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Genetic Identification of a Hantavirus Associated with an Outbreak of Acute Respiratory Illness

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A mysterious respiratory illness with high mortality was recently reported in the southwestern United States. Serologic studies implicated the hantaviruses, rodent-borne RNA viruses usually associated elsewhere in the world with hemorrhagic fever with renal syndrome. A genetic detection assay amplified hantavirus-specific DNA fragments from RNA extracted from the tissues of patients and deer mice (*Peromyscus maniculatus*) caught at or near patient residences. Nucleotide sequence analysis revealed the associated virus to be a new hantavirus and provided a direct genetic link between infection in patients and rodents.

An outbreak of a respiratory illness with high mortality was recently reported in the shared border region (Four Corners) of New Mexico, Arizona, and Colorado in the southwestern United States (1). Patients were defined as having unexplained adult respiratory distress syndrome (ARDS) or acute bilateral pulmonary interstitial infiltrates in the presence or absence of prodromal symptoms (2). Mortality in confirmed patients has been in excess of 75%, frequently in previously healthy adults between 20 and 40 years of age. Serologic surveys of patients failed to detect evidence of agents normally associated with severe respiratory illness but did detect immune cross-reactivity with previously characterized hantavirus antigens (1). This finding was unexpected because hantaviruses had not previously been associated with outbreaks of acute human disease in North America nor had hantaviruses found elsewhere in the world been associated with a

severe, predominantly respiratory illness.

Hantaviruses are rodent-borne viruses belonging to the family Bunyaviridae. They possess a negative sense, single-stranded RNA genome consisting of three segments, designated large (L), medium (M), and small (S), which encode the virus polymerase protein (L), the glycoproteins G1 and G2, and the nucleocapsid (N) protein, respectively (3–7). At least four distinct virus serotypes have been clearly defined that differ in their overall geographic distribution, rodent host, and degree of pathogenicity for humans (8). The Hantaan (HTN) serotype viruses, associated with field mice (*Apodemus agrarius*) and found predominantly throughout Korea, China, and far eastern Russia, cause severe hemorrhagic fever with renal syndrome (HFRS). The Seoul (SEO) serotype viruses are probably found worldwide, which reflects the range of their primary host, *Rattus norvegicus*. The SEO viruses have been associated with a generally more moderate form of HFRS, particularly in Korea and China. Recently, three cases of mild HFRS disease associated with SEO-related virus were described in the United States, although the SEO virus was probably introduced into the United States by colonization of its Eurasian rat host (9). The Puumala (PUU) serotype viruses, found throughout Scandinavia and Europe west of the Ural Mountains, are associated with a relatively mild form of HFRS. The primary

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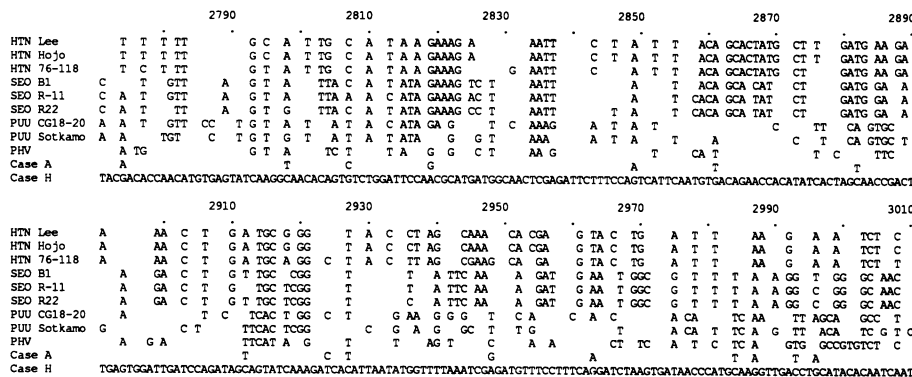


Fig. 1. Comparison of nucleotide sequences of a hantavirus associated with ARDS in the Four Corners region. The nucleotide sequence of 241 bp of the PCR product derived from the two representative hantaviruses detected in New Mexico (case H) and reported in Arizona ARDS cases was compared with the same genome region of previously characterized hantaviruses (11). Sequence differences are shown relative to the case H sequence. All sequences are in the viral complementary DNA (+) sense and numbered relative to the overall hantavirus sequence alignment (nucleotides 1 to 3722, including gaps).

rodent host is the bank vole (*Clethrionomys glareolus*). The fourth serotype, Prospect Hill (PH), is known from only two virus isolates, both of which are from indigenous North American rodent species. The original isolate was from meadow voles (*Microtus pennsylvanicus*) in Frederick, Maryland, and the other from Minnesota. PH cross-reactive antibodies have been found in other North American microtine and cricetid rodents and in humans involved in trapping mammals (10), but PH has not been associated with any human disease. Other, less well characterized hantaviruses, such as Thottapalayam from a shrew in India, may represent additional serotypes.

