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Cloning and Sequencing of the Matrix Protein (M) Gene of Turkey Rhinotracheitis Virus Reveal a Gene Order Different from That of Respiratory Syncytial Virus

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Received December 13, 1990; accepted October 10, 1991

Several biochemical properties and the sequence of the fusion glycoprotein (F) have indicated that turkey rhinotracheitis virus (TRTV) is a pneumovirus, subfamily Pneumovirinae of the Paramyxoviridae family. As TRTV was known to generate polycistronic mRNAs, cDNA was generated from TRTV strain UK/3BV/85-infected Vero cell mRNAs using an oligonucleotide primer corresponding to a region of the F gene. Sequencing of four cDNAs revealed that the gene adjacent to the beginning (3' end) of the F gene was that for the matrix (M) protein, i.e., that TRTV had the partial gene order 3'-M-F-5'. This was unexpected as human respiratory syncytial (RS) virus, the type species of the genus Pneumovirus, has the partial gene order 3'-M-SH-G-F-5', where SH and G are the small hydrophobic protein and attachment glycoprotein, respectively. Instead TRTV resembled the Morbillivirus and Paramyxovirus genera of the Paramyxoviridae (subfamily Paramyxovirinae) which have the partial gene order 3'-M-F-5'. Two further oligonucleotides, one corresponding to a sequence near the end of the M gene and the other (oligo 6) to a sequence near the beginning of the F gene, with their 5' ends spaced 300 nucleotides apart on the basis of the cDNA sequence, were used in a polymerase chain reaction (PCR) using genomic RNA as template. Only a PCR product of 0.3 kb was obtained. The same sized product was also obtained using these oligonucleotides and genomic RNA from three other TRTV strains (W/91/78, UK/8544/85, and SA/2381/88) which had been grown in chicken tracheal organ cultures. In addition PCR was performed using genomic RNA from TRTV3BV and SA/2381/88 with oligo B and another oligonucleotide near the 5' end of the gene upstream from M, spaced 1141 nucleotides apart on the basis of the sequence data. Only a 1.14-kb PCR product was obtained. Larger products would have been expected if another gene had been situated between M and F. The absence of such larger products, plus the demonstration that infected cells contained M-F dicistronic mRNAs, supported the conclusion that in the TRTV genome the M gene is adjacent to the F gene in the order 3'-M-F-5'. The 5'termini of the M and F mRNAs were confirmed by mRNA mapping. The TRTV M gene encoded a protein of 254 amino acids, very similar to that of RS virus (256 residues; 37% amino acid identity) but very different from that of the morbillviruses and paramyxoviruses (approximately 350 residues). Thus, on the basis of the sequences of both the M and the F genes, TRTV is more closely related to the pneumoviruses than to the paramyxoviruses and morbillviruses.

INTRODUCTION

Turkey rhinotracheitis virus (TRTV), as the name indicates, was first isolated from turkeys with respiratory disease, although it has subsequently been shown that chickens and other domestic birds can also be infected (McDougall and Cook, 1986; Jones et al., 1987; Gough et al., 1988; and references in Yu et al., 1991). Biochemical analysis of the TRTV-induced proteins (Collins and Gough, 1988; Ling and Pringle, 1988; Cavanagh and Barrett, 1988) and RNAs (Cavanagh and Barrett, 1988) suggested that this virus was a member of the Paramyxoviridae family and more closely related to the genus Pneumovirus (subfamily Pneumovirinae; Pringle, 1991) than to the other two genera in this family, Morbillivirus and Paramyxovirus (subfamily Paramyxovirinae). Our recent cloning and sequencing of the TRTV fusion (F) protein gene has supported this view. Thus the TRTV F protein had 39% amino acid identity with that of human respiratory syncytial (RS) virus, a pneumovirus, but only about half that with members of the other two genera (Yu et al., 1991). In order to identify the gene upstream (3' proximal) from the F gene, we have used an oligonucleotide, based on the F gene sequence, to prime cDNA synthesis in the hope that some clones would be generated from polycistronic mRNA which comprised F and the 3' proximal gene. This was indeed the case, although the gene adjacent to the 3' terminus of the F gene was not that which had been expected from knowledge of the gene order of RS virus and pneumonia virus of mice (PVM) (Collins et al., 1984; Chambers et al., 1990).

