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Sequencing of prototype viruses in the Venezuelan equine encephalitis antigenic complex

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Received 16 March 1999; received in revised form 4 June 1999; accepted 4 June 1999

Abstract

The 5’ nontranslated region (5’NTR) and nonstructural region nucleotide sequences of nine enzootic Venezuelan equine encephalitis (VEE) virus strains were determined, thus completing the genomic RNA sequences of all prototype strains. The full-length genomes, representing VEE virus antigenic subtypes I–VI, range in size from 11.3 to 11.5 kilobases, with 48–53% overall G+C contents. Size disparities result from subtype-related differences in the number and length of direct repeats in the C-terminal nonstructural protein 3 (nsP3) domain coding sequence and the 3’NTR, while G+C content disparities are attributable to strain-specific variations in base composition at the wobble position of the polyprotein codons. Highly-conserved protein components and one nonconserved protein domain constitute the VEE virus replicase polyproteins. Approximately 80% of deduced nsP1 and nsP4 amino acid residues are invariant, compared to less than 20% of C-terminal nsP3 domain residues. In two enzootic strains, C-terminal nsP3 domain sequences degenerate into little more than repetitive serine-rich blocks. Nonstructural region sequence information drawn from a cross-section of VEE virus subtypes clarifies features of alphavirus conserved sequence elements and protease recognition signals. As well, whole-genome comparative analysis supports the reclassification of VEE subtype-variety IF and subtype II viruses. © 1999 Elsevier Science B.V. All rights reserved.

Keywords: Alphavirus; VEE; Nonstructural proteins; Nucleotide sequence

1. Introduction

Venezuelan equine encephalitis (VEE) viruses are arthropod-borne, Western Hemisphere alphaviruses in the family Togaviridae (Walton and Grayson, 1988). Individual VEE virus isolates are
designated as either epizootic or enzootic based on origin. Epizootic strains, isolated from sporadic equine epizootics and epidemics, multiply to high titer in non-immune equines and are transmitted indiscriminately by suitable mammalophilic or ornithophilic mosquito vectors to humans, domestic livestock, wild mammals, and birds (Sudia and Newhouse, 1975). The only interepizootic ‘reservoirs’ identified thus far have been in the form of incompletely formalin-inactivated vaccines (Kinney et al., 1992a). In contrast, enzootic strains are readily isolated in discrete geographic foci sharing common elevations, climate, and vegetation (Johnson and Martin, 1974). Elegant early field work implicated *Culex* (primarily subgenus *Melanoconion*) mosquitoes and wild rodent or marsupial hosts in enzootic transmission cycle maintenance (Chamberlain et al., 1964; Galindo et al., 1966; Grayson and Galindo, 1968).

The VEE antigenic complex (Shope et al., 1964), as initially defined by a short-incubation hemagglutination inhibition assay using spiny rat antisera (Young and Johnson, 1969) and later refined and expanded by hemagglutination and neutralization tests (France et al., 1979; Calisher et al., 1982; Kinney et al., 1983; Calisher et al., 1985), currently consists of six subtypes (I–VI). Subtype I is divided into five varieties (IAB, IC, ID, IE, IF) and subtype III is divided into three varieties (IIIA, IIIB, IIIC).

Prior to this decade, VEE viruses isolated from epizootics–epidemics were invariably subtype IAB or IC strains. Not surprisingly, subtype IAB and IC strains are experimentally equine virulent, although virulence varies among strains (Mackenzie et al., 1976). The more divergent enzootic strains occupy remaining antigenic classifications, and are for the most part equine benign. However, intrathecally inoculated subtype ID and IE strains cause a fulminant encephalitis in horses (Dietz et al., 1978), and recent equine outbreaks in southern Mexico with 40–50% case-fatality rates yielded subtype IE strains (Oberste et al., 1998).

This confirmation of the previously suspected involvement of certain enzootic VEE virus strains in outbreaks (Morilla-Gonzalez and Mucha-Macias, 1969), along with the isolation of long-quiescent subtype IC lineages from outbreaks in Venezuela and Columbia (Rico-Hesse et al., 1995; Weaver et al., 1996), raised or renewed questions concerning determinants of equine virulence and sources of equine-virulent strains.

Our laboratory has been engaged in the nucleotide sequencing of prototype strains in the VEE complex to examine genetic relationships and develop improved serologic (Hunt et al., 1990, 1991; Roehrig et al., 1991; Hunt and Roehrig, 1995) and polymerase chain reaction (PCR) reagents (Pfeffer et al., 1997, 1998) for VEE diagnostics. We have previously reported the full-length genomic sequences of subtype IAB, IC, and ID viruses (Kinney et al., 1989, 1992a,b). The complete sequence of a subtype IE virus, strain 68U201, has also been determined (Oberste et al., 1996).

The organization of the VEE virus positive-sense, single-stranded RNA genome is 5′-methylated cap-nontranslated region (5′NTR)-nonstructural protein 1 (nsP1)-nsP2-nsP3-nsP4-26S mRNA junction region-capsid-E3-E2-6K-E1-3′NTR-polyadenylated tail. Analogous to other alphaviruses, nonstructural replicase polyproteins are likely translated directly off the plus strand after viral uncoating, orchestrating complementary minus-strand synthesis and (following additional intracellular processing) subgenomic and genomic plus-strand RNA synthesis (de Groot et al., 1990; Strauss and Strauss, 1994; Lemm et al., 1998).

