The gene encoding the nucleocapsid protein: sequence analysis in murine hepatitis virus type 3 and evolution in *Coronaviridae*

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Summary. The nucleoprotein-encoding gene (N) of murine hepatitis virus type 3 (MHV 3), from the Mill Hill strain, was cloned and sequenced. It was compared to gene N from other murine coronaviruses and was found to share more similarities with N sequences from MHV 1 and MHV JHM strains than with the published MHV 3 N sequence which is almost identical to MHV A59. We suggest that the evolution of some MHV N sequences resulted from a double recombination phenomenon between two ancestors. Furthermore, comparison of protein N from avian and mammalian coronaviruses leads to the hypothesis that horizontal transfer events of the virus from one host species to another have occurred.

Introduction

Murine hepatitis viruses (MHV) are enveloped RNA viruses belonging to the family *Coronaviridae*. Their genome is a polyadenylated, non-segmented, single stranded RNA [20] coding for three to four major structural proteins according to the antigenic group: the spike glycoprotein (S), the HE antigen, the membrane associated matrix glycoprotein (M), and the internal nucleocapsid protein (N), in addition to non-structural proteins such as an RNA-dependant RNA polymerase [18]. The order of the genes from 5' to 3' is: RNA polymerase, HE, S, M, N [9]. During replication, a full-length negative-stranded RNA and six polyadenylated messenger RNAs are synthesized in addition to the full length positive-sense RNA. The subgenomic RNAs form a nested set, all overlapping with the 3' end of the genomic RNA [10].

Among the MHV group, different strains induce various pathologies [27].
Intraperitoneal injection of MHV 3 from the Mill Hill strain (MHV3 MH) in adult mice results in fulminant hepatitis or in chronic infection of brain [23, 26]. In order to better characterize MHV3 MH and assess its genetic origin, we cloned and sequenced the 3' end of MHV3 MH RNA, i.e., the nucleoprotein-encoding gene. By comparing nucleotide sequences, we suggest that N genes from murine hepatitis viruses evolved by a double recombination phenomenon between two ancestors. We also report the phylogeny of protein N from avian and mammalian Coronaviridae.

Materials and methods

Viruses

MHV3 MH belongs to the family Coronaviridae, genus Coronavirus, murine hepatitis virus type 3 from the Mill Hill strain. The virus used throughout this work was derived from a single isolate of plaque purification [24] and was stored at −80°C.

Cloning of cDNA

Viral particles were purified [21] and the RNA extracted [10]. Viral RNA was used for cDNA synthesis and the cDNA fragments were cloned into the Eco RI site of λ gt 11 phage [28]. MHV 3 recombinant clones were detected by immunodetection of the fusion protein using polyclonal antibodies against MHV3 MH which were prepared by immunizing a rabbit with purified virus [21] and were revealed by a rabbit-PAP system (DAKO) with 3-amino-9-ethylcarbazole as substrate.

DNA and RNA sequencing

Restriction fragments of MHV 3 inserts were cloned in the multisite of M 13 mp 18 and 19 replicative forms. DNA sequencing was performed by the dideoxy chain termination method [15]. RNA sequencing was done [4] on purified virion RNA with a synthetic oligonucleotide (GGCTGATTCCCTCCTGCTC) complementary to positions 146 to 128 (Fig. 1) as primer.

Sequence analyses

Computer analyses were performed using the Bisance facilities [3] on the VAX of CITI2 in Paris and on PC microcomputers using the MUST package [14]. Amino acid sequences were first aligned using the Clustal program [6] and corrected by hand. For phylogenetic analyses, PAUP version 2.4.1. [22] was used which deduces phylogenetic relations on the basis of maximum parsimony. The following N protein sequences from the Swiss-Prot Database Rel. 23 were used: avian infectious bronchitis virus, IBV (M 21515); turkey enteric coronavirus, TCV (P26020); murine hepatitis viruses: MHV1 (P18446); MHV3 PM (P18447), MHV A59 (P18448), MHV JHM (P03417); porcine respiratory coronavirus: PRC (P24411), porcine transmissible gastroenteritis viruses: TGEV fs (P05991), TGEV purdue (P04134); bovine coronaviruses: BCV f15 (P19902), BCV mebus (P10527); feline infectious peritonitis virus, FIP (P25909); human coronaviruses: HCV 229E (P15130), HCV OC43 [7].
Sequence and evolution of the gene N from MHV 3

Results

Sequence of MHV 3 MH gene N

The cDNA library constructed from murine hepatitis virus RNA of the Mill Hill strain was screened by immunodetection (data not shown). A positive cDNA was subcloned in M13 vectors and sequenced. The sequence corresponded to the nucleoprotein-encoding gene but lacked the 5' end. Viral RNA was then sequenced directly using a synthetic oligonucleotide designed from the cDNA sequence just determined. The resulting composite sequence is shown in Fig. 1 beginning at the initiation codon (accession number X63538 in the EMBL Database).

