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Sequence Analysis of the Nucleocapsid Protein Gene of Human Coronavirus 229E

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Human coronaviruses are important human pathogens and have also been implicated in multiple sclerosis. To further understand the molecular biology of human coronavirus 229E (HCV-229E), molecular cloning and sequence analysis of the viral RNA have been initiated. Following established protocols, the 3'-terminal 1732 nucleotides of the genome were sequenced. A large open reading frame encodes a 389 amino acid protein of 43,366 Da, which is presumably the nucleocapsid protein. The predicted protein is similar in size, chemical properties, and amino acid sequence to the nucleocapsid proteins of other coronaviruses. This is especially evident when the sequence is compared with that of the antigenically related porcine transmissible gastroenteritis virus (TGEV), with which a region of 46% amino acid sequence homology was found. Hydrophathy profiles revealed the existence of several conserved domains which could have functional significance. An intergenic consensus sequence precedes the 5'-end of the proposed nucleocapsid protein gene. The consensus sequence is present in other coronaviruses and has been proposed as the site of binding of the leader sequence for mRNA transcriptional start. This region was also examined by primer extension analysis of mRNAs, which identified a 60-nucleotide leader sequence. The 3'-noncoding region of the genome contains an 11-nucleotide sequence, which is relatively conserved throughout the Coronavirus family and lends support to the theory that this region is important for the replication of negative-strand RNA.

INTRODUCTION

Human coronavirus 229E (HCV-229E) belongs to one of two major antigenic groups of human coronaviruses (MacNaughton, 1981). It shares antigenic relationships with other coronaviruses, such as porcine transmissible gastroenteritis virus (TGEV), feline infectious peritonitis virus (FIPV), and canine coronavirus (CCV). The other well-characterized human coronavirus, HCV-OC43, is in a separate antigenic group which includes mouse hepatitis virus (MHV) and bovine coronavirus (RCV). Both human coronaviruses are mainly respiratory pathogens and have been estimated to cause up to 25% of common colds (McIntosh et al., 1974; Wege et al., 1982). They have also been implicated in gastrointestinal diseases (Resta et al., 1985). Furthermore, the isolation of coronaviruses bearing an antigenic relationship to HCV-OC43 from the central nervous system of two patients with multiple sclerosis has suggested a possible etiologic relationship between human coronaviruses and multiple sclerosis (Burks et al., 1980). This possibility is supported by the observation that neurotropic strains of MHV cause demyelination in the central nervous system of rodents (Weiner and Stohlman, 1978). Thus, human coronaviruses are important human pathogens.

The structural and biochemical properties of several coronaviruses, particularly MHV and avian infectious peritonitis virus (IBV), have been well characterized (Lai et al., 1987; Boursnell et al., 1987). The virion contains a single stranded, positive sense RNA molecule (molecular weight 6–8 x 10^6 Da) (Lai and Stohlman, 1978) associated in a helical conformation with nucleocapsid proteins (N). The viral nucleocapsid is enclosed by an envelope, in which are embedded at least two types of viral proteins, the peplomer (E2) and matrix (E1) glycoproteins. Coronavirus RNA replication occurs in the cytoplasm of infected cells and is mediated by a virus-encoded RNA-dependent RNA polymerase (Brayton et al., 1982). The virus-specific mRNA in infected cells comprises a genomic-sized RNA plus six subgenomic mRNA species. These mRNAs are arranged in a nested-set structure, which is characterized by RNAs having common 3'-termini but extending for varying lengths in the 5' direction (Lai et al., 1981). Only the 5' proximal regions of each mRNA are translated (Rottier et al., 1981). A unique feature of the structure of coronavirus is the existence, at the 5'-end of each mRNA, of an identical leader sequence. This sequence is derived from the 5'-end of the genomic RNA and is of approximately 70 nucleotides in length (Lai et al., 1983, 1984). Recent evidence has supported a role for the leader sequence in mediating a novel type of discontinuous transcription of genomic RNA (Baric et al., 1985; Makino et al., 1986; Shieh et al., 1987).