The present study reports the 5′NTR and nonstructural region sequences of nine enzootic VEE virus strains, which, when combined with the 26S mRNA sequences of these strains (Kinney et al., 1998), completes the genomic sequences of all currently identified prototype strains in the VEE virus antigenic complex.

### 2. Materials and methods

#### 2.1. Viruses

Information concerning strain name, abbreviation, classification, origin, and GenBank accession
number is shown in Table 1. All viruses except 68U201 (Oberste et al., 1996) are plaque-purified stocks that have been used to establish antigenic relationships and virus phenotypes, as summarized previously (Kinney et al., 1998).

2.2. Reverse transcriptase-polymerase chain reaction and sequencing

Extraction of viral genomic RNA, amplification of cDNA by reverse transcriptase-polymerase chain reaction (RT-PCR), and agarose gel purification and sequencing of cDNA were performed as previously described (Kinney et al., 1998). RT-PCR utilized degenerate amplimers designed to anneal to regions of conserved amino acid sequence in VEE virus and other alphaviruses. Initially, the majority of the nonstructural region was amplified using three overlapping sense/anti-sense amplimer combinations (designations include the 3’ terminal nucleotide genomic position in TRD virus, standard ambiguity code is R = A/G, Y = C/T, H = A/C/T, M = A/C, D = A/G/T, K = G/T, V = A/C/G, B = C/G/T, N = A/C/G/T):

| Virus | Abbreviation | Subtype | Origin | Sequence reference/isolation reference | GenBank Accession Number |
|-------|--------------|---------|--------|----------------------------------------|--------------------------|
| TRD   | TRD          | IAB     | Trinidad, 1943 | Kinney et al., 1989 | L01442 |
| TC-83 | TC-83        | d       |         | Kinney et al., 1989 | L01443 |
| 71-180| 71-180       | IAB     | Texas, USA, 1971 | Kinney et al., 1992a | AF069903 |
| P676  | P676         | IC      | Venezuela, 1963 | Kinney et al., 1992b | L04653 |
| 3880  | 3880         | ID      | Panama, 1961  | Kinney et al., 1992b | L00930 |
| Men I | Mena II      | IE      | Panama, 1962 | Galindo et al., 1966 | AF075252 |
| 68U201| 68U201       | IE      | Guatemala, 1968 | Oberste et al., 1996 | U34999 |
| 78V-3531| 78V-3531 | IIF     | Brazil, 1976 | Calisher et al., 1982 | AF075257 |
| Everglades Fe3-7c | EVE | II | Florida, USA, 1963 | Chamberlain et al., 1964 | AF075251 |
| Mucambo BeAn 8 | MUC | IIIA | Brazil, 1954 | Causey et al., 1961 | AF075253 |
| Tonate CaAn 410d | TON | IIIB | French Guiana, 1973 | Digoutte and Girault, 1976 | AF075254 |
| 71D-1252 | 71D-1252 | IIIC | Peru, 1971 | Scherer and Anderson, 1975 | AF075255 |
| Pixuna-BeAr35645 | PIX | IV | Brazil, 1961 | Shope et al., 1964 | AF075256 |
| Cabassou CaAr 508 | CAB | V | French Guiana, 1968 | Digoutte and Girault, 1976 | AF075259 |
| AG80-663 | AG80-663 | VI | Argentina, 1980 | Calisher et al., 1985 | AF075258 |

### Table 1

Reference information for Venezuelan equine encephalitis (VEE) virus strains

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**Abbreviation**: Karabatsos, 1985.

**Subtype**: Variety classification scheme for the VEE antigenic complex (Young and Johnson, 1969; France et al., 1979; Calisher et al., 1982; Kinney et al., 1983; Calisher et al., 1985; Roehrig and Mathews, 1985; Roehrig et al., 1991).

**Sequence reference/isolation reference**: Previously sequenced VEE viruses in **bold**, isolation references included in sequencing reports referenced.

**GenBank Accession Number**: Vaccine strain derived from TRD virus by passage in tissue culture (Berge et al., 1961).

**Strain name abbreviation**: Two subtype IE strains were isolated from the same patient in consecutive years, the latter in 1962 (M.A. Grayson, personal communication).
VE-2008 5'-CAYYGHGGRCNYTGAAY ACNGAYGARGARTACTA or
VE-2518 5'-GGDGAYCCKAARCTGTYG GYTTYTYYAYATGATGTG

cVE-4506 5'-TCTVCVAGATRCATAT CTCYTCBRYNGCCTCYCT
VE-4477 5'-GCGRATGTRGCGYATTYT GYMGRGAYAARAARTGGGA
cVE-7155 5'-ACYACBGCRCTCDATRATYT TNACYTCCATRTTYARCCA

Additional degenerate and virus-specific primers (sequences available on request) were designed to fill in sequence gaps. The infrequency of invariant neighboring wobble position nucleotides in the nonstructural region required synthesis of more degenerate primers than were used for the 26S mRNA region (Kinney et al., 1998). All initial priming sites in the genome were derived as internal sequences in other amplicons, and both strands of the amplified cDNA were sequenced.

