VIROLOGY 209, 337-346 (1995)

Evolution of Mouse Hepatitis Virus (MHV) during Chronic Infection: Quasispecies Nature of the Persisting MHV RNA

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Received December 24, 1994; accepted March 14, 1995

Coronavirus infection of mice has been used extensively as a model for the study of acute encephalitis and chronic demyelination. To examine the evolution of coronavirus RNA during chronic demyelinating infection, we isolated RNA from intracerebrally inoculated mice at 4, 6, 8, 13, 20, and 42 days postinfection and used reverse transcription-polymerase chain reaction amplification methods (RT-PCR) to detect viral sequences. RNA sequences from two viral structural genes, the spike gene and the nucleocapsid gene, were detected throughout the chronic infection. In contrast, infectious virus was not detectable from brain homongenates beyond 13 days postinfection. These results indicate that coronavirus RNA persists in the brain at times when infectious virus is not detected. To determine if genetic changes were occurring during viral replication in the host, we cloned and sequenced the RT-PCR products from the spike and nucleocapsid regions and analyzed the sequences for mutations. Sequencing of the cloned products revealed that a variety of mutant forms of viral RNA persisted in the CNS, including point mutants, deletion mutants, and termination mutants. The mutations accumulated during persistent infection in both the spike and the nucleocapsid sequences, with greater than 65% of the mutations encoding amino acid changes. These results show that a diverse population or quasispecies consisting of mutant and deletion variant viral RNAs (which may not be capable of producing infectious virus particles) persists in the central nervous system of mice during chronic demyelinating infection. The implications of these results for the role of persistent viral genetic information in the pathogenesis of chronic demyelination are discussed. © 1995 Academic Press, Inc.

INTRODUCTION

Mouse hepatitis virus (MHV) is an enveloped virus with a 31-kb single-stranded RNA genome (Lai, 1990; Spaan et al., 1990). The virion particle is composed of three glycoproteins (S, spike; M, matrix; and HE, hemagglutinin-esterase) and the nucleocapsid protein (N), which is associated with the MHV RNA. MHV replicates in the cytoplasm of infected cells using a viral RNA-dependent RNA polymerase that is translated from the genomic RNA. In infected rodents, MHV is able to replicate in the central nervous system (CNS), resulting in acute encephalitis and chronic demyelination. Infection of rodents with strains of MHV which induce predominantly chronic demyelination has been used as a model system for studying human demyelinating diseases such as multiple sclerosis.

Many laboratories have used the MHV model system to investigate the role of the virus and the immune system in chronic demyelination. These studies have shown that viral structural genes such as the spike and hemagglutinin-esterase are important in determining the neuropathogenicity (encephalitis versus chronic demyelination) of the virus and in the ability of the virus to spread through the CNS (reviewed by Kyuwa and Stohlman, 1990). However, it has been difficult to assess the role of the virus in chronic infection because infectious virus is rarely isolated after 20 days postinfection (Knobler et al., 1982; Dalziel et al., 1986; Yokomori et al., 1993) and may not be representative of the persistent viral RNA population. Therefore, we elected to analyze the evolution of MHV RNA by examining specific regions of the RNA throughout the chronic infection. The specific regions examined were the S1 portion of the spike glycoprotein gene and a portion of the nucleocapsid gene. These regions were selected because they encode viral structural proteins important in the pathogenesis of the disease. The S1 region of the spike glycoprotein has previously been characterized as "hypervariable" (Banner et al., 1990), whereas the nucleocapsid region is more conserved among MHV strains.

In this study, we report the evolution of coronavirus RNA during persistent infection. We infected mice with a strain of MHV which consistently caused demyelination (Fleming et al., 1986) and analyzed the viral RNA isolated from the brain at various time points after infection. The viral RNA was detected by reverse transcription-polymerase chain reaction amplification methodology. By se-

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quence analysis of the amplified products, we found that MHV RNA persists in the brains of infected animals in multiple mutant/defective forms. The majority of the mutations we detected were biased transitions (U to C and A to G) and resulted in amino acid changes. These results demonstrate the rapid evolution of coronavirus RNA during persistent infection and suggest that a diverse population of coronavirus RNA persists in the central nervous system where it may contribute to the pathogenesis of chronic demyelination.

MATERIALS AND METHODS

Virus

MHV-JHM antigenic variant 2.2-V-1 was propagated in DBT cells and quantitated by plaque assay as previously described (Fleming *et al.*, 1986). This low virulence variant was selected by resistance to neutralization by an antibody directed against the spike glycoprotein (Fleming *et al.*, 1986) and causes predominantly chronic demyelination in mice (Fleming *et al.*, 1986). Relative to the parental virus, JHM-DL, from which it was selected, MHV-JHM 2.2-V-1 encodes a single nucleotide change in the S2 region of the spike gene (nucleotide 3340) which results in an amino acid change from leucine to phenylalanine (Wang *et al.*, 1992).

Animal experiments

Six-week-old male C57BL/6J mice (Jackson Laboratories, Bar Harbor, ME) were used in all experiments. Upon arrival, selected mice were screened by enzymelinked immunosorbent assay (Fleming and Pen, 1988) and found to be seronegative for murine coronaviruses. Mice were inoculated intracerebrally with 10^3 plaque forming units (PFU) of MHV-JHM 2.2-V-1 diluted in 30 μ l Dulbecco's modified minimal essential medium. At the indicated times, mice were sacrificed by CO_2 inhalation, and brains were removed aseptically, immediately frozen on dry ice, and stored at -70° . In representative animals, viral titers were determined on one-half of the fresh, unfrozen brain by plaque assay as previously described (Fleming *et al.*, 1986).

