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Murine Coronavirus Spike Glycoprotein Mediates Degree of Viral Spread, Inflammation, and Virus-Induced Immunopathology in the Central Nervous System

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The mouse hepatitis virus (MHV) spike glycoprotein is a major determinant of neurovirulence. We investigated how alterations in spike affect neurovirulence using two isogenic recombinant viruses differing exclusively in spike. S4R, containing the MHV-4 spike gene, is dramatically more neurovirulent than SA59R, containing the MHV-A59 spike gene (J. J. Phillips, M. M. Chua, E. Lavi, and S. R. Weiss, 1999, J. Virol. 73, 7752–7760). We examined the contribution of differences in cellular tropism, viral spread, and the immune response to infection to the differential neurovirulence of S4R and SA59R. MHV-4 spike-mediated neurovirulence was associated with extensive viral spread in the brain in both neurons and astrocytes. Infection of primary hippocampal neuron cultures demonstrated that S4R spread more rapidly than SA59R and suggested that spread may occur between cells in close physical contact. In addition, S4R infection induced a massive influx of lymphocytes into the brain, a higher percentage of CD8⁺ T cells, and a higher frequency of MHV-specific CD8⁺ T cells relative to SA59R infection. Despite this robust and viral-specific immune response to S4R infection, infection of RAG1⁻/⁻ mice suggested that immune-mediated pathology also contributes to the high neurovirulence of S4R.

INTRODUCTION

The severity of viral infections depends on the extent of tissue destruction and cellular dysfunction mediated by a combination of direct virus infection and immune-mediated destruction. The relative contribution of these two components differs depending on a number of virus and host factors including viral tropism, rate of viral spread, and specificity of the immune response.

Infection of mice with the murine coronavirus, mouse hepatitis virus (MHV), provides a model for studying acute virus-induced neurological disease. We have previously demonstrated that the spike (S) gene is a major determinant of MHV neurovirulence. Recombinant viruses containing the spike gene of the highly neurovirulent MHV-4 strain, S4R, exhibit a dramatically more neurovirulent phenotype (3 log₁₀ decrease in intracranial LD₅₀) than isogenic recombinants containing the spike gene of the mildly neurovirulent MHV-A59 strain, S₅₉₉₃R (Phillips et al., 1999). (S₅₉₉₃R and S₄R have all other genes derived from MHV- A59.) By defining the contribution of direct virus-mediated pathology and immune-mediated pathology to the differential neurovirulence of S₄R and S₅₉₉₃R, we can better elucidate the mechanisms by which the MHV-4 spike confers high neurovirulence.

The MHV spike glycoprotein is required for viral entry and spread (Boyle et al., 1987; Collins et al., 1982; Sturman and Holmes, 1981). Expressed on the virion surface, spike is responsible for binding to the viral receptor and mediating virus–cell fusion, and subsequent to infection spike, expressed on the host cell membrane, can mediate cell–cell fusion. The spike is also vital for the immune response to infection as it is able to induce both a cell-mediated and a humoral-mediated immune response (Bergmann et al., 1996; Castro and Perlman, 1995; Collins et al., 1982). Thus the high neurovirulence conferred by the MHV-4 spike may be due to alterations in a number of aspects of infection including viral entry, viral spread, or the immune response to infection.

Although MHV infects many cell types in the brain, including neurons, astrocytes, and oligodendrocytes (Knobler et al., 1981; Lavi et al., 1984, 1988; Weiner, 1973), it has been suggested that neuronal tropism is a major determinant of MHV neurovirulence (Dubois-Dalcq et al., 1982; Fleming et al., 1986; Knobler et al., 1981). The high neurovirulence of S₄R is associated with extensive spread of virus in the brain (Phillips et al., 1999). Increased viral spread could either reflect increased infection of a particular cell-type, such as neurons, or alternatively increased infection of multiple cell types in the CNS.

In the CNS, clearance of infectious MHV requires multiple components of the immune response including CD8⁺ and CD4⁺ T cells and B cells. Adoptive transfer experiments in combination with depletion experiments have demonstrated that both CD8⁺ and CD4⁺ T cells are critical for normal viral clearance (Korner et al., 1991;
Stohlman et al., 1986, 1995, 1998; Sussman et al., 1989; Williamson and Stohlman, 1990; Yamaguchi et al., 1991). Furthermore, the peak of T lymphocyte infiltration into the CNS is coincident with falling titers of infectious virus in the CNS (Williamson et al., 1991). Relative to infection with S4R, infection with S4R results in a dramatic infiltration of inflammatory cells into the CNS (Phillips et al., 1999). Differences in either the composition of these infiltrates or their virus-specific activity could have a profound influence on neurovirulence.

The MHV spike is a major determinant of MHV neurovirulence. To identify the critical parameters of infection mediating the differential neurovirulence of S4R and S49R, we characterized the cellular tropism and composition and function of the lymphocytic response to infection with S4R and S49R. Our results suggest that the high neurovirulence conferred by alterations in the MHV spike are mediated by both increased viral spread in multiple cell types in the brain and immune-mediated pathology.

