Genetic Characterization of Venezuelan Equine Encephalitis Virus from Bolivia, Ecuador and Peru: Identification of a New Subtype ID Lineage

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Abstract

Venezuelan equine encephalitis virus (VEEV), a member of the family Togaviridae genus Alphavirus, has been responsible for hundreds of thousands of human and equine cases of severe disease in the Americas. A passive surveillance study was conducted in Peru, Bolivia and Ecuador to determine the arboviral etiology of febrile illness. Patients with suspected viral-associated, acute, undifferentiated febrile illness of <7 days duration were enrolled in the study and blood samples were obtained from each patient and assayed by virus isolation. Demographic and clinical information from each patient was also obtained at the time of voluntary enrollment. In 2005–2007, cases of Venezuelan equine encephalitis (VEE) were diagnosed for the first time in residents of Bolivia; the patients did not report traveling, suggesting endemic circulation of VEEV in Bolivia. In 2001 and 2003, VEE cases were also identified in Ecuador. Since 1993, VEEV has been continuously isolated from patients in Loreto, Peru, and more recently (2005), in Madre de Dios, Peru. We performed phylogenetic analyses with VEEV from Bolivia, Ecuador and Peru and compared their relationships to strains from other parts of South America. We found that VEEV subtype ID Panama/Peru genotype is the predominant one circulating in Peru. We also demonstrated that VEEV subtype ID strains circulating in Ecuador belong to the Colombia/Venezuela genotype and VEEV from Madre de Dios, Peru and Cochabamba, Bolivia belong to a new ID genotype. In summary, we identified a new major lineage of enzootic VEEV subtype ID, information that could aid in the understanding of the emergence and evolution of VEEV in South America.

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Introduction

Venezuelan equine encephalitis virus (VEEV), a member of the family Togaviridae genus Alphavirus, has been responsible for outbreaks involving hundreds-of-thousands of equine and human cases of severe disease in the Americas [1]. At least 14 varieties within the six VEE subtype complex of alphaviruses have been recognized. Only subtype I varieties AB and C have caused major epizootics/epidemics, whereas subtypes II through VI and subtype I varieties D, E and F are enzootic strains that are generally avirulent in horses, but capable of causing human disease [2].

In Ecuador, VEE was first confirmed in 1944 when the virus was isolated from the blood of a sick horse [3]. However, clinical cases that were compatible with VEEV infection had been observed as early as 1940. In 1958, VEEV neutralizing antibodies were found in sera of inhabitants of the Pacific coastal region of Ecuador [4]. In 1968–1969, a large outbreak involving more than 30,000 equids was reported, and in 1975–1977, field work investigations in Ecuador yielded VEEV isolates that were characterized genetically as the Southwestern Colombia/Ecuador ID genotype [5,6].

In Peru, VEEV was first isolated in the 1940s when subtype IAB caused epizootics and epidemics along the Peruvian coast [7,8]. Field work investigations were conducted afterwards in the Amazon region of Peru, resulting in the isolation of 11 VEE complex alphavirus strains [9,10]. Ten isolates were later identified as subtype ID VEEV, whereas one strain was identified as a (new) subtype IIIC virus [9,10,11]. Evidence of VEEV human infections was only obtained in 1993–1995, when subtype ID was isolated from febrile patients residing in the Amazon region of Peru [12,13].

In 2000, in collaboration with the Ministries of Health of Bolivia, Ecuador and Peru, a passive surveillance study was initiated with the purpose of investigating the etiology of febrile illness. As part of the surveillance activities in Peru, several VEEV strains were obtained from febrile patients. Genetic analyses of
Author Summary

Venezuelan equine encephalitis virus (VEEV) has been responsible for hundreds of thousands of human and equine cases of severe disease in the Americas. In 2005–2007, cases of Venezuelan equine encephalitis (VEE) were diagnosed for the first time in residents of Bolivia; the patients did not report traveling, suggesting endemic circulation of VEEV in Bolivia. In 2001 and 2003, VEE cases were also identified in Ecuador. We characterize recent VEEV from Bolivia, Ecuador and Peru and compared their relationships to strains from other parts of South America. We found that most VEEV from Peru grouped within a particular genetic lineage known to circulate in Panama and Peru whereas the VEEV circulating in Ecuador belong to a genetic lineage that circulates in Colombia and Venezuela. Importantly, the VEEV from Madre de Dios, Peru and Cochabamba, Bolivia belong to a new genetic lineage. This finding could aid in the understanding of the emergence and evolution of VEEV in South America and underscores the need for continuous monitoring for VEEV activity.