In contrast to other coronaviruses, the molecular biology of human coronaviruses is relatively poorly understood. The genomic RNA of both HCV-229E and HCV-OC43 has a molecular weight of approximately 6

Sequence data from this article have been deposited with the EMBL/GenBank Data Libraries under Accession No. J04419.

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The six subgenomic RNA species appear to have lower molecular weights than those of the corresponding MHV RNAs (Weiss and Leibowitz, 1981). The structure of these mRNAs is not yet known. Analysis of purified HCV-229E virions has revealed three major polypeptides: a glycosylated protein with a molecular weight of 180 kDa, a phosphorylated nucleocapsid protein of 50 kDa, and a family of polypeptides with molecular weights of 25, 23, and 21 kDa (Kemp et al., 1984). In addition, several minor nonstructural polypeptides of 107, 92, and 39 kDa have been identified (Kemp et al., 1984). The functions of these proteins have not yet been characterized.

To further understand the molecular biology of HCV-229E, we have initiated molecular cloning and sequence analysis of HCV-229E RNA. In this paper we report the sequence analysis of the gene encoding the nucleocapsid protein of HCV-229E. In addition, the mRNA leader sequence was also identified. The results are compared with sequences of other coronaviruses including MHV, BCV, IBV, and TGEV.

### MATERIALS AND METHODS

#### Virus and cells

HCV-229E (obtained from Dr. J. Fleming, University of Southern California) was propagated at low multiplicities of infection in human fetal lung cells L132 (Kennedy and Johnson-Lussenberg, 1975/1976), using Dulbecco’s modified Eagle’s medium (DMEM) supplemented with 10% fetal calf serum.

#### Virus purification and preparation of virion RNA

Following a virus adsorption period of 1 hr at 37°C, HCV-229E-infected L132 monolayers were incubated at 37°C for 24 to 48 hr, at which time the cell culture fluid was harvested. Viruses were precipitated from 2 liters of culture fluid with 50% ammonium sulfate and centrifuged at 8000 rpm for 30 min. The pellet was resuspended in NTE buffer and centrifuged at 8000 rpm for 30 min. The supernatant of culture fluid with 50% ammonium sulfate was collected and treated with proteinase K (0.2 mg/ml) for 20 min at 37°C, followed by 1% SDS for 30 min at 37°C. Genomic RNA was extracted with phenol and then with phenol/chloroform, and precipitated with ethanol.

The diluted virus suspension was pelleted at 26,000 rpm for 13 hr at 4°C in a Beckman SW28.1 rotor. The virus band at the interface between discontinuous sucrose gradient consisting of 60, 50, and 30% sucrose was collected and diluted three-fold with NTE buffer. The diluted virus suspension was centrifuged on a linear sucrose gradient at 26,000 rpm for 13 hr at 4°C in a Beckman SW28.1 rotor. The virus band at the interface between 50 and 30% sucrose was collected and diluted three-fold with NTE buffer. The diluted virus suspension was centrifuged on a linear sucrose gradient at 26,000 rpm in an SW28.1 rotor for 4 hr at 4°C. The virus band was collected and treated with proteinase K (0.2 mg/ml) for 20 min at 37°C, followed by 1% SDS for 30 min at 37°C. Genomic RNA was extracted with phenol and then with phenol/chloroform, and precipitated with ethanol.

### Preparation of intracellular RNA

Monolayers of L132 cells grown in 100 X 20-mm culture dishes were infected with HCV-229E. Cells were incubated in phosphate-free DMEM containing 1% diazoyed fetal calf serum 4 hr prior to RNA extraction. Actinomycin D (1 μg/ml) (Sigma) and [3H]-orthophosphate (70 μCi/ml) (ICN Radiochemicals) were added at 3 and 2 hr, respectively, prior to RNA extraction at 1 hr post-infection (p.i.). Cells were collected in cold phosphate-buffered saline and centrifuged at 5000 rpm for 3 min at 4°C. The pellet was mixed with cold 0.5% Nonidet P-40 in NTE buffer, incubated for 10 min at 4°C, and then centrifuged at 5000 rpm for 3 min. The supernatant was transferred to a fresh tube containing 1/10 vol of 10% SDS at room temperature and vortexed briefly. Intracellular RNA was extracted with phenol and phenol/chloroform and precipitated with ethanol. Poly(A)-containing RNA was selected by oligo(dT)-cellulose chromatography as previously described (Makino et al., 1984).