2.3. 5' nontranslated region sequence determination

5'NTR sequences were obtained using the 5' RACE System kit (GIBCO BRL, Bethesda, MD) according to supplied protocol. The antisense oligomer cVE-963 (5'-GTBACYTT-GCARCACAAGATWCCCTCGGRTGCAT-3', where W = A/T) primed synthesis of the first strand cDNA. Following RNA degradation, the purified cDNA from each strain was divided into two aliquots, and the aliquots tailed with either dATP or dCTP. The dATP-tailed cDNA was amplified using a synthesized poly-T primer (5'-CACAG-CTGCGAGCTTGATCGTACCTTTTTTT-3') and cVE-256 (5'-GYGR-CARAYRCTGRTAYTRKHTHHTHART-ACATC-3'), a VEE virus-specific antisense primer. The dCTP-tailed cDNA was amplified using the supplied anchor primer and cVE-256. Comparison of sequences determined from the alternatively-tailed cDNAs assured the identity of the 5' terminal nucleotide.

2.4. Published partial or complete Venezuelan equine encephalitis virus nonstructural region sequences

We previously reported partial N-terminal nsP1 and C-terminal nsP4 coding sequences of enzootic strains included in this study (Pfeffer et al., 1997; Kinney et al., 1998). Full-length nucleotide sequences have been determined for five naturally-isolated VEE virus subtype I strains and for TC-83, the TRD-derived vaccine strain (Kinney et al., 1989, 1992a,b; Oberste et al., 1996). In the course of this study, an A→T artifact in the TRD virus sequence at nt 4809 was discovered. As originally reported (Kinney et al., 1989), the nucleotide at this position using plaque-purified TRD virus as starting template for sequence determination was T. Resequencing using cDNA amplified from low-passage (equine-1/mouse-1, 1961 seed) virus that had not been plaque-purified revealed an A at this position, as is found in TC-83 and every VEE virus strain. A partial MENA virus nsP3-nsP4 coding sequence, GenBank accession no. U34978 (Oberste et al., 1996), is identical to sequence obtained using our laboratory stock of MENA virus. Conflicts with a previous partial C-terminal nsP4 coding sequence determination, ranging from one nucleotide difference in MUC virus to 11 differences in 78V-3531 virus, are likely due to sequencing methods employed (Weaver et al., 1992).

2.5. Sequence analysis

Nucleotide and amino acid sequence analysis was performed using programs available in the GCG package, version 8.0 (Devereux et al., 1984), with default settings used in all cases. Phylogenetic analysis, including bootstrap resampling (Felsenstein, 1985), was performed using maximum parsimony algorithms implemented by PAUP V3.1.1 (Swofford, 1993). Characters were either unweighted or were unweighted in codon first and second positions when codon third position characters were given zero weight. Throughout this study, all comparative or phylogenetic analyses involving aligned VEE virus deduced polyprotein P1234 sequences exclude the C-termi-
Fig. 1. (A) Schematic VEE virus 5’NTR and nonstructural region subgenomic map (not drawn to scale), with ranges of 5’NTR and nsP coding sequence lengths above and deduced amino acid sequence lengths below individual nsP designations. The C-terminal nsP3 domain is shaded. Asterisks indicate stop codons. (B) Total genome lengths and nonstructural region landmarks of prototype VEE viruses and representative alphaviruses. Nucleotide positions for the ends of the nsP3 and nonstructural polyprotein coding sequences include respective stop codons. Polyprotein codon position and total genomic %G+C content are also listed (right). The C-terminal nsP3 domain coding sequence is excluded from nonstructural region %G+C content calculations. For strain abbreviations, see Table 1.

### 3. Results

#### 3.1. Genome organization

A schematic VEE virus subgenome and accompanying chart (Fig. 1) provide nonstructural region landmarks for TRD virus and the nine enzootic strains sequenced in this study — Mena II (antigenic subtype-variety IE), 78V-3531 (IF), Everglades Fe3-7c (II), Mucambo BeAn 8 (IIIA), Tonate CaAn 410d (IIIB), 71D-1252 (IIC), Pixuna BeAr 35645 (IV), Cabassou CaAr 508 (V) and AG80-663 (VI) — as well as for North American Eastern equine encephalitis (EEE) virus strain 82V-2136 (Weaver et al., 1993) and Sindbis virus (SIN) strain HRsp (Strauss et al., 1984).

The actual N-termini of the VEE virus nonstructural polyproteins or processed proteins have not been directly confirmed by protein sequencing. The favorable context for the putative initiation codon (Kozak, 1981, 1987) and shared amino acid identity with the cognate alphavirus nsP ter-
mini along with the known specificity of the papain-like proteinase (Strauss and Strauss, 1994), suggest these are the most likely termini. The VEE virus nsP2 is divided (dotted line) into N- and C-terminal domains after the isoleucine residue at position 456 (numbering based on TRD virus), corresponding to the final residue in the alignment of SIN nsP2 with certain single-stranded RNA plant viral helicases (Ahlquist et al., 1985) and the region where the N-terminus of the SIN nsP2 proteinase domain was mapped using infectious clone deletion mutants (Hardy and Strauss, 1989). Comparative analysis of VEE virus sequences (see below) indicates that selection pressures are different in N- and C-terminal portions of this protein. The VEE virus N-terminal nsP3 domain ends at residue 330, a conserved aliphatic residue following an invariant tyrosine residue. Only strains 78V-3531 and AG80-663 contribute to the variation in N-terminal nsP3 domain size.