RNA extraction

Brains were thawed, and each half brain was manually disrupted in a Tenbrock homogenizer containing 2 ml phenol/chloroform (1/1) and 2 ml of lysis buffer (0.5% SDS, 0.025 M EDTA, 0.074 M NaCl). RNA was subsequently extracted once with lysis buffer, and proteins removed by extraction with phenol/chloroform (1/1) and with chloroform. RNA was precipitated first in the presence of 3.0 M ammonium acetate (final concentration) and incubated overnight at -20° . The RNA was pelleted by centrifugation for 15 min at 10,000 g and the pellet was resuspended in DEPC-treated water. The RNA was

reprecipitated in the presence of 0.3 *M* sodium acetate and 2.5 volumes of ethanol. RNA from four to six brains were combined in order to prepare representative pools from each time point. RNA quantity was determined by optical density measurements at 260 nm.

Input virus RNA was isolated by mixing 5×10^6 PFU of MHV-JHM 2.2-V-1 with one-half of an uninfected mouse brain and immediately subjecting the mixture to RNA extraction as described above.

Reverse transcription-polymerase chain reaction (RT-PCR) of brain RNA

Two micrograms of total brain RNA was denatured with 20 mM methylmercury hydroxide for 5 min at room temperature, after which β -mercaptoethanol was added at a final concentration of 0.14 M. For cDNA synthesis, the RNA was incubated at 42° for 60 min in a reaction containing 50 mM Tris-HCl (pH 8.3), 50 mM KCl, 10 mM MgCl₂, 10 mM DTT, 0.5 mM spermidine, 1.25 mM of each dNTP, 40 units RNasin, 1 μ g random hexamer oligonucleotide primers, and 4 units avian myeloblastosis virus reverse transcriptase (Promega, Madison, WI). The mixture was incubated at 95° for 3 min, cooled on ice for 1 min, and incubated at 42° for 60 min after the addition of another 4 units of reverse transcriptase.

The synthesized cDNA was amplified by PCR using a method adapted from Saiki et al. (1988) and Chang et al. (1993) to minimize Tag errors. Briefly, 20 μ l of reverse transcription mixture was mixed with 80 μ l of a master mix buffer containing 7.5 mM Tris-CI (pH 9.0), 37.5 mM KCl, 0.075% Triton X-100, 0.84 mM MgCl₂, 0.25 μ M synthetic oligonucleotide primers, and 2.5 units of Tag polymerase (Promega). The reaction mixture was overlaid with 50 μ l mineral oil and subjected to 30 (Fig. 2B) or 35 (Fig. 2A) cycles of amplification with each cycle consisting of 94° for 1 min, 60° for 1 min, and 72° for 1 min. At the end of this period, an additional extension step at 72° for 7 min was performed. MHV-JHM nucleotide sequences for oligonucleotide PCR primers are indicated in Table 1. RT-PCR products were resolved by electrophoresis on a 1.2% agarose gel in 1× TAE buffer (0.04 M Tris-acetate, pH 8.5, 0.002 M EDTA) and stained with ethidium bromide.

cDNA cloning and sequencing

PCR-amplified products generated by 35 cycles of amplification of the S1 and N genes were purified using the Magic PCR Preps DNA purification system (Promega) and ligated into the pGEM-T vector (Promega). The ligation buffer contained 30 mM Tris-Cl, pH 7.8, 10 mM MgCl₂, 10 mM DTT, and 0.5 mM ATP and 3 units of T4 DNA ligase. The ligation product was used to transform competent *Escherichia coli* strain DH5 α (BRL). Plasmid DNA was isolated from the bacteria by Magic Miniprep purification resin (Promega) and sequenced by the

TABLE 1
Oligonucleotide Primers Used to Amplify MHV Sequences

Primer	Sequence (5' to 3') ^a	Position ⁶
SI-1	TATGAATTCTACGTTATGTCCAGGCTGAGTC	1084-1106
SI-2	TATGGATCCATAGAGGTCATATCTGACGC	1895-1916
\$1-3	ATGTTGCCTACGCCCAGC	1391-1408
SI-4	CACTCACGATGAATTGTGCC	1510-1529
AM8-1	CCAAGCTTCTGCACCTGCTAGTCGATCTG	672-693
AM8-2	CCGGTACCACCATCTTGATTCTGGTAGGC	1215-1237
AM8-3	ATGAATTCAGCGCAAGCCTGCCTCTACTG	747-768
AM8-4	<u>ATGGATCCT</u> GAATATTGCAGCTCATACAC	1122-1144

^e Additional nucleotides (underlined) provide restriction endonuclease recognition sites.

Sanger dideoxy chain termination method (Sanger et al., 1977) using Sequenase Enzyme 2.0 (Sequenase DNA sequencing kit, version 2.0, U.S. Biochemicals) with reverse and —40 primers. Internal primers for the spike 1 region and nucleocapsid region (SI-3, SI-4, AM8-3, and AM8-4, see Table 1) were used in some sequencing reactions. Reaction products were analyzed on 5% Hydrolink (J. T. Baker) gels containing 7 M urea in TBE.

