A combination of mutations in the S1 part of the spike glycoprotein gene of coronavirus MHV-A59 abolishes demyelination

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The A59 strain of coronavirus, mouse hepatitis virus (MHV), produces acute hepatitis, meningoencephalitis, and chronic demyelination. The authors have previously shown that the spike (S) glycoprotein gene of MHV contains determinants of virulence, hepatitis, and demyelination. They then identified viruses containing mutations in the S gene that exhibit alterations in viral pathogenesis. In the present study, the authors produced new recombinant viruses with each one of these S gene mutations by site-directed mutagenesis and targeted recombination and studied the effect of each individual mutation on the pathogenesis of the virus. They identified a combination of mutations in the S1 gene (I375M and L652I) that abolishes demyelination. Individual mutation and other combinations of mutations in the S gene only interfere with virulence and hepatitis and only reduce demyelination (I375M), but do not abolish demyelination completely. Thus, demyelination determinants exist within genomic regions on both sides of the hypervariable region, downstream from the receptor-binding domain in the S1 part of the MHV spike glycoprotein gene. The structure and precise function of these regions awaits further investigation. Journal of NeuroVirology (2004) 10, 41–51.

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Introduction

Mouse hepatitis virus (MHV) is a member of the coronavirus genus, which belongs to the Coronaviridae family of the Nidovirales order (Cavanagh, 1997; Lai and Cavanagh, 1997). Nidoviruses produce a variety of hepatic, enteric, and neurologic diseases in animals, and upper respiratory and enteric diseases in humans (Lai and Cavanagh, 1997; Lavi et al, 1999; Lavi and Weiss, 1989). Infection of MHV-A59 in 4-week-old C57Bl/6 (B6) mice produces a biphasic disease with acute hepatitis and meningoencephalitis followed by chronic inflammatory demyelinating disease (Lavi et al, 1984, 1986). MHV infection in mice (with both A59 and JHM strains) serves as an excellent laboratory model for virus-induced demyelination, similar to multiple sclerosis (MS) in humans (Houtman and Fleming, 1996; Knobler et al, 1981; Lavi et al, 1984b; Perlman et al, 1990; Sorensen et al, 1984; Wege et al, 1982; Weiner, 1973). Because viral infection is considered as a potential trigger for demyelination in MS (Allen and Brankin, 1993), understanding the mechanism of MHV-induced demyelination and the upstream events that lead to the autoimmune phenomenon may provide important insight into the mechanism of interaction between viruses and the nervous system, especially with respect to
the role of local and systemic immune system in viral neurotropism.

To understand the molecular basis of MHV neurotropism, we investigated a strain of MHV (MHV-2) that is closely related to MHV-A59 but is only weakly neurotropic (Hirano et al., 1974, 1981; Wege et al., 1981; Yamada et al., 1997; Yokomori et al., 1991). We sequenced the entire genome of MHV-2 and studied its detailed pathologic properties (Das Sarma et al., 2001b). We found that MHV-2 produces acute hepatitis and meningitis, without encephalitis, without demyelination, and without persistence of virus in the central nervous system (CNS) (Das Sarma et al., 2001b). We then replaced the S gene of A59 with that of MHV-2 by targeted recombination and produced recombinant viruses that have a persistence-positive, demyelination-negative phenotype. We concluded that demyelination determinants, independent of viral persistence, map to the S gene of MHV (Das Sarma et al., 2000). Additional studies identified molecular determinants of pathogenesis in the S gene of MHV (Leparc-Goffart et al., 1998; Navas et al., 2001; Phillips et al., 1999, 2001).