RESULTS

S4R and S49R exhibit similar cellular tropism

We previously observed that at the peak of virus replication, day 5 postinfection, the more neurovirulent S4R infects a significantly greater number of cells in the basal

![FIG. 1. Identification of S4R- and S49R-infected neurons in the brain 5 days after intracranial inoculation. Representative pictures are shown of double-immunofluorescent labeling of sagittal brain sections taken from S4R-inoculated (A and B), and S49R-inoculated (C and D) mice. A and C show immunofluorescent labeling of viral antigen in the basal forebrain (red). B and D show the corresponding labeling of MAP2b in neurons (green). Note MAP2b-positive cells exhibit the characteristic morphology of neurons. Arrows identify cells double-positive for viral antigen, and MAP2b in both S4R and S49R inoculated brains. Note the greater total number of viral antigen-positive cells following inoculation with S4R. No viral antigen staining was observed in mock-infected mice, when sections were incubated in preimmune sera, or when the primary antibody was omitted (data not shown). Magnification, X380.]

![FIG. 2. Quantification of viral antigen positive cells in the brain following inoculation with S4R and S49R. The total number of viral antigen-positive cells per brain section following infection with S4R (open squares), and S49R (open diamonds) was determined on days 3 and 5 postinfection. On day 5 postinfection two brain sections per mouse were examined, and the mean number of viral antigen-positive cells per sagittal section was determined (n = 5). For the quantification of viral antigen-positive cells on day 3 a single sagittal section was examined for each infected mouse (n = 4). The mean number of viral antigen-positive cells per brain section (horizontal line) following infection with S4R was significantly greater than with S49R on days 3 and 5 postinfection (two-tailed t test, P < 0.05, and P < 0.001, respectively).]
forebrain, hippocampus, and cingulate gyrus than S4R (Phillips et al., 1999). Increased infection could reflect differences in viral spread in a single cell type or in multiple cell types. To quantitatively compare the cellular tropism of S4R and S49R, we performed double-label immunofluorescence on sagittal brain sections for viral

### TABLE 1

| Virus | Animal | Section 1 | Section 2 |
|-------|--------|-----------|-----------|
|       | No. of viral antigen-positive cells | No. of cells double-positive with AP14 (%) | No. of viral antigen-positive cells | No. of cells double-positive for GFAP (%) |
| S4R   | 1      | 599       | 328 (54.8) | 478       | 132 (27.6) |
|       | 2      | 1016      | 493 (48.5) | 1096      | 111 (10.1) |
|       | 3      | 883       | 469 (53.1) | 765       | 148 (19.3) |
|       | 4      | 1192      | 597 (50.1) | 713       | 202 (28.3) |
|       | 5      | 249       | 143 (57.4) | 738       | 99 (13.4) |
|       | Group mean (52.8) | | | Group mean (19.7) |
| S49R  | 1      | 529       | 291 (55.0) | 288       | 61 (21.2) |
|       | 2      | 97        | 52 (53.6)  | 330       | 44 (13.4) |
|       | 3      | 239       | 134 (56.1) | 259       | 37 (14.3) |
|       | 4      | 403       | 208 (51.6) | 616       | 170 (27.6) |
|       | 5      | 326       | 163 (50.0) | 282       | 80 (28.4) |
|       | Group mean (53.3) | | | Group mean (21.0) |

* Mice were inoculated intracranially with 10 PFU of either S4R or S49R. Animals were sacrificed on day 5 postinfection and perfused, and the brain was fixed in formalin, embedded in paraffin, and sectioned sagittally. Section 1 and 2 are adjacent.

* The slides were double-immunostained with anti-MHV-A59 serum and AP14, a monoclonal antibody to MAP2b, or a mouse anti-GFAP antibody. Sagittal sections were coded and systematically examined in a blinded fashion (magnification ×190), and the total number of viral antigen-positive cells and double-positive cells per section was determined. The (%) is the percentage of the total number of viral antigen positive cells counted that were also positive for the indicated marker.

**FIG. 3.** Viral spread in primary hippocampal neuron cultures. Neurons were cultured for 4 days and then infected with S4R (A and B) or S49R (C and D). Representative pictures on day 3 postinfection are shown. Colocalization of viral antigen (A and C) and neuronal cell markers (B and D) were demonstrated using double-immunofluorescence for viral antigen (green) and MAP2 (red). The arrows indicate viral antigen-positive neurons, while the arrowheads indicate viral antigen-positive neurites. Note the large focus of viral antigen-positive cells following infection with S4R (A). Magnification, ×190.
antigen and MAP2b, a neuron-specific marker (De Camilli et al., 1984; Riederer et al., 1995), or glial fibrillary acidic protein (GFAP), an astrocyte-specific marker. Mice were inoculated intracranially with 10 PFU of S4R or S A59R and animals were sacrificed at 3 and 5 days postinfection.

The sections were systematically scanned (magnification, ×190) in a blinded fashion, and the total number of viral antigen-positive and double-positive cells was determined. Representative pictures of double-immunofluorescence for viral antigen and MAP2 (Fig. 3). Viral antigen-positive cells were often located in discrete foci. Moreover, viral antigen-positive neurites could often be observed between infected cells or between foci of infected cells, suggesting that perhaps one means of viral spread in these cultures was along neurites. To examine the pattern of viral spread in these cultures, we determined the total number of viral antigen-positive cells and the total number of foci of infected cells per coverslip at different time points (Table 2). On day 1 postinfection the average number of foci and the total number of antigen-positive cells was similar following infection with S4R and S A59R. On day 3 postinfection, although the average number of foci was similar, the average number of viral antigen-positive cells was significantly greater following infection with S4R than S A59R (two-tailed t test, P < 0.01).