3.2. 5′ nontranslated region

Individual VEE virus 5′NTR sequences, shortest among the alphaviruses, vary in length by 0–8 nucleotides (Fig. 2A). Despite an overall lack of sequence conservation, alphavirus 5′NTRs are predicted to form stable secondary structures (Ou et al., 1983; Niesters and Strauss, 1990a). An older method of calculating the free energy of duplex formation is used (Tinoco et al., 1973), relying on a strict interpretation of Table 1 in this reference without modifications proposed in the text. $\Delta G = \text{free energy at 25°C.}$

![Diagram](image)

**Fig. 2.** VEE virus conserved sequence elements. (A) 5′NTR sequence alignment, with both conserved nucleotides (bold) and consensus sequence (CONS) derived from nucleotides present in at least 5 of 6 VEE virus subtypes. A dash indicates a gap used to improve alignment. 5′NTR sequences of 71–180 and 3880 viruses are identical to TRD virus (Kinney et al., 1992a,b). Previously reported sequences are underlined. See Table 1 for strain abbreviations. (B) Proposed secondary structures for the VEE virus 51-nt CSE (left) and additional VEE virus nsP1 coding region CSEs (right). The 51-nt CSE nucleotide positions and putative stem sequence are based on TRD virus. Remaining non-stem 51-nt CSE sequence is a composite of all VEE virus strains (Y = A/U, Y = A/G/C, H = A/C/U, N = A/C/G/U). The C-G pairs replaced by U-A pairs in certain strains are boxed. The shaded boxes indicate additional pairs which could hydrogen bond in subtype IE MENA virus. For the putative hairpin beginning at TRD nt-67, the sequence is that of 71D-1252 virus, although a similar structure with slightly lower stability can be modeled for other VEE virus strains. TRD virus sequence is used for the other two hairpins. To facilitate comparison with earlier studies (Ou et al., 1983; Niesters and Strauss, 1990a), an older method of calculating the free energy of duplex formation is used (Tinoco et al., 1973), relying on a strict interpretation of Table 1 in this reference without modifications proposed in the text. $\Delta G = \text{free energy at 25°C.}$
Table 2

| nsP | N-term. nsP2 | C-term. nsP2 | N-term. nsP3 | nsP4 |
|-----|--------------|-------------|-------------|------|
|     | %nt(%aa) | %nt(%aa) | %nt(%aa) | %nt(%aa) | %nt(%aa) |
| TRD, TC-83, 71–180, P676, 3880, EVE | 89 (98) | 84 (96) | 84 (96) | 85 (98) | 86 (98) |
| +MENA, 68U201 | 77 (93) | 70 (93) | 68 (88) | 70 (92) | 70 (93) |
| +MUC, TON, 71D-1252 | 67 (87) | 61 (86) | 56 (78) | 59 (82) | 60 (87) |
| +CAB, PIX, 78V-3531, AG80-663 | 60 (79) | 55 (79) | 48 (66) | 55 (75) | 55 (81) |
| +EEE<sup>a</sup> | 49 (60) | 52 (72) | 38 (48) | 44 (56) | 50 (72) |
| +SIN<sup>a</sup> | 41 (48) | 44 (56) | 27 (30) | 35 (40) | 45 (63) |

<sup>a</sup> NA EEE virus strain 82V-2136 sequence, SIN strain HR<sub>sp</sub> sequence.

et al., 1983) potentially required for ribosome orientation or (via the 3′ complement on the minus strand) promotion of plus-strand RNA synthesis (Niesters and Strauss, 1990b; Pardigon and Strauss, 1996). Putative VEE virus 5′NTR hairpins are less stable and structurally less complex than stem-loop structures modeled for other alphavirus 5′NTR sequences. A secondary structure previously modeled for TRD virus (Dubuisson et al., 1997) has a calculated free energy of −4.4 kcal by the method used here (Tinoco et al., 1973). A 5′ terminal hairpin formed from the conserved UGGGCGG heptamer (circled in the consensus sequence) starting at TRD virus nt-2 and either the conserved GCCCA (nt-21) or the conserved CUACCCA (nt-36) has a calculated free energy of −11.2 kcal using the former combination and −7.2 kcal (−8.2 kcal on the minus strand) using the latter combination.

In contrast to conserved sequence element (CSE) putative stem–loop structures noted below, the VEE virus 5′NTR stems are not supported by compensatory pairs of nucleotide changes (Noller and Woese, 1981; Pace et al., 1989; Niesters and Strauss, 1990b) in sequences of other VEE virus strains.

### 3.3. Nonstructural region

Overall VEE virus nonstructural region deduced amino acid sequence identity is 77%, compared to 60% in the structural region (Kinney et al., 1998). Table 2 details the stepwise approach to the nonstructural region consensus sequence, starting with a consensus determined for selected subtype I and II viruses and adding increasingly more divergent strains. EEE and SIN genotypes (Strauss et al., 1984; Weaver et al., 1993) are added to the overall VEE virus consensus sequence to emphasize certain trends. For any particular VEE virus nsP coding sequence, the majority of nucleotide changes are silent, occurring initially at the codon wobble positions and continuing until the wobble positions are essentially saturated, as noted previously in other alphavirus sequence comparisons (Strauss and Strauss, 1994). For example, the nsP4 coding sequences of subtype IAB, IC, ID, and II strains share only 86% nucleotide identity yet are virtually identical at the deduced amino acid level. It is clear from every generated consensus sequence that individual nsPs (and protein domains) diverge at different rates. The VEE virus N-terminal nsP2 domain, for example, is much more conserved than the C-terminal nsP2 domain. This divergence is even more striking when SIN and EEE virus nsP2 sequences are included in the comparison.