To further explore which parts of the S gene are directly responsible for the demyelinating phenotype, we studied a set of recombinant viruses that were produced by infection with both MHV-2 and LA-7, a temperature-sensitive (ts) mutant of MHV-A59 (Keck et al., 1988). These recombinant viruses were all found to be nondemyelinating. Sequencing the S genes of these recombinant viruses revealed that the recombinants' S gene was either derived from MHV-2 or from MHV-A59. All the recombinant viruses with an S gene derived from MHV-A59 had three identical amino acid substitutions within the S gene (Das Sarma et al., 2001a). Because these recombinant viruses were produced by dual infection of cultures with MHV-2 and LA-7, we could not rule out that they also contained other mutations in addition to the mutations in the S gene. We therefore used site-directed mutagenesis and targeted recombination to produce new recombinant viruses, with selective S gene mutations that were associated with the demyelination-negative phenotype.

**Results**

**Generation of the S gene recombinant viruses by targeted RNA recombination**

We have previously identified three amino acid substitutions in the S gene (Figure 1) that are potentially associated with demyelination (I375M, L652I, and T1087N). By using the site-directed mutagenesis kit (see Materials and methods), we mutated three different sites individually or jointly. Targeted recombination was carried out between synthetic RNAs transcribed from recombinant vectors and recipient virus Alb4 (Penn99-1, 2) or fMHV (all other recombinants) according to previously described methods (Peng et al., 1995; Das Sarma et al., 2000). Alb4 is thermostable and displays small plaque morphology at 39°C; thus putative recombinants were selected as large plaque viruses after treatment for 24 h at 40°C and plating on L2 cell at 39°C. The use of the virus with feline spike protein (fMHV) also provided a

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**Structure of the MHV-A59 spike glycoprotein (S) gene**

![Figure 1](image.png)

Figure 1 Schematic representation of the map of the spike glycoprotein gene with emphasis on the S1 portion of the gene. RBD, receptor binding domain; HVR, hypervariable region; TCE, CD8+ T-cell epitope H-2Kb: RCQIFANI; TMD, transmembrane domain; SDR1, S demyelination region 1; SDR2, S demyelination region 2.
useful method to select recombinant viruses against the background of parental viruses. Because fMHV does not grow in mouse cells, the recombination process with the transfected plasmid could be achieved in feline cells (FCWF). Recombinant viruses with S genes derived from mutated mouse S gene regained the ability to grow in mouse 17Cl-1 cells. By switching from feline cells to mouse cells that did not support the growth of parental fMHV, we were able to select the recombinant viruses because only recombinant viruses with a mouse S gene were present in these cultures. The potential recombinants were plaque purified twice and screened by using polymerase chain reaction (PCR) sequencing to detect the expected mutations made in the S gene. To verify that no secondary site mutations had been introduced into the S gene, we sequenced the entire S gene of the new recombinant viruses.

In vitro analysis of new recombinant viruses
We analyzed the ability of the new recombinant viruses to grow in L2 cells by growth-curve kinetics analysis, and compared it with the MHV-A59 kinetics. We also performed growth analysis of the new viruses in different temperatures (33°C, 37°C, 39°C) to rule out a ts phenotype. The growth kinetics of the viruses at 37°C is shown in Figure 2. It demonstrates that all viruses have similar growth properties. In addition, none of the viruses was ts.

In vivo analysis of new recombinant viruses
We analyzed the virulence and pathologic properties of the new recombinant viruses and compared it to that of wtMHV-A59 and the control recombinant virus wtR-A59. The virulence of the viruses is shown in Table 1. The T1087N mutation in S2 reduced viral virulence by fivefold. However, all other individual and combination of mutations in the S1 gene reduced virulence significantly (as assessed by LD50), and rendered the recombinant viruses nonlethal at the maximum titer of virus injected (50,000 plaque-forming units [PFU]). Reduced virulence correlated with reduction in the amount of hepatitis, mostly seen in the viruses that had mutations in the S1 gene.