RESULTS

Persistence of MHV-JHM RNA

To examine the evolution of coronavirus RNA during chronic demyelinating infection, we extracted brains from mice at 4, 6, 8, 13, 20, and 42 days postinfection and analyzed them for infectious virus and viral RNA. We were able to detect production of infectious virus until 13 days post infection (data not shown) by plaque assay of brain homogenates. To determine whether coronavirus RNA persists in mouse brains beyond times when infectious virus is detected, we performed RT-PCR using primers specific to the S1 region of the spike glycoprotein and to the nucleocapsid gene (Fig. 1). We were able to amplify both the spike (Fig. 2A) and the nucleocapsid (Fig. 2B) sequences from total RNA isolated from MHVinfected mouse brain at 4, 6, 8, 13, 20, and 42 days. PCR products of the expected 832-bp size for the S1 region and of the 565-bp size for the nucleocapsid region were detected. No PCR product was detected in the reagents alone control (Figs. 2A and 2B, lane C) or uninfected brain RNA (Figs. 2A and B, Iane U). As shown in Figs. 2A and 2B, the RT-PCR products of the S1 and the nucleocapsid genes were detected at 42 days postinfection without resorting to additional cycles of amplification or the use of nested primers. Clearly the amount of viral RNA decreases significantly after 13 days postinfection; however, these results demonstrate that coronavirus RNA persists in the brains of infected mice as long as 42 days postinfection. Indeed, preliminary results suggest that coronavirus RNA may persist in the brain of infected mice for the life of the animal (Pooley and Fleming, unpublished).

Sequencing of viral cDNA PCR products

To confirm the identity of the PCR products and to examine the evolution of viral RNAs during chronic infection, we cloned and sequenced the S1 and nucleocapsid PCR products from brain samples taken at 4, 8, 13, 20, and 42 days p.i. To control for mutations generated during the RNA isolation and RT-PCR amplification process, we mixed input virus particles (5 × 10⁶ PFU) with uninfected brain and immediately subjected the mixture to RNA extraction and RT-PCR amplification (see Materials and Methods). The PCR products were ligated into the plasmid pGEM-T as described under Materials and Methods and individual clones were sequenced by the Sanger dideoxy chain termination method (Sanger et al., 1977). The mutations detected in each clone are listed in Table 2 and summarized in Table 3. Sequencing of eight clones representing input virus showed that seven of the eight clones encoded the wild-type sequence. A single mutation (A to G) was detected in one clone (S0-8). These results reflect the "master" sequence (wild type) typically found in the virus population derived from tissue culture cells (Steinhauer and Holland, 1987). In contrast, sequencing of individual clones at 4, 8, 13, 20, and 42 days postinfection showed that multiple point mutants and deletion mutants are detectable in the persistent RNA population (Table 3). These results are consistent with an increase in the diversity of the quasispecies in the persistent MHV RNA population.

A total of 20 individual clones were sequenced from PCR products generated at Day 4 postinfection. There was an average of 1 mutation per clone, with 5 of the 20 clones containing wild-type sequences. In the samples generated from RNA isolated at 42 days postinfection, we detected an increase in the number of mutations (55 mutations in 20 clones), with only 1 of 20 clones encoding wild-type sequence. The mutations were spread throughout the S1 region (Fig. 3). There was an average of 2 to 3 mutations per clone (range 0 to 6) and 1 clone contained a deletion from nucleotides 1529 to 1618. The same deletion was detected in a Day 8 clone, suggesting that this deletion variant may be generated early during the course of infection. It is striking that this particular deletion mutant is present in independently isolated brain RNA samples (Day 8 and Day 42). Interestingly, several

⁶ Nucleotide position of SI according to Parker et al. (1989). Nucleotide position of N according to Skinner et al. (1983).

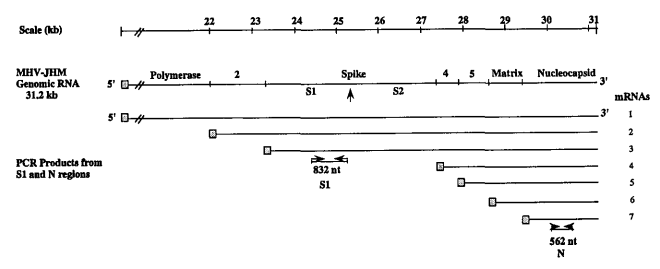


FIG. 1. Schematic representation of the 31-kb MHV-JHM genome and seven mRNAs. The shaded boxes represent the viral leader sequences that are at the 5'-end of each of the mRNAs. The 22-kb polymerase gene is not drawn to scale. Arrows exemplify the primers used to amplify specific regions of spike and nucleocapsid sequences.

other attenuated variants have been described in which this region of the S1 has been deleted (Dalziel et al., 1986; Fleming et al., 1986; Morris et al., 1989; Parker et al., 1989; La Monica et al., 1991; Wang et al., 1992). This region (from nt 1500 to nt 1959) has in fact been termed "hypervariable" (Banner et al., 1990; Wang et al., 1992). However, the mutations we detected were not limited to the hypervariable region, but were distributed throughout the S1 region (Fig. 3).

To further investigate the evolution of the MHV RNA, we examined a second region of the virus, the nucleocapsid region. The sequence analysis of the nucleocapsid PCR products revealed a pattern similar to that seen in the S1 gene, that is, an increased number of mutations in the persistent RNA (Tables 4 and 5). At Day 4 post infection, 11 of 20 clones maintained the wild-type sequence and the remaining clones encoded one or two mutations per clone. In contrast, only 8 of the 20 clones randomly isolated at 42 days

post infection maintained the wild-type nucleic acid sequence. These results indicate that mutations may potentially occur throughout the RNA genome.