MATERIALS AND METHODS

Cells and virus

The TRTV strain used for cloning was 3BV, plaque-purified clone R1.1, this being the UK/3B/85 field iso-
late (McDougall and Cook, 1986) after passage in cell culture (Yu et al., 1991). This virus was propagated in Vero cells (ATCC No. CCL 81; Flow Laboratories, Irvine, Scotland). The South African TRTV strains SA/91/78 and SA/2381/88 and strain UK/8544/85 were grown in chick embryo tracheal organ cultures (Cook et al., 1976). The coronavirus infectious bronchitis virus (IBV), strain Beaudette, was grown in Vero cells.

Preparation of TRTV-induced RNAs

Flasks (75 cm²) of Vero cells were inoculated with 1 ml of undiluted TRTV strain 3BV working stock which resulted in the infection of less than 1% of the cells. After 72 hr at 37 °C actinomycin D (1 μg/ml; Sigma) was added and 24 hr later the cells were lysed by the guanidinium isothiocyanate method as described by Chomczynski and Sacchi (1987) except that isopropanol was substituted by ethanol. Poly(A)+ RNA was selected using Hybond mAP affinity paper according to the manufacturer’s instructions (Amersham International). RNA of strains SA/91/78, UK/8544/85, and SA/2381/88 was extracted from 60 tracheal organ culture rings, using guanidinium isothiocyanate, which had been infected 3 days previously. Virions of TRTV-3BV were prepared by centrifugation of clarified medium from infected Vero cells at 35,000 gmax for 90 min in a Sorvall TFA-20.250 angle rotor. Nucleocapsids were sedimented from Nonidet P-40-produced extracts of infected Vero cells (Ward et al., 1983) at 100,000 gmax for 40 min in a Sorvall TH641 swing-out rotor. The genomic RNA of both virions and nucleocapsids was extracted using guanidinium isothiocyanate. IBV-Beaudette mRNAs were extracted from Vero cells by the same method.

Preparation of cDNA

cDNA synthesis was performed with a cDNA synthesis kit (Amersham) using the poly(A)+ RNA from approximately two 75-cm² flasks of infected cells. Reverse transcription was primed using a synthetic oligonucleotide corresponding to nucleotides 531–547 of the TRTV F gene (Vu et al., 1991).

Cloning and sequencing

cDNA was cloned into EcoRV-cut phagemid pBluescript KS+ (Northumbria Biologicals, Cramlington, Northumbria, UK). Colonies containing TRTV-specific inserts were identified by probing with a 32P-labeled, random hexanucleotide (Pharmacia)-primed probe corresponding to a region (398 nucleotides long) near the 3' part of the F gene, generated by Psfl restriction of clone F17 (Yu et al., 1991). The clones, in pBluescript, were sequenced by dideoxy chain termination following the generation of a nested set of deletions, as described by Yu et al. (1991). Briefly, the recombinant phagemid was digested within the polylinker either with BstXI and XbaI or with Apal and HindIII (Boehringer-Mannheim) to generate deletions from opposite ends and deletions generated with exonuclease III (Northumbria Biologicals). Sequences were analyzed on a MicroVAX 3600 computer system using the Wisconsin program GAP.

Polymerase chain reaction (PCR)

The PCR was performed essentially as described by Sambrook et al. (1989) using genomic RNA from virions and nucleocapsids in the case of strain UK/3BV/85 and total infected tracheal organ culture RNA for the other three TRTV strains. Reverse transcription was done with 2 units of AMV reverse transcriptase (Anglia Biotechnology, Colchester, UK); cDNA was amplified using 2.5 units of Taq polymerase (Boehringer-Mannheim) in 25 cycles, the polymerization step lasting for 5 or 15 min, as described under Results. Two microliters from the 80-μl reaction volume was analyzed in 1% agarose gels, run at 100 V for 2 hr, and stained with ethidium bromide (5 μg/ml).

Oligonucleotides

The positions of the 25-mer oligonucleotides (Severn Biotech, Kidderminster, UK) used for priming DNA synthesis are shown in Fig. 2. Oligonucleotide A (positive sense) corresponded to nucleotides 604–629 of the M sequence (Fig. 3). Oligonucleotide B (negative sense) corresponded to nucleotides 53–78 in the F gene (Vu et al., 1991). Oligonucleotide C (positive sense), which corresponded to part of the gene upstream (3') from the M gene, had its 5' terminus 237 nucleotides from the beginning of the M gene. Oligonucleotide D (negative sense) corresponded to nucleotides 54–79 of the M sequence (Fig. 3).