The effect of the three mutations on the pathologic picture of encephalitis and hepatitis is shown in Figures 3 and 4. Generally, none of the mutations or combination of mutations affected the extent of encephalitis. Hepatitis was abolished in all the viruses that contained the I375M mutation individually or with L652I (Penn01-1, 2; Penn2K-3, 4; and Penn2K-1, 2). Hepatitis was reduced in Penn01-3, 4 (L652I individual mutation), but interestingly, hepatitis was not significantly changed in combinations of the T1087N mutation with each one of the other S1 mutations (Penn01-5, 6, and Penn01-7, 8). The effect of the three mutations on viral replication in the infected organs (brain and liver) is shown in Figure 5. Viral titers in the brain and liver were consistent with the pathologic picture. Because viruses were injected into the brain, the brain titers on day 1 generally represent residual injected virus.

The effect of the three mutations on demyelination is shown in Table 2. Demyelination was not affected by the reduced virulence and most nonlethal viruses were still able to cause demyelination. Only viruses containing both I375M and L652I mutations in the S1 gene were nondemyelinating. Therefore, the two combinations that abolished demyelination were the one that included the I375M and L652I mutations (Penn2K-3 and Penn2K-4) and the one that included all three mutations I375M, L652I, and T1087N (Penn2K-1 and Penn2K-2). Neither one of these individual mutations was able to abolish demyelination completely. The I375M individual mutation and the combination of L652I and T1087N partially reduced demyelination, especially at the lower doses of viral injections. Interestingly, the combination of I375M and T1087N mutations did not reduce demyelination at the same level as the individual I375M mutation.

Analysis of viral RNA persistence
Viral RNA persistence of all recombinant and parental viruses in mouse organs (brain, spinal cord and liver) was analyzed by reverse transcriptase

Table 1 Virulence of recombinant viruses

| Viruses             | LD50     |
|---------------------|----------|
| MHV-A59             | 4000 PFU |
| WtR-A59             | 5000 PFU |
| Penn2K-1, 2 (I375M, L652I, and T1087N) | >50,000 PFU |
| Penn2K-3, 4 (I375M and L652I) | >50,000 PFU |
| Penn01-1, 2 (I375M) | >50,000 PFU |
| Penn01-3, 4 (L652I) | >50,000 PFU |
| Penn99-1, 2 (T1087N) | 20,000 PFU |
| Penn01-5, 6 (I375M and T1087N) | >50,000 PFU |
| Penn01-7, 8 (L652I and T1087N) | >50,000 PFU |