To determine whether there was a bias for any type of mutations during persistent infection, we categorized the mutations as transitions or transversions (Table 6). Analysis of the type of the nucleotide changes detected during coronavirus persistent infection revealed that the mutations consisted predominantly of U to C and A to G transitions (Table 6, 85 U to C and A to G mutations/135 total mutations; 63%). Biased mutations, most dramatically seen in persistent measles infection (Cattaneo et al., 1988), have been proposed to occur due to modification of double stranded RNA intermediates that form during viral RNA transcription and replication by the cellular helicase activity (Bass et al., 1989, see Discussion). Overall, the persisting MHV RNA contains a variety of mutations, the majority (65%) of which encode amino acid

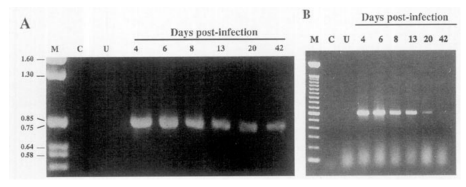


FIG. 2. Long-term persistence of MHV-JHM 2.2-V-1 RNA in brains of C57/B16 mice. Brain RNA was isolated at 4, 6, 8, 13, 20, and 42 days postinoculation and subjected to RT-PCR amplification of 35 cycles using spike-specific primers (A) and 30 cycles using nucleocapsid-specific primers (B). Lane C shows no products were amplified from reagents alone; Iane U shows no products were amplified from uninfected brain RNA. Lane M indicates DNA standards (A) and 100 bp marker DNA (B). Products were analyzed by electrophoresis on a 1.2% agarose gel and visualized by staining with ethicium bromide.

TABLE 2

Mutations Detected in the Spike SI Region during Persistent MHV-JHM Infection

Clone	Nucleotide	Codon change	Amino acid	Clone	Nucleotide	Codon change	Amino acid
	clones sequenced, 7	clones encoded wild	type sequence	Day 42: 20) clones sequenced,	, 1 clone encoded wil	d-type sequence
8-03	1127 A → G	GAU → GGU	Asp → Gly	S42-2	1416 U → C	UUU → UUC	No change
D 4 00		- 1	1 -		1472 C → U	ACU → AUU	Thr → Ile
Day 4: 20	ciones sequencea,	5 clones encoded will	a-type sequence		1546 C → U	CCC → UCC	Pro → Ser
84-1	1335 U → C	ncn → ncc	No change	S42-3	1271 C → U	UCA → UUA	Ser → Leu
	1878 A → G	GAA → GAG	No change		1480 A → G	ACC → GCC	Thr → Ala
34-4	1783 G → C	GUA → CUA	Val → Leu	S42-1	1688 G → A	GGU → GAU	Gly → Asp
, ,	1734 U → C	DGU → UGC	No change		1764 G → A	UGG → UGA	Trp → TERM
34-5	1118 A → G	AAU → AGU	Asn → Ser	S42-7	1266 C → U	GCC → GCU	No change
34-7	1741 G → A	GCC → ACC	Ala → Thr		1280 U → C	CUG → CCG	Leu → Pro
34-8	1767 A → G	UCA → UCG	No change		1313 A → G	AUA → GUA	lle → Asp
54-0 54-9	1667 G → A	GGC → GAC	Gly → Asp		1518 U → C	AUU → AUC	No change
			•		1800 A → U	CAA → CAU	Gln → His
64-10	1205 A → G	GAU → GGC	Asp → Gly	S42-13	1215 U → A	AUU → AUA	No change
	1675 U → C	UGU → CGU	Cys → Arg	012 10	1821 A → G	UUA → UUG	No change
34-11	1167 A → G	UCA → UCG	No change	S42-9	1119 U → C	AAU → AAC	No change
	1841 C → U	ACU → AUU	Thr → Ile	04E-0	1199 A → G	CAA → CGA	Gin → Arg
34-12	1299 G → A	AAG → AAA	No change	040.10		GAU → GAA	Asp → Glu
	1509 A → G	ACA → ACG	No change	S42-12	1173 U → A		Ash → Giu
34-14	1230 U → C	NOO → OOC	No change		1529-1618	Deletion	1
84-16	1560 G → A	UCG → UCA	No change		1649 A → G	AAA → AGA	Lys → Arg
	1797 C → U	UGC → UGU	No change		1737 U → C	UCU → UCC	No change
34-17	1336 U → C	UGG → CGG	Trp → Arg		1872 U → C	AAU → AAC	No change
	1869 U → C	CCN → CCC	No change	S42-15	1446 A → G	GCA → GCG	No change
34-18	1261 G → C	CCn → CCn	Ala → Pro		1592 A → G	GAG → GGG	No change
34-19	1232 U → C	UUG → UCG	Leu → Ser	S42-16	1197 U → C	CGU → CGC	No change
34-20	1869 U → C	CCN → CCC	No change		1277 A → G	CAG → CGG	GIn → Arg
			· ·		1472 C → U	ACU → AUU	Thr → Ile
	Day 8: 4 o	clones sequenced			1592 A → G	gag → ggg	Glu → Gly
S 8- 2	1432 U → C	UAU → CAU	Tyr → His		1781 U → C	UUA → UCA	Leu → Ser
3 <i>6-2</i> 38-3	1697 A → G	GAA → GGA	Glu → Gly	S42-17	1090 U → C	UAU → CAU	Tyr → His
		DELETION	Giu - Giy		1148 U → A	AUG → AAG	Met → Lys
S8-4	1529-1618		No obooga		1167 A → G	UCA → UCG	No change
S8-6	1233 G → A	UUG → UUA	No change	S42-18	1212 A → G	CAA → CAG	No change
	1332 G → A	UCG → UCA	No change		1339 A → U	AAU → UAU	Asn → Tyr
	Day 13: 4	clones sequenced			1890 C → G	GGC → GGG	No change
	,	********		S42-19	1159 A → U	AGU → UGU	Ser → Cys
S13-1	1349 U → C	UAU → CAU	Try \rightarrow His		1263 U → C	GUU → GCU	Val → Ala
	1468 U → C	UGC → CGC	Cys → Arg		1764 G → A	UGG → UGA	Trp → TERN
S13-4	1288 U → C	CUU → CUC	No change	S42-20	1167 A → G	UCA → UCG	No change
	1592 A → G	GAG → GGG	Glu → Gly	0 12 20	1169 U → C	GUU → GCU	Val → Ala
	1760 G → A	GGA → GAA	Gly → Glu		1172 A → G	GAU → GGU	Asp → Glv
S13-5	1285 U → C	AGU → AGC	No change		1177 U → C	UUU → CUU	Phe → Leu
	1595 G → A	UGC → UAC	Cys → Try		1616 U → C	UUU → UCU	Phe → Ser
	1763 G → A	UGG → UAG	Trp → TERM		1875 U → C	ACU → ACC	No change
S13-6	1543 G → U	GGG → UGG	Gly → Trp	042.21	1870 A → G	AAU → GAU	_
	1726 A → U	AAG → UAG	Lys → TERM	S42-21			Asn → Asp
		7 - 12		S42-22	1109 U → C	UUG → UCG	Leu → Ser
	Day 20: 4	clones sequenced			1127 A → G	GAU → GGU	Asp → Gly
220 E	1204 ∐ → △	GUU → GCU	Val → Clu	0.40.00	1841 C → A	ACU → AAU	Thr → Asn
S20-5	1394 U → C		Val → Gly	S42-23	1755 U → C	UCU → CCU	Ser → Pro
000.0	1367 G → A	GGU → GAU	Gly → Asp	S42-24	1156 G → A	GGU → AGU	Gly → Ser
S20-6	1192 A → G	AGC → GGC	Ser → Gly		1311 C → U	ACC → AUC	Thr → Ile
	1413 U → C	UGU → UGC	No change		1631 U → G	CUC → CGC	Leu → Arg
S20-7	1289 U → C	AGU → AGC	No change	S42-24	1682 G → A	GGU → GAU	Gly → Asp
	1628 A → G	GAU → GGU	Asp → Gly	S42-25	1307 U → C	GUU → GCU	Val → Ala
S20-8	1109 U → C	AAA → AAC	No change		1640 G → A	GGG → GAG	Gly → Glu
					1127 A → G	GAU → GGU	Asp → Gly