Primer extension

The 5' terminus of the M mRNA was determined by primer extension using a 25-mer oligonucleotide (oligonucleotide D) complementary to nucleotides 54–79 of the M mRNA sequence shown in Fig. 3. The primer was 5' end-labeled with polynucleotide kinase and [γ-32P]ATP. The primor (100 ng, 5 × 104 cpm) was added to 15 μg of total infected cell extract RNA or 4 μg of poly(A)+ RNA from infected cells. Reverse transcription was performed as described previously (Cavanagh and Davis, 1988). The same oligonucleotide was used to prime a sequencing reaction (Yu et al., 1991) using pBluescript containing insert MF36 as substrate. All reactions were analyzed in the same sequencing gel.
so that the 5' terminus of the M mRNA could be deduced directly by comparison with the sequence determined from clone MF36. The 5' terminus of the F mRNA was mapped in the same way using oligonucleotide B, whose 5' terminus was located 78 nucleotides from the presumed start of the F mRNA, and using pBluescript containing insert MF37.

Detection of polycistronic mRNAs

Polycistronic mRNAs of TRTV were detected by Northern blot analysis (Sambrook et al., 1989) using a restriction fragment in the case of M and a PCR product of F to make random hexanucleotide-primed 32P-labeled probes. IBV-Beaudette mRNAs, all of which have been sequenced (Boursnell et al., 1987), were used as RNA size markers. The IBV-specific probe comprised a complete cDNA copy of the nucleocapsid (N) gene of IBV, produced by the PCR. All IBV mRNAs were detected by this probe because all the mRNAs contain the N gene at their 3' end (Boursnell et al., 1987). TRTV mRNAs were electrophoresed before and after selection of poly(A)+ RNA; IBV RNAs were used without poly(A)+ selection. Before transfer of the RNA to nitrocellulose, the gel was exposed to uv light (302 nm) (Ultra-Violet Products, Inc., U.S.A.) for 2 min to nick the RNA sufficiently to improve the transfer of genome and anti-genome TRTV RNA and the large IBV mRNAs.

Sequence databases

The sequence of the M mRNA (nucleotides 1–824 in Fig. 3) will appear in the DDBJ, EMBL, and Genbank Nucleotide Sequence Databases under Accession No. X58639.

RESULTS

Cloning and sequencing of the gene 3' proximal to the F gene

In order to identify the gene upstream (3' proximal) from the F gene, an F gene-specific oligonucleotide was used to generate cDNA from poly(A)+-infected cell RNA, and clones were selected using a radiolabeled F gene restriction fragment. This strategy was based on the knowledge, gained from probing Northern blots of infected cell RNA with F gene-specific probes, that polycistronic mRNAs were generated during transcription (Fig. 1, lanes 3 and 5) and the hope that some of these mRNAs contained F and the upstream gene. Six clones were obtained, two of which did not extend beyond the primer in the direction of the 5' end of the F gene, to nucleotide position 1170 in the F gene sequence (Vu et al., 1991). Every nucleotide of the gene 3' proximal to F was sequenced from at least two clones; 93% were sequenced from three clones. The entire sequence was determined from both strands of the cDNA and 96% of nucleotides were determined 5 to 16 times and the remaining 4% on 3 or 4 occasions.

The M gene is adjacent to the 3' end of the F gene of TRTV

As described above the identity of the gene sequences contained within each clone was ascertained by sequencing. Clones MF35 and 36 encoded a large ORF of 254 amino acids. The next longest ORF would encode an ORF of only 51 residues, other ORFs being less than half that size. Comparison of the sequence of the longest ORF with a databank of Paramyxoviridae sequences revealed that it had 37% amino acid identity with the M protein of RS virus which comprises 256 residues (Fig. 3). The calculated molecular weights of
The solid areas show those parts of the clones that were se-
the sequence just upstream from the F ORF (nucleo-
tides 827-835, doubly underlined in Fig. 3) which we
the ORF was the sequence GGGACAAGU (nucleotides
that nucleotides l-9 and 827-835 (Fig. 3) marked the
the sequence GGGACAAGU. This was direct evidence
the tension showed that the lower band of the doublet co-
tective mRNAs having been reached (Fig. 4). The same
sequences (Vu et al., 1991). Confirmation that these sequences
did mark the beginnings of the M and F mRNAs was
migrated with a G residue which was the beginning of
the M protein for TRTV and RS virus were 27,595 and
28,714, respectively.