Figure 2 Viral titer kinetics of the newly constructed recombinant viruses in a 24-h analysis of samples from infected L2 cells, as evaluated by agarose-based plaque assays at 37°C on L2 cells following infection of 1 PFU per cell (m.o.i = 1). Only one virus of each pair of identical recombinant viruses is shown. All viruses have growth patterns similar to MHV-A59.
Figure 3  Pathologic analysis of recombinant viruses during acute infection. Mice were injected IC with 1000 PFU per mouse with each virus. Only one recombinant virus from each set of duplicate viruses is shown; however, similar findings were found for both viruses. Each time point represents the pathologic analysis of two to three mice. Dark bars represent encephalitis whereas white bars represent hepatitis. The intensity of the pathologic damage of encephalitis was based on a 3-level scale: 0 = no inflammatory infiltrates in the brain; 1 = focal inflammatory infiltrate; 2 = multifocal inflammatory infiltrates; 3 = diffuse inflammatory infiltrates in the entire brain (panencephalitis). The intensity of the hepatitis was evaluated based on a 4-level scale: 0 = no hepatitis; 1 = one to two foci of necrosis in each section of liver; 2 = three to six necrotic foci in each liver section; 3 = more than six foci of nonbridging necrosis in each liver section; 4 = diffuse bridging necrosis of the liver with islands of preserved parenchyma.
Figure 4  Histopathology of mouse organs following infection with MHV-A59 and demyelination-negative recombinant viruses. (A) Four-week-old C57Bl/6 mice were injected IC with 1000 PFU of MHV-A59 and sacrificed 5 days post infection. Liver section shows necrotic lesions with cellular debris, inflammatory cells (polymorphonuclear leukocytes, lymphocytes and macrophages), and focal hemorrhage, representing acute hepatitis (H&E ×200). (B) Four-week-old C57Bl/6 mice were injected IC with 1000 PFU of MHV-A59 and sacrificed 5 days post infection. Brain section shows perivascular lymphocytic infiltration microglial proliferation and pyknotic neurons representing acute encephalitis (H&E ×200). (C) Four-week-old C57Bl/6 mice were injected IC with 5000 PFU of the MHV-A59 and sacrificed 30 days post infection. Spinal cord section shows multiple inflammatory demyelinating lesions in the white matter of the posterior and lateral columns (Luxol-Fast-Blue and cresyl violet ×50). (D) Four-week-old C57Bl/6 mice were injected IC with 1000 PFU of Penn2K-1 and sacrificed 5 days post infection. Liver section shows normal histology (H&E ×200). (E) Four-week-old C57Bl/6 mice were injected IC with 1000 PFU of Penn2K-1 and sacrificed 5 days post infection. Brain section shows perivascular lymphocytic infiltration microglial proliferation and pyknotic neurons representing acute encephalitis (H&E ×200). (F) Four-week-old C57Bl/6 mice were injected IC with 5000 PFU of the Penn2K-1 and sacrificed 30 days post infection. Spinal cord section shows normal histology (Luxol-Fast-Blue and cresyl violet ×40). (G) Four-week-old C57Bl/6 mice were injected IC with 1000 PFU of Penn2K-3 and sacrificed 5 days post infection. Liver section shows normal histology (H&E ×200). (H) Four-week-old C57Bl/6 mice were injected IC with 1000 PFU of Penn2K-3 and sacrificed 5 days post infection. Brain section shows perivascular lymphocytic infiltration microglial proliferation and pyknotic neurons representing acute encephalitis (H&E ×200). (I) Four-week-old C57Bl/6 mice were injected IC with 5000 PFU of the Penn2K-3 and sacrificed 30 days post infection. Spinal cord section shows normal histology (Luxol-Fast-Blue and Cresyl Violet ×40).

(RT)-PCR. The results of RNA persistence in the spinal cord are shown in Figure 6. Both MHV-A59 and recombinant virus wtR-A59 showed a prominent band consistent with MHV RNA. Similar bands were detected following infections with Penn99-1, Penn01-1, Penn01-3, and Penn01-5, all of which are demyelinating viruses. The demyelinating-negative virus Penn2K-1 showed no RNA persistence in the cord; however, the other demyelinating-negative virus Penn2K-3 showed a band similar to demyelinating-positive viruses. The demyelination-positive virus Penn01-7 showed a weaker positive band. Thus, although some degree of RNA persistence may be required for demyelination, the presence of persistence is insufficient for the development of demyelination. RNA persistence in the brain was similar to the spinal cords but with a slightly weaker signal. RNA persistence in the liver was minimal and was only detected in MHV-A59 and wtR-A59 infections (data not shown).
Discussion

The MHV genome is the largest among RNA viruses. Although the generation of an infectious MHV clone has been recently reported (Yount et al., 2002), the new construct appears to be attenuated in vivo. Thus, molecular pathogenesis studies continue to require a complex manipulation of the MHV genome in order to introduce targeted changes into specific locations. To date, the best technique available for this purpose is targeted 3' end heterologous recombination, originally introduced by Paul Masters (Peng et al., 1995).
Table 2  Demyelinating properties of recombinant viruses