TABLE 3
Summary of Mutations Detected in the Spike Region during Persistent MHV-JHM Infection

Days postinfection		Total mutations detected		Туре о			
	Clones sequenced		S	С	Del	Term	Mutations per thousand nucleotides sequenced
0 (input)	8	1	0	1	0	0	0.2
4	20	22	12	10	0	0	1.4
8	4	5	2	2	1	0	1.7
13	4	10	2	6	0	2	3.4
20	4	7	3	4	0	0	2.4
42	20	55	15	37	1	2	3.7

^a S, silent; C, change of amino acid; Del, deletion; Term, termination codon.

changes. The diversity of the quasispecies in coronavirus persistent infection may have important implications for chronic pathogenicity.

DISCUSSION

In this study, we examined the evolution of coronavirus RNA during persistent infection. We found that a variety of mutant/defective viral RNAs emerge throughout chronic viral infection in mouse brain and that mutant viral RNAs persist in the brain at times when infectious virus is not detected (after 13 days postinfection). These findings emphasize the diversity and rapid evolution of coronavirus RNA. As early as 8 days postinfection, multiple mutants and a deletion variant were detected. The mutations in the viral RNA were detected in two regions that were examined, the spike S1 and the nucleocapsid region. By 42 days postinfection, the majority of the clones we sequenced contained multiple mutations and/ or deletions. Previous studies of chronic MHV infection have shown that infectious viruses with deletions or mutations in the spike gene (Dalziel et al., 1986; Fleming et al., 1986; Morris et al., 1989; Parker et al., 1989; La Monica et al., 1991; Wang et al., 1992) and/or hemagglutininesterase gene (Yokomori et al., 1993) can be isolated. In this study, we demonstrate that a mutant or defective viral RNA population, which may not be capable of producing infectious virus, persists in the brain. This defective viral RNA population could potentially contribute to the pathogenesis of chronic demyelination by perturbing normal cell functions and/or by chronically stimulating the immune system (Oldstone, 1989, 1991).

An increase in the heterogeneity in the MHV sequences isolated during the chronic infection is clear for both the spike S1 and the nucleocapsid regions of MHV RNA (Tables 3 and 5). The frequency of mutations we detected in the spike region rose from 1.4 mutations per thousand nucleotides sequenced at Day 4 to 3.7 mutations per thousand nucleotides sequenced at Day

42 (Table 3). The sequencing of individual cDNA clones representing persistent viral RNAs was essential for this study. As shown in Fig. 3, an average of 9 of 10 clones contained the wild-type sequences at any given position. Therefore, sequencing of uncloned PCR products would have revealed an overall wild-type or consensus sequence. By sequencing cloned PCR products, we unveiled a previously undetected diversity in the persistent viral RNA population. This diversity or heterogeneity in the persistent MHV infection is consistent with what has been seen in other persisting RNA viral infections such as measles virus (Baczko, et al., 1986; Cattaneo, et al., 1988), hepatitis delta virus (HDV) (Chao et al., 1990; Lee et al., 1992), and hepatitis C virus (HCV) (Martell et al., 1992; Higashi et al., 1993). In the case of HDV, rapid evolution and microheterogeneity of the viral RNA were demonstrated by isolating, cloning, and sequencing HDV RNA from the serum of a single patient over a 2-year period of chronic infection (Lee et al., 1992). The heterogeneity of HCV has been well documented both in individual patients and in isolates from different geographical regions (Choo et al., 1991; Martell et al., 1992; Higashi et al., 1993). These observations of diversity in persisting RNA virus populations are consistent with the models developed to describe the rapid evolution of RNA viruses and the existence of RNA quasispecies (Steinhauer and Holland, 1987; Eigen, 1993). The evolution of quasispecies in RNA virus infections appears to be a common theme in the development of "smoldering" and chronic viral infections (Martell et al., 1992).

