O	NA	–	–	NA
27	NA	O	NA	–	–	NA
28	NA	O	NA	+	–	NA
29	NA	O	NA	–	–	NA
30	NA	O	NA	–	–	NA
31	NA	O	NA	–	–	NA
32	NA	O	NA	+	–	NA
33	NA	O	NA	–	–	NA
34	M/50	I	NA	–	–	NA
35	NA	O	NA	–	+	NA
36	NA	O	NA	–	+	NA
37	NA	O	NA	–	–	NA
38	NA	O	NA	–	–	NA
39	NA	O	NA	–	–	NA
40	NA	O	NA	–	–	NA
41	NA	O	NA	–	–	NA
42	NA	O	NA	–	–	NA
43	NA	O	NA	–	–	NA

Case	Sex/age, y	Inpatient/ outpatient	Clinical diagnosis and parameters	IFA		Neutralization, titer
				TAMV IgM	TAMV IgG	
44	NA	O	NA	-	-	NA
45	F/21	I	Urticaria	-	-	NA
46	M/60	I	Gastritis	-	-	NA
47	M/41	O	Hyperlipidemia, hyperglycemia, coronary heart disease, renal dysfunction	-	+	NA
48	NA	O	NA	-	+	NA
49	NA	O	NA	-	-	NA
50	F/1	O	Rickets	-	-	NA
51	F/20	O	Suspected hepatitis	-	-	NA
52	M/45	O	Hypertension	-	-	NA
53	M/39	O	Hyperlipidemia, hyperglycemia	-	-	NA
54	F/30	I	Suspected chickenpox	-	-	NA
55	M/50	O	NA	-	-	NA
56	F/35	O	NA	+	-	NA
57	M/71	O	NA	-	-	NA
58	F/75	O	Hypertension	+	-	32
59	F/55	I	NA	-	-	NA
60	NA	O	NA	-	+	NA
61	NA	O	NA	-	+	16
62	NA	O	NA	-	-	NA
63	M/49	I	burn	-	-	NA
64	M/53	O	NA	-	-	NA
65	F/20	O	NA	-	-	NA
66	F/60	O	NA	-	-	NA
67	NA	O	NA	-	-	NA
68	F/40	O	Cholecystitis	-	-	NA
69	NA	I	NA	-	+	NA
70	F/56	I	NA	-	+	32
71	M/15	I	NA	-	+	NA
72	M/70	I	NA	+	+	64
73	M/62	I	NA	-	-	NA
74	F/15	I	NA	-	-	NA
75	NA	I	NA	-	-	NA
76	F/61	I	NA	-	+	NA
77	F/51	I	NA	-	-	NA
78	F/42	I	NA	-	+	NA
79	F/20	I	NA	-	+	NA
80	NA	I	NA	-	+	NA
81	NA	O	NA	-	+	NA
82	NA	O	NA	-	+	16
83	F/55	I	NA	-	+	NA
84	NA	O	NA	-	+	NA
85	NA	O	NA	-	-	NA
86	NA	I	NA	-	-	NA
87	NA	I	NA	-	-	NA
Total	20 M, 29 F, 38NA/1-75	26 I, 61 O	NA	17 (19.5%)	21 (24.1%)	6 (6.9%)

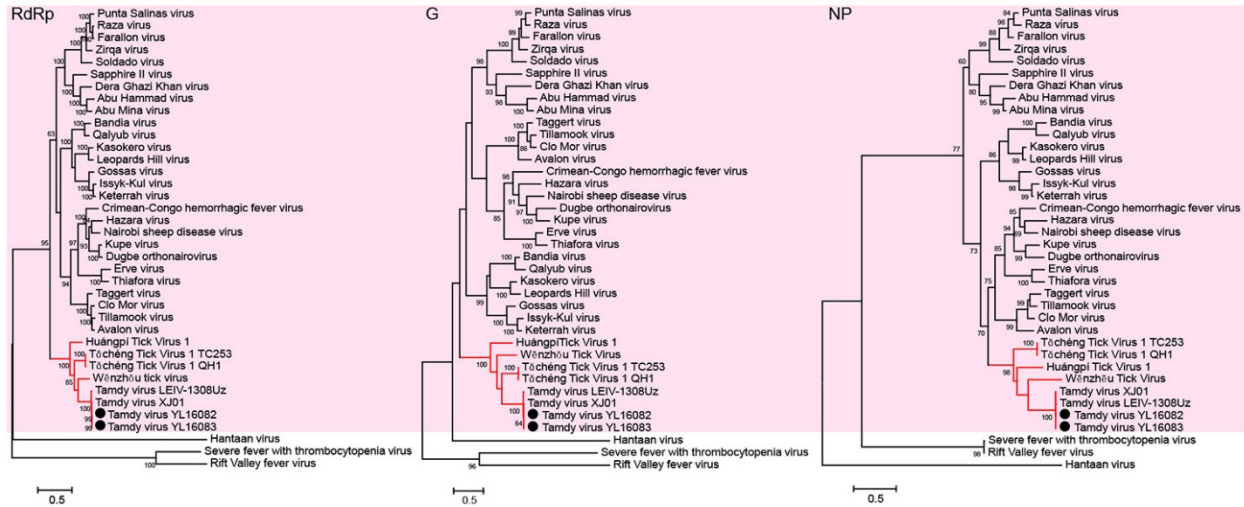
*IFA, immunofluorescence assays; NA, not available; I, inpatient; O, outpatient; TAMV, Tamdy virus; +, positive for antibodies; -, negative for antibodies