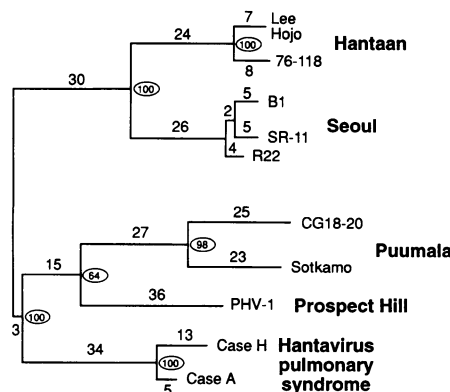


Fig. 2. Phylogenetic relation of the ARDS-associated hantavirus from the Four Corners region (bottom) to previously characterized hantaviruses. Phylogenetic analysis of the virus sequence differences within 241 bp of the amplified fragment was performed by the maximum parsimony method (12). The horizontal distances represent the number of nucleotide step differences (indicated adjacent to the lines) present between branch nodes and taxa (that is, viruses). Bootstrap confidence limits exceeding 50% are indicated in ovals next to each branch node.

The two pairs of hantavirus serotypes, HTN and SEO and PUU and PH, each share considerable nucleotide sequence similarity (approximately 70%). Precise regions of sequence conservation within the G2 protein coding region of the M segment of the virus genome were identified, and deoxyoligonucleotide primers were designed for detection of small amounts of hantavirus of either pair of serotypes by a nested reverse transcriptase-polymerase chain reaction (RT-PCR) assay (11). The first-round primers were expected to amplify a DNA fragment that contained predicted conserved sequence targets that could be used in a nested PCR by a second pair of PCR primers to give additional specificity and sensitivity to the detection assay. Specific nested PCR DNA products of the correct size [278 base pairs (bp)] were obtained with only the PUU-PH primer set

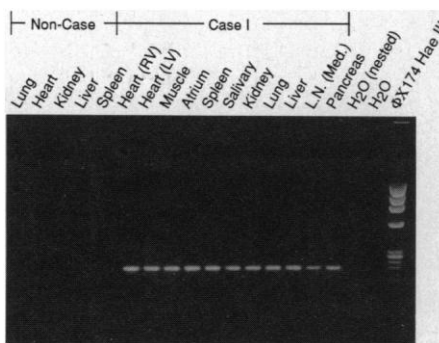


Fig. 3. Agarose gel electrophoresis analysis of hantavirus PCR DNA products from nested RT-PCR assays of total RNA extracted from ARDS patient autopsy specimens. Shown is a hantavirus-specific DNA band, 172 bp in length, amplified from total RNA from various tissues from case I. RV and LV, heart right and left ventricles, respectively; L.N. (Med.), medial lymph nodes.

when used with total RNA extracted from tissues from several of the outbreak patients (that is, nested HTN-SEO primer reactions were negative, as were control autopsy materials). Amplified DNA from the tissues of two representative ARDS patients (case H in New Mexico and case A in Arizona) was extracted from the agarose gel and analyzed with an automated thermocycle sequencing technique (Applied Biosystems, Foster City, California) with the same primers used for PCR product synthesis.

The DNA bands were found to contain hantavirus-like sequences (Fig. 1), but their nucleotide sequence differed from that of any of the known hantaviruses by at least 30%. A 7% nucleotide difference was also observed between PCR bands from case H and those from case A. Phylogenetic analysis of the sequence differences by the maximum parsimony method (12) indicated that the hantavirus associated with ARDS in the Four Corners region was novel, representing a distinct lineage, and was most closely related to PH, the only other hantavirus recovered from rodent species indigenous to North America (Fig. 2). This relation is supported by the serologic cross-reactivity obtained with convalescent-phase sera from patients (13).

On the basis of sequences derived from the two representative hantaviral ARDS cases, we designed a new second-round (nested) pair of PCR primers [+2816 AA-GGTAACACAGT(G/C)TCTGGATTC and -2955 GGTTATCACTTAGATC(C/T)TGAAAGG]. A broader screening of autopsy specimens was then carried out by an RT-PCR assay with the original first-round primers followed by the new second-round primers, which generate a specific, 172-bp DNA band. Frozen-tissue autopsy specimens were available from 10 patients who met the case definition (2). When tested with the new primer set, all 10 patients were found to be PCR positive (Fig. 3). Lung, heart, liver, kidney, and spleen tissues were frequently found to be positive. At the time of death, the virus was apparently distributed extensively throughout the body. Currently, the site of virus replication is unknown, and no precise quantitation of virus in different tissues has been performed. However, immunohistochemical analysis of tissues with a monoclonal antibody that cross-reacts with conserved hantavirus nucleoprotein epitopes revealed widespread endothelial involvement, with deposition of antigen in lung, kidney, heart, pancreas, adrenals, and skeletal muscle (14). The pathophysiologic role of the endothelial involvement in the increased vascular permeability, which is a hallmark of this syndrome, remains to be elucidated.