Almost immediately upstream from the beginning of
the ORF was the sequence GGGACAAGU (nucleotides
1–9, doubly underlined in Fig. 3) which was identical to
the sequence just upstream from the F ORF (nucleo-
tides 827–835, doubly underlined in Fig. 3) which we
have suggested is the start sequence for the F mRNA
(Yu et al., 1991). Confirmation that these sequences
did mark the beginnings of the M and F mRNAs was
obtained by mapping the 5’ end of the respective
mRNAs. Gene-specific oligonucleotides were 5’ end-
labeled and then used for primer extension. Both the M
and the F primer-extended products terminated to yield
a double band, consistent with the ends of the respective
mRNAs having been reached (Fig. 4). The same
end-labeled oligonucleotides were used in sequencing
reactions, with [35S]ATP present, with pBluescript con-
taining either the M or the F cDNA inserts. Electropho-
resis of the reaction products in the same gels as those
used for analysis of the cDNA generated by primer ex-
tension showed that the lower band of the doublet co-
migrated with a G residue which was the beginning of
the sequence GGGACAAGU. This was direct evidence
that nucleotides 1–9 and 827–835 (Fig. 3) marked the
beginning of the M and F mRNAs, respectively. The
upper band of the doublet, which was sometimes of
lower intensity, was considered most likely to be an
artifact, as proposed previously (Jorgensen et al., 1987).
The presence of partial cross-banding in the
pBluescript-generated sequence was a consequence
of the presence of some n − 1 oligonucleotide in the primer preparations.

At the end of the TRTV F mRNA is the sequence
AGUUA followed three nucleotides later by the poly(A)
tail (Yu et al., 1991). Immediately prior to the start of
the F gene, i.e., near the end of the M sequence, was the
similar sequence AGUCA (nucleotides 812 to 816, sin-
gly underlined, in Fig. 3) followed two nucleotides later
by AAAAAAUU. This observation, coupled with the
identification of the 5’ terminus of the F mRNA (Fig. 4B),
indicates that these sequences mark the end of the M
mRNA, the UU residues being the M-F intergenic re-
region (in the mRNA sense). Thus nucleotides 1 and 824
in Fig. 3 define the termini of the TRTV M mRNA.

**Northern blot analysis**

Northern blots of TRTV-infected cell RNA, before
and after selection of poly(A)+ RNA, were probed with
F- and M-specific probes. In addition to the monocis-
tronic F mRNA the F probe hybridized to an mRNA of
approximately 2.63 kb (Fig. 1, lanes 3 and 5; size esti-
ated using IBV mRNAs as markers), a mRNA of this
length also being revealed by the M-specific probe
(Fig. 1, lanes 7 and 9). The size estimate of 2.63 kb is
close to that of 2.56 kb predicted for a M-F dicistronic
mRNA based on the sequences for M and F of TRTV
and assuming a poly(A) tail of approximately 100 nu-
cleotides. The probes also detected small amounts of
several other mRNA species and some genome-sized
RNA; Afzal et al. (1990) detected both genome and
anti-genome in poly(A)+-selected mumps virus RNAs.
TRTV clones MF35 and MF36, which extended into
the gene upstream from M, could have been generated
from one or more of these additional RNA species.

**PCR evidence that the TRTV M gene is adjacent to
the F gene**

From the Northern blot analysis it was most likely
that clones MF34–37 had been generated by reverse
transcription of polycyclstronic mRNAs which faithfully
reflected the gene order of the M and F genes in the
genome. However, it was possible that in the TRTV
genome there were one or more genes between M and
F and that during the synthesis of our cDNA clones
there might have been a fault leading to the deletion of
intervening genes. Alternatively the virus RNA prepara-
tion may have contained deletion genomes which pro-
duced defective RNA from which the cDNAs were gen-
erated, although we have no evidence for the exis-
tence of such RNA. To investigate these possibilities
we have used the PCR to generate cDNA, using UK/
3RV/85 virion RNA as template, covering the proposed
M-F junction. First oligonucleotides A and B (Fig. 2)
were used to prime cDNA synthesis. If the gene order M-F was correct, the only product generated by the PCR would be 300 nucleotides in length. This was indeed the case, irrespective of whether the elongation time in the PCR was 5 min (Fig. 5, lane b) or 15 min (Fig. 5, lane c). The long incubation time was included to permit the amplification of any genes which might have been located between M and F; no long PCR products corresponding to such a RNA population were observed. Sequencing of the 0.3-kb PCR product after cloning confirmed that the sequence was the same as that obtained from sequencing clones MF35 and 36. Finally, the PCR was performed using oligonucleotides C and B with an extended polymerization time of 15 min. This produced only a single product of approximately 1.14 kb (Fig. 5, lane f) which was that predicted from the distance between oligonucleotides C and B based on the sequencing of clones MF35 and 36. The same result was obtained with RNA extracted from Vero cells which had been infected at very low multiplicity with TRTV strain 3BV which was only two passages beyond the plaque-purification stage (Fig. 5, lane g). In addition the PCR produced only one product, of 1.14 kb, when RNA of the tracheal organ culture-passaged strain SA/2381/88 was used (Fig. 5, lane h).