Analysis of the mutations detected in the persistent coronavirus sequences revealed several interesting features. First, the majority of mutations detected in the persistent viral sequences are predicted to encode amino acid changes (65%). The prevalence of mutations that result predominantly in amino acid changes during coronavirus persistence suggests that MHV RNA evolution may be driven by positive selection. One of the most

	CTACGTTATGTCCAGGCTGAGTCTTTGTCGTGTAATAATATGATGCGTCCAAAGTGTATGGTATGTGTTGTTGTTCTCAGTTGATAAGTTTGCTATCCCCCGGAAGCCGTCAAATTGA

S42-7	***************************************
542-9	
S42-12	·····

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	C
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1206	TTTACARATTGGCRACTCCGGATTTTTGCARACGGCTAATTATARGATTGATACCGCTGCCACATCATGTCRGCTGTATTACAGTCTTCCTAAGAATAATCTTACCATARATAACTATAACCACACATCATGTCRGCTGTATTACAGTCTTCCTAAGAATAATCTTACCATARATAACTATAACCACACACACACAC
F42_1	
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	C

642-15	
394-11	***************************************
1222	CCTCGTCTTGGAATAGGAGGTATGGTTTTAATGATGCTGGTGTGTTTTGGCAAAAGTAAACATGATGTTCCCTACGCCCAGCAATGTTTACTGTGCGACCTAGCTATTGTCCGTGTGCAAA
	CCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCC

	g

S42-17	***************************************
	CCGGACRTAGTTAGCGCTTGCACTAGTCAGACCAAACCCATGTCTGCTTATTGCCCCACAGGCACAATTCATCGTGAGTGTTCTCTTTGGAATGGGCCCCATTTGCGCTCGGCACGTGTAGG

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542-12	[Deletion
S42-13	

542-16	······································
S42-17	***************************************
	TTCCGGCACGTACACGTGTGAGAGCACTTGTAAACCCAATCCATTGATACGTATGATCTCCGCTGTGGGCAAATTAAAACTATTGTTAATGTGGGCGATCATTGTAAAGGTCTGGGTGTTT

S42-12	Deletion]
542-13	
842-15	

1694	THGANGATANATGTGGC NATAGCGATCCACATAAGGGCTGTTCTTGTGCCAATGATTCTTTATCGGATGGTCACATGACACTTGTTTAGTANATGATCGCTGCCAAATTTTTGCTAACATA
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\$42-1 \$42-2	······································
\$42-1 \$42-2	
\$42-1 \$42-2 \$42-3	***************************************
\$42-1 \$42-2 \$42-3 \$42-7	······································
542-1 542-2 542-3 542-7 542-9	
542-1 542-2 542-3 542-7 542-9 542-12	······································
\$42-1 \$42-2 \$42-3 \$42-7 \$42-9 \$42-12 \$42-13	
\$42-1 \$42-2 \$42-3 \$42-7 \$42-9 \$42-12 \$42-13	
\$42-1 \$42-2 \$42-3 \$42-7 \$42-9 \$42-12 \$42-13 \$42-15	
\$42-1 \$42-2 \$42-3 \$42-7 \$42-9 \$42-12 \$42-13 \$42-15	
\$42-1 \$42-2 \$42-3 \$42-7 \$42-9 \$42-12 \$42-13 \$42-15 \$42-17	
\$42-1 \$42-2 \$42-3 \$42-7 \$42-9 \$42-12 \$42-13 \$42-15 \$42-17	
\$42-1 \$42-2 \$42-3 \$42-7 \$42-7 \$42-12 \$42-13 \$42-15 \$42-16 \$42-1 \$42-1	T
\$42-1 \$42-2 \$42-3 \$42-7 \$42-9 \$42-13 \$42-15 \$42-17 1816 \$42-1 \$42-2 \$42-2	TTGTTARATGGCATTAATAGTGGGACTACGTGTTCCACAGATTTACAATTGCCTAATACTGAAGTGGCCACTGGCGTTTGCGTCAGATATGACCTCTATGG
\$42-1 \$42-2 \$42-3 \$42-7 \$42-9 \$42-13 \$42-15 \$42-17 1816 \$42-1 \$42-2 \$42-3 \$42-7	TIGTTARATGCATTARTAGTGGGACTACGTGTTCCACAGATTTACAATTGCCTAATACTGAAGTGGCCACTGGCGTTTGCGTCAGATATGACCTCTATGG
\$42-1 \$42-2 \$42-3 \$42-7 \$42-9 \$42-13 \$42-15 \$42-16 \$42-1 \$42-1 \$42-3 \$42-3 \$42-9	T
\$42-1 \$42-2 \$42-3 \$42-7 \$42-9 \$42-13 \$42-15 \$42-16 \$42-17 1816 \$42-1 \$42-2 \$42-7 \$42-7 \$42-7 \$42-7	TTGTTARATGGCATTARTAGTGGGACTACGTGTTCCACAGATTTACAATTGCCTAATACTGAAGTGGCCACTGGCGTTTGCGTCAGATATGACCTCTATGG C.
\$42-1 \$42-2 \$42-3 \$42-7 \$42-9 \$42-15 \$42-15 \$42-17 1816 \$42-17 \$42-1 \$42-2 \$42-3 \$42-7 \$42-3 \$42-3 \$42-1 \$42-3 \$42-1	TIGTTARATGCATTARTAGTGGGACTACGTGTTCCACAGATTTACAATTGCCTAATACTGAAGTGGCCACTGGCGTTTGCGTCAGATATGACCTCTATGG C
\$42-1 \$42-3 \$42-3 \$42-7 \$42-7 \$42-12 \$42-13 \$42-16 \$42-16 \$42-1 \$42-2 \$42-2 \$42-2 \$42-13 \$42-13 \$42-13 \$42-13 \$42-13	TOTALANTIGGENTTANTAGTGGGACTACGTGTTCCACAGATTTACAATTGCCTAATACTGAAGTGGCCACTGGCGTTTGCGTCAGATATGACCTCTATGG G
\$42-1 \$42-2 \$42-3 \$42-7 \$42-7 \$42-1 \$42-15 \$42-16 \$42-16 \$42-1 \$42-2 \$42-1 \$42-1 \$42-1 \$42-1 \$42-1 \$42-1 \$42-1 \$42-1 \$42-1 \$42-1 \$42-1 \$42-1 \$42-1 \$42-1	TIGTTARATGCATTARTAGTGGGACTACGTGTTCCACAGATTTACAATTGCCTAATACTGAAGTGGCCACTGGCGTTTGCGTCAGATATGACCTCTATGG C