A systematic rodent trapping effort initiated this year in the Four Corners region

allowed a calculation of the relative density of the rodent population. The deer mouse (*Peromyscus maniculatus*) was the dominant rodent species, accounting for 63% of all captures. Initial serologic testing (by immunoglobulin G enzyme-linked immunosorbent assay) of all small rodents captured at or near case and control houses indicated an extremely high seropositivity to PUUPH antigens (approximately 33%) among *P. maniculatus* (15). Of 54 seropositive *P. maniculatus* tested, 44 (82%) gave a positive PCR band, which indicates a high persistence of the virus in this rodent reservoir even in the presence of a substantial humoral immune response. Similar results were recently obtained with PCR analysis of seropositive, SEO virus-infected rats (16). Indeed, the presence of virus with persistent shedding in excreta, concurrent with the presence of circulating antibodies, is a hallmark of hantaviral infection in other rodent species (17). In addition, of 45 seronegative *P. maniculatus* tested, 14 (31%) were PCR positive. These likely represent recently infected individuals that have not yet mounted an immune response or, alternatively, may indicate a greater sensitivity of detection of virus infection by PCR. These data suggest that the virus can be maintained and actively transmitted in this rodent species and implicate *P. maniculatus* as an important virus reservoir. Hantaviral infection of other rodent species (*P. truei*, *P. boyleyi*, *Tamias dorsalis*) in this region could also be detected but to a much lesser extent (15).

Nucleotide sequence analysis was performed on PCR products obtained from each of the PCR-positive patients and from PCR-positive rodents captured at or in the vicinity of the residences of the patients. Despite the fact that the region of the genome (within the G2 coding region) targeted for amplification is generally highly conserved among hantaviruses and that virus-infected samples were collected over a period of less than 2 months from a limited geographic area, a surprisingly high level of genetic diversity was observed among the samples (Fig. 4). However, identical sequences were consistently obtained from independent RNA preparations from the same tissue or PCR products from different tissues within an individual.

Comparison of the human and rodent sequences demonstrated a direct genetic link between the virus in infected *Peromyscus* rodents and the virus in the human hantaviral ARDS cases. Viral sequences from two of the cases (G and H, from Pinehill/Ramah, New Mexico) were genetically closely clustered. The viral sequence of case G was identical to that obtained from three *P. maniculatus* captured at or near the patient's residence. Similarly, the viral sequence of case H was identical to those of two *P. maniculatus* trapped at or near the patient's residence. These results suggest that these individuals may have acquired infection in or near their residences. A third genetically closely related patient (case J) was from 35 miles northeast, in the Crownpoint area. However, an

additional patient (case I), from Black Mesa, Arizona, was also contained within this genetic cluster.

Two other patients (cases B and C) were from southwestern Colorado. Their viral sequences were closely related to one another and to those from rodents captured nearby but were distinct from the New Mexico clusters. Case A was officially recorded as an Arizona case. This individual had resided in the vicinity of Springerville, Arizona, 11 days before the onset of illness in nearby Snowflake, Arizona (and death 3 days later), but had resided in Hesperus, in southwestern Colorado, before this trip. The viral sequence from this patient was found to be closely related to that obtained from other human and rodent samples from Colorado and identical to that of a *P. maniculatus* captured near the patient's residence in Hesperus, which suggests that this individual acquired the infection in Colorado before traveling to Arizona.

On the basis of the limited sequence region analyzed, cases D, E, and F and additional rodents do not appear to form a geographic cluster. A clearer picture may emerge as these studies are extended to other regions and segments of the virus genome. However, these initial results of extensive virus genetic diversity obtained over a small sampling period, together with examples of geographic clustering of virus variants, suggest that this virus may have been independently maintained for some time in rodent populations in multiple areas of the southwestern United States. More extensive studies of virus and rodent populations will be needed to confirm this hypothesis and to gain a better understanding of the epidemiology and public health impact of this disease. As *P. maniculatus* represents one of the most abundant small mammals in North America, it is of particular importance to determine the potential role it may play as a reservoir for hantaviruses associated with severe ARDS.