Materials and Methods

Study subjects

The study protocols were approved by the Ministry of Health of the participant countries and the Naval Medical Research Center Institutional Review Board (protocols NMRC.D.2000.0006, NMRC.D.2001.0002, NMRC.D.2000.0008) in compliance with all applicable Federal regulations governing the protection of human subjects. The study subjects were patients who presented with a diagnosis of acute, febrile undifferentiated illness in their home or at military or civilian outpatient clinics at the specimen collection study sites described below. A signed consent form was obtained from each volunteer after they were informed about the study and a standardized questionnaire was used to obtain demographic and clinical information from each patient at the time of voluntary enrollment. Travel history information was also recorded. The criteria for inclusion in the program have been described previously and consist of fever 38°C or higher and no more than 7 days duration accompanied by headache, myalgia, or other nonspecific symptoms such as ocular and/or joint pain, generalized fatigue, cough, nausea, vomiting, sore throat, rhinorrhea, difficulty breathing, diarrhea, bloody stool, jaundice, dizziness, disorientation, stiff neck, petechiae, ecchymoses, bleeding gums or nose [12,13]. During the acute phase of illness blood samples were obtained from each patient, and when possible, convalescent samples were obtained 10 days to 4 weeks later for serological studies.

Study sites

Human VEEV isolates included in this study were obtained from the specimen collection sites in Peru located in the city or around Iquitos and Yurimaguas in the Department of Loreto and Puerto Maldonado in Madre de Dios. Iquitos is a city of about 400,000 residents located in the Amazon River basin in the Department of Loreto approximately 120 meters above sea level. Yurimaguas is located at the confluence of the Huallaga and Paranapura rivers in the steamy rainforests of northeastern Peru with a population of about 63,000 inhabitants.

Puerto Maldonado is one of the most important cities of the southern jungle. It sits on the banks of the Madre de Dios river, which connects it with Rivera Alto in Bolivia and with Assis, in Brazil (242 km). Puerto Maldonado is a city of approximately 25,000 inhabitants and is located about 256 meters above sea level.

VEEV isolates were also obtained from Eterazama, Cochabamba department in Bolivia. Eterazama is located approximately 450 meters above sea level. At the time of the 2001 census, it had a population of about 2,500 people.

Two VEEV strains were obtained from Shell and Puyo located in the Pastaza province of the Amazon River basin of Ecuador. Puyo is located approximately 950 meters above sea level and in 2006 it had a population of about 25,000 inhabitants. Shell is a town of 5,000 people located 5 km from Puyo. Figure 1 shows the geographic distribution of the sites with confirmed VEE cases identified as part of this febrile surveillance study.

Virus isolation

Patient specimens were sent to the biosafety level 3 containment laboratory at NMRC. Sera were diluted 1:10 in Eagle's minimum essential medium (EMEM) supplemented with 2% fetal bovine serum, 200 μg of streptomycin, and 200 U/ml of penicillin. Diluted sera were then inoculated into monolayers of confluent African green monkey kidney epithelial cells (Vero) and Aedes albopictus mosquito (C6/36) cells. Slides were prepared from the infected cells and an immunofluorescence assay (IFA) was performed using polyclonal antibodies against several arboviruses endemic in Peru [12,13,14,15,16,17]. A variety of arboviruses such as dengue, Oropouche, Mayaro, group C, yellow fever and guaroa were isolated from these samples, including the human VEEV isolates listed in Table 1 that were selected for sequencing.

RNA extraction, RT, and PCR amplification

Viral RNA was extracted using the QIAamp viral RNA mini kit (Qiagen, Valencia, CA) or Trizol reagent (Invitrogen, Carlsbad, California) following the manufacturer's protocols. The reverse transcription (RT) reaction was done using 1× RT buffer, 0.2 mM dNTPs, 1 μM of primers, 80 U of Rnasin ribonuclease inhibitor (Promega, Madison, WI), 1 mM of dithiothreitol (DTT), 200 U of SuperScript reverse transcriptase (Invitrogen), and 5 μl (1/10th) of the extracted RNA. The reactions were incubated at 42°C for 1 hr. The polymerase chain reaction (PCR) included 1× PCR buffer, 0.2 mM dNTPs, 1 μM of primers, 3 mM of MgCl2, 2.5 U of GoTaq DNA polymerase (Promega, Madison, WI) and 20% of RNA. The conditions for the PCRs included incubation at 95°C for 2 min, 35 cycles of 95°C for 30 sec, 48°C for 30 sec, 72°C for 1 min. A final extension of 72°C for 10 min was used to ensure complete double-stranded DNA synthesis. The primers used for the PCR amplification and sequencing reaction have been previously described [14,18].
Sequencing and phylogenetic analyses

