The Immunoevasive Function Encoded by the Mouse Cytomegalovirus Gene m152 Protects the Virus Against T Cell Control In Vivo

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Summary

Cytomegaloviruses encode numerous functions that inhibit antigen presentation in the major histocompatibility complex (MHC) class I pathway in vitro. One example is the mouse cytomegalovirus (MCMV) glycoprotein gp40, encoded by the m152 gene, which selectively retains murine but not human MHC class I complexes in the endoplasmic reticulum–Golgi intermediate compartment/cis-Golgi compartment (Ziegler, H., R. Thäle, P. Lucin, W. Muranyi, T. Flohr, H. Hengel, H. Farrell, and U.H. Koszinowski. 1997. Immunity. 6:57–66). To investigate the in vivo significance of this gene function during MCMV infection of the natural host, we constructed recombinants of MCMV in which the m152 gene was deleted, as were the corresponding virus revertants. We report on the following findings: Deletion of the m152 gene has no effect on virus replication in cell culture, whereas after infection of mice, the m152-deficient virus replicates to significantly lower virus titers. This attenuating effect is lifted by reinsertion of the gene into the mutant. Mutants and revertants grow to the same titer in animals deprived of the function targeted by the viral gene function, namely in mice deficient in β2-microglobulin, mice deficient in the CD8 molecule, and mice depleted of T cells. Upon adoptive transfer of naive lymphocytes into infected mice, the absence of the m152 gene function sensitizes the virus to primary lymphocyte control. These results prove that MHC-reactive functions protect CMVs against attack by CD8+ T lymphocytes in vivo.

Key words: cytomegalovirus • immune evasion • virus mutants • MHC class I • CD8 T lymphocytes

The T cell–mediated immune response is decisive for control and clearance of most viral infections. CD8+ T cells limit virus infections by secretion of cytokines with antiviral activity and by direct cytolysis of infected cells. To achieve effective surveillance and elimination of virus-infected cells, CD8+ T cells need to recognize viral peptides in the context of MHC class I molecules at the surfaces of infected cells for maturation to CTLs or for reactivation from memory (1). Presentation of viral peptides via this pathway requires degradation of viral proteins by the proteasome and the translocation of the peptides into the endoplasmic reticulum (ER) by transporters associated with antigen presentation for loading into the binding groove of nascent MHC class I molecules and subsequent egress of MHC complexes to the cell surface (for review see references 2 and 3).

Certain viruses make use of specific and unique genes to thwart this pathway of virus peptide presentation (for review see reference 4). For example, the adenovirus E3-19K protein binds and arrests MHC class I molecules in the ER, and the herpes simplex virus type I–infected cell protein (ICP)47 inhibits transport of peptides into the ER by competing for the peptide binding site (5–8). Remarkably, CMVs use multiple genes to interfere with the MHC class I pathway of antigen presentation. The human (H)CMV encodes at least four viral polypeptides, each of which can independently and by different molecular mechanisms interfere with MHC class I antigen presentation to inhibit efficient recognition of infected cells by CTLS (9–15). In mouse (M)CMV, at least three genes affect MHC class I molecules (16, 17, 18). The function of the viral proteins is usually of selective specificity for the target proteins of the natural host (16). Detailed studies of the molecular mechanisms by which individual viral polypeptides act at various steps of the antigen presentation pathway in vitro are contrasted by the pau-
city of data concerning the relevance of their function in vivo. Only the adenovirus E3-19K and herpesvirus ICP47 proteins have been investigated in mice so far (19, 20). However, mice do not represent the natural host of these viruses, and, given the species specificity of the functions, such studies might not completely reflect their physiological role for virus control.

Therefore, the question of the biological significance of the individual viral evasion mechanisms should be addressed in the natural host of the virus. MCMV offers the unique opportunity to study the biological impact of herpesviral immune evasion genes in vivo. Comparable to HCMV, MCMV already tightly controls the presentation of viral antigens at early (E) phases of infection (21). The effect is achieved by blocking transport of MHC class I molecules to the cell surface (22). We have identified the m152 gene responsible for inhibition of MHC class I antigen presentation in the E phase of virus gene expression. The MCMV E glycoprotein gp40 encoded by the m152 gene blocks the export of MHC class I complexes from the ER/cis-Golgi compartment and thereby prevents the presentation of viral peptides to CTLs (16). Similar to HCMV, additional MCMV functions exist that also control antigen presentation in the MHC class I pathway (23). Two additional proteins we have identified are the products of the m04 and m06 genes, which form complexes with MHC class I molecules (17, 18).