To examine the kinetics of viral mRNA synthesis, intracellular RNA was extracted from virus-infected L132 monolayers in 60 X 15-mm culture dishes at 7, 21, 29, 46, and 58 hr postinfection.

#### cDNA cloning

cDNA cloning was performed using a modified method of Gubler and Hoffman (1983). The poly(A)-containing RNA extracted from 229E-infected L132 monolayers was precipitated, dried, and resuspended in 6.72 μl of autoclaved water. The RNA was incubated with 10 mM methylmercuric hydroxide in an 8 μl total volume for 10 min at room temperature. First-strand cDNA synthesis was carried out in a 50-μl reaction mixture containing 60 units RNasin (Promega Biotec), 10 mM MgCl₂, 100 mM KCl, 50 mM Tris–HCl (pH 8.3 at 42°C), 10 mM DTT, 1.25 mM dNTPs, 40 μCi [α-32P]dATP (3000 Ci/mmol), 28 mM β-mercaptoethanol, and 10 ng oligo(dT)₁₂₋₁₈ primer. After 5 min at room temperature, 40 units of AMV reverse transcriptase (Life Science) was added and the mixture was incubated for 1 hr at 42°C. The reaction was stopped by adding 4.4 μl of 250 mM EDTA. The products were extracted with phenol/chloroform and precipitated with ethanol containing 0.3 M ammonium acetate. For second-strand synthesis, the 100-μl reaction mixture contained 5 mM MgCl₂, 100 mM KCl, 20 mM Tris–HCl (pH 7.5), 50 μg/ml bovine serum albumin (BSA), 10 mM ammonium sulfate, 0.15 mM β-NAD, 100 μM dNTPs, 25 units of Escherichia coli DNA polymerase I, 2 units of E. coli DNA ligase, and 0.8 units of RNase H. Sequential incubations were for 1 hr at 12°C and 1 hr at 22°C. The reaction was stopped by the addition of 8.7 μl of 250 mM EDTA and the products were extracted with phenol/
chloroform and precipitated with ethanol in the presence of 0.3 M ammonium acetate. Homopolymeric tailing of double-stranded cDNA with poly(C) was carried out in a 12-μl reaction mixture containing 10 units of terminal transferase, 200 mM potassium cacodylate, 0.5 mM CaCl₂, 25 mM Tris—HCl (pH 6.9), 2 mM DTT, 250 μg/ml BSA, and 50 μM dCTP at 37°C for 4 min. The dC-tailed double-stranded DNA was annealed to 200 μg of dG-tailed PstI-cut pBR322 plasmid in 20 μl of a buffer containing 10 mM Tris—HCl (pH 7.4), 100 mM NaCl, and 0.25 mM EDTA. The mixture was incubated for 5 min at 68°C and then cooled slowly overnight. The annealed molecules were used to transform E. coli MCl061 as described (Dagert and Erhlich, 1979).

**Colony hybridization**

Colonies grown on LB/tetracycline plates were incubated at 37°C for 12 hr and transferred to Colony/Plaque Screen disks (New England Nuclear). Bacterial lysis and DNA fixation were carried out according to the methods previously described (Grunstein and Hogness, 1975). The disks were prehybridized in a solution containing 0.2% polyvinylpyrrolidone (MW 40,000), 0.2% Ficoll (MW 400,000), 0.2% BSA, 0.06 M Tris—HCl (pH 7.5), 1% SDS, 1 M NaCl, 10% dextran sulfate, and 100 μg/ml denatured salmon sperm DNA at 65°C for 6 hr. Fragments derived from either the 5’- or 3’-ends of gene 7 were labeled with 32P by nick-translation and added to the solution. Hybridization was carried out for 20 hr at 65°C. The disks were then washed twice in 2X SSC (0.3 M NaCl, 30 mM sodium citrate) at room temperature, twice in 2X SSC containing 1% SDS for 30 min at 65°C, and twice in 0.1X SSC at room temperature for 30 min. The disks were air-dried and exposed to X-ray film at -70°C.