FIG. 3. Multiple mutations detected in the S1 region amplification products synthesized from RNA isolated from mice 42 days after inoculation with MHV-JHM. PCR products were cloned into plasmid pGEM-T and sequenced as described under Materials and Methods. Mutations detected in 10 individual clones are indicated.

TABLE 4

Mutations Detected in the Nucleocapsid Region during Persistent
MHV-JHM Infection

Clone	Nucleotide	Codon change	Amino Acid						
Day 4: 20 C	Day 4: 20 Clones sequenced, 11 clones encoded wild-type sequence								
N4-1	916 A → C	AAC → ACC	Asn → Thr						
N4-3	1004 U → C	AGU → AGC	No change						
N4-5	857 A → G	CAA → CAG	No change						
	1135 A → U	CAA → CUA	$GIn \rightarrow Leu$						
N4-6	1103 U → C	GCU → GCC	No change						
N4-11	956 C → U	GGC → GGU	No change						
N4-13	1005 G → A	GAU → AAU	Asp → Asn						
N4-20	696 U → C	UCG → CCG	Ser → Pro						
N4-25	849 A → G	ACA → GCA	Thr → Ala						
N4-29	840 A → G	AAG → GAG	Lys → Glu						
	977 C → U	GGC → GGU	No change						
Day 20: 4 (Clones sequenced,	2 clones encoded wild	d-type sequence						
N20-6	1087 A → G	AAG → AGG	Lys → Arg						
	1163 U → A	AGU → AGA	Ser → Arg						
N20-7	990 A → G	$AAA \rightarrow GAA$	Lys → Glu						
	767 U → C	ACU → ACC	No change						
Day 42: 20	Clones sequenced,	8 clones encoded will	d-type sequence						
N42-1	1166 U → C	ACU → ACC	No change						
N42-2	1116 A → U	$AAA \rightarrow UAA$	Lys → TERM						
	1150 U → C	GUU → GCU	Val → Ala						
N42-3	767 U → C	ACU → AAC	No change						
N42-4	847 U → C	GUA → GCA	Val → Ala						
	1002 A → G	AGU → GGU	Ser → Glu						
N42-6	771 A → G	$AAA \rightarrow GAA$	Lys → Glu						
	871 U → C	GUC → GCC	Val → Ala						
	1024 U → C	CUU → CCU	Leu → Pro						
N42-7	740 U → C	AGU → AGC	No change						
	934 A → C	CAG → CCG	$GIn \rightarrow Pro$						
	1146 G → A	GCA → ACA	Ala \rightarrow Thr						
N42-8	1034 G → A	UUG → UUA	No change						
N42-9	949 A → U	AAG → AUG	Lys → Met						
	1061 U → C	AAA → AAC	No change						
N42-10	941 U → C	UGU → UGC	No change						
	1049 U → C	GGU → GGC	No change						
N42-11	868 A → G	gaa → agg	Lys → Arg						
N42-12	792 A → G	AUU → GUU	lle → Val						
	911 U → C	ACU → GGG	No change						
	1109 A → G	GGA → GGG	No change						
N42-18	912 C → U	CCA → UCA	Pro → Ser						

striking examples of positive Darwinian evolution in viruses is that of Influenza A virus (Fitch et al., 1991). In the case of influenza A virus, at least two significant factors contribute to rapid evolution of the virus; the high mutation rate of the viral polymerase, and the selection of mutations which allow the virus to escape immune surveillance. Like Influenza virus, coronaviruses are replicated by a viral RNA-dependent RNA polymerase with no known proofreading activity and may evolve rapidly through random polymerase errors (Lai, 1990). In the MHV-infected animal, the immune system may play a role in neutralizing the wild-type virus and allowing viruses which have altered or deleted epitopes to propagate. The strong selection for specific S1 region deletion variants has been shown previously by isolating and characterizing attenuated variants of coronavirus (Dalziel et al., 1986; Fleming et al., 1986; Morris et al., 1989; Parker et al., 1989; La Monica et al., 1991; Wang et al., 1992). These attenuated variants have been shown to have significant deletions or point mutations in the spike glycoprotein gene, predominantly in the S1 "hypervariable" region (Banner et al., 1990; Wang et al., 1992). Our results indicate that mutations which encode amino acid changes are not limited to the S1 hypervariable region but also occur in the nucleocapsid region and potentially throughout the genome. Mutations are probably generated by errors introduced randomly throughout the genome (Steinhauer and Holland, 1987). Our results are consistent with the observations of Yokomori et al., who showed that multiple HE protein-defective mutants are generated during both acute and subacute phases of viral infection in the mouse brain (Yokomori et al., 1993).

