Malpais spring virus is a new species in the genus vesiculovirus

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Abstract

**Background:** Malpais Spring virus (MSPV) is a mosquito-borne rhabdovirus that infects a variety of wild and feral ungulates in New Mexico, including horses and deer. Although, initial serologic tests and electron microscopy at the time of isolation nearly 25 years ago provided evidence that MSPV is a novel virus, possibly related to vesiculoviruses, the virus still has not been approved as a new species.

**Findings:** Use of the illumina platform allowed us to obtain the complete genome of MSPV. Analysis of the complete 11019 nt genome sequence of the prototype 85-488NM strain of MSPV indicates that it encodes the five common rhabdovirus structural proteins (N, P, M, G and L) with alternative ORFs (> 180 nt) in the N, M and G genes, including a 249 nt ORF in the G gene predicted to encode a 9.26 kDa highly basic transmembrane protein. Although antigenically very distant, phylogenetic analysis of the L gene indicates that MSPV is most closely related to Jurona virus, also isolated from mosquitoes in Brazil, as well as a number of other vesiculoviruses.

**Conclusions:** In sum, our analysis indicates MSPV should be classified as a member of the genus *Vesiculovirus*, family *Rhabdoviridae*. The complete genome sequence of MSPV will be helpful in the development of a reverse genetics system to study the unique aspects of this vesiculovirus in vivo and in vitro, and will assist development of specific diagnostic tests to study the epidemiology of MSPV infection.

**Keywords:** Malpais spring virus (MSPV), Family *rhabdoviridae*, Genus *vesiculovirus*, Genome sequence, Phylogeny

Background

Malpais Spring virus (MSPV) is a negative-sense RNA virus currently assigned to the family *Rhabdoviridae* in the order *Mononegavirales* [1]. MSPV was first isolated from pools of *Aedes campestris* and *Psorophora signipennis* mosquitoes collected during an investigation of unexplained deaths among feral horses living near a spring in the Malpais Lava Flow on the White Sands Missile Range in New Mexico, USA [2]. Initial serologic tests and electron microscopy provided evidence that MSPV is a novel rhabdovirus, possibly related to vesiculoviruses [2]. MSPV is lethal to newborn mice following intracerebral (i.c.) inoculation and causes a cytopathic effect (CPE) in Vero cells, forming sizeable (3–4 mm in diameter) plaques six days post-inoculation [2]. Plaque reduction neutralization tests conducted using sera from wild mammals living in the area where the virus was recovered, indicated a relatively high prevalence (30-100%) of specific MSPV neutralizing antibodies among feral horses, coyotes, mule deer, gemsbok (*Oryx gazella*) and pronghorn, although the virus could not be clearly associated with the horse deaths [2].

Findings

**Methodology**

**Virus source**

The prototype strain of MSPV (85-488NM) was obtained from the World Reference Center for Emerging Viruses and Arboviruses at the University of Texas Medical Branch. The virus stock used had been passaged four times by i.c. inoculation of newborn mice and three times in Vero cells.
Next generation sequencing
The complete genome of MSPV (85-488NM) was determined by de novo sequencing using the Illumina platform.

Library construction
Viral RNA (0.05-1.7 μg) was fragmented by incubation at 94° C for eight (8) minutes in 19.5 μl of fragmentation buffer (Illumina 15016648). First and second strand synthesis, adapter ligation and amplification of the library were performed using the Illumina TruSeq RNA Samplec Preparation kit under conditions prescribed by the manufacturer (Illumina). Samples were tracked using the “index tags” incorporated into the adapters as defined by the manufacturer.

Sequence analysis
Cluster formation of the library DNA templates was performed using the TruSeq PE Cluster Kit v3 (Illumina) and the Illumina cBot workstation using conditions recommended by the manufacturer. Paired end 50 base sequencing by synthesis was performed using TruSeq SBS kit v3 (Illumina) on an Illumina HiSeq 1000 using protocols defined by the manufacturer. Cluster density per lane was 645–980 k/mm² and post filter reads ranged from 148–178 million per lane. Base call conversion to sequence reads was performed using CASAVA-1.8.2. Virus assembly was performed using SeqMan Lasergene software (DNASTAR). In certain cases, pre-filtering of reads to remove host sequence enhanced the assembly process. Assembly was carried out using a fasta file of hamster ribosomal RNA sequences to remove host DNA from the assembly, thus reducing the number of contigs present.

Nucleotide sequence accession number
The GenBank accession number of MSPV prototype 85-488NM strain is KC412247.