Here, we constructed mutants of MCMV that lacked the m152 gene as well as the corresponding virus revertants. We investigated the susceptibility of these recombinant viruses to host immune control. We demonstrate that the deletion of the m152 gene results in high susceptibility of the virus to CD8 T cell control. We conclude that even the deletion of a single viral gene from a group of genes that interfere with the MHC class I presentation pathway affects the fitness of CMV in vivo.

Materials and Methods

Cells and Viruses. Mice. NIH 3T3 cells (American Type Culture Collection [ATCC] CRL1658) were grown in DMEM supplemented with 10% newborn calf serum. Primary mouse embryonic fibroblasts (MEFs) prepared from BALB/c; mice and B12 cells (24) were grown in MEM with 10% FCS. The Smith strain of MCMV was grown in MEM with 10% newborn calf serum. Primary mouse embryonic fibroblasts (MEFs) prepared from BALB/c; mice and B12 cells (24) were grown in MEM with 10% FCS. The Smith strain of MCMV was grown in MEM with 10% newborn calf serum.

Construction of Recombination Plasmids and Recombinant Viruses. Plasmid constructions were performed by standard methods (25). Plasmid p152KO was used for generating m152− recombinant viruses. The selection strategy of a 5-kb NotI–BamHI fragment comprising a loxp-flanked lacZ cassette (26) into the XhoI/NheI-hdigested plasmid pEcoO ΔM B (all sites were blunt-ended by treatment with Kl enow DNA polymerase). Plasmid pEcoO ΔM B contains a 5.0-kb EcoRI–MluI fragment of the MCMV genome (MCMV nucleotides 209,756–214,714) encompassing the m152 gene (27). To generate recombinant plasmid pm152gpt, the Escherichia coli gpt gene was flanked with loxp sites and inserted into an XhoI site of plasmid pEcoO ΔM B at the 3′ end of the m152 gene.

Recombinant viruses were generated by homologous recombination in NIH 3T3 as described previously (26). LaZ + recombinants were identified by 5-bromo-4-chloro-3-indolyl-β-d-galactopyranoside (X-gal) staining and isolated by at least five rounds of limiting dilution passage. Recombinant viruses carrying the gpt marker were first enriched by selection in medium that was supplemented with 12.5 μg/ml mycophenolic acid (GIBCO BRL) and 100 μg/ml xanthine (Sigma Chemical Co.) (28) and further purified by limiting dilution. LaZ− and gpt− mutants were generated by a single passage through the recombinase Cre− cell line, N2 (26). LaZ− recombinants were identified as white plaques after X-gal color screening and purified by limiting dilution. Gpt− mutants were selected on STO cells (ATCC CR-L1503) in medium containing 20 μg/ml 6-thioguanine (Sigma Chemical Co.) as described previously (29). To characterize the recombinant virus genomes, viral DNA was isolated from infected cells and analyzed by Southern blot analysis (26).

Characterization of Viral Proteins. B12 cells were infected with wild-type MCMV or m152 recombinant viruses. Cells were pulse labeled at 37°C for 60 min with 500 μCi/ml [35S]methionine (1,200 Ci/mmol; Amersham) in methionine-free MEM supplemented with 5% dialysed FCS and chased in the presence of 10 mM nonlabeled methionine for 2 h. Labeled cells were washed in ice cold PBS and disrupted in lysis buffer (140 mM NaCl, 20 mM Tris-HCl, pH 7.6, 5 mM MgCl2, 1% NP-40, and 1 mM PM SF). Cytoplasmic extracts were precleared by incubation with normal mouse serum, antiactin mAb (Boehringer Mannheim), and protein A-coupled Sepharose (Pharmacia). Immunoprecipitations were performed with anti-K b-sIgM MA-215 ascitic fluid, and immune complexes were retrieved using protein A-coupled Sepharose. Endoglycosidase H (Endo H; Boehringer Mannheim) digestion and SDSPAGE were performed as described previously (23).

Cytolytic Activity. Target cells were labeled for 90 min with Na2 CrO4 and a 4-h standard release assay was performed with 103 target cells and a graded number of effector cells in fivefold dilution steps as described (21, 30). In short, for selective and enhanced expression of immediate-early (IE) genes, MEFs were infected with 0.5 PFU of recombinant viruses or wild-type MCMV per cell by centrifugation (800 g, 30 min). Infection was performed in the presence of cycloheximide (50 μg/ml), which was removed 3 h later by washing with medium containing actinomycin D (5 μg/ml). Limited E gene expression after CH treatment was achieved by removal of cycloheximide using inhibitor-free medium and by adding actinomycin D to the final concentration of 5 μg/ml after 1.5 h. To generate pp89-specific polyclonal CTLs, MCMV-infected spleen cells were restimulated with pp89-derived antigenic peptide (21), and recombinant IL-2 (100 U/ml) was added 5 d later. Cultures were restimulated with gamma-irradiated syngeneic MEFs and by adding actinomycin D at a concentration of 10−8 M. Data represent the mean percentage of specific lysis from three replicate cultures (see Fig. 2 B).