The sequence of Fig. 1 was translated in the 3 possible reading frames and 2 long open reading frames (ORF) were found. The smaller one (positions 26 to 685) encoded a hypothetical protein of 220 amino acids, 24,009 Da in molecular mass, rather basic (11.4 and 8.2% of basic and acid amino acids, respectively) and serine-rich and leucine-rich (8.6 and 17% of total amino acids, respectively).

The larger ORF (Fig. 1) was 1371 bases long, the 3' untranslated region contained 295 bases including the first A of the polyA tail, with no canonical polyadenylation signal but containing a 10 base motif (position 1593 to 1602) which is relatively conserved among coronaviruses [16]. The predicted 457 amino acid protein of 50,065 Da is similar to other murine hepatitis nucleocapsid proteins whereas it was 55 kDa in molecular mass when estimated by "Western" blotting (data not shown). The nucleocapsid protein is composed by 14.7% basic amino acids (33 lysine, 30 arginine, and 4 histidine) and 10.3% acidic residues (25 aspartic acid, 22 glutamic acid), giving a basic protein, a property expected for a nucleic acid binding protein. Basic amino acid residues are clustered in the central part of the protein and acidic ones in the carboxy terminus.

Evolution of the nucleoprotein-encoding gene in murine hepatitis viruses

In order to analyze the variability of the nucleoprotein-encoding gene of MHV, all available sequences were aligned (Fig. 1) and compared pairwise (Table 1). The following sequences were used: MHV 1 [13], MHV JHM [17], MHV A59 [1, 13], MHV 3 PM [13], and MHV 3 MH (this work). The sequence of gene N from MHV S, which was shown to be a recombinant one [13], was not considered. The N sequences from the two MHV 3 strains appeared highly divergent: the gene from MHV 3 MH (position 1 to 1371) was longer by 9 bases than the MHV 3 PM sequence and was different at 90 positions whereas MHV 3 PM and MHV A59 sequences were almost identical (2 bases were different in the coding sequence leading to one different amino acid) as already stated [13]. Such a low level of sequence variation between 2 different strains
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### Table 1. Nucleoprotein-encoding gene sequence similarity among the different MHV strains (%)

|        | 3 MH | 1   | JHM | A 59 | 3 PM |
|--------|------|-----|-----|------|------|
| 3 MH   |      |     |     |      |      |
| 1      | 96.3 |     |     |      |      |
| JHM    | 94.5 |     |     |      |      |
| A 59   | 94.3 |     |     |      |      |
| 3 PM   | 94.1 |     |     |      |      |

Numbers above the diagonal represent the per cent similarities among nucleotides (from the initiation codon to the first A of the polyA tail; in parentheses: coding sequence). Numbers below the diagonal are per cent similarities at the amino acid level.

is within experimental error so that MHV3 PM and MHV A59 nucleocapsid sequences were considered to be the same one.

We then turned to an analysis of relationships between strains by identifying informative positions. Informative positions for the parsimony analysis are defined as those at which 2 different nucleotides are present at least twice each. With 4 sequences (without MHV A59), there are 23 informative positions, spread over the whole sequence (Fig. 2 A). A graphical representation of the informative positions and of their character state is provided in Fig. 2 B. This juxtaposition of the nucleotides in the informative positions can be interpreted in terms of associations. The informative positions at the two ends of the sequence group MHV 1 with MHV JHM while those in the central region group MHV 1 with MHV3 MH.
Fig. 2. Informative positions of four MHV N sequences. A Informative positions (two different bases at a given position, present at least twice each) are listed in the order (upper line) by which they appear in the sequences of Fig. 1. Gaps are indicated by a dash. Vertical lines indicate the limits of the 3 domains. B Schematic representation. Blank and shaded boxes indicate identity of nucleotides, respectively, when read vertically. By convention, the nucleotides of the first sequence were all shaded without any implication as to whether the nucleotide association is parental within this sequence.