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11. The complete M genome segment nucleotide se-

Case	Rodent	Location									
1	A	Snowflake	AZ*	g	t	t	c				
2	1 (A)	Hesperus	CO	g	t	t	c				
3	2	Grand Junction	CO	g	t	t	c				
4	B	Cortez	CO	g	t	t t	g				
5	C	Grand Junction	CO	g	c	t	t	g	g		
6	3	Grand Junction	CO			t	t	g	g	c	
7	D	Sedan	NM	g	a						
8	E	Farmington	NM	g	a						
9	4 (H)	Pinehill/Ramah	NM			t	c				
10	5	Springerville	AZ			t	c				
11	6	Springerville	AZ			t	c				
12	F	Alpine Ranchero	AZ	g		t	c				
13	7	Black Mesa	AZ	g			c				
14	G	Pinehill/Ramah	NM			g t					
15	8 (G)	Pinehill/Ramah	NM			g t					
16	9 (G)	Pinehill/Ramah	NM			g t					
17	10 (G)	Pinehill/Ramah	NM			g t					
18	H	Pinehill/Ramah	NM			g	c	c	a		
19	11 (H)	Pinehill/Ramah	NM			g	c	c	a		
20	12 (H)	Pinehill/Ramah	NM			g	c	c	a		
21	13 (G)	Pinehill/Ramah	NM			g	c	c			
22	I	Black Mesa	AZ			g t	c	c			
23	J	Crownpoint area	NM			t g	c	c			
24	14	Black Mesa	AZ	g	a	t	a	c	t	g	
				Consensus: A C T C A C G T C C A A T A T T T G T							
				2	2	2	2	2	2	2	2
				8	8	8	8	8	8	9	9
				1	3	4	4	5	6	7	7
				9	5	3	6	9	5	1	3
				6	2	8	6	2	8	6	2
				1	4	7	5	8	4	5	8

Fig. 4. Nucleotide sequence differences among the PCR products derived from hantaviral ARDS patients from the Four Corners and from rodents trapped in the region. Shown is a genetic cluster analysis (18) of 139 bp of virus sequences relative to the consensus sequence. Nucleotide difference positions are numbered as in Fig. 1. The asterisk indicates that the infection was acquired in Colorado (see text).

quences of hantaviruses Hantaan strains 76118 (GenBank numbers M14627 and Y00386), Lee (D00377), and Hojo (D00376); Seoul strains B-1 (X53861), SR-11 (M34882), and R22 (S68035); Puumala strains CG18-20 (M29979) and Sotkamo (X61034); and Prospect Hill strain PHV-1 (X55129) were aligned with the GAP, PILEUP, and LINEUP programs of the GCG software package (Genetics Computer Group, Madison, WI) run on a VAX computer. Predicted conserved positions for the synthesis of nested RT-PCR oligonucleotide primers for HTN-SEO viruses or PUU-PH viruses were as follows. HTN-SEO first-round primers: +2548 GATATGAATGATTG(T/C)TTTGT and -2859 CCATCAGGGTCT(T/C)TCCA; second-round: +2590 TGTATAATTGGGAC(T/A)GTATCTAA and -2751 GCAAAGTTACATTT(T/C)TTCCT (position numbering of the oligonucleotide 3' terminus was relative to the total aligned and gapped sequence length of 3722 nucleotides); PUU-PH first-round primers: +2671 TTTAAGCAATGGTG(C/T)ACTAC(T/A)AC and -3108 CCATAACACAT(AT)GCAGC; second-round: +2770 AGAAAGAAATGTGCATTTGC and -3012 CCTGAACCCCATGC(AT/C)CCA-TC. Because of the hazardous nature of the agent,

all steps of the homogenization of autopsy tissue samples and the total RNA extraction and purification were performed under biosafety level 3 conditions. RNA extraction, first-round RT-PCR reactions, and subsequent product DNA gel electrophoresis analysis were performed essentially as described [L. L. Rodriguez, G. J. Letchworth, C. F. Spiropoulou, S. T. Nichol, *J. Clin. Microbiol.* **31**, 2016 (1993)], except the following cycle profile run on a Perkin-Elmer 9600 thermocycler was used: 41°C for 1 hour, followed by 40 cycles at 94°C for 40 s, at 38°C for 45 s, and at 72°C for 60 s. Second-round reaction conditions used 3% of the first-round reaction product, no RT step; the following profile was used: 35 cycles at 94°C for 40 s, at 43°C for 45 s, and at 72°C for 60 s.

12. PAUP: Phylogenetic Analysis Using Parsimony, version 3.0 s; D. L. Swofford (Illinois Natural History Survey, Champaign, 1991). The software was run on a SUN SPARC 10 workstation. Two equally parsimonious trees were obtained (with minor branch order differences among the SEO viruses) with the use of the BANDB option. Bootstrap confidence limits were obtained with 10,000 repetitions, and the ALLTREES option was used to

examine the tree-length frequency distribution.

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