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Evolutionary Relatedness of the Predicted Gene Product of RNA Segment 2 of the Tick-Borne Dhori Virus and the PBl Polymerase Gene of Influenza Viruses

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The complete nucleotide sequence of the second largest RNA segment of Dhori/India/1313/61 virus was determined and the deduced amino acid sequence was compared with the polymerase (P) proteins of influenza A, B, and C viruses. RNA segment 2 (2224 nucleotides) of Dhori virus contains a single long open reading frame that can encode a 716-amino acid polypeptide (81.3 kDa). The predicted polypeptide shares between 27 and 31% sequence identities with the PBl polypeptides of influenza A, B, and C viruses. Among the regions most highly conserved are the sequences around the Asp-Asp motif common to many RNA polymerases. In spite of the high level of sequence identity between the Dhori RNA segment 2 gene product and the influenza A, B, and C virus PBl proteins the amino acid composition of the Dhori protein indicates an acidic charge feature at pH 7.0 in contrast to the basic nature of the PBl proteins of the influenza viruses. We suggest that the Dhori PBl-like protein be designated the Pp protein of this virus.

Members of the Dhori virus serogroup are as yet unclassified tick transmitted viruses that share structural and genetic properties with the Orthomyxoviridae (Clerx et al., 1983). Dhori viruses have been isolated from a variety of tick and vertebrate species (Anderson and Casals, 1973; Filipe and Casals, 1979; Karabatsos, 1985; Williams et al., 1973). The genome of the virus consists of seven unique segments of single-stranded RNA with a total size of approximately 11.9 kb. The viral RNAs have been shown to encode information in the negative-sense (Clerx et al., 1983; Fuller et al., 1987; Freedman-Faulstich and Fuller, 1990) and it was previously shown that the Dhori nucleoprotein (encoded by RNA segment 5) shares conserved amino acid sequences with the influenza A, B, and C virus nucleoproteins (Fuller et al., 1987). The sequence of RNA segment 4 encodes the single envelope protein of Dhori virus (Freedman-Faulstich and Fuller, 1990). The envelope protein does not share sequence identity with any orthomyxovirus envelope protein. We have also determined the nucleotide sequence of the Dhori segment 6 RNA and this segment most likely encodes the viral matrix protein (unpublished data). The 3' and 5' nontranslated ends of RNA segments 4, 5, and 6 are very similar to the conserved ends of the genes of the recognized members of the Orthomyxoviridae (Clerx et al., 1983; Fuller et al., 1987; Freedman-Faulstich and Fuller, 1990) and segment 3 of Thogoto virus (Staunton et al., 1989).

A second group of tick-transmitted viruses, the Thogoto viruses, possess a segmented genome with six or seven segments of single-stranded RNA (Clerx et al., 1983; Staunton et al., 1989). Thogoto virus has been shown to replicate in Rhipicephalus appendiculatus ticks and to be transmitted to laboratory animals (Davies et al., 1986). The sequence of the third largest segment of Thogoto/SiAr/126/72 virus was recently determined (Staunton et al., 1989) and shown to encode in the minus sense a protein that shares 15-20% sequence identity with the influenza A and B virus PA proteins. Recently the sequence of the influenza C virus PA-like protein (designated P3) was determined (Yamashita et al., 1989) and the Thogoto PA-like protein shares a 19.9% sequence identity with the influenza C virus P3 protein. Based on this finding and the fact that the nontranslated ends of the Thogoto virus RNAs are similar to those of the Orthomyxoviridae genes, it is suggested that Thogoto virus is evolutionarily related to the Orthomyxoviridae. It is not yet known, however, to what degree Dhori and Thogoto viruses are related to each other.

We report here the nucleotide sequence of the second largest segment of the Dhori/India/1313/61 virus. The sequence of this RNA segment predicts one gene product which while likely to represent the equivalent of the influenza virus polymerase protein PBl, is...
unique in having a net negative charge unlike the net positive charge of the corresponding A, B, and C virus proteins. Despite these differences the data reported here establish that the influenza viruses and Dhori virus share a common evolutionary ancestor.

MATERIALS AND METHODS

Virus

Dhori/India/1313/61 virus was grown in Green monkey kidney (GMK-Vero) cells as previously described (Clerx et al., 1983). Virus purification and RNA extraction have also been described previously (Clerx et al., 1983).