| Viruses     | No. of mice with demyelination/total no. of mice (%) | Demyelinated quadrants/total number of quadrants (%) |
|-------------|------------------------------------------------------|------------------------------------------------------|
|             | 500 PFU  | 4000–5000 PFU | 50,000 PFU | 500 PFU  | 4000–5000 PFU | 50,000 PFU |
| MHV-A59     | 5/5 (100%) | 9/9 (100%) | N/A | 33/84 (39%) | 98/128 (77%) | N/A |
| Penn09-2    | 8/8 (100%) | 6/6 (100%) | 2/2 (100%) | 27/44 (61%) | 32/52 (62%) | 8/12 (67%) |
| Penn01-1    | 3/9 (33%)  | 2/10 (20%) | 9/10 (90%) | 7/160 (4%)  | 2/184 (1%)  | 24/188 (13%) |
| Penn01-3    | 10/10 (100%) | 10/10 (100%) | 10/10 (100%) | 50/168 (30%) | 54/184 (30%) | 64/200 (32%) |
| Penn2K-3    | 0/10 (0%)  | 0/10 (0%)  | 1/10 (10%) | 0/116 (0%)  | 0/120 (0%)  | 1/120 (0.8%) |
| Penn01-5    | 10/10 (100%) | 9/10 (90%) | 10/10 (100%) | 83/200 (42%) | 33/111 (30%) | 97/148 (66%) |
| Penn01-7    | 3/5 (60%)  | 1/5 (20%)  | 10/10 (100%) | 12/100 (12%) | 21/96 (22%)  | 36/200 (18%) |
| Penn2K-1    | 0/10 (0%)  | 0/10 (0%)  | 0/10 (0%)  | 0/120 (0%)  | 0/124 (0%)  | 0/100 (0%)  |

Note. MHV-A59 and representative recombinant viruses of each pair were analyzed for their ability to cause spinal cord demyelination following IC injections with three different doses of the virus. The other virus of each pair presented a similar phenotype. MHV-A59 was not evaluated at the highest dose because there are no surviving mice at this dose on day 30 post infection.

We used this technique in our previous studies, in which we identified the demyelination determinants in the S gene (Das Sarma et al., 2000). We continued our investigation in the present study to better define the regions of genomic control of demyelination within the S gene.

Although incidental mutations during coronavirus replication are not uncommon, we used strict precautions to reduce the chance that incidental mutations might interfere with molecular pathogenesis studies. We sequenced the entire S gene of each new recombinant virus immediately before it was used in pathogenesis studies to confirm the presence of the desired mutations and to exclude the possibility of incidental undesired mutations. The pathogenesis of new recombinant viruses was tested immediately after the generation of the viruses to avoid unnecessary passages of the viruses in culture, which tend to increase the chance for mutations. We generated two identical recombinant viruses for each mutation in two independent recombination procedures. Both viruses were tested in mice to ensure an identical in vivo phenotype. These precautions reduced the chance for incidental mutations to a negligible level.

Using these principles, we found that a combination of mutations in the S1 gene abolishes demyelination only when introduced simultaneously. The I375M mutation, which appears to be the dominant and the more important one, reduces the amount of demyelination but does not abolish it completely. However, I375M in combination with L652I completely abolishes demyelination. Thus, although this may not be the only genomic site that controls demyelination, the region in the S1 gene downstream from the receptor-binding domain contains important determinants of demyelination (Figure 1).
The coronavirus spike protein is a type I membrane protein with approximately a 1300-residue ectodomain, an 18-residue transmembrane portion, and a 38-residue cytoplasmic tail (Schmidt et al., 1987). The full-length spike glycoprotein oligomerizes rapidly after synthesis and then cleaves by a furin-like protease into two similar-sized fragments: a peripheral S1 and a membrane-anchored S2 (Sturman et al., 1985). S1 and S2 are associated by noncovalent interactions. S1 is responsible for binding to cellular receptors whereas S2 has the primary responsibility for membrane fusion. The receptors for MHV are members of the family of carcinoembryonic antigen cell adhesion molecules (CEACAM), of the immunoglobulin superfamily (Devksler et al., 1991, 1993a, 1993b). Dimers of S1 bind to a single molecule of CEACAM. The binding sites on S1 are presumably juxtaposed, causing a steric hindrance or global conformational changes that preclude additional interactions (Lewicki and Gallagher, 2002). After binding to receptor, the spike protein undergoes conformational changes, including alternative disulfide linkages and separation of S1 from S2, which may increase the propensity of S to fuse membranes (Taguchi and Matsuyama, 2002).