Aminos and Infection Conditons. BALB/c (H-2b haplotype) and C57BL/6 mice (H-2b haplotype) were bred at the Central Animal Facilities at the Medical Faculty, University of Rijeka. Mice were housed under specific pathogen-free conditions in the Whitehead Institute of Biomedical Research, Cambridge, MA. Data represent the mean percentage of specific lysis from three replicate cultures (see Fig. 2 B).
CD8+ T lymphocytes (33). Mice homozygous for the deletion of the gene encoding the CD8 molecule (CD8−−) were obtained from the Centre de Développement des Techniques Avancées pour l’Expérimentation Animale, Institut de Transgenose, Orleans, France. The absence of CD8+ T lymphocytes in βm−− and CD8−− mice was verified by flow cytometry as described previously (34). Neonatal mice, 24 h and 4 d postpartum, were injected intraperitoneally with recombinant viruses or wild-type MCMV. 6–8-wk-old mice were injected either in the posterior footpad or i.p. with 2 × 10^6 PFU of virus in a volume of 50 and 500 μl of diluent, respectively, as described (35).

Detection of Infectious MCMV in Tissues and Statistical Evaluation. Plaque assays were performed in MEF as described previously (36, 37). Statistical significance of differences between the experimental groups was determined by the Mann-Whitney exact rank sum test. Virus titers (x and y) were considered significantly different for P (x versus y) < alpha = 0.05 (one sided), where P is the observed probability value and alpha is a selected significance level.

In Vivo Depletion of Lymphocyte Subsets. In vivo depletion of CD4+ and CD8+ T lymphocyte subsets was performed by intraperitoneal injection of mAbs (rat anti–mouse) to CD4 (YTS 191.1) and/or CD8 (YTS 169.4) molecules (38). Adult and newborn mice received 1 mg and 250 μg of antilymphocyte antibodies, respectively, at the time of injection and every fifth day throughout the experiment. The efficacy of T lymphocyte depletion was >95%, as assessed by cytofluorometric analysis of spleen cells using FITC- or PE-conjugated antibodies directed against mouse CD4 and CD8 molecules (Becton Dickinson; nos. 1333 and 1447).

Adoptive T Lymphocyte Transfer. Donor T lymphocytes were harvested from spleens of uninfected (nonprimed) or latently infected (MCMV-primed) mice. Recipient mice were injected with 2 × 10^6 nylon wool-purified cells were injected intravenously into recipient mice. Mice that did not receive cell transfer were used as negative controls. Mice were killed on day 14 after infection, and tissues were harvested for virus titer determinations.

Results

Generation of Recombinant MCMV. To investigate the significance of the m152 gene product in the course of infection, a targeted deletion of the m152 gene and subsequent reintroduction of this gene into the MCMV genome was performed (Fig. 1 A). The recombinant virus ΔM 69.21 was generated by homologous recombination between the wild-type MCMV genome and the recombinant plasmid p152K O. In this plasmid, a 1.2-kb X ho l-N hel fragment containing the m152 gene was replaced by a loxP-flanked E. coli lacZ gene. The lacZ marker was excised by passing the ΔM 69.21 recombinant through the recombinase Cre+ cell line N 2 (26) to create the m152−lacZ− deletion mutant ΔM 69.24. To generate a revertant virus, the m152 gene, together with the loxP-flanked gpt gene, was reinserted by homologous recombination into the ΔM 69.21 genome. After positive selection (28) of the m152−gpt+ virus rM 69.26, the gpt marker gene was again removed by passing the virus through the recombinase Cre+ cell line N 2 to generate the m152−gpt− revertant virus (designated rM 96.27).