Isolation of infected cell mRNA

Dhori virus-infected cellular RNA was isolated from Vero cells at 48 hr postinfection, phenol extracted, ethanol precipitated, and purified by oligo(dT)-cellulose chromatography as previously described (Collins et al., 1982; Fuller et al., 1983).

Cloning of virus-specific DNA and identification of clones

The Dhori/India/1313/61 virus strain was used to infect GMK-Vero cells. Infected cell mRNA was reverse transcribed with an oligo(dT)12-18 primer. Second-strand cDNA synthesis was performed by the method of Gubler and Hoffman (1983). The blunt-ended double-stranded cDNAs were ligated to pBR322 which had been cut by PvuII and dephosphorylated. Clones bearing Dhori virus specific inserts were identified by colony hybridization using radiolabeled cDNA which had been obtained using viral RNA as template and the oligonucleotide 5′ AGCAA(A/T)AACAGCATG 3′ as primer. This oligonucleotide is complementary to sequences common to the 3′ ends of the seven viral RNA segments (Clerx et al., 1983). Virus-specific clones were organized by cross-hybridization into several pools. Viral RNA was extracted from cesium chloride gradient (12–42% (w/w)) purified Dhori virus and resolved on a 1.5% agarose gel containing 10 mM methylmercuric hydroxide. The viral RNAs were transferred from the agarose gel to Biotrans nylon membranes (ICN Inc., Irvine, CA) and hybridized with nick-translated plasmid DNA.

Sequencing of cloned cDNA

Both strands of insert DNA of pD50-40 were directly sequenced using specific synthetic oligonucleotide primers and the modified T7 DNA polymerase Sequenase (United States Biochemical Corp.). All oligonucleotides were synthesized using an Applied BioSystems 380A synthesizer (Applied BioSystems, Foster City, CA). For plus-strand as well as for minus-sense sequencing specific primers were synthesized to permit the reading of new sequences over a distance of 150–200 nucleotides. In order to obtain sequences located at the 3′ end terminus of the RNA segment 2, polymerase chain reactions were performed using purified viral RNA and the following oligonucleotides, 5′AGCAA(A/T)AACAGCATG 3′ and 5′ATCTCTGAGAC 3′. The sequence located at the 5′ end terminus of the RNA segment 2 was also obtained by polymerase chain reaction performed on purified viral RNA and the following oligonucleotides, 5′AGTAA(A/C)ATCAAGCA 3′ and 5′ACCTCTTTGTGGAAG 3′. The resulting dsDNAs were cloned into M13mp18RF and directly sequenced using universal primer.

RESULTS AND DISCUSSION

An analysis of the Dhori virus genome was attempted by DNA cloning of the poly(A)-containing RNA of virus-infected GMK-Vero cells. Virus-specific cDNA clones were obtained and by cross-hybridization were grouped into seven pools. Nick-translated DNAs of plasmids pD21-39, pD50-40, pD43-22, pD68-30, pD22-39, pD26-16, and pD12a-35 hybridized to RNAs 1, 2, 3, 4, 5, 6, and 7, respectively in a Northern blot analysis of Dhori virus RNA (Fig. 1). The clone pD12a-
FIG. 2. Complete nucleotide sequence of RNA segment 2 of Dhori/India/1313/61 virus and the deduced amino acid sequence. Sequencing of segment 2-specific cDNA from pD50-40 was accomplished using Sequenase (United States Biochemical Corp., Cleveland, OH), and specific oligonucleotides (see Materials and Methods). The 25 3′-terminal and the 16 5′-terminal sequences (underlined) of the gene were determined as described in the text.

35, contains a small insert of segment 1 as well as an insert for segment 7 (unpublished results). Plasmid pD50-40 was selected for further study since it had the largest insert among a total of nine different clones belonging to the group that hybridized to RNA segment 2.

The insert of pD50-40 was then sequenced using specific synthetic oligonucleotide primers and the mod-
The RNA is 2224 nucleotides in length and the cDNA for longer than 64 amino acids and the longest open reading frame in the genome sense is 109 amino acids. The RNA encodes a single long open reading frame that could be initiated by an AUG codon at nucleotides 25 through 27. No other reading frame is open in the antigenome sense (see Table 1 above diagonal) between Dhori and influenza PB1 proteins yield alignment scores of 25–26 which clearly indicate an evolutionary relationship between the Dhori virus PB1-like protein and the influenza PB1 proteins. A similar comparison between the Dhori segment 2 gene product and the influenza A/PR/8/34 PB2 protein and PA protein gave alignment scores of 0.25 and 0.015, respectively.