We hypothesize that the I375M and L652I mutations mark potential S1 genomic sites that control demyelination. These mutations are in the middle of two regions of yet undesigned name or function on either side of the hypervariable region of S1. As shown in Figure 1, we provisionally labeled these regions as S demyelination region 1 (SDR1) and S demyelination region 2 (SDR2). There are no known T- or B-cell epitopes in these regions; however, these S1 regions may influence spike protein functions in several ways. Binding of S1 to receptor is unlikely to be significantly affected by sites outside the first 330 amino acids because the CEACAM binding site can exist wholly in the amino terminal 330 residues of S1. However, SDR1 and SDR2 may still have influence on S1-CEACAM binding kinetics, which may affect pathogenesis. Similarly, the SDR1 and SDR2 regions may alter the kinetics and/or extent of the fusion activity of the spike protein, or the kinetics and/or extent of separation of S1 from S2. Either one of these changes may have an effect on the pathogenesis of the virus. The fact that two mutations are necessary for the abolishment of demyelination may rely on the three-dimensional spatial structure and folding of the S1 part of the molecule and on potential chemical interactions between the two sites.

Both I375M and L652I mutations also reduce hepatitis and therefore reduce the overall virulence of the viruses resulting in a nonlethal phenotype. However, the reduced virulence per se is not enough to abolish demyelination. We have shown here and in a previous publications (Das Sarma et al., 2000, 2001a, 2001b) that viruses that lack the ability to produce demyelination can be either less virulent (Penn2K-1, and Penn2K-2) or significantly more virulent (Penn98-1 and Penn98-2, and MHV-2) than MHV-A59. Moreover, the individual L652I mutation has a nonlethal phenotype but no effect on demyelination. In addition, ability of the virus to produce encephalitis may be a prerequisite, but is insufficient for the development of subsequent demyelination. For example, Penn2K-3 and penn2K-4 produce encephalitis similar to demyelinating viruses but do not produce demyelination. This study, therefore, provides further evidence that the ability of a virus to produce chronic demyelination is not in direct correlation with the virulence of the virus. Similarly, viral persistence may be a prerequisite but is insufficient for the development of demyelination. The property of demyelination is therefore a unique additional neurotropic property, which appears to be at least in part encoded by the S1 gene of the MHV genome.

Materials and methods

Viruses and cells

The following plaque-purified viruses were used in this study: MHV-A59 (Budzilowicz et al, 1985; Lavi et al., 1984a, 1984b), Alb4 (obtained from Paul Masters, Albany, NY), and fMHV (Kuo et al, 2000) (obtained from Paul Masters, Albany, NY). Stock viruses had titers of 10⁷ to 10⁸ PFU/ml. Viruses were propagated and titered on murine L2 cells, 17Cl-1 cells, and feline FCWF cells in Dulbecco’s minimum essential medium (DMEM) with 10% fetal bovine serum (FBS).

Plasmids and PCR mutagenesis

The pMH54 plasmid (obtained from Paul Masters) contains the 1.1-kb, 3′ end of MHV-A59 genome from the HE gene downstream (Das Sarma et al, 2000; Kuo et al, 2000). The sequence of the S gene in pMH54 is identical to that of MHV-A59 except for the introduction of AvrII and sbfi restriction sites, which allowed the insertion of different S genes into the background of pMH54, and enabled the identification of recombinant viruses containing full length recombinant S gene. Site-directed mutagenesis experiments were performed using the Stratagene QuickChange kit (Cedar Creek, TX).