Despite the difference in the absolute number of infected cells, S4R had a similar cellular tropism as S A59R. As shown in Table 1, we found that the percentage of viral antigen-positive cells that were also positive for neuronal or astrocytic markers was remarkably similar for S4R and S A59R. On day 5 postinfection, neurons appeared to account for approximately 53% of either S4R- or S A59R-infected cells, while astrocytes accounted for approximately 20% or 21% of S4R- or S A59R-infected cells, respectively. Thus the increased neurovirulence of S4R correlated with increased infection of multiple CNS cell types, including both neurons and astrocytes.

Viral spread of S4R and S A59R differs in primary neuronal cultures

The extent of neuronal infection is thought to be a determinant of MHV neurovirulence (Dubois-Dalcq et al., 1982; Fleming et al., 1986; Knobler et al., 1981). As an in vitro model of CNS infection and to examine the ability of S4R and S A59R to spread in neurons, we infected primary hippocampal neuron cultures obtained from B6 mice with the two viruses. Immunostaining for MAP2 and GFAP demonstrated that the cultures consisted primarily of neurons with less than 8% of the cells exhibiting GFAP immunoreactivity (data not shown). On days 0, 1, and 3 after infection, supernatants were titered for infectious virus and the number of viral antigen-positive cells was determined by immunofluorescence. Neurons did not exhibit cytopathic effect for the duration of the experiment as assessed by bright-field microscopy.

To compare the number of cells infected with each virus, we used immunofluorescence to detect viral antigen expression. S4R and S A59R infection of neurons was confirmed by cell morphology and by double-immunostaining for viral antigen and MAP2 (Fig. 3). Viral antigen-positive cells were often located in discrete foci. Moreover, viral antigen-positive neurites could often be observed between infected cells or between foci of infected cells, suggesting that perhaps one means of viral spread in these cultures was along neurites. To examine the pattern of viral spread in these cultures, we determined the total number of viral antigen-positive cells and the total number of foci of infected cells per coverslip at different time points (Table 2). On day 1 postinfection the average number of foci and the total number of antigen-positive cells was similar following infection with S4R and S A59R. On day 3 postinfection, although the average number of foci was similar, the average number of viral antigen-positive cells was significantly greater following infection with S4R than S A59R (two-tailed t test, P < 0.01).

Thus, on day 3 the number of viral antigen-positive cells per focus was greater for S4R, 5.79 ± 1.15, than S A59R, 3.17 ± 1.59 (mean ± SD). The difference in the number of infected cells, despite a similar number of foci, between S4R and S A59R, suggested that S4R spread faster via cell–cell spread than S A59R. Thus, the results in primary neuronal cultures were consistent with the results in vivo and suggested that the MHV-4 spike conferred an inherent ability to spread rapidly from cell to cell in the CNS.

Consistent with previous studies in which primary neuronal cultures were infected with MHV (Dubois-Dalcq et al., 1982; Pasick et al., 1994), the amount of infectious virus released into the media was minimal. Following infection with S4R or S A59R, the titers of infectious virus in the supernatants initially dropped, day 0 to day 1 postinfection, and then remained the same from day 1 to day 3 postinfection (Table 2). The inactivation of infectious virus in the residual inoculum most likely accounted for the precipitous drop in

### Table 2

| Virus | Day | Viral titers log (PFU/mL) mean ± SD | No. of foci mean ± SD | No. of viral antigen-positive cells mean ± SD |
|-------|-----|------------------------------------|-----------------------|---------------------------------------------|
| S4R   | 0   | 3.79 ± 0.39                        | 0                     | 0                                           |
|       | 1   | 2.13 ± 0.34                        | 60 ± 17               | 367 ± 108                                   |
|       | 3   | 2.28 ± 0.22                        | 127 ± 52              | 842 ± 31                                    |
| S A59R| 0   | 4.21 ± 0.12                        | 0                     | 0                                           |
|       | 1   | 3.59 ± 0.15                        | 122 ± 79              | 302 ± 165                                   |
|       | 3   | 3.38 ± 0.19                        | 162 ± 34              | 471 ± 109                                   |

* Primary hippocampal neuron cultures were infected on day 4 post-explant at an m.o.i. of 5. Coverslips containing infected neurons were fixed in 2% paraformaldehyde and immunostained for viral antigen (mouse antinucleocapsid).

a Supernatants from infected cultures were collected at the indicated times and viral titers were determined on L2 cells. The titers are the mean titers from two independent experiments (n = 6).

b A group of viral antigen-positive cells that appeared to exhibit cell–cell contact was defined as a focus. The data shown represent the results from two independent experiments (n = 4).

c The total number of viral antigen-positive cells per coverslip was determined (n = 4, except on day 3 following infection with S4R, n = 3). On day 3 the mean number of viral antigen-positive cells was greater following infection with S4R than S A59R (two-tailed t test, P < 0.01).
viral titers from day 0 to 1 postinfection. On days 1 and 3, the titers of infectious virus were lower in S4R-infected cultures than S598R-infected cultures.

Differences in the composition of the lymphocytic response to infection with S4R and S598R

The MHV-4 spike appears to mediate more efficient spread in cells of the CNS than the MHV-A59 spike. Previously we observed a difference in the immune response to infection with S4R and S598R by H&E staining. S4R infection induced a greater infiltration of inflammatory cells into the brain than S598R infection (Phillips et al., 1999). The immune response to MHV infection is complex, and multiple components of the immune response are required for optimal clearance of infectious virus from the CNS. Therefore, both quantitative and qualitative differences in the antiviral immune response may also contribute to the differential neurovirulence of S4R and S598R.