A ranking of invariant nsP amino acid residues, arranged by total numbers of invariant alanine, arginine, etc. residues in the deduced polyprotein P1234 sequence, was generated, and ratios of total invariant residues/total residues for each amino acid were calculated for every sequenced strain (data not shown). In general, four-codon amino acid families are more prevalent in the VEE con-
sensus sequence than two-codon families, and amino acids with specific, irreplaceable roles in protein structure or enzymatic activity, i.e. glycine, cysteine, and tryptophan, are well-conserved and seldom extraneous.

Previously identified alphavirus nsP or cognate ‘Sindbis-like supergroup’ protein functions and features are listed in Table 3, along with invariant residue correlates in VEE virus sequences. Only one Sindbis-like supergroup invariant residue in the replicase complex is not strictly conserved by all VEE virus strains. This nsP2 helicase domain valine residue (Gorbalenya et al., 1989) is an isoleucine in strain 71D-1252.

3.4. Conserved sequence elements

The VEE virus nsP1 deduced amino acid consensus sequence is less conserved than the nsP4 sequence (Table 2, line 4), while the corresponding nsP1 coding sequence is more conserved. One reason for this is the presence of CSEs, most notably the alphavirus 51-nt CSE, in the nsP1 coding region. The 51-nt CSE has been well-characterized in SIN (Niester and Strauss, 1990a) and may represent a promoter for RNA minus-strand synthesis. The VEE virus 51-nt CSE is shown in Fig. 2B, along with three other nsP1 coding region CSEs.

Unlike the putative VEE virus 5’NTR stem–loop structure, proposed RNA secondary structures for these CSEs are supported by pairs of compensatory nucleotide changes (Noller and Woese, 1981; Pace et al., 1989) in different VEE virus strains. The boxed C-G pair in the stem of the second hairpin of the 51-nt CSE (Fig. 2) is a U-A pair in all three of the subtype III strains and in strains 78V3531 and AG80-663. Similarly, the boxed C-G pair in the stem of the 20-nt near-perfect palindrome beginning at TRD virus nt-1118 is a U-A pair in Eve and MUC viruses. Compensatory changes are lacking in the proposed stems.

Table 3

| Nonstructural protein or protein domain | (putative) Function(s) | Conserved residues/motifs/domains<sup>a</sup> | Comments |
|----------------------------------------|------------------------|---------------------------------------------|----------|
| nsP1<sup>b</sup>                        | membrane-associated methyltransferase, guanylytransferase | H-37, D-89, R-92, Y-248 [VEE virus specific: N-terminal 55 amino acid (aa) residues] | VEE virus C-terminal residues vary — C-terminal truncations well-tolerated in SIN and SF virus constructs; 'Invariant' nsP2 V-389 is isoleucine in strain 71D-1252 |
| N-terminal nsP2 domain                 | RNA helicase, ATPase, GTPase | 6–7 conserved segments, including GVPGSGKS and DEAF NTP-binding signature motifs, 14 ‘invariant’ aa residues | Other plus strand RNA viral papain-like proteinases have flexible catalytic dyad spacing; Only aa similarity is with rubella virus and coronavirus proteins of undetermined functions |
| C-terminal nsP2 domain                 | Papain-like proteinase | Catalytic dyad residues C-477 and H-546 (C-481 and H-558 in SIN), W-547 | |
| N-terminal nsP3 domain                 | Phosphoprotein necessary for minus-strand, 26S mRNA synthesis | | |
| nsP4                                  | RNA-dependent RNA polymerase | N-terminal Y residue, GDD motif, 8 conserved domains | |

<sup>a</sup> Invariant VEE virus residues in <strong>b</strong> number, based on TRD virus nonstructural protein sequences.

<sup>b</sup> References: nsP1 (Mi et al., 1989; Mi and Stollar, 1991; Rozanov et al., 1992; Laakkonen et al., 1994; Peranen et al., 1995; Wang et al., 1996; Ahola et al., 1997; Pfleger et al., 1997); nsP2 (Gorbalenya et al., 1988, 1989, 1991; Hodgman, 1988; Ding and Schlesinger, 1989; Hardy and Strauss, 1989; Strauss et al., 1992; Rikkenon et al., 1994); nsP3 (Gorbalenya et al., 1991; LaStarza et al., 1994a,b); nsP4 (Kamer and Argos, 1984; Koonin, 1991; Shirako and Strauss, 1998).
of the two additional VEE virus-specific CSEs. However, equivalent calculated free energies and proximity to the 51-nt CSE double hairpins support the involvement of these CSEs in a large 5% terminal secondary structure, as has been modeled for SIN (Niesters and Strauss, 1990b).

3.5. Proteinase sites

Proteinase recognition sites have been identified in other alphaviruses by N-terminal sequencing of processed nsPs and by mutations which abolish in vitro cleavage (Strauss and Strauss, 1994). By alignment, the putative substrates for the VEE virus papain-like proteinase are predicted to be (residues given as P4P3P2P1, P1, P2, P3, P4; variable residues in brackets): nsP1/nsP2 — EAGA/G[S,T]VE; nsP2/nsP3 — EAG[C,S,T]/APSY; and nsP3/nsP4 — [D,E]AGA/YIFS. The P3, P2, and P1 residues constitute the major recognition signal for alphavirus proteinase cleavage (de Groot et al., 1990; Strauss and Strauss, 1994), with the P3 residue generally alanine, the P2 residue invariably glycine, and the P1 residue generally alanine or glycine, although any non-bulky residue may be tolerated, as indicated by the P1 residues allowed at the putative VEE virus nsP2/nsP3 cleavage site. The P4 residue is acidic in all VEE virus strains. The VEE virus proteinase may require an acidic P4 residue or a particular 3-dimensional conformation for cleavage, if for no other reason than the presence of additional AGA motifs in the VEE virus polyprotein P1234 consensus sequence besides those at putative cleavage sites.