Targeted RNA recombination and selection of recombinants

Targeted RNA recombinations were carried out between Alb4 (for Penn99-1, 2), fMHV (for all other recombinant viruses), and synthetic RNAs. Alb4 is thermodabile and displays small plaque morphology at 39°C. fMHV contains an S gene from FIPV on an MHV-A59 background (Kuo et al, 2000) and can only grow in feline cells FCWF. The synthetic RNAs were supplied by the pMH54 plasmid containing the entire 3′ end of the MHV genome from gene 2 (HE) to 7 (N). For recombination, L2 cells spinner culture (for Penn99-1, 2 only) or FCWF (all other recombinants)
was harvested and infected at multiplicity of infection (m.o.i.) = 1 of Alb4 (at 33°C), or fMHV (at 37°C), respectively. Cells were then transfected by electroporation with the synthetic RNA by using two consecutive pulses from Bio-Rad Gene Pulser apparatus set at 0.3 kV and 960 μF. Infected and transfected cells were plated onto a monolayer of 17Cl-1 cells, and the released viruses were harvested at approximately 24 to 48 h post infection. For selection of the recombinant viruses, a two-step plaque purification was performed and PCR sequencing of the S gene was used to confirm the presence of the desired mutations. For each mutation, we produced two identical viruses in two independent recombination events. This strategy was designed to reduce the possibility that random unrecognized mutations would affect the phenotype. The new recombinant viruses were designated as the following (also in Table 1): (1) Penn2K1 and Penn2K2 contain an MHV-A59 S gene with the I375M, L652I 2-point mutations; (2) Penn2K3 and Penn2K4 contain an MHV-A59 S gene with the I375M and L652I 2-point mutations; (3) Penn99-1 and Penn99-2 contain an MHV-A59 S gene with the T1087N single-point mutation (Das Sarma et al., 2001c); (4) Penn01-1 and Penn01-2 contain an MHV-A59 S gene with the I375M single-point mutation; (5) Penn01-3 and Penn01-4 contain an MHV-A59 S gene with the L652I single-point mutation; (6) Penn01-5 and Penn01-6 contain an MHV-A59 S gene with the I375M and T1087N two-point mutations; (7) Penn01-7 and Penn01-8 contain an MHV-A59 S gene with the L652I and T1087N 2-point mutations. Finally, a control virus, wtR-A59, was generated by targeted recombination in which the S gene of MHV-A59 was combined back into the MHV-A59 background, to eliminate the possibility that any changes in pathogenesis might derive from the recombination process itself.

Viral growth curves
L2 cell cultures were prepared in 12-well plates in DMEM with 10% FBS. Confluent monolayers were infected with each virus (m.o.i. = 1) in duplicate wells and incubated for 1 h at 37°C. After adsorption, the cells were washed with PBS three times and then fed with 1.5 ml medium. At the indicated times, the cells were lysed by three cycles of freeze-thawing, and the supernatants were removed and titered by plaque assay on L2 cells as previously described (Lavi et al., 1984b).

Histologic analysis
For organ analysis, mice were sacrificed at various time points post infection. After perfusion with PBS, brains and livers were removed. Half of the brain and a portion of liver were fixed in 10% buffered formalin, paraffin-embedded, and stained with hematoxylin and eosin (H&E) for a blinded histologic analysis. Samples of brain and liver were frozen in −80°C, to be used later for assessment of viral titers by plaque assays. For assessment of demyelination, mice were sacrificed at 30 days post infection. Mice were perfused with PBS followed by 10% buffered formalin. Spinal cords were removed, formalin-fixed, and paraffin embedded. Four to six transverse sections of cervical, thoracic, and lumbar levels of the cord were stained with Luxol fast blue (LFB) and the number of quadrants of cord containing demyelination were counted as a fraction of the total number of cord quadrants evaluated in each individual mouse (Lavi et al., 1984b, 1986, 1988, 1990).
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