Purified PCR products were sequenced directly, and sequencing analyses of the PCR products was performed using an Applied Biosystems (Foster City, CA) Prism automated DNA sequencing kit according to the manufacturer’s protocol. Deduced amino acid sequences were aligned using the ClustalW algorithm in the MacVector version 9.0 software package (MacVector, Inc., Cary, NC), and the nucleotide sequences were aligned manually based on codon positional homology and compared to VEEV sequences from previously published studies available in the genbank database [5,14,15,19]. Phylogenetic analyses were performed using the neighbor joining, maximum parsimony, and maximum likelihood algorithms implemented in the PAUP* version 4.0 software package [20,21]. The outgroup consisted of homologous sequences of 4 major lineages of eastern equine encephalitis virus (EEEV). For the neighbor joining analysis, the HKY85 distance formula was used, and bootstrap analyses [22] were performed with 1,000 replicates to place confidence values on the nodes within trees. For the maximum parsimony analysis, the heuristic algorithm was employed.

For maximum likelihood analysis, the general time-reversible (GTR) model of nucleotide substitution was used, with a proportion of 0.255 nucleotide sites being invariable and a gamma distribution among-site rate variation (alpha shape parameter) of 0.726. The starting tree in the analysis was found using neighbor joining, which was followed by successive rounds of tree bisection reconstruction branch-swapping, identifying the maximum likelihood substitutions parameter at each stage until the tree of highest likelihood was found. Bootstrapping was subsequently performed to assess the robustness of tree topologies using 1,000 replicate neighbor joining trees under a maximum likelihood substitution model.

Antigenic characterization

To evaluate antigenic differences between the VEEV subtype ID Panama/Peru and Peru/Bolivia genotypes, convalescent sera
from patients from Madre de Dios (infected with subtype ID Peru/ Bolivia genotype) and Loreto, Peru (infected with subtype ID Panama/Peru genotype) and Cochabamba, Bolivia (infected with subtype ID Peru/Bolivia genotype) were tested against homologous and heterologous strains of VEEV. Most convalescent sera were obtained 2 to 4 weeks after infection; however, one sample was collected 7 years after VEE infection.

Samples were processed using a previously described plaque reduction neutralization test (PRNT) [23]. Briefly, sera were heat-inactivated at 56°C for 30 min and 2-fold dilutions were mixed with 100 PFU of virus and incubated at 4°C overnight. The mix was added onto a monolayer of Vero cells and incubated at 37°C for 1 hr before adding an overlay of 0.4% of agarose in EMEM. After 48 hr, plates were stained with 0.25% crystal violet in 20% methanol and plaques were counted. The PRNT titer presented in Table 2 is the reciprocal of the highest serum dilution capable of neutralizing 80% of approximately 100 plaque-forming units (PFU) of virus. The traditional serological criteria was used to determine whether the viruses were antigenically distinguishable (i.e. at least fourfold difference between the homologous and heterologous titers of both, or one but not both of the two sera tested) [24].
Genetic characterization of VEEV isolates

In order to determine the genetic relationship among the new VEEV isolates from Bolivia, Ecuador, and Peru and the strains from other areas in South America including Colombia and Venezuela, RT-PCR amplifications and sequencing of the partial PE2 gene (815 bp) were carried out. This genome region was chosen because there is an extensive GenBank database of sequences available for comparison, and because this area undergoes critical amino acid substitutions that are associated with VEE epizootic/epidemic emergence [14,15,18,26,27]. Recent VEEV strains were selected for sequencing based on date of collection and geographic origin. Overall, the phylogenetic trees that were generated in this study, using neighbor joining, maximum parsimony, and maximum likelihood methods, had identical topologies except for some groupings between subtypes III and V (Figure 2). Twenty-four isolates from Peru grouped with the Panama/Peru ID genotype with strong bootstrap support, and no evidence of circulation of the Colombia/Venezuela ID genotype in Peru was observed among the recent Peruvian isolates.

Interestingly, the 12 strains from Puerto Maldonado, Madre de Dios in Peru and Eterazama (Cochabamba) in Bolivia formed a distinct clade within subtype ID that is a sister to the Colombia/Venezuela and the Peru/Panama genotypes. Sequence comparisons between the Panama/Peru and Bolivia/Peru ID genotype viruses revealed about a 3% difference at the nucleotide level and a 0.4% difference at the amino acid level whereas the Colombia/Venezuela and Peru/Panama genotypes are known to differ by about 5% at the nucleotide level and 0.8% at the amino acid level [14]. A 1.7% nucleotide difference was observed between the Colombia/Venezuela and Bolivia/Peru genotypes.