3.6. C-terminal nsP3 domain

Because of large duplications and amplification of small serine-rich blocks in different strains, the deduced VEE virus C-terminal nsP3 domain ranges in size from 174–234 amino acids, with only 27 amino acid residues invariant (Fig. 3). Of the four nsP3 carboxyl region domains previously identified by comparison of five subtype I strains (Oberste et al., 1996), only domain 4 remains inviolate in the overall VEE virus alignment. Although divergence in this region is so great that any proposed alignment likely contains errors, the majority of VEE virus C-terminal nsP3 domain sequences coalesce around two motifs corresponding to truncated domains 2 and 3. The domain 2 SXWSXPXASDF motif (where 'X' indicates a variable amino acid residue) is conserved by all strains except PIX. Only strains 78V-3531 and AG80-663 do not preserve at least one copy of the imperfect PXPAPRT repeat in domain 3. One copy of this repeat is also found in EEE virus strain 82V-2137 (Weaver et al., 1993).

Duplication events appear to be common in the alphavirus C-terminal nsP3 domain (Strauss et al., 1988), although the variety and size of direct repeats in VEE virus sequences have not been previously noted. A 34-amino acid duplication is present in all VEE virus subtype IAB, IC, ID, and II strains. One copy of this duplication was deleted during propagation of a TRD virus infectious clone, with no demonstrable effect on viability (Davis et al., 1989). Both subtype IE strains possess a larger upstream duplication obscured by a subsequent deletion event. TON virus has two duplications of 34 and 20 amino acids, neither of which is found in other subtype III strains. The region corresponding to domain 2 has been duplicated at least once (Fig. 3) in subtype IV PIX virus. Beyond domain 2, sequences of strains 78V-3531 and AG80-663 comprise little more than short serine, alanine, and/or arginine-containing repeats. Translation in alternative reading frames rules out the possibility that sequencing errors or recent indels mask motif conservation in these strains.

The relative ages of these direct repeats can be determined by nucleotide alignment. The two copies of the 20 amino acid duplication in TON virus, for example, share 59 of 60 nucleotides. Using an estimate of $3 \times 10^8$ substitutions/nucleotide (Strauss and Strauss, 1986), this duplication occurred $< 100$ generations prior to isolation.

The opal termination codon is preserved by all VEE virus strains, as are 12 of 14 amino acid residues immediately upstream (Fig. 3). Amino acid sequence conservation upstream of the opal codon is atypical for alphaviruses (Strauss and Strauss, 1994).
Fig. 3. Alignment of VEE virus C-terminal nsP3 domain sequences (single letter code) from the residue corresponding to TRD virus nsP3 V-331 to the putative nsP3
nsP4 proteinase cleavage site. A dash indicates a gap introduced to improve alignment. Asterisk indicates nsP3 opal stop codon. Strains with
\[80\% \text{ nsP3 amino acid sequence identity to strain(s) selected for alignment} \text{ are omitted. Consensus sequence (CONS) includes residues conserved in at least 7 of the 9 strains selected for alignment, with boxed}
residues conserved by all sequenced VEE virus strains. Domains 1–4 are as designated by Oberste et al. (1996). The direct repeats
present in the various strains are indicated on the right, with lines drawn to the downstream copy of the direct repeat in the
alignment (underlined) the upstream copy of the direct repeat is omitted from the alignment. Degenerate, repetitive portions
78V-3531 virus and AG80-663 virus C-terminal nsP3 domain sequences excluded from the alignment (flanked by tildes) are given
on the lower right. For strain abbreviations, see Table 1. IABCDII = Present in all VEE virus subtype IAB, IC, ID, and II strains.
IE = present in all subtype IE strains.

3.7. Phylogenetic analysis

Nonstructural region codon third position nucleotides contribute the overwhelming majority of informative characters to VEE virus phylogenetic
analysis. Of the more than 3000 variant nucleotide positions in the VEE virus nonstructural
region alignment, 70\% are in the third position, 21\% in the first position, and 9\% in the second position of the codon. Because multiple substitutions
at the same position increase as more divergent VEE virus strain sequences are added to the
alignment, codon wobble position nucleotides become misleading and contribute little more than
background noise to parsimony-based phylogenetic analysis. This is manifested in inconsistent
branching orders and low bootstrap values for phylogenetic trees inferred from individual or
combined nsP coding sequences. Substituting R or Y for codon wobble position purines or pyrimidines is of little benefit, due to the predominance
of four-codon amino acid families in VEE virus nsP sequences. Thus, the only appropriate VEE virus character sets for parsimony-based analysis
Fig. 4. (Continued)

P1234 sequences or corresponding codon first and second position nucleotides produce the same topology as that shown in Fig. 4. The tree in Fig. 4 is derived from combined nonstructural and structural region codon first and second position nucleotides, with EEE virus strain 82V-2136 (Weaver et al., 1993) serving as the outgroup. This tree is well-supported by bootstrap resampling (Felsenstein, 1985), as only two partitions are present in fewer than 70% of resampled trees. An identical branching pattern is again reproduced when combined deduced nonstructural and structural region amino acid sequences are used, although bootstrap proportions in this case are somewhat lower (data not shown).