**Northern hybridization**

Intracellular RNA from virus-infected cells was denatured by glyoxal treatment and separated by electrophoresis on a 1% agarose gel containing 10 mM sodium phosphate (pH 7.0) as described previously (McMaster and Carmichael, 1977). RNA transfer to Biodyne nylon filters (ICN Radiochemicals) and subsequent hybridization were performed according to the method described by Thomas (1980).

**Primer extension**

A synthetic oligodeoxyribonucleotide was 5’-end-labeled with [γ-32P]ATP by polynucleotide kinase (Pedersen and Haseltine, 1980). The total amount of poly(A)-containing RNA extracted from 229E-infected cell monolayers in three 150 mm culture dishes was incubated in 8 μl of distilled water containing 10 mM methylmercuric hydroxide for 10 min at room temperature. A further incubation was carried out in a 50-μl reaction volume containing 60 units of RNasein (Promega), 10 mM MgCl₂, 100 mM KCl, 50 mM Tris—HCl (pH 8.3 at 42°C), 10 mM DTT, 1.25 mM dNTPs, 28 mM β-mercaptoethanol, 5’-end-labeled synthetic oligodeoxyribonucleotides, and 20 units of AMV reverse transcriptase (Life Science) for 1 hr at 42°C. Reaction products were extracted with phenol/chloroform, precipitated with ethanol, and then analyzed by electrophoresis on a 6% polyacrylamide gel containing 8.3 M urea. The primer-extended product was identified by autoradiography and eluted from the gel according to the published procedure (Maxam and Gilbert, 1977).

**DNA sequencing**

Sequencing was carried out by the dideoxyribonucleotide chain termination method (Sanger et al., 1977) as well as the chemical modification procedure (Maxam and Gilbert, 1977). In the first method, fragments of cDNA inserts generated by various restriction endonucleases were cloned into the M13 vectors mp18 and mp19 (Messing and Vieira, 1982). [α-32P]-dATP was used as a label. Sequence data were also obtained by chemical modification (Maxam and Gilbert, 1977) of various cDNA fragments subcloned into the pT7-3 vector (Tabor and Richardson, 1985). In the second method, cDNA fragments were 3’-end-labeled with Klenow fragment at internal restriction sites or, alternatively, at the polylinker cloning site of pT7-3. End-labeled cDNA restriction fragments were separated by electrophoresis on preparative polyacrylamide gels (Maxam and Gilbert, 1980) and purified as described previously (Hansen et al., 1980; Hansen, 1981). Sequencing of the primer-extended product of mRNA7 was performed by the chemical modification procedure (Maxam and Gilbert, 1977). Sequence analysis was performed by the Intelligenetics and Seqaid programs. Hydrophathy profiles were constructed using the PepPlot program of the University of Wisconsin Computer Genetics Group, which employs both the Kyle-Doolittle (KD) and Goldman, Engelman, Steitz (GES) algorithms.

**RESULTS**

**Kinetics of HCV-229E mRNA synthesis**

To determine the optimum time for extracting 229E-specific mRNAs, we first studied the kinetics of virus-specific mRNA synthesis. Intracellular RNA was extracted from infected L132 monolayers at specified times p.i. The RNA was separated by agarose gel electrophoresis (Fig. 1). As can be seen, viral mRNA synthesis could be detected as early as 7 hr p.i. and reached maximum at 28 hr p.i. Thereafter, total RNA
synthesis gradually declined. By 46 hr p.i. only the most abundant mRNA species were evident. The number and size of these mRNA species are comparable to those of MHV mRNAs and are in agreement with previously published results (Weiss and Leibowitz, 1981). Significantly, mRNA 2a, which was previously found only in BCV-infected cells and proposed to encode hemagglutinins (King et al., 1985; Keck et al., 1988), was not present. This is consistent with the finding that HCV-229E does not have hemagglutinating activity (Hierholzer, 1976). The relative amounts of the mRNA species were the same throughout the replication cycle. Therefore, in all of our subsequent experiments, the virus-specific intracellular RNAs were extracted at 15 hr p.i.