Phylogenetic analyses
The L protein sequence of MSPV was compared with those of 44 other rhabdoviruses downloaded from GenBank (members of the genera Cytorhabdovirus, Novirhabdovirus and Nucleorhabdovirus were excluded because their excessive divergence reduced phylogenetic resolution). All protein sequences were aligned using MUSCLE [3] under default settings, and ambiguously aligned regions were then removed using the Gblocks program [4]. This resulted in a final sequence alignment of 1136 amino acid residues. The phylogenetic relationships among these sequences were determined using the maximum likelihood (ML) method available in PhyML 3.0 [5], employing the WAG+Γ model of amino acid substitution and subtree pruning and regrafting (SPR) branch-swapping. The robustness of each node was determined using 1,000 bootstrap replicates utilizing nearest neighbor interchange (NNI) branch-swapping.

Antigens and immune reagents
Antigens used in complement fixation (CF) tests and for immunizing animals were prepared from infected newborn mouse brains by the sucrose/acetone extraction method [6]. Hyperimmune mouse ascitic fluids were prepared against each of the 21 rhabdoviruses listed in Table 1. The immunization schedule consisted of four intraperitoneal injections given at weekly intervals. Immunogens consisted of 10% crude suspensions of homogenized infected mouse brain mixed with equal volumes of complete Freund’s adjuvant just prior to inoculation. Sarcoma 180 cells were also given intraperitoneally after the final immunization in order to induce ascites formation. All animal experiments were carried out under an animal protocol approved by the University of Texas Medical Branch IACUC committee.

| Table 1 Antigenic relationships of MSPV with confirmed and tentative members of the genus Vesiculovirus |
|-------------------------------------------------|-------------------------------------------------|------------------|
| Genus | Viral antigen | MSPV antibody* | Homologous antibody |
|-------|---------------|----------------|---------------------|
| Carajas (CJSV) | 0 | 128 |
| Chandipura (CHPV) | 0 | 256 |
| Cocal (COCV) | 0 | 256 |
| Isfahan (ISFV) | 0 | 64 |
| Maraba (MARV) | 0 | 512 |
| Piry (PIRV) | 0 | 64 |
| Moreton (MORV) | 0 | 256 |
| VSV Indiana (VSIV) | 0 | 256 |
| VSV New Jersey (VSNJ) | 0 | 32 |
| Boteke (BTKV) | 0 | 256 |
| Farmington (FARV) | 0 | 256 |
| Gray Lodge (GLOV) | 0 | 8 |
| Jurona (JURV) | 0 | 1024 |
| Klamath (KLAV) | 0 | 256 |
| Kwatta (Kwav) | 0 | 64 |
| La Joya (LV) | 0 | 512 |
| Perinet (PERV) | 0 | 512 |
| Porton (PORV) | 0 | 256 |
| Radi (RADV) | 0 | 512 |
| Jug Bogdanovac (YBV) | 0 | 256 |
| Malpais Spring (MSPV) | 512 | - |

* Antibody titers expressed as reciprocal of highest positive antibody dilution. 0 = <8.
Serologic testing

Complement fixation tests were conducted according to a microtechnique described previously [7], using 2 full units of guinea-pig complement. Titers were recorded as the highest dilutions giving 3+ or 4+ fixation of complement on a scale of 0–4+.

Results

The complete genome of MSPV (85-488NM) showed that the 11,019 nt MSPV genome exhibits typical rhabdovirus organization and consists of 3’- and 5’- terminal non-coding regions (approximately 63 nt and 97 nt, respectively) and 5 coding regions (genes), each bounded by putative transcriptional regulatory sequences (Figure 1A). Similar to those of most other animal rhabdoviruses, the transcription initiation (AU[U/C]GUC) signal and transcription termination/polyadenylation ([G/U]UAC[U]) signal are relatively conserved (Figure 1B). Gene 1 (N), encodes a 426-aa protein (47.4 kDa); gene 2 (P), encodes a 274-aa acid protein (29.8 kDa; pl 4.75); gene 3 (M), encodes a 226-aa protein (25.6 kDa); gene 4 (G), encodes a 522-aa protein, which is a class 1 transmembrane glycoprotein; and gene 5 (L), encodes a 2096-aa protein (238.7 kDa), which the RNA-dependent RNA polymerase (RdRp).