We used flow cytometry to quantitatively compare the composition of the immune response in the brain to infection with S4R and S598R. Total brain mononuclear cells were isolated on days 5 and 7 postinfection and were examined for surface expression of CD45 (a marker of nucleated cells of hematopoietic lineage), CD8, CD4, and B220.

At the peak of inflammation, day 7 postinfection, the difference in the total number of inflammatory cells in the brain between S4R and S598R was dramatic. The number of immune cells, as determined by CD45 positivity, isolated from each brain following S4R and S598R infection was between 1.2 x 10^6 and 1.1 x 10^7 cells and was on average fivefold greater following infection with S4R than S598R. Representative data are shown in Fig. 4. Flow analysis of brain-derived lymphocytes revealed that the percentage of CD8+ T cells was also greater following infection with the more neurovirulent S4R as compared with S598R (Fig. 4B). In contrast, the percentage of cells positive for CD4 or B220 (a B cell marker) was similar following infection with S4R and S598R. We next examined the ratio of CD8+ to CD4+ T cells for S4R and S598R from four independent experiments. The mean ratio of CD8+ to CD4+ T cells following infection with S4R, 2.83 ± 0.26 (mean ± SEM), was significantly greater than the ratio following infection with S598R, 1.73 ± 0.06 (mean ± SEM) (two-tailed t test, P < 0.01) (n = 4). Due to the difference in total cells isolated per brain, the estimated numbers of CD45+, CD4+, CD8+, and B220+ cells per mouse brain were between three- and fivefold greater following infection with S4R than with S598R (Fig. 4C). Thus, the high neurovirulence of S4R relative to S598R was correlated with a greater total number of mononuclear cells in the brain, and a significant increase in the percentage of CD8+ T cells.

On day 5 following infection with either S4R or S598R, the magnitude of the inflammatory response to infection in the brain was much less than on day 7 postinfection as previously observed by Haring et al. (2001) and Williamson et al. (1991). In addition, the numbers of CD8+ or CD4+ T cells in the brain were just above the level of detection. Even on day 5 postinfection, however, the number of immune cells in the brain following infection with S4R was still greater than following S598R infection (data not shown).

Functional differences in the T cell response to infection with S4R and S598R

In addition to differences in the magnitude and in the composition of the immune response to infection with S4R and S598R, we hypothesized that there might be differences in the function of MHV-specific T cells. To compare the efficiency of functionally activated, MHV-specific CD8+ and CD4+ T cells in the CNS following infection with S4R or S598R, we performed intracellular IFN-γ assays. Brain-derived T cells on day 7 postinfection were isolated and challenged with peptides containing the immunodominant or sub-immunodominant CD8+ T cell epitope, S510 or S598, or the immunodominant CD4+ T cell epitope, M133, and stained for intracellular IFN-γ as described previously (Murali-Krishna et al., 1998). Representative results are shown in Fig. 5. Following S4R infection, approximately 13.6 and 9.3% of the CD8+ T cell population in the CNS was S510-specific and S598-specific, respectively. S510 and S598 are the only two known CD8+ T cell epitopes for MHV; therefore, the total frequency of MHV-specific CD8+ T cells in the CNS following S4R infection was approximately 22.8%. The MHV-A59 spike lacks the immunodominant S510 epitope (Luytjes et al., 1987). Thus, following infection with S598R the MHV-specific CD8+ T cell response was S598-specific and accounted for approximately 11.8% of the CD8+ T cell response.

Within the population of cells isolated from the CNS, a proportion of CD4+ cells exhibited intracellular IFN-γ in the absence of peptide (Fig. 5B). The reason for this background level of IFN-γ expression is not known; however, it has been suggested that infected microglia and macrophages in the CNS cell preparation may stimulate CD4+ T cells (Haring et al., 2001). The frequency of M133-specific CD4+ T cells following infection with S4R or S598R was similar, accounting for approximately 16.0 and 15.4% of the CD4+ T cell response, respectively. As this may overestimate the frequency of MHV-specific cells in the CNS, we also subtracted the percentage of IFN-γ-expressing cells in the absence of peptide from the total percentage of IFN-γ-expressing cells after peptide stimulation. The calculated frequency of M133-specific CD4+ T cells following infection with S4R, 10.6%, and S598R, 8.9%, was also similar. Thus, infection with either S4R or S598R resulted in a robust CD8+ and CD4+ T cell response.
response that was both epitope-specific and functional. The major difference between the frequency of virus-specific T cells induced by S4R and S A59R was the increased frequency of functionally activated CD8\(^+\) T cells following infection with the more neurovirulent S4R. Note that due to the increased numbers of inflammatory cells in the brain following S4R infection relative to S A59R infection, the total number of virus-specific CD8\(^+\) and CD4\(^+\) T cells in the brain was greater following infection with S4R than S A59R.

Contribution of lymphocytes to differential neurovirulence

S4R infection resulted in a greater number of lymphocytes and a higher frequency of virus-specific immune
cells in the CNS than S_{as9}R infection. To evaluate the contribution of lymphocytes to the differential neurovirulence observed with S_{4}R and S_{as9}R, we compared the survival of infected RAG1^{-/-} mice (B6 background), which lack functional T and B cells (Mombaerts et al., 1992), and immunocompetent B6 mice. RAG1^{-/-} mice infected with S_{as9}R experienced 100% mortality while immunocompetent mice survived following challenge with S_{as9}R (Fig. 6A). Thus, S_{as9}R-infected immunocompetent B6 mice are protected from death by the activated immune response, and without this response the mice succumb to a lethal infection. In contrast, RAG1^{-/-} mice infected with S_{4}R exhibited prolonged survival relative to immunocompetent B6 mice; the median survival time for S_{4}R-infected mice was 13 days in RAG1^{-/-} and 10 days in B6 mice (statistically different, Mann–Whitney rank sum test, \(P < 0.0001\)) (Fig. 6B). Prolonged survival in S_{4}R-infected RAG1^{-/-} mice suggested that T or B cells contributed to pathology.