A second PCR was done using oligonucleotides C and D whose 5' ends were spaced 316 nucleotides apart (Fig. 2). In agreement with the analysis of the 3' (genome sense) portion of clones MF35 and 36, the PCR produced only the predicted product of approximately 0.3 kb (Fig. 5, lanes d and e), confirming that the sequence upstream from the beginning of M was as had been deduced from sequencing clones MF35 and 36. Finally, the PCR was performed using oligonucleotides C and B with an extended polymerization time of 15 min. This produced only a single product of approximately 1.14 kb (Fig. 5, lane f) which was that predicted from the distance between oligonucleotides C and B based on the sequencing of clones MF35 and 36. The same result was obtained with RNA extracted from Vero cells which had been infected at very low multiplicity with TRTV strain 3BV which was only two passages beyond the plaque-purification stage (Fig. 5, lane g). In addition the PCR produced only one product, of 1.14 kb, when RNA of the tracheal organ culture-passaged strain SA/2381/88 was used (Fig. 5, lane h).
DISCUSSION

The deduced sequence of the M protein of TRTV confirms the notion, based initially on superficial observations of TRTV-induced proteins and RNAs and more recently on the sequence of the F protein, that of the three genera within the Paramyxoviridae family, TRTV most closely resembles the Pneumovirus genus. First, TRTV and RS virus, the type species of the Pneumovirus genus, have M proteins of almost identical length, comprising 254 and 256 residues, respectively, in contrast to the much longer M polypeptides (335–377) possessed by paramyxoviruses (Chambers et al., 1986; McGinnes and Morrison, 1987; Sato et al., 1987; Blumberg et al., 1984; Spriggs et al., 1987; Luk et al., 1987; Galinski et al., 1987; Sakai et al., 1987; Sheshberadaran and Lamb, 1990; Elliott et al., 1989; Elango, 1989) and morbilliviruses (Bellini et al., 1986; Limo and Yilma, 1990). Second, the M protein hydrophathy plots for TRTV and RS virus are remarkably similar (Fig. 6).

The Wisconsin program GAP revealed that the M protein of TRTV had 37% amino acid identity with that of RS virus and 17–21% with members of the other two genera. However, this figure of 17–21% tends to exaggerate the extent to which the M proteins of the pneumoviruses resemble those of the other two genera since 7–11 gaps had to be inserted in order to produce even this low level of alignment.

The M protein of paramyxoviruses forms an ordered array at the inner surface of the virion envelope (Heggeness et al., 1982; Buechi and Bachi, 1982) and plays an important role in the virus maturation process, the protein being believed to interact with the viral ribonucleoprotein (RNP) (Mountcastle et al., 1974; Shimizu and Ishida, 1975; Yoshida et al., 1976) the inner surface...
TRTV and RS virus have a highly hydrophobic terminus of the M protein of the morbilliviruses which proteins of paramyxoviruses and morbilliviruses are basic and acidic residues, respectively, in this region. However, this is not a general observation for pneumoviruses as RS virus has almost equal numbers of acid and basic residues (8 and 7, respectively) in the first third of the molecule (Satake and Venkatesan, 1984). Parainfluenza virus 3/Sendai virus group of paramyxoviruses, whereas within the morbillivirus genus the amino acid identity is about 75% (Rima, 1989). This suggests that the evolutionary pathways of TRTV and

![Hydropathy profiles for the (A) TRTV and (B) RS virus M proteins. Positive values indicate hydrophobic regions and negative values indicate hydrophilic regions. The midpoint line represents a grand average of the hydropathy of the amino acid composition of a large number of sequenced proteins (Kyte and Doolittle, 1982). Each point on the profile represents the average hydropathy of a span of five residues.](image)