The two Ecuadorian strains that were obtained from patients in 2001 and 2003 grouped within the Colombia/Venezuela ID genotype, which also includes the epidemic/epizootic subtype IAB and IC strains. Previous isolates of ID genotype viruses from Ecuador have grouped with the Southwest Colombia/Ecuador ID genotype [5]. However, these isolates were collected on the western side of the Andes mountains, while the two new isolates were collected from the eastern side of this mountain range, in the Amazon basin.

Antigenic characterization of the Panama/Peru and Peru/Bolivia ID genotypes

Although the Panama/Peru and the new Bolivia/Peru ID genotypes differ by only 3% at the nucleotide level, we examined the possibility that they are antigenically distinguishable. To test this possibility, convalescent sera from patients infected with either the Panama/Peru ID genotype or the new Bolivia/Peru ID genotype strain were tested for their ability to neutralize homologous and heterologous viruses. The results demonstrated that the convalescent sera from these patients equally neutralize both genotype strains (Table 2), suggesting that these strains are not antigenically different. Because the convalescent sera were obtained 2 weeks to a month after infection, we also included in the analyses serum from a volunteer who contracted VEE infection (with the Panama/Peru genotype strain) 7 years before. The results obtained with this serum also failed to antigenically discriminate between the Panama/Peru vs the Peru/Bolivia genotypes. Overall, the VEE neutralizing titers range between 20 to 640 (Table 2).

The lack of convalescent sera from the Colombia/Venezuela genotype strains prevented us from performing similar testing. We did not attempt to produce antibodies in animals to test antigenic differences.

Discussion

VEEV continues to cause sporadic outbreaks of severe febrile disease in South America. In 2005, cases of VEE were detected for the first time in Eterazama, Cochabamba department in Bolivia.
Prior to this report, there was no evidence of VEEV circulation in this country, and more importantly, there was no proof that VEEV was responsible for human illness in Bolivia. The continuous isolation of VEEV from Bolivian (2005–2007) patients with no report of traveling suggests endemic circulation of the virus in Cochabamba. Because our surveillance activities in Cochabamba, Bolivia were initiated only in 2005, it is difficult to assess how long the virus has been present in this area. In addition, the limited extent of our surveillance activities in Bolivia that includes only the Beni, Cochabamba and Santa Cruz departments may be preventing us from detecting VEEV cases in other areas in Bolivia.

Surveillance activities in Madre de Dios, Peru began in 2004 but it was not until 2005 that the first VEE human cases were reported. The construction of the Interoceanic Highway that began in 2000 and is scheduled for completion in 2010 has disrupted the ecology in Madre de Dios (where most of the construction activities are currently undergoing) and cause an increase in forest disturbance [28]. The highway, which creates a coast to coast trucking route between the coastal cities of Ilo, Matarani and Marcona in Peru and the Brazilian ports of Rio de Janeiro and Santos, is therefore possibly causing the emergence of new viruses in the area. In addition to the highway, gold mining extraction practices have also intensified during recent years, which is probably another key factor for the emergence of VEE and other arboviruses in Madre de Dios. Only between 2000 and 2005, Peru had the world sixth highest loss of old-growth forests, losing 224,600 hectares per year [28].

In Ecuador, VEEV was first detected in 1944 when the virus was isolated from the blood of a sick horse [3]. In 1968–1969, a large outbreak involving more than 30,000 equines was reported in Ecuador, which later extended to Central America, Mexico and...
in these areas.

Examine the possibility of emergence of epizootic viruses.

VEEV subtype ID, the Bolivia-Peru ID genotype. This information could aid in the understanding of the emergence and evolution of VEEV in South America. Further ecological and surveillance activities are needed in Madre de Dios, Peru and Bolivia to identify the vectors and reservoir host(s) involved in transmission and to determine the public health impact and distribution of VEEV in the region. In addition, field investigations are needed to examine the possibility of emergence of epizootic strains of VEEV in these areas.

Supporting Information

Alternative Language Abstract S1 Translation of the abstract into Spanish by PVA. Found at: doi:10.1371/journal.pntd.0000514.s001 (0.03 MB DOC)

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The sequences generated in this study were deposited in GenBank (accession numbers GQ336464-GQ336482 and GQ358223-GQ358224).

Author Contributions

Conceived and designed the experiments: PVA APA VS LB JV SM SCW TJK. Performed the experiments: PVA APA AD XL. Analyzed the data: PVA APA LB JV SCW. Contributed reagents/materials/analysis tools: VS LB JV VF SM JF WRE YR SCW TJK. Wrote the paper: PVA APA SCW TJK.

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