Evolution of coronavirus protein N in birds and mammals

We compared all N protein sequences of avian and mammalian coronaviruses available in the Swiss-Prot Database Rel. 23 plus HCV OC43 [7]. Amino acid sequences were first aligned using the Clustal program [6]; 222 positions can be aligned unambiguously. In a first step, we verified by PAUP [22] that the N sequences of viruses infecting the same species (mouse, pig, and cattle) were clustered together on the tree. We chose one sequence for pig and cattle viruses and two for MHV (MHV 1 and 3), the most distant ones. For avian and turkey viruses and among human viruses which are not closely related on the tree, individual sequences were used. The phylogenetic tree of the various N proteins constructed by the parsimony method is shown in Fig. 3 A, while the cladogram of birds and mammals based on morphological characters [12] is provided in Fig. 3 B. First, we found that human OC43, bovine, and turkey enteric coronaviruses give a monophyletic group. The distances between these 3 viruses were smaller than between MHVs. Second, porcine and feline viruses were more closely related than their respective hosts.

The same analysis was performed on protein M sequences of the same coronaviruses (when available) and an identical most parsimonious tree was obtained (data not shown).

Discussion

We sequenced the 3' part of the MHV3 MH genome and showed that it contained 2 open reading frames. The smallest one was also detected in other MHV N sequences [13], although these ORF are shorter (207 aa). The only exception is for the MHV JHM sequence in which a premature stop codon is found. This
hypothetical protein has not yet been detected. The longer ORF which codes for the nucleocapsid protein presents a discrepancy between the calculated and estimated molecular mass which could be explained by post-translational modifications such as phosphorylation of serine residues, which account for 8.5% of total amino acids. Phosphorylation of MHV nucleoproteins has been described [19].

We showed that the nucleotide sequences of gene N of 2 viruses which were classified under the name MHV 3 are different. The N gene of the virus used in our laboratory (Mill Hill strain) seemed original by comparison with the MHV3 PM [13] which is almost identical to MHV A59. The identity of these 2 sequences could be explained by genetic recombination between the 3' termini of MHV 3 and A59 genomes leading to a common N gene. Such recombination processes have been observed among laboratory strains of murine coronaviruses, under special laboratory conditions [5, 8, 11]. Alternative hypotheses can also be considered such as a mixture of viruses in the infected animal from which MHV 3 was first isolated or a mixture of the two strains in the cultured cells.

A double recombination event between two ancestors is suggested by the distribution of the informative positions in three regions, two of which are of the same kind (MHV 1 is the same as MHV JHM at the ends and MHV 1 is the same as MHV3 MH in the central part). Only 2 positions (108 and 895) support grouping MHV 1 with MHV3 PM in the left and central portion of the sequence, respectively, but this can be interpreted as convergence in 2 out of 23 positions which does not blur the phylogenetic information. It is not
possible to determine the polarity of recombination or the nature of the ancestors.

The junctions of the putative recombination events would be between positions 254 and 260 and between positions 1149–1194, as indicated by the dashed lines in Fig. 2. It is interesting to note that a three domain structure for the nucleoprotein has been suggested, with spacers A (positions 420 to 486) and B (positions 1143 to 1215) defined as hot spots of variability between domains [13]. Spacer B corresponds to the second putative recombination region that we have identified, but spacer A is not located within the first recombination region. If instead of comparing nucleotide sequences, we converted the sequence into amino acids, very little information appears since most nucleotide informative positions were silent at the amino acid level (only 6 informative positions remain at the amino acid level).

As coronaviruses were shown to have a marked host specificity [9, 20], it is generally assumed that the virus and its host evolved in parallel, the position of the former on the phylogenetic tree mimicking the position of the latter. We checked this hypothetical co-evolution between virus and host by constructing the phylogenetic tree of M and N protein sequences. The M and N genes are located next to each other and it would have been better to use 2 distant genes, one at the beginning of the genomic RNA, the N gene being located at the 3’ end, but the sequences are not all available. Nevertheless, the fact that we obtain the same tree with the sequences of 2 different viral proteins is a strong argument in favor of the generality of the tree.

The N sequences of turkey, bovine, and human OC43 coronaviruses are so closely related [25] that they are less polymorphic than the MHV N sequences, leading to the conclusion that the interspecific diversity is smaller than the intraspecific polymorphism. The HE proteins of BCV and HCV OC43 were also shown to be very similar [29]. It is possible that very recent host changes have occurred as a result either of domestication or of some laboratory forced infections.

The phylogenetic tree cannot be explained without one horizontal transfer of viruses between feline and porcine hosts on a large evolutionary scale (over million years) because it is well known that Artiodactyla (pig and cattle) are not closely related to rodents or carnivores [2, 12] (see Fig. 3 B). Thus, we could infer from the phylogenetic tree of N proteins that very recent transfer phenomena have occurred in parallel with horizontal transfer and co-evolution.

In conclusion, our analyses suggest that co-evolution between the virus and its host and horizontal transfer both participate in the evolution of *Coronaviridae*. The relative importance of the 2 phenomena could be studied if more “wild” viral strains were available from “wild” hosts.

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