The many interactions of M are believed to involve complex interactions of both hydrophobic and ionic natures (Caldwell and Lyles, 1986; Faaberg and Peeples, 1988). As for morbilliviruses and paramyxoviruses (Chambers et al., 1986; Gelfinski et al., 1987, Limo and Yilma, 1990) the hydropathy plots for TRTV and RS virus indicate that the M protein is moderately hydrophobic. While none of the sequences are obvious candidates for membrane-spanning structures, both TRTV and RS virus have a highly hydrophobic sequence of 14 residues centering around amino acid 200 (Figs. 3 and 6). This stretch in particular might be membrane associated. Bellini et al. (1986) have drawn attention to a sequence of 13 to 14 largely nonpolar residues situated about 90 residues from the carboxy terminus of the M protein of the morbilliviruses which might interact with the lipid environment of the virus envelope. With respect to ionic interactions the M proteins of paramyxoviruses and morbilliviruses are basic in nature, basic and acidic residues accounting for 12–14 and 8%, respectively, of M residues. Many (12–16) of the basic residues occur in pairs. In contrast, TRTV and RS virus M proteins have only a small preponderance of basic over acidic residues, these residues accounting for 9–11 and 8–9%, respectively, of total residues. TRTV and RS virus have only 6 and 2 basic residues, respectively, in pairs (Satake and Venkatesan, 1984). TRTV M protein does resemble that of the other two genera in that the amino terminal one-third of M has an acidic character, TRTV M having 7 and 2 acidic and basic residues, respectively, in this region. However, this is not a general observation for pneumoviruses as RS virus has almost equal numbers of acid and basic residues (8 and 7, respectively) in the first third of the molecule (Satake and Venkatesan, 1984).

All paramyxoviruses and morbilliviruses examined to date have the partial gene order 3′-M-F-(SH)-HN/H-5′; SH is in parentheses to indicate that this gene is not possessed by all viruses in these genera. In contrast two pneumoviruses, human RS virus and PVM, have the equivalent genes in the order 3′-M-SH-G-F-5′; i.e., they have, unlike the other two genera, the SH and cell attachment protein gene (G) on the 3′ side of the F gene (Collins et al., 1984; Chambers et al., 1990). It was a surprise, therefore, when we found that TRTV had the M gene immediately adjacent to the 3′ side of the F gene. There was no evidence from probing Northern blots that TRTV UK/3BV/85 had generated defective RNA which might have had genes deleted between M and F. Even if such defective RNA had been present, then in order to generate clones MF34–37 the deletion event would have had to be such as to delete genes in their entirety while at the same time preserving completely intact the M gene and the beginning of F. We have confirmed the beginning of both the M and the F mRNAs by 5′ mapping and have shown that M has mRNA start and end sequences very similar to those of F. The PCR data supported the gene order deduced from clones MF34–37. Although the PCR would tend to favor the amplification of small products, e.g., the 300-nucleotide product obtained using oligonucleotide pairs A/B, over larger ones, e.g., the product which would have been obtained if there had been a gene between M and F, there was no sign at all of larger PCR products to substantiate the view that there was a gene between M and F. Moreover, the gene order M-F was further supported by the PCR data obtained with three other TRTV strains, representing isolates from two continents, isolated over a 10-year period, which had different passage histories, and which had been propagated in chicken tracheal organ cultures.

Although the M protein of TRTV is clearly more closely related to that of RS virus than to that of the other two genera, the amino acid identity was only 37%. This is similar to the amino acid identity shared between the morbilliviruses and the human and bovine parainfluenza virus 3/Sendai virus group of paramyxoviruses, whereas within the morbillivirus genus the amino acid identity is about 75% (Rima, 1989). This suggests that the evolutionary pathways of TRTV and
RS virus diverged before those of members of the morbillivirus genus.

The subfamily Pneumovirinae is of particular interest because, based on the analysis of the genomes of human RS virus and PVM, the member viruses contain several genes that are not present in species of the Paramyxovirinae subfamily (Collins et al., 1984; Chambers et al., 1990). Given that TRTV has some pneumovirus characteristics but appears to have diverged a considerable time ago from the other known members of this genus, further analysis of the number, nature, and order of TRTV genes will be instructive with regard to whether those genes currently known to be unique to the pneumovirus genus are essential for replication. In addition, study of TRTV may help to clarify the evolutionary relationships between the genera of the Paramyxoviridae family.

ACKNOWLEDGMENTS

We thank Rachel Purvis for technical assistance, Dr. Jane Cook for providing isolates of TRTV, and Dr. Tom Barrett for advice during the course of this work. This work was supported by the Ministry of Agriculture, Fisheries and Food.

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