We examined virus titers in the brain and other organs of RAG1^{-/-} mice. At five days postinfection, the titers of infectious virus in the brains of RAG1^{-/-} mice were similar to those in B6 mice, following S_{as9}R infection 6.0 vs 5.4 and following S_{4}R infection 5.1 vs 5.1 [log_{10}(PFU/g)] (\(n = 2\)), respectively. In addition, similar to the infection in immunocompetent mice the level of infectious virus was minimal in the liver and undetectable in the spleen, kidney, lung, heart, and a portion of the small intestine.

FIG. 5. MHV-specific CD8^{+} and CD4^{+} T cells following infection with S_{4}R and S_{as9}R. Mononuclear cells were isolated from the brain of S_{4}R- and S_{as9}R-infected mice on day 7 postinfection, and the frequency of CD8^{+} and CD4^{+} T cells specific for MHV-specific epitopes were determined by intracellular IFN-\(\gamma\) staining. Cells were cultured with or without peptide as indicated above each column. The numbers in the upper right quadrants indicate the percentage of CD8^{+} T cells (A) or CD4^{+} T cells (B) that are positive for the intracellular IFN-\(\gamma\) stain following infection with either S_{4}R (top row) or S_{as9}R (bottom row). The numbers in the right lower quadrants represent the percentage of CD8^{+} T cells (A) or CD4^{+} T cells (B) that are negative for the intracellular IFN-\(\gamma\) stain. Brain mononuclear cells were pooled from six to eight animals, and these data are representative of two to five experiments.
S4R died by 43 days postinfection. No deaths were observed in B6 mice. Although not shown on the graph, all RAG1/H11002/H11002 mice inoculated with S4R infected and survived without apparent pathology. We therefore conclude that the vanishing virus in the CNS, suggesting that immune-mediated neuropathogenesis using S4R and SA59R, two isoforms of S4R.

Effect of a functional T and B cell response on mortality following intracranial inoculation with S4R (A) or SA59R (B). B6 (open symbols) or RAG1/H11002/H11002 (closed symbols) mice were inoculated with 10 PFU of virus and observed daily for mortality. The percentage survival and median time to death in days (number to right of line) is shown. Following inoculation with SJ7 (B), there was a significant difference in the median time to death between RAG1/H11002/H11002 and B6-infected mice (13 and 10 days, respectively, Mann–Whitney rank sum test, P < 0.0001). Although not shown on the graph, all RAG1/H11002/H11002 mice inoculated with S4R died by 43 days postinfection. No deaths were observed in B6 mice after 19 days postinoculation. The data shown represent the results from two independent experiments from a total of 14 RAG1/H11002/H11002 mice per virus, 11 B6 mice inoculated with SA59R, and 15 B6 mice inoculated with S4R.

Following inoculation with SA59R (A) or S4R (B). B6 (open symbols) or RAG1/H11002/H11002 (closed symbols) mice were inoculated with 10 PFU of virus and observed daily for mortality. The percentage survival and median time to death in days (number to right of line) is shown. Following inoculation with SJ7 (B), there was a significant difference in the median time to death between RAG1/H11002/H11002 and B6-infected mice (13 and 10 days, respectively, Mann–Whitney rank sum test, P < 0.0001). Although not shown on the graph, all RAG1/H11002/H11002 mice inoculated with S4R died by 43 days postinfection. No deaths were observed in B6 mice after 19 days postinoculation. The data shown represent the results from two independent experiments from a total of 14 RAG1/H11002/H11002 mice per virus, 11 B6 mice inoculated with SA59R, and 15 B6 mice inoculated with S4R.

DISCUSSION

The spike glycoprotein is a major determinant of MHV neurovirulence. We examined how alterations in spike affect neuropathogenesis using S4R and SA59R, two isoforms of S4R. Compared to SA59R, S4R infection of neuronal cultures resulted in greater viral antigen spread, however, the tropism of S4R and SA59R for neurons and astrocytes was identical. Thus differences in viral spread, not differences in neuronal tropism per se, contributed to the differential neurovirulence of S4R and SA59R.

Despite the similarity in neuronal tropism, however, the number of infected neurons was greater following infection with S4R than SA59R due to the greater total number of cells infected. To compare the ability of S4R and SA59R to spread in cells from the CNS, we examined the infection of primary hippocampal neuron cultures. S4R infection of neuronal cultures resulted in greater viral spread than SA59R infection, and the pattern of spread suggested that S4R spreads more rapidly from cell to cell than SA59R. Viral spread along neuronal processes could contribute to rapid dissemination of virus to multiple regions of the brain. Future studies will address how the MHV-4 spike confers rapid cell to cell spread, and the effect of infection on neuronal viability.

In addition to increased viral spread, S4R mediated a greater influx of immune cells into the CNS than SA59R. To identify how this increased immune response to infection might influence the differential neurovirulence of S4R and SA59R, we compared the composition and the functional features of the immune response to infection. Our studies suggest that differences in viral spread in multiple cell types, differences in the magnitude of the immune response to infection, and immune-mediated pathology contribute to the differential neurovirulence of S4R and SA59R. We have shown previously (Phillips et al., 1999, 2001) that, during acute infection, the titers of infectious virus in the brains of animals infected with S4R and SA59R are similar. Thus, the spread of viral antigen, neurovirulence, and mortality are not determined by the level of infectious virus in the CNS.