Southern blot analysis of the recombinant virus genomes confirmed the recombination events at the expected positions (Fig. 1 B). In the mutant virus genomes, the original 23.3-kb HindIII E fragment and the 5.9-kb EcoRI O fragment were removed, and the m152 gene product in the course of infection, a targeted deletion of the m152 gene and subsequent reintroduction of this gene into the MCMV genome was performed (Fig. 1 A). The recombinant virus ΔM 69.21 was generated by homologous recombination between the wild-type MCMV genome and the recombinant plasmid p152K O. In this plasmid, a 1.2-kb X ho l-N hel fragment containing the m152 gene was replaced by a loxP-flanked E. coli lacZ gene. The lacZ marker was excised by passing the ΔM 69.21 recombinant through the recombinase Cre+ cell line N 2 (26) to create the m152−lacZ− deletion mutant ΔM 69.24. To generate a revertant virus, the m152 gene, together with the loxP-flanked gpt gene, was reinserted by homologous recombination into the ΔM 69.21 genome. After positive selection (28) of the m152−gpt+ virus rM 69.26, the gpt marker gene was again removed by passing the virus through the recombinase Cre+ cell line N 2 to generate the m152−gpt− revertant virus (designated rM 96.27).

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Figure 1. Characterization of m152 recombinant viruses (A) Genome structure of recombinant viruses. The HindIII cleavage map of the MCMV strain Smith genome is shown at the top, and the expanded HindIII E region of wild-type and recombinant viruses is shown below, with the HindIII (H) and EcoRI (E) cleavage sites indicated. The open box with the arrow depicts the position and orientation of the m152 gene, and the shaded boxes represent viral sequences that were used for homologous recombination. The positions of the loxP sites are indicated by asterisks (*). The marker genes used for selection, lacZ and gpt, are indicated. The probe used for Southern blot analysis is represented by a bar. The expected sizes of the HindIII and EcoRI fragments are indicated by arrows. (B) Southern blot analysis of the recombinant virus genomes. DNA was isolated from infected NIH 3T3 cells and digested with restriction enzymes HindIII and EcoRI, respectively. Sizes of the DNA fragments are indicated in kb.
with CTLs specific for the MCMV antigen pp89 at the indicated E/T ratios in a 4-h 51Cr-release assay.

The altered glycosylation pattern of newly synthesized molecules can be used to locate the export block of nascent MHC class I molecules in MCMV-infected cells. Therefore, the majority of MHC class I molecules from cells infected with wild-type MCMV exhibit high mannose N-linked glycans typical for this compartment that are sensitive to Endo H and migrate faster in gels after digestion with Endo H (Fig. 2 A). In contrast, MHC class I complexes in cells infected with the m152 deletion mutant ΔM C95.21 as well as in uninfected cells acquire Endo H-resistant glycans, indicating the normal egress from the ER GIC/cis-Golgi compartment. As expected, the MHC class I transport was affected again in cells infected with the revertant virus rMC96.27, demonstrated by the reappearance of molecules sensitive to Endo H digestion.

Deletion of the m152 gene restores presentation of viral antigens to CTLs. The transport arrest of MHC class I molecules by the MCMV m152 gene product at early (E) times of virus replication prevents surface expression of these molecules and thus the recognition and lysis of infected cells by specific CTLs (1). A deletion of this gene should restore the recognition of infected cells by CTLs under the experimental conditions. To test this, MEFs were infected with the m152 deletion mutant ΔM C95.21, the revertant virus rMC96.27, or wild-type MCMV. Infected cells were arrested in the IE or E phase of the MCMV replication cycle and used in a CTL assay with MHC class I–restricted CTLs specific for the MCMV antigen pp89 (21, 30, 36). As expected, recognition and cytolysis were equivalent for cells infected with all three viruses during the IE phase of the viral replication cycle, a time at which the m152 gene product is not yet expressed (Fig. 2 B). However, recognition was impaired during the E phase when cells infected with wild-type or revertant virus were used as targets. In contrast, efficient recognition of ΔM C95.21-infected cells was seen, confirming that retention of MHC class I molecules and the associated block in antigen presentation is mediated under these conditions exclusively by the m152 gene.

Replication of m152 deletion mutant and revertant virus in cell culture. Multistep growth curves of recombinant