**Molecular cloning of HCV-229E genomic RNA and intracellular virus-specific mRNAs**

cDNA cloning was initially performed using virion genomic RNA as a template. The size of inserts in the resultant cDNA clones ranged from 0.2 to 0.5 kb in length. One clone, A34, contained a 0.45-kb insert, which was subsequently characterized by restriction mapping and Northern blot analysis. The 0.45-kb fragment was labeled with $^{32}$P by nick-translation and hybridized with intracellular RNA from 229E-infected cells. The result, shown in Fig. 2, revealed that the fragment hybridized to each of the mRNA species. This result suggested that the HCV-229E subgenomic mRNAs possess a nested-set structure similar to other coronaviruses (Lai, 1988) and that A34 represented a cDNA clone of either the 3' end of the genomic RNA or the leader sequence.

Cloning was subsequently carried out using intracellular RNA from 229E-infected cells as a template. The
resulting cDNA clones were screened by colony hybridization using the 0.45-kb fragment from clone A34 as a nick-translated probe (Fig. 3). Several positive colonies were identified and characterized further. Clone L8 contained a 3.6-kb insert but lacked a 3'-poly(A) tail. Clone L37, which contained an insert of 1.7 kb, overlapped L8 but was 0.1 kb shorter at the 3'-end. This clone also lacked a poly(A) sequence (see below). Therefore, additional cDNA clones were isolated using a 0.24-kb Bal I–EcoRI fragment of L0 (Fig. 3a) as a probe. These latter clones were further characterized by Southern blot analysis. Clone S10 contained an insert of 0.8 kb which overlapped the 3'-ends of the two previous clones and extended another 0.4 kb in that direction. Figure 3b shows the orientation and sizes of clones L8, L37, S10, and A34 with reference to the viral genome. Restriction enzyme sites used for sequencing are also shown.

Sequencing of the cDNA clones

To determine the sequence of the 3'-end of HCV-229E genome, various restriction fragments of L8, L37, and S10 were subcloned into M13 vectors. For L8, only the 1.2-kb fragment extending from an internal PstI site toward the 3'-end was sequenced. Clone L37 was also sequenced in part. Figure 3c shows the cDNA fragments and strategy used in sequencing. Each region
was verified by dideoxy chain termination sequencing of both strands or by the chemical modification method. Clone S10 was found to have a poly(A) stretch of 34 bases. Figure 4 shows the complete DNA sequence with a translation of the main open reading frame (ORF) in one-letter amino acid code. This ORF extends from base 147 to base 1313 and predicts a 389 amino acid protein with a molecular weight of 43,366 Da. This predicted molecular weight is slightly smaller than the measured molecular weight of the nucleocapsid protein of HCV-229E, which is 50 kDa as determined by SDS-polyacrylamide gel electrophoresis (MacNaughton, 1980). The difference is probably due to phosphorylation or other modification of the protein.

Figure 5 shows a schematic diagram of the possible ORFs obtained by translating the nucleotide sequence. The ORF in frame 3 is likely the one which encodes the nucleocapsid protein. In frame 2, the 5'-flanking region probably contains part of the sequence of the matrix protein encoded by gene 6. This possibility is sup-

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**FIG. 4.** The primary nucleotide sequence of the 3'-end of HCV-229E RNA and the deduced amino acid sequence of the nucleocapsid protein.

A primer extension study was carried out using a synthetic oligodeoxyribonucleotide complementary to an 18-mer sequence underlined near the 5'-end of the gene. The 3' noncoding region contains a conserved sequence which is shown by the double line. The intergenic conserved sequence, TCTAAACT, is also shown (dotted line) was verified by dideoxy chain termination sequencing of both strands or by the chemical modification method. Clone S10 was found to have a poly(A) stretch of 34 bases. Figure 4 shows the complete DNA sequence with a translation of the main open reading frame (ORF) in one-letter amino acid code. This ORF extends from base 147 to base 1313 and predicts a 389 amino acid protein with a molecular weight of 43,366 Da. This predicted molecular weight is slightly smaller than the measured molecular weight of the nucleocapsid protein of HCV-229E, which is 50 kDa as determined by SDS-polyacrylamide gel electrophoresis (MacNaughton, 1980). The difference is probably due to phosphorylation or other modification of the protein. The predicted protein shares features with the nucleocapsid proteins of TGEV, MHV, BCV, HCV-OC43, and IBV (Kapke and Brian, 1986; Armstrong et al., 1983; Lapps et al., 1985, 1984). Namely, the protein is highly basic and rich in serine residues. Sixty percent of the amino acid residues are basic and 12% are acidic. There are 39 serine residues (10% of total), which are presumed to be sites of phosphorylation (Stohlman and Lai, 1979). When compared to TGEV, with which HCV-229E shares antigenic properties, both N proteins have identical amounts of basic and acidic amino acids and serine residues and similar molecular weights (Kapke and Brian, 1986).