Neuronal infection has been proposed to be a major determinant of MHV neurovirulence (Dubois-Dalcq et al., 1982; Fleming et al., 1986; Knobler et al., 1981). Using double-immunohistochemistry, we quantitatively compared the tropism of S4R and SA59R for neurons and astrocytes. We determined that on days 3 and 5 postinfection, S4R infected a greater total number of cells than SA59R throughout the brain. Despite the difference in viral spread, however, the tropism of S4R and SA59R for neurons and astrocytes was identical. Thus differences in viral spread, not differences in neuronal tropism per se, contributed to the differential neurovirulence of S4R and SA59R.

To identify the parameters of MHV infection mediated by spike and important in neurovirulence, we compared the cellular tropism of S4R and SA59R and both quantitative and functional features of the immune response to infection. Our studies suggest that differences in viral spread in multiple cell types, differences in the magnitude of the immune response to infection, and immune-mediated pathology contribute to the differential neurovirulence of S4R and SA59R. We have shown previously (Phillips et al., 1999, 2001) that, during acute infection, the titers of infectious virus in the brains of animals infected with S4R and SA59R are similar. Thus, the spread of viral antigen, neurovirulence, and mortality are not determined by the level of infectious virus in the CNS.

Neuronal infection has been proposed to be a major determinant of MHV neurovirulence (Dubois-Dalcq et al., 1982; Fleming et al., 1986; Knobler et al., 1981). Using double-immunohistochemistry, we quantitatively compared the tropism of S4R and SA59R for neurons and astrocytes. We determined that on days 3 and 5 postinfection, S4R infected a greater total number of cells than SA59R throughout the brain. Despite the difference in viral spread, however, the tropism of S4R and SA59R for neurons and astrocytes was identical. Thus differences in viral spread, not differences in neuronal tropism per se, contributed to the differential neurovirulence of S4R and SA59R.

Despite the similarity in neuronal tropism, however, the number of infected neurons was greater following infection with S4R than SA59R due to the greater total number of cells infected. To compare the ability of S4R and SA59R to spread in cells from the CNS, we examined the infection of primary hippocampal neuron cultures. S4R infection of neuronal cultures resulted in greater viral spread than SA59R infection, and the pattern of spread suggested that S4R spreads more rapidly from cell to cell than SA59R. Viral spread along neuronal processes could contribute to rapid dissemination of virus to multiple regions of the brain. Future studies will address how the MHV-4 spike confers rapid cell to cell spread, and the effect of infection on neuronal viability.

In addition to increased viral spread, S4R mediated a greater influx of immune cells into the CNS than SA59R. To identify how this increased immune response to infection might influence the differential neurovirulence of S4R and SA59R, we compared the composition and the function of the lymphocytic response to infection. At the peak of inflammation, day 7 after infection, the more neurovirulent S4R induced the infiltration of a greater number of T cells and B cells into the CNS with a specific increase in the percentage of CD8+ T cells as compared to SA59R.

We also compared the virus-specific activity of CD4+ and CD8+ T cells from the CNS of animals infected with S4R and SA59R by measuring intracellular IFN-γ secretion...
in response to specific viral epitopes. Both S₄R and S₅₉₋R induced robust and virus-specific immune responses; however, there were differences in the specificity of the CD₈⁺ T cell response. The two H-2b CD₈⁺ T cell epitopes recognized in B6 mice are located in the spike glycoprotein at residues 510–518 (S₅₁₀) and 598–605 (S₅₉₈) (Bergmann et al., 1996; Castro and Perlman, 1995). A 52-amino-acid deletion in the spike of MHV-A59 relative to the spike of MHV-4 removes the more immunodominant 510–518 epitope (Luujjes et al., 1987; Parker et al., 1989). Consistent with this difference in epitopes, the frequency of S₅₁₀-specific CD₈⁺ T cells in the CNS of S₄R and SA₅₉₋R was similar. The two viruses also shared a similar frequency of response to the dominant CD₄⁺ T cell epitope (Xue and Perlman, 1997), M₁₃₃, located in the viral matrix protein.

Thus, expression of the S₅₁₀ epitope was responsible for the greatest difference in the frequencies of virus-specific, functionally activated T cells following infection with S₄R and S₅₉₋R. S₄R infection induced a greater percentage of CD₈⁺ T cells in the CNS than S₅₉₋R infection. The response to the S₅₁₀ epitope in S₅₉₋R did not appear to induce a significantly greater immune response to S₅₉₈, however, as the frequency of S₅₉₈-specific CD₈⁺ T cells in the CNS following infection with S₄R or S₅₉₋R was similar. The two viruses also shared a similar frequency of response to the dominant CD₄⁺ T cell epitope (Xue and Perlman, 1997), M₁₃₃, located within the viral matrix protein.