Comparative analysis of the deduced amino acid sequences revealed highly conserved regions in the PB1 proteins of influenza A, B, and C viruses (Kawacka et al., 1989; Yamashita et al., 1989). Although the functional domains of the PB1 proteins of influenza viruses have not been determined, the conserved regions may be important for basic functions of the PB1 proteins, such as initiation and chain elongation (Braam et al., 1983). It has also been observed that there are four highly conserved motifs among RNA-dependent RNA polymerases (plus- or minus-strand virus polymerases) as described by Poch et al. (1989) and the Dhori PA protein shares all of these motifs (Fig 3). A comparison of the Dhori virus protein with the PB1 proteins shown in Fig. 3 also reveals regions of highly conserved sequences. A 15-residue sequence motif (a double aspartic acid core flanked by hydrophobic residues) is found in many DNA and RNA polymerases (Argos, 1988; Poch et al., 1989). The PB1 gene in influenza A, B, and C viruses (Yamashita et al., 1989) and they are most likely required for nucleotide addition during viral RNA synthesis (Braam et al., 1983). A pairwise comparison of the amino acid sequence identities of the Dhori virus protein and the influenza A, B, and C virus PB1 proteins are shown in Table 1. The optimized alignments were made by the method of Needleman and Wunsch (1970) and indicate that the Dhori PB1-like protein shares sequence identities of 30.7, 26.9, and 29.8% with the PB1 proteins of influenza A/PR/8/34, influenza B/Ann Arbor/1/66, and influenza C/J/J/50 virus, respectively (Table 1). In addition, we compared the sequence of the Dhori PB1-like protein to the PB1 proteins of influenza A, B, and C viruses using the PCOMPARE (Intelligenetics) program. This program determines the probability that the similarity between two sequences could occur by chance. The score is expressed as the number of standard deviation units by which the maximum score for the real sequences exceeds the average maximum score for 100 randomizations of the sequence. An alignment score greater than 5 (probability of similarity occurring by chance \( < 3 \times 10^{-7} \)) implies an evolutionary relationship between proteins (Dayhoff et al., 1983).

**Table 1**

| Amino Acid Sequence Identites and ALIGN Scores of the Dhori PA Protein and the PB1 Proteins of Influenza A, B, and C Viruses |
|--------------------------------------------------|
| Dhori/India/1313/61                               |
| A/PR/8/34                                        |
| B/Ann Arbor/1/66                                 |
| C/J/J/50                                         |
| Dhori/India/1313/61                               |
| 1313/61                                          |
| 30.7%                                            |
| 26.9%                                            |
| 29.9%                                            |
| A/PR/8/34                                        |
| 30.7%                                            |
| 26.3%                                            |
| 26.9%                                            |
| 60.9%                                            |
| 26.9%                                            |
| 39.2%                                            |
| B/Ann Arbor/1/66                                 |
| 30.7%                                            |
| 26.3%                                            |
| 26.9%                                            |
| 60.9%                                            |
| 39.2%                                            |
| C/J/J/50                                         |
| 29.9%                                            |
| 26.9%                                            |
| 29.8%                                            |
| 60.9%                                            |
| 29.8%                                            |
| 64.3%                                            |
| 70.0%                                            |
| 30.7%                                            |
| 26.9%                                            |
| 29.8%                                            |

Proteins and the PB1 Proteins of Influenza A, B, and C Viruses

| Percent amino acid identity (below diagonal) based on an alignment of the two sequences by the method of Needleman and Wunsch (1970). |
|------------------------------------------------------------------------------------------------------------------------|
| Dhori/India/1313/61                               | A/PR/8/34*                                                  | B/Ann Arbor/1/66 | C/J/J/50*                                |
| 1313/61                                          | 26.3*                                                      | 25.8            | 25.7                                     |
| A/PR/8/34                                        | 30.7%                                                      | 152.1           | 64.3                                     |
| B/Ann Arbor/1/66                                 | 26.9%                                                      | 60.9%           | 12.1%                                    |
| C/J/J/50                                         | 29.9%                                                      | 39.2%           | 12.1%                                    |