Figure 5 shows a schematic diagram of the possible ORFs obtained by translating the nucleotide sequence. The ORF in frame 3 is likely the one which encodes the nucleocapsid protein. In frame 2, the 5'-flanking region probably contains part of the sequence of the matrix protein encoded by gene 6. This possibility is sup-
reported by the finding that reading frame 2 remains open at the extreme 5'-end. Furthermore, the sequence TCTAAACT, which is found in the intergenic regions of several other coronaviruses (Kapke and Brian, 1986; Skinner and Siddell, 1984; Armstrong et al., 1983; Lapps et al., 1987; Kamahora et al., 1988; Budzilowicz et al., 1985), is also present between the presumed initiation codon of the main ORF and the 3'-end of gene 6. This sequence is the proposed site of fusion of the leader sequence with the mRNA coding region (Shieh et al., 1987; Makino et al., 1986; Budzilowicz et al., 1985).

The 3'-noncoding region contains the sequence TGGAAGAGCCA, 75 nucleotides from the 3'-end (Fig. 4), which is relatively conserved among coronaviruses and is found at approximately the same location in all of these viral genomes (Kapke and Brian, 1986; Skinner and Siddell, 1984; Armstrong et al., 1983; Lapps et al., 1987; Kamahora et al., 1988; Boursnell et al., 1985) (Table 1). There is only one nucleotide difference in this conserved sequence when it is compared with that of TGEV, BCV, and HCV-OC43. Two and three nucleotide differences are found in IBV and MHV, respectively. This conservation of sequence and location suggests that it may be important for viral RNA replication.

In frame 1, there are several additional ORFs of at least 30 amino acids. Some of these, including one found in the 3'-noncoding region, lack appropriate translation start sites. Another long internal ORF is found from base 322 through 693. This contains an appropriate initiation sequence and encodes a hypothetical protein of 13,974 Da, which is rich in leucine residues (17%). The significance of this ORF remains to be defined.

Leader sequence of HCV-229E

The mRNAs of coronaviruses contain a stretch of leader sequence which is derived from the 5'-end of the viral genome and exhibits homology with the intergenic consensus sequence (Shieh et al., 1987; Budzilowicz et al., 1985). Since our cDNA clones did not appear to contain leader sequences, we used primer extension studies to determine the sequence of the HCV-229E leader RNA. A synthetic oligodeoxyribonucleotide which was complementary to an 18-mer sequence located near the 5'-end of the gene (Fig. 4) was end-labeled and used in a primer extension study with poly(A)-selected intracellular mRNA as a template. The reaction products, separated by agarose gel electrophoresis, revealed six bands (data not shown). Since these bands were most likely to represent the primer-extended products of the individual mRNA species, the smallest and most abundant band, corresponding to the primer-extended product of mRNA7, was eluted and sequenced by the chemical modification method (Maxam and Gilbert, 1977). The sequence of the 3'-end of the leader RNA was identical to the L8 sequence from nucleotides 129 to 171. At nucleotide 128, immediately 5' to the proposed leader mRNA fusion site, the sequence diverged from the L8 sequence and revealed a putative 60-base leader sequence which is shown in Fig. 6. The figure also shows a degree of homology with the leader sequence of IBV. Considerably less homology exists between the leader sequence of HCV-229E and those of HCV-OC43 and MHV-JHM (data not shown).