To examine the role of the T and B cell response in the differential neurovirulence of S₄R and S₅₉₋R, we compared the virulence of these viruses in the presence and absence of a functional T and B cell response. Despite prolonged replication of infectious S₄R in the CNS of RAG⁻/⁻ mice, these mice exhibited prolonged survival relative to immunocompetent mice. In contrast, the T and B cell response to S₅₉₋R infection was primarily protective. These data suggest that immune-mediated pathology contributes to the high neurovirulence of S₄R. Interestingly, previous studies with MHV have also suggested that, in addition to its role in protection, the inflammatory response, particularly the T cell response, can be immunopathologic. Prolonged survival of perforin-deficient mice infected with a JHM strain of virus suggested that CD₈⁺ CTLs might contribute to disease (Lin et al., 1997). Moreover, CD₄⁺ T cells have also been implicated in immunopathology; both adoptive transfer experiments with CD₄⁺ T cells and infection of CD₄⁺ knockout mice have suggested that CD₄⁺ T cells in the CNS may increase disease morbidity (Lane et al., 2000; Wu et al., 2000). To determine whether CD₈⁺ or CD₄⁺ T cells were primarily responsible for the prolonged survival in RAG⁻/⁻ mice, we compared the survival of CD₈⁻/⁻, CD₄⁻/⁻, and immunocompetent B6 mice infected with S₄R; however, the absence of either T cell subset alone did not confer significantly increased survival (data not shown).

Previous studies have demonstrated the complexity of the immune response to MHV and the requirement for multiple components for clearance (Korner et al., 1991; Stohlman et al., 1986, 1995, 1998; Sussman et al., 1989; Williamson and Stohlman, 1990; Yamaguchi et al., 1991; Bergmann et al., 2001). Our results suggest that MHV-induced immunopathology is also complex and involves multiple components of the immune response to infection. Studies with LCMV and HIV have suggested that virus strains that exhibit rapid viral spread are associated with increased immune-mediated pathology (Wodarz and Krakauer, 2000; Moskophidis et al., 1995). Future studies will delineate which components of the immune response to S₄R contribute to immunopathology and examine how increased viral spread may induce such a response.

Increased neurovirulence conferred by the MHV-4 spike correlates with both rapid viral spread in the CNS and immune-mediated pathology. By studying two isogenic viruses with defined alterations in spike in vivo models of MHV infection, we can begin to dissect the different roles of the spike glycoprotein in neurovirulence. Furthermore, we can examine the contribution of virus- and immune-mediated pathology to neurovirulence.

**MATERIALS AND METHODS**

**Virus and cells**

The recombinant viruses, referred to as S₄R (containing the MHV-4 spike gene) or S₅₉₋R (containing the MHV-A59 spike gene), were generated using targeted RNA recombination and are in the MHV-A59 background (Fischer et al., 1997). We have in the past described independent isolates of both S₄R (S₄R22 and S₄R29) and S₅₉₋R (S₅₉₋R13 [wtR13] and S₅₉₋R16) (Phillips et al., 1999, 2001). Since the group of recombinant viruses expressing the MHV-4 spike gene are phenotypically indistinguishable from each other, they will be referred to collectively as S₄R; similarly, since the group of viruses
expressing the A59 spike gene are phenotypically indistinguishable from each other, they will be referred to collectively as S$_{A59}$R. All viral stocks were grown in 17Cl-1 cells, and infectious virus was quantified by plaque assay, using murine L2 cells as indicator cells.

Inoculation of mice

Four-week old male C57Bl/6 (B6) mice were purchased from the National Cancer Institute (Bethesda, MD), and four-week old male RAG1$^-/-$ mice were purchased from the Jackson Laboratory (Bar Harbor, ME). No mature B or T lymphocytes are produced in RAG1$^-/-$ mice (Mombaerts et al., 1992). All mice were certified MHV-free. Mice were anesthetized with Isoflurane (Abbott Laboratories), and injections were performed in the left cerebral hemisphere. Each injection contained 10 PFU of virus diluted to a volume of 20 μl per mouse in PBS containing 0.75% bovine serum albumin. Mock-infected controls were inoculated similarly but with an uninfected 17Cl-1 cell lysate at a comparable dilution.

Double-label indirect immunofluorescence on brain sections

At various times after infection with S$_R$ or S$_{A59}$R, mice were sacrificed and perfused with 10 ml of PBS, and the brains were removed. The right half of the brain was fixed in Formalin for 2 days, embedded in paraffin, sectioned, and left unstained for immunofluorescence. Adjacent sagittal brain sections close to midline, 10–20 μm apart, were examined. Sections were blocked with 2% goat serum in Tris–HCl and 0.1% Triton X-100 for 1 h, incubated with primary antibodies for 2 h, and incubated with secondary antibodies for 1 h. All incubations were carried out in a humidified chamber at 37°C, and after each step the sections were washed (three times for 5 min each) in PBS. For localization of viral antigen, sections were incubated with a 1:100 dilution of rabbit anti-MHV-A59 serum, made against detergent-disrupted MHV-A59, obtained from Mark Denison (Vanderbilt University, TN). Neurons were identified with a 1:200 dilution of biotin-conjugated horse anti-mouse immunoglobulins (Vector Laboratories). Murine antibodies were detected with a 1:200 dilution of fluorescein avidin D (FITC) for 30 min, purchased from Vector Laboratories.

Control slides were incubated in parallel with preimmune rabbit serum, and sections from mock-infected animals were incubated with rabbit anti-MHV-A59 serum. For each double-immunostaining scheme, it was verified that the secondary antibody did not bind in the absence of the primary antibody or cross-react with primary antibodies of different species.