4. Discussion

Comparative analysis is often the best initial experimental method for determining secondary structures of RNA molecules (Noller and Woese, 1981; Pace et al., 1989), and is especially appropriate for examining alphavirus genomic elements potentially involved in formation of secondary structures required for host or viral protein interactions. Essential features of the alphavirus 51-nt CSE structure, for example, are confirmed or clarified by 51-nt CSE sequences reported here. Substitutions in putative stem structures of enzootic VEE virus strains represent natural experiments identical to SIN mutants constructed in vitro (Niesters and Strauss, 1990a). The requirement for strict maintenance of stem length in the second stem–loop of the 51-nt CSE as demonstrated by anti-pairing changes in O’Nyong-nyong virus (Levinson et al., 1990) is apparently violated.

Fig. 4. Maximum-parsimony phylogenetic tree for VEE virus prototype strains derived from branch-and-bound search in PAUP V3.1.1 (Swofford, 1993) using codon first and second position nucleotides from combined nonstructural (excluding the C-terminal nsP3 domain coding sequence) and structural regions, rooted with EEE virus strain 82V-2136 (Weaver et al., 1993) as the outgroup (7056 total characters, 1255 parsimony-informative characters). Percentages to the left of internal nodes indicate bootstrap support for 1000 pseudoreplicates (Felsenstein, 1985), using 10 random-addition heuristic searches per pseudoreplicate.

are deduced amino acid sequences or codon first and second position nucleotides. Maximum parsimony analysis (Swofford, 1993) using polyprotein
by MENA virus, which could form a stem lengthened by two base pairs (Fig. 2B). An equivalent calculated free energy for the MENA virus hairpin, due to the presence of a non-canonical G-U pair within the putative helix (Tinoco et al., 1973), may indicate that overall stability, rather than absolute stem length, determines possible nucleotide changes.

Secondary structure estimations using cross-species comparisons of homologous RNA molecules define a covariation that preserves pairing (such as a U-A pair mutating to a C-G pair) in a putative helical region as supportive of a stem, with two independent covariations taken as ‘proof’ of that stem (Pace et al., 1989). While this definition is more appropriate for purely structural RNA molecules, the alphavirus 51-nt CSE second stem is ‘proven’ and the VEE virus 20-nt palindrome stem supported by available sequences. In VEE virus, both CSEs are located in the only regions of the genome characterized by high concentrations of invariant wobble position nucleotides, and both may be part of more extensive secondary structures. The proposed involvement of the 51-nt CSE in a large 5' terminal secondary structure (Niester and Strauss, 1990a) has been mentioned. The 20-nt palindrome is within the region (nt 735-1255) corresponding to the putative SIN packaging signal (Weiss et al., 1989). Experiments using SIN RNA transcribed from progressively truncated cDNA clones rule out the palindrome as the exclusive packaging signal (Weiss et al., 1989), but capsid binding may require a secondary structure which includes this palindrome.

The proposed VEE virus 5'NTR secondary structures are less stable than those of other alphaviruses (Ou et al., 1983), and are not supported by covariation of potential stem-forming pairs. Because it would be helix-disruptive, the G-to-A mutation at nt 3 contributing to attenuation of the TC-83 strain (Fig. 2) indirectly supports a putative VEE virus 5'NTR stem (Dubuisson et al., 1997). However, while the contribution of this mutation to TC-83 attenuation is well-established (Kinney et al., 1993), the contribution of this mutation to attenuation as a result of 5'NTR secondary structure disruption awaits demonstration. As evidence from SIN 5'NTR and 51-nt CSE mutants indicates (Niester and Strauss, 1990a,b), nucleotide changes can influence a secondary structure model and a viral genome disproportionately. Preservation of ‘proximal’ nucleotides in the putative SIN 5'NTR stem (Niester and Strauss, 1990b), or preservation of the SIN 51-nt CSE linear nucleotide sequence over and above preservation of amino acid sequence and stem–loop free energy (Niester and Strauss, 1990a), may be more important determinants of viral fitness than secondary structure preservation. For another member of the Togaviridae, preservation of secondary structure may not even be required. The rubella virus 51-nt CSE homolog conserves many alphavirus wobble position nucleotides and deduced amino acid residues despite lacking an apparent alphavirus-like secondary structure (Domínguez et al., 1990).

Comparative analysis is the foundation for much of our current understanding of the less tractable, cellurally less plentiful alphavirus nonstructural proteins. Many of the experiments which helped define functions or functional residues for these proteins were prompted by or directed by sequence comparisons (Ahliquist et al., 1985; Hardy and Strauss, 1989; Rozanov et al., 1992; Strauss et al., 1992; Wang et al., 1996; Ahola et al., 1997; Shirako and Strauss, 1998), and support for additional proposed nsP properties relies on shared identity with viral or nonviral proteins of known function (Gorbalenya et al., 1988; Rikkonen et al., 1994). Potential VEE virus epizootic residues have been identified in structural proteins by sequence comparison (Powers et al., 1997; Kinney et al., 1998; Oberste et al., 1998). Functional attributes of ‘epizootic nsPs’ must be defined for similar comparisons in the nonstructural region to have meaning (LaStarza et al., 1994a; Oberste et al., 1996).