**DISCUSSION**

This report presented the primary sequence of the nucleocapsid gene and leader sequence of HCV-229E. When compared to the known sequences of other coronaviruses (Kapke and Brian, 1986; Skinner and Siddell, 1984; Armstrong et al., 1983; Lapps et al., 1987; Kamahora et al., 1988; Boursnell et al., 1985), common features of coronavirus nucleocapsid proteins emerged; namely, they are highly basic and have a high proportion of serine residues, which have been shown to be important for acquisition of the leader sequence. These features are also consistent with the observation that the nucleocapsid gene is highly conserved among coronaviruses.

**TABLE 1**

| Virus       | 3' conserved sequence |
|-------------|-----------------------|
| HCV-229E    | TGGAAGAGCCA (75)      |
| TGEV        | TGGAAGAGCTA (76)      |
| BCV         | GGGAAAGGC (79)        |
| HCV-OC43    | GGGAAAGGC (79)        |
| IBV         | GGGAAAGGCTA (81)      |
| MHV-JHM     | GGGAAAGGTC (82)       |
| MHV-A59     | GGGAAAGGTC (82)       |

Note. Number in parenthesis indicates distance, in nucleotides, of the conserved sequence from the poly(A) region.
to be sites of phosphorylation (Stohlman and Lai, 1979). The relationship between the nucleocapsid genes of HCV-229E and TGEV is particularly interesting since the viruses are antigenically related (MacNaughton, 1981). The predicted molecular weights of the N protein and the number of potential phosphorylation sites of both viruses are almost identical. Although these two viruses have little nucleotide sequence homology between their nucleocapsid genes, the amino acid sequences are homologous within a limited region. Amino acid sequence analysis revealed several structural features common to both viruses, which may have functional significance. For instance, there is a region of 46% homology within the amino-terminal one-third of the protein which extends from residues 29 to 134 in HCV-229E, and 41 to 146 in TGEV. Furthermore, approximately 10 amino acids downstream from the homologous region in both proteins lies an area which is abundant in serine residues, suggesting that this may be an important functional domain of the molecule. To further examine such functional homology between the two proteins, hydropathy profiles were constructed (Fig. 7). The contour of these plots suggests that a certain degree of functional homology exists within the first and last one-third of each molecule, with an additional region around position 200.

![Hydropathy profiles of coronavirus N proteins. Both the K-D (solid line) and GES (dashed line) curves are depicted with scales on the right and left, respectively.](image-url)
The peak around position 200 occurs just after the serine-rich region of the molecule. The relative conservation of these regions suggests a possible role in the interaction of the N protein with the viral genome. Similar structural features exist among the N proteins of HCV-229E, IBV, MHV, HCV-OC43, and BCV (Skinner and Siddell, 1984; Lapps et al., 1987; Kamahora et al., 1988; Bourssell et al., 1985). This is demonstrated by the hydropathy profiles of these proteins, which are also shown in Fig. 7. Further studies are required to reveal the functional significance of the conserved domains.

Another interesting finding is the open reading frame internal to the main coding region of the HCV-229E N gene. Thus far, two other coronaviruses, BCV and MHV-JHM, have been found to contain internal ORFs in gene 7 (Skinner and Siddell, 1984; Lapps et al., 1987) which are preceded by optimum translation initiation signals according to Kozak's consensus sequence (Kozak, 1983). The predicted amino acid sequences could encode hypothetical proteins of molecular weights 13,973; 14,842; and 23,057 for HCV-229E, MHV-JHM, and BCV, respectively. Interestingly, all three sequences are abundant in leucine residues (17 to 19%). HCV-OC43 also has two smaller internal ORFs encoding potential leucine-rich proteins of 8830 and 16,297 molecular weights (Kamahora et al., 1988). Further studies to determine whether this hypothetical protein can be detected in 229E-infected cells or by in vitro translation of a full-length cDNA clone (i.e., L8) are in progress.

Finally, the 3'-noncoding conserved sequence of gene 7 lends additional support to a common ancestry for coronaviruses, regardless of antigenic subgroup. This sequence has been proposed as a recognition site for the virus-encoded RNA-dependent RNA polymerase prior to negative-strand synthesis (Kapke and Brian, 1986). Certainly future studies must focus on examining the role of this conserved region in the viral replication cycle.

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