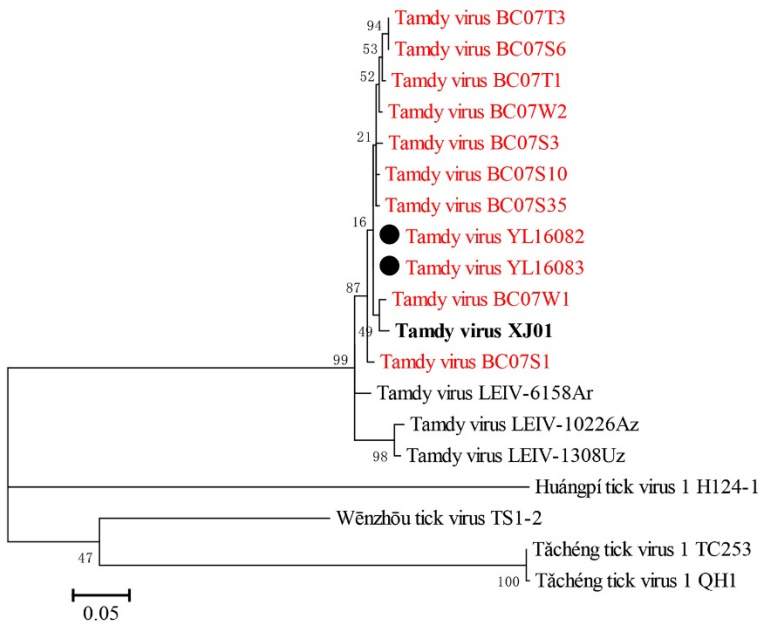
Appendix Figure 1. Isolation of 2 new TAMV strains (YL16082 and YL16083) by incubating clarified homogenates of suckling mouse brains in Vero E6 cells. We monitored productive virus replication using immunofluorescence assays for each passage (P1–P4) with a TAMV-NP antiserum.