Alternative ORFs (< 180 nt) also occur in the N gene (198 nt), M gene (189 nt) and G gene (249 nt) encoding potential polypeptides for which BlastP searches revealed no significant sequence relationships to known proteins (Figure 1A). The putative proteins encoded in the N and M genes are unremarkable in structure. The putative 83-aa product encoded in the G gene is predicted (Phobius and HMMTOP) to be a highly basic transmembrane protein (9.26 kDa; pl 11.36) with a 53-aa N-terminal cytoplasmic domain, a 24-aa helical transmembrane domain and a very short (6-aa) C-terminal ectodomain (Figure 1C). ORFs encoding similar small hydrophobic proteins of unknown function have been identified in the genomes of other rhabdoviruses [8]. The MSPV G protein contains 12 cysteine residues in the ectodomain that form six disulphide bridges that are common to all vesiculovirus G proteins and three predicted N-glycosylation sites, two of which align with those of the vesicular stomatitis Indiana virus (VSIV) G protein [9].

Serologic studies conducted at the time of MSPV isolation indicated a weak antigenic relationship with another probable vesiculovirus Jurona virus (JURV) by indirect immunofluorescence, but none with VSIV, vesicular stomatitis New Jersey, Cocal, Piry or Chandipura viruses (CHPV) [2]. We could not demonstrate an antigenic relationship in CF tests between MSPV and an expanded set of confirmed and tentative vesiculovirus species, including JURV (Table 1).

Phylogenetic analysis based on an alignment of 1136

Figure 1 Genome organization of MSPV. A. Schematic representation of the genome organization of MSPV. Block arrows indicate the location of long ORFs, including overlapping ORFs in the N, M and G genes (light shading). B. Sequences of MSPV gene junctions illustrating potential transcription initiation sequences (shaded), conserved transcription termination/polyadenylation sequences (solid underlined) and initiation codons (dotted underline). C. Schematic representation of the predicted membrane topology of a small, basic transmembrane protein encoded in the alternative ORF in the G gene.
amino acid residues of the L protein sequence of 45 rhabdoviruses indicated that MSPV is most closely related (although with weak bootstrap support) to JURV, which has been isolated from Haemagogus spp. mosquitoes in the state of Para, Brazil in 1962 [10,11], as well as to Isfahan, CHPV and Perinet viruses (Figure 2). These five viruses form a well-supported (100% bootstrap support) phylogenetic cluster within the genus Vesiculovirus.

Discussion
The genome organization and phylogenetic relationships indicate that Malpais Spring virus should be classified as a new species in the genus Vesiculovirus. This is also consistent with the previous report of a distant antigenic relationship with JURV, which also falls phylogenetically within the vesiculovirus cluster. Evidence of alternative ORFs in the MSPV N, M and G genes is unique amongst vesiculovirus genomes sequenced to date but overlapping ORFs appear to occur commonly in rhabdoviruses [8], including VSIV in which an alternative ORF in the P gene encodes the 55-aa and 65-aa C and C’ proteins which are known to be expressed in infected cells from alternative initiation codons [12,13]. The complete genome sequence of MSPV will be helpful in the development of a reverse genetics system to study the unique aspects of this vesiculovirus in vivo and in vitro, and will

Figure 2 Phylogenetic tree of L protein sequences from 45 rhabdoviruses. The position of MSPV is shaded and bootstrap support values (>70%) are shown for key nodes. All horizontal branch lengths are drawn to a scale of amino acid substitutions per site, and the tree is rooted in the position observed in a broader analysis of the Rhabdoviridae (i.e. including members of the Cytorhabdovirus, Novirhabdovirus and Nucleorhabdovirus; not shown). (*) Genera not formally approved by the International Committee on Taxonomy of Viruses (ICTV).
assist development of specific diagnostic tests to study the epidemiology of MSPV infection.

**Abbreviations**

MSPV: Malpais Spring virus; CPE: Cytopathic effect; i.c.: Intracerebral; ML: Maximum likelihood; NNI: Nearest neighbor interchange; CF: Complement fixation; RdRp: RNA dependent RNA polymerase; VSV: Vesicular stomatitis Indiana virus; JURV: Jurona virus; CHPV: Chandipura virus.

**Competing interests**

The authors do hereby declare that they have no competing interests in this scientific work.

**Authors’ contributions**

NV, SW, TG, RBT performed the laboratory experiments. APATdR performed the serologic assays. ECH performed the phylogenetic analyses; PJW performed the genomic analyses. NV, ECH, PJW, RBT contributed in finalizing the manuscript. All authors have read and approved the final manuscript.

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