The second feature of the mutations detected in the persistent coronavirus sequences is that the mutations are biased, with 63% of the mutations being U to C or A to G transitions. This type of "biased hypermutation" has been previously described in association with persistent infection of measles virus which results in measles inclusion body encephalitis (Cattaneo *et al.*, 1988). Although the mechanism for generating biased mutations is not known, it has been proposed to occur during inefficient or partially blocked stages in viral replication (Bass *et al.*, 1989). Such inefficient replication may occur if double-

TABLE 5
Summary of Mutations Detected in the Nucleocapsid Region during Persistent MHV-JHM Infection

Days postinfection				Туре			
	Clones	Total mutations detected	s	С	Del	Term	Mutations per thousand nucleotides sequenced
4	20	11	5	6	0	0	1,1
20	4	4	1	3	0	0	2.0
42	20	23	8	14	0	1	2.4

^a S, silent; C, change of amino acid; Del, deletion; Term, termination codon.

TABLE 6
Type of Nucleotide Change Occurring in the S and N RNA during Persistent Infection

	Transition			Transversion								
	∩→C	C → U	G → A	A → G	U → A	A → U	G → C	C → G	U → G	G→U	A→C	C → A
S — Day 4	8	2	4	6	0	0	2	0	0	0	0	0
S — Day 8	1	0	2	1	0	0	0	0	0	0	0	0
S — Day 13	4	0	3	1	0	1	0	0	0	1	0	0
S — Day 20	4	0	1	2	0	0	0	0	0	0	0	0
S — Day 42	17	6	6	15	3	3	0	1	2	0	0	1
S — Total	34	8	16	25	3	4	2	1	2	1	0	1
N — Day 4	3	2	1	3	0	1	0	0	0	0	1	0
N — Day 20	1	0	0	2	1	0	0	0	0	0	0	0
N — Day 42	1 1	1	2	6	0	2	0	0	0	0	1	0
N — Total	15	3	3	11	1	3	0	0	0	0	2	0

stranded RNAs (dsRNA) are stable intermediates of replication. For replication to occur from a dsRNA, a putative viral or cellular helicase activity is required to unwind the RNA. Such a dsRNA unwinding/modifying activity has been characterized in human cells (Bass and Weintraub, 1988). This enzyme can modify adenosine residues to inosine residues, resulting in a destabilizing of the double-stranded RNA and unwinding, thereby allowing elongation by the RNA polymerase (Wagner et al., 1989). The result of the modification of adenosine to inosine is a predominance of A to G and U to C mutations during the replication of plus- and minus-strand RNA (Bass et al., 1989). Clearly, biased hypermutations are much more prevalent in measles virus persistent infection as compared to coronavirus persistent infection (there are greater than 50% U to C transitions and over 20% of the U residues in certain regions are changed to C in persistent measles RNA, Wong et al., 1991). However, it is interesting to speculate that a similar mechanism for generating biased mutations may contribute to the mutations seen in two different types of viral persistent CNS infections, measles and coronavirus.

The mechanism by which the mutant viral RNAs are generated and the importance of selective pressures (immune system, neuron or glial cell environment, and viral replication competency) exerted on the persistent viral RNA remains to be determined. The mutations may be introduced randomly by the viral RNA-dependent RNA polymerase during replication. Error rates for viral RNA polymerases have been estimated to be on the order of 10^{-4} to 10^{-5} (Steinhauer and Holland, 1987; Parvin *et al.*, 1986; De La Torre *et al.*, 1992). The high error rate during replication allows for rapid evolution (Steinhauer and Holland, 1987; Holland *et al.*, 1982). Mutations may also be introduced by the dsRNA unwinding/modifying activity as described above (Bass *et al.*, 1989; Wong *et al.*, 1991; Rataul *et al.*, 1992). The mutant or defective viral ge-

nomes generated during this process may have an advantage in the persistently infected cell by being unable to synthesize infectious virus and therefore escaping immune recognition. The diverse population or "quasispecies" (Eigen, 1993) of RNAs that we detect in coronavirus persistent infection may reflect the multiple pathways that are used to generate and maintain the persistent viral RNA.

The role of persistent coronavirus RNA in the pathogenesis of chronic demyelination remains to be elucidated. The persistent viral RNA may be translated at low levels, resulting in chronic antigenic stimulation at sites of viral infection. Alternatively, persistent viral RNA sequences could result in abnormal neuronal or glial cell function, potentially disrupting normal CNS functions. Reactivation of latent or defective virus may be another mechanism contributing to pathogenesis in persistent coronavirus infection. Studies with the Sindbis virus (SV) model system for acute encephalitis infection of mice have demonstrated that SV RNA persists in neurons of infected mice for at least 17 months after infection (Levine and Griffin, 1992). Interestingly, the persistent Sindbis virus can be reactivated under conditions of waning antibody titer (Levine and Griffin, 1992). Further study of coronavirus persistence in the absence of immune selection in vivo (in SCID mice) may shed light on the mechanisms involved in the generation of mutant viruses and the role of persistent viral RNA in the pathogenesis of chronic demyelination.

ACKNOWLEDGMENTS

We thank Colette Ruedin for excellent technical assistance. This work was supported by Public Health Service Research Grant Al32065 from the National Institutes of Health, a Junior Faculty Research Award from the American Cancer Society (to S.C.B.), and National Multiple Sclerosis Society Research Grant RG2283-A-2 (to J.O.F.).

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