Isolation of primary hippocampal neurons

Primary hippocampal neurons were obtained from day 14 to day 16 embryonic mice, as described previously (Banker and Goslin, 1991; Pasick et al., 1994; Rall et al., 1997), except the cells were maintained in neurobasal medium (Gibco/BRL) containing 4 μg/ml glutamate, in the absence of an astrocyte feeder layer. The mice used in these experiments were C57Bl/6 transgenic mice expressing the human MV receptor, CD46 (Rall et al., 1997). Cells were plated at a density of 2–7 × 10^5 cells per glass coverslip.

Infection of primary hippocampal neuron cultures and virus detection

On day 4 postexplant, cells were infected at a multiplicity of infection (m.o.i.) of 5 with S$_R$ or S$_{A59}$R or mock-infected with 17Cl-1 cell lysate. After 1 h at 37°C with gentle agitation, the inoculum was removed; conditioned neurobasal medium was added back to the cultures, and the cultures were maintained at 37°C in 5% CO$_2$ until harvesting. In each of two independent experiments three coverslips were infected per virus per time point.

The levels of infectious virus in the supernatants as a function of time following infection were determined by plaque-assay on L2 cell monolayers. For the localization of viral antigen coverslips containing infected cells were washed once with PBS, fixed in 1% paraformaldehyde, and stored at 4°C. Coverslips were blocked with 2% goat serum in Tris–HCl and 0.1% Triton X-100 for 15 min, incubated with primary antibodies for 2 h, and incubated with secondary antibodies for 1 h. Between each step the coverslips were washed in Tris–HCl and 0.1% Triton X-100 six times, and all incubations were carried out at room temperature in a humidified chamber. To detect viral antigen coverslips were incubated with a 1:20 dilution of mouse antinucleocapsid antibody, obtained from Julian L. Leibowitz (Texas A&M University, TX). This was followed by incubation with the secondary antibody, a 1:200 dilution of biotin-conjugated horse anti-mouse immunoglobulins (Vector Laboratories), and then incubation with a 1:200 dilution of fluorescein avidin D (FITC) (Vector Laboratories) for 30 min. MAP2 and GFAP were detected with a 1:300 dilution of rabbit anti-MAP2 obtained from M.-Y. Lee (University of Pennsylvania, PA) and a 1:250 dilution of rabbit anti-cow glial fibrillary acidic protein (GFAP) (Dako Corp.), respectively. Rabbit antibodies were detected with a 1:40 dilution of TRITC-conjugated swine anti-rabbit immunoglobulins (Dako
Isolation of monocytes from the CNS

Monocytes were prepared from the CNS as previously described by Pewe et al. (1999) on days 5 and 7 after intracranial inoculation with 10 PFU of $S_R$ or $S_{ASR}$ or mock-infected with/7CI cell lysate. Two to six brains were pooled per sample. In brief, animals were sacrificed and perfused with 10 mL of PBS, and a single-cell suspension from the brain was obtained by passing cells through a nylon mesh bag (64-μm pore diameter). Percoll (Pharmacia) was added to a final concentration of 30%, and the lysate was centrifuged at 1300 g for 30 min at 4°C. The cell pellet was resuspended and washed. The cells were then layered over 2 mL Lympholyte-M (Cedarlane Laboratories) and centrifuged at 1300 g for 20 min at room temperature. Cells were removed from the interface, washed once, and counted.

Flow analysis

Expression of cell-surface markers was determined by staining cells with monoclonal antibodies specific for CD8α (clone 53–6–7), CD4 (clone RM4-5), CD45 (leukocyte common antigen, clone 30-F11, and B220 (CD45R/B220, clone RA3-6B2) purchased from PharMingen. For flow cytometry approximately $1 \times 10^6$ cells were washed three times with PBS 1% BSA and blocked with 50 μg of rat IgG and a 1:200 dilution of anti-CD16/CD32 (FcγRII/II receptor, clone 2.4G2). Cells were then surface stained for the appropriate markers for 1 h at 4°C, washed three times with PBS 1% BSA, and fixed in 2% paraformaldehyde. Cells were analyzed using a FACSscan flow cytometer (Becton–Dickinson). The total number of cells positive for each marker per mouse was determined by multiplying the fraction of cells positive for the marker by the total number of live cells isolated per brain. As controls, monocytes were incubated with isotype controls and monocytes from the brains of mock-infected mice were incubated with antibodies to cell markers.

Intracellular IFN-γ staining

Intracellular IFN-γ expression in response to peptide stimulation was performed as previously described (Murrali-Krishna et al., 1998). We cultured $1 \times 10^6$ brain-derived monocytes/well for 5 h at 37°C in 200 μl of RPMI 1640 complete supplemented with 5% FCS, 10 units human recombinant IL-2, and 1 μg/ml Brefeldin A (Golgi-stop, PharMingen) either with or without epitope peptides. The peptides were used at a concentration of 0.1 μg/ml. Cells were then stained as above for surface expression of CD8 or CD4 and incubated overnight at 4°C. For intracellular IFN-γ staining cells were then fixed and permeabilized using the Cytotof/Cytoperm kit (PharMingen), and stained with a FITC-conjugated monoclonal rat anti-mouse IFN-γ antibody (clone XMG 1.2, PharMingen).

Survival assays

RAG1−/− and B6 mice were inoculated intracranially with 10 PFU of $S_R$ or $S_{ASR}$. In two independent experiments, a total of 14 RAG1−/− mice were analyzed per virus, and in parallel a total of 11 B6 mice inoculated with $S_{ASR}$ and 15 B6 mice inoculated with $S_R$ were analyzed. Mice were examined for signs of disease or death on a daily basis up to 21 days postinfection, after which they were examined every fourth day, and the percentage survival was calculated.

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