The variety of alphavirus 5'NTR and nonstructural region mutations leading to attenuation for cell lines or laboratory animals (Niester and Strauss, 1990b; Kuhn et al., 1992; Kinney et al., 1993; Rikkonen, 1996; Dryga et al., 1997) make it unwise to dismiss identified VEE virus nucleotide or amino acid differences as irrelevant in the
absence of functional studies. This is especially true for the 5’NTR, which is emerging as a major determinant of viral replication and pathogenesis (Kinney et al., 1993; Dubuisson et al., 1997).

On the other hand, the overall 77% amino acid sequence identity in the VEE virus polyprotein P1234 (excluding the C-terminal nsP3 domain) does not contain conservative amino acid changes or changes near the ends of processed proteins, where structure is almost certainly less constrained. Engineered into a TRD virus genetic background, many enzootic strain nsP conservative amino acid substitutions would likely have negligible attenuating effect. An example of the probable lesser (or independently insufficient) role of naturally-occurring VEE virus nonstructural region mutations in attenuation or epizootic strain emergence is provided by the recent equine outbreak in Chiapas, Mexico.

Isolation of a 68U201-like subtype IE strain from this outbreak (Oberste et al., 1998) indicates the epizootic phenotype can be maintained over a range of nonstructural region sequences. More than 120 amino acid residues in the polyprotein P1234 sequence differ between epizootic IAB or IC strains and strain 68U201. Providing the Mexican VEE virus isolate is not a recombinant with a IAB or IC strain nonstructural region and a 68U201-like structural region [the near-identity of partial nsP3 coding sequences of the Mexican isolate and strain 68U201 (Oberste et al., 1998) make this unlikely], replicase modules at least 5% dissimilar are capable of equivalent equine virulence.

VEE virus sequence comparisons provide insight into the C-terminal nsP3 domain coding sequence and the polyprotein codon wobble position nucleotides, two of the least conserved ‘regions’ of the genome. In vitro, the alphavirus nsP3 tolerates C-terminal domain deletions, duplications, or linker insertions (Davis et al., 1989; LaStarza et al., 1994a) provided the reading frame is preserved. Given the plasticity of alphavirus C-terminal nsP3 domain size (Strauss et al., 1988; LaStarza et al., 1994a), the infrequency of deletion events and predominance of duplication events in VEE virus C-terminal nsP3 domains is noteworthy. A TRD virus C-terminal nsP3 domain direct repeat secondary structure has been proposed (Davis et al., 1989), and related structures can be drawn for the duplications found in enzootic VEE virus strains. The mechanism for the generation of duplications may be related to the sequences themselves, as G + C content analysis or secondary structure modeling of C-terminal nsP3 coding sequences stripped of direct repeats fail to reveal sequence qualities peculiar to this region that would favor polymerase slippage or template switching.

Conservation of certain motifs by most VEE virus strains and a high serine concentration by all strains (Fig. 3) suggests a C-terminal region function beyond that of separating proteins in the replication complex. The hypothesis that nonconserved portions of the C-terminal nsP3 domain may determine host protein interactions or vector competence (Oberste et al., 1996) is not disproved by the additional sequences reported here, since there are nonconserved portions of the C-terminal domain specific to each VEE virus subtype and each strain. However, such a model would have to explain the finding of widely divergent deduced C-terminal nsP3 domain sequences (e.g. CAB and 78V-3531 virus sequences) in strains with the same identified mosquito vector (Digoutte and Girault, 1976; Calisher et al., 1982), or of essentially the same sequence (e.g. P676 and EVE virus sequences) in strains with different vectors and hosts (Chamberlain et al., 1964; Mackenzie et al., 1976).

While the lack of nucleotide sequence conservation renders the C-terminal nsP3 coding region unreliable for RT-PCR diagnosis of VEE virus infection using nondegenerate primers (Brightwell et al., 1998), we had no difficulty amplifying the C-terminal regions of all strains using a single degenerate primer pair hybridizing to neighboring conserved regions. Because nucleotide sequences of the C-terminal nsP3 region distinguish between even closely-related strains, the diagnostic utility of this region is obvious.

Few non-methionine, non-tryptophan codon third position nucleotides are invariant in the VEE virus nonstructural region. Provided changes are silent and preserve RNA secondary structure, codon third position nucleotides may simply drift,
or may be selected for based on prevailing vector or host genome codon biases. The high wobble position G + C content in PIX virus, for example, could represent a quasispecies minority population founder effect or a specific adaptation to indigenous fauna.

The use of data sets uncorrected for superimposed changes at codon third positions as more divergent VEE virus sequences are added can produce misleading phylogenies. Even without this correction, the monophyly of VEE virus subtype I and subtype II strains (excluding subtype IF 78V-3531 virus) and the monophyly of subtype III strains are absolute, regardless of the region analyzed or algorithm used (Powers et al., 1997; Kinney et al., 1998). The phylogenetic clustering of EVE virus with subtype IAB, IC, and ID strains has been previously noted (Powers et al., 1997; Kinney et al., 1998), as has the clustering of 78V-3531 virus with subtype VI AG80-663 virus. A reclassification of VEE virus subtype IF and subtype II strains has been proposed (Kinney et al., 1998), to which the complete nucleotide sequences reported here provide further support.

Acknowledgements

We would like to thank Ryan J. Barrett and Kevin M. Myles for technical assistance. All synthetic oligonucleotide primers used in this study were supplied by the Biotechnology Core Facility Branch, Centers for Disease Control and Prevention, Atlanta, Georgia.

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