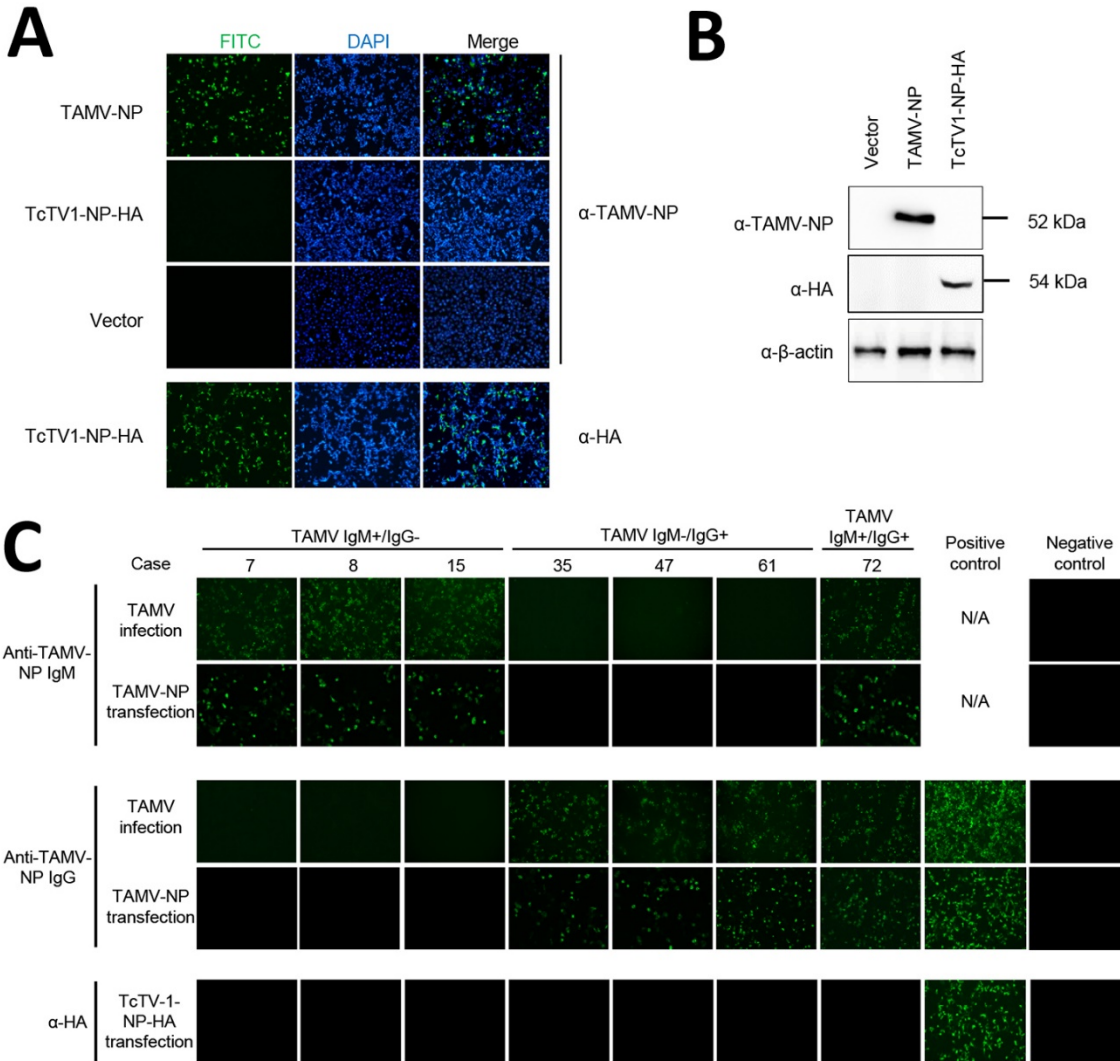
Appendix Figure 2. Infection susceptibility of different cells revealed by immunofluorescence staining. These include 3 different human cell lines and 1 each from monkeys, sheep, cattle, swine, dogs, and mice.



Appendix Table 3. Maximum likelihood phylogenetic trees based on the complete amino acid sequences of the RNA-dependent RNA polymerase (RdRp), glycoprotein (G), and nucleoprotein (NP) of virus members in the genus *Orthonairovirus* (shaded in pink), family *Nairoviridae*. The clade of *Tamdy orthonairovirus* species is shown by red lines. The 2 new TAMV isolates are labeled by black dots. Trees were constructed by bootstrap method of 1000 replicates.



Appendix Figure 4. Maximum likelihood tree based on the partial (423 nt) sequences of the L segment of species *Tamdy orthonairovirus*. TAMV strains identified in this study are indicated by red characters; the 2 isolates (YL16082 and YL16083) are indicated by black dots. TAMV strain XJ01, identified in a recent report (7), is indicated in bold. Trees were constructed using a bootstrap method of 1000 replicates.



Appendix Figure 5. Examination of potential cross-reaction between TAMV NP polyclonal antibodies and Táchéng tick virus 1 (TcTV-1) NP using A) immunofluorescence assays (IFA) and B) western blot. We transfected Vero E6 cells with TAMV-NP expression plasmid, TcTV-1-NP-HA expression plasmid, and control plasmid. At 48 hours after infection, we immune-stained fixed cells using α-TAMV-NP. We immune-stained cells transfected with TcTV-1 NP-HA expression plasmid using α-HA as a control. Transfected cells were harvested and detected by western blot using α-TAMV-NP, α-HA, and α-β-actin. C) Examination of reactivity of serum samples from febrile patients toward TAMV NP and TcTV-1 NP by IFA using virus-infected cells or plasmid-transfected cells. Representative images of serum samples from 7 febrile patients, with 3 IgM-positive samples (case-patients 7, 8, and 15), 3 IgG-positive samples (case-patients 35, 47, and 61), and 1 sample positive for both IgG and IgM (case-patient 72), determined using IFA with TAMV-infected cells. The upper panel shows images from the 7 samples being examined by α-TAMV-NP IgM using both TAMV-infected cells and TAMV NP transfected cells as antigens; the middle panel shows images from the examination of α-TAMV-NP IgG; and the bottom panel shows images from detection of TcTV-1 NP antibodies using TcTV-1-NP-HA transfected cells. NA, not applicable.