* PA protein 716 amino acids (81.3 kDa) (this article).
* PB1 protein 757 amino acids (86.6 kDa) (Winter and Fields, 1982).
* PB1 protein 752 amino acids (84.3 kDa) (DeBorde et al., 1987).
* P3 protein 754 amino acids (85.9 kDa) (Yamashita et al., 1989).
* Alignment score (above diagonal) determined by the program PCOMPARE (Intelligenetics) using the unitary matrix, a bias of 0, a gap penalty of 3, and 100 random runs. A score of 5 or greater is considered significant (Dayhoff, 1983).
* Percent amino acid identity (below diagonal) based on an alignment of the two sequences by the method of Needleman and Wunsch (1970).
FIG. 3. Comparison of the deduced amino acid sequences of the PBl proteins of influenza A/PRI8/34 (Winter and Fields, 1982) B/Ann Arbor/66 (DeBorde et al., 1988) and C/JJ/50 (Yamashita et al., 1990) with the PCU protein of Dhori/India/Gl313/61 virus. The alignment was obtained by a compilation of the pairwise comparisons by the method of Needleman and Wunsch (1970). An asterisk (*) denotes identity with the amino acid of the Dhori P Bl protein at that position and a dash (-) indicates a gap introduced in the sequence for optimal pairing. The four domains designated by the lines above and below the sequence represent those highly conserved regions defined in RNA polymerases by Poch et al. (1989).

B, and C shares this motif (W/FWD/TGLQSSDDFA/VLI/FV/A), at residues 435/438 to 449/452 (domain 3, Fig. 3). In the Dhori virus protein, we found a similar sequence at residues 419 to 433 (TGDHVESSDDFIHFF). All rules of this Asp—Asp polymerase motif are fulfilled except for position 7 which contains a serine instead of the traditional Gly, Met, Cys, Val, or Leu residue (Aragos, 1988). Thus it is likely that the Dhori virus protein also possesses a polymerase function similar to that of the PBl proteins of influenza viruses.

The influenza virus PBl protein is transported to the nucleus independently of other viral proteins (Smith et al., 1987; Akkina et al., 1987). Nath and Nayak (1990) have recently shown that the nuclear localization sig-
nal of influenza virus (A/WSN/33) PB1 protein was located within residues 180–252 of PB1. The study suggested that two discontinuous regions of the PB1 both containing a stretch of basic amino acids were required for its nuclear localization. Homologous sequences, however, were not found in the Dhori virus protein.

A comparison of some of the physical properties of the Dhori and influenza PB1 proteins indicates that the Dhori protein is the smallest in this group (716 amino acids; M, 81,292) and unlike the influenza PB1 proteins which have a strong positive charge at neutral pH (+23-A/PR/8/34; +12-B/AA/1/66; +14-C/JJ/50) the Dhori protein is slightly acidic with a charge of −10 (PEPTIDESORT; Devereux et al., 1984). Since the nomenclature of the influenza virus polymerase proteins is based on their charge characteristics a different name for the related protein of Dhori virus may be appropriate and we thus suggest the designation Pa protein. This nomenclature is proposed for the designation of the Dhori polymerase proteins [Paα-PB1, Pbβ-PB2, and Pbγ-PAI] since the charge characteristics do not appear to be constant across virus groups. For example, the PA-like polymerase protein of influenza C/JJ/50 virus (Yamashita et al., 1989) encodes a weakly basic protein designated P3 (segment 3 gene product). The third largest RNA segment of Thogoto virus encodes a PA-like protein with only a very weak negative charge (−2.5 at pH 7.0; Staunton et al., 1989).

Genes of other viruses also share sequence identity with influenza virus segments. It has been previously demonstrated that some members of the coronavirus group and influenza C virus have a hemagglutinin-esterase (HE) glycoprotein that recognizes the same N-acetyl-9-O-acetylmuramic acid receptors on cells (Vlasak et al., 1988). The influenza C virus HE glycoprotein shares an approximately 30% amino acid sequence identity with a coronavirus HE glycoprotein present in bovine coronavirus (Kienzle et al., 1990) and mouse hepatitis virus (Luytjes et al., 1988). This example of significant identity between a plus-strand RNA and a minus-strand RNA virus supports the concept of a cassette model of evolution for the HE gene and is most likely an example of a recombinational event. In contrast, we feel that the relationship of the Dhori virus group to the orthomyxovirus group is most likely an example of divergent evolution in light of the similarities in genome structure and common replication strategies and of the observation that several Dhori virus gene products share significant identity with orthomyxovirus genes.

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