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**Figure 2.** Functional characterization of the m152 deletion mutants. (A) Normal maturation of newly synthesized MHC class I molecules in cells infected with the m152 deletion mutant. B12 cells were either mock infected or infected with wild-type MCMV or ΔM C95.21 and rMC96.27 recombinants 6 h after infection, cells were pulse labeled for 1 h with [35S]methionine, and newly synthesized molecules were chased for 2 h. *K* MHC class I complexes were precipitated from cell lysates with anti-K*κ* mAb MA-215. Half of the precipitates were digested with Endo H or mock treated before separation by 12.5% SDS-PAGE. The different glycosylation forms of the MHC class I heavy chains with regard to Endo H sensitivity are denoted as r, Endo H resistant; s, Endo H sensitive; or d, Endo H digested. MWM, molecular weight marker. (B) Restoration of MHC class I antigen presentation in cells infected with the m152 deletion mutant. BALB/c MEFs were infected with wild-type (w.t.) MCMV, the m152 deletion mutant ΔM C95.21, and the revertant virus rMC96.27 under conditions that allowed expression of only IE (∆) or IE and E viral proteins (●). Antigen presentation was tested with CTLs specific for the MCMV antigen pp89 at the indicated E/T ratios in a 4-h 51Cr-release assay.
and wild-type viruses served to assess whether the deletion of the m152 gene affects virus growth in cell culture. After infection of NIH 3T3 fibroblasts at a multiplicity of infection of 0.1 PFU per cell, replication of the m152 deletion mutant and revertant were indistinguishable from that of MCMV wild-type virus (Fig. 3). Identical results were obtained by comparing the replication capacity of the m152 deletion mutants, the revertant virus, and MCMV wild type on primary MEFs (not shown), indicating that the m152 gene product is completely dispensable for virus growth in fibroblasts.

Replication of the m152 Deletion Mutant In Vivo. Considering the fact that three different MCMV genes affect nascent MHC molecules and that m152 merely represents the gene that is expressed first, it was not clear whether or not the deletion of this gene would have any detectable impact on the susceptibility of the virus to immune control in vivo. Whereas adult mice control the infection with tissue culture-derived wild-type MCMV effectively, young mice allow virus replication to high titers (39, 40). To detect even minor differences in virulence due to deletion of the single m152 gene, we assayed virus replication in neonatal mice. To avoid the potential influence of marker gene products on the biological properties of mutant viruses, the in vivo experiments were performed mainly with the m152 deletion mutant ΔMC95.24 and the revertant virus rMC96.27, although the other mutants gave comparable results (data not shown). Neonatal mice were injected with 100 PFU of the m152 deletion mutant, the revertant virus, or wild-type MCMV and monitored for 30 d. After infection with wild-type MCMV or revertant virus, 53 and 75%, respectively, of animals succumbed to infection (Fig. 4 A). In contrast, infection with the m152 deletion mutant was survived by the majority of mice (25% mortality). With respect to clinical signs, all three groups of mice exhibited during the first week of infection significant runting and a general failure to thrive compared with mock-infected controls. By 14–20 d after infection, however, most animals that survived the infection with the m152 deletion mutant had recovered. In contrast, clinical signs persisted throughout the course of observation for wild-type MCMV and revertant virus-infected mice. The different disease courses correlated with the body weights of infected mice. On day 26 after infection, the av-
Average body weight of mice that survived infection with ΔmC95.24 was comparable to that of the control group (9.79 ± 1.86 and 10.9 ± 1.16 g, respectively), whereas mice infected with the revertant virus still appeared runted (7.04 ± 1.70 g; data not shown).

To assess whether the differences were due to an altered tissue tropism associated with the m152 deletion, virus titers were determined for lungs, spleen (Fig. 4 B), and salivary glands (data not shown). The mutant ΔmC95.24 yielded lower titers in the spleen and lungs as compared with wild-type MCMV and the revertant virus. Although the differences in virus titers in tissues of neonatal mice did not exceed 1–2 log10 steps, this finding was reproducible both in MCMV-infect mouse strains. In the salivary glands, this observation could not be made. In this organ, the virus titer yielded by the m152 deletion mutant was indistinguishable from that of the wild-type and revertant virus. In this context, it is of interest to note that we have demonstrated earlier that the salivary gland represents the only organ in which MCMV replication is exempt from CD8+ T cell control (41). Altogether, the lack of the m152 gene results in an attenuated course of infection and in restricted virus growth.

The attenuation of the m152 deletion mutant is a function of T cells. Immunodeficient mice were used to assess whether the attenuated phenotype of the m152 deletion mutant indeed reflected an enhanced sensitivity to T cell control. BALB/c mice were immunodepleted by gamma irradiation and by injection with cytolytic antibodies to T lymphocytes and NK cells. In immunodepleted animals, all three viruses replicated to high titers without significant titer differences (data not shown). This demonstrated already that the attenuated phenotype of the m152 deletion mutant is caused by an increased sensitivity to immune control mechanisms. The m152 deletion mutant replicates to lower virus titers than the revertant virus (Fig. 5 A, left panels; P < 0.005) in undepleted BALB/c as well as C57BL/6 mice (Fig. 5 A, top and bottom panels, respectively). This growth restriction was abrogated after depletion of T lymphocyte subsets (Fig. 5 A, right panels), indicating that the attenuated phenotype of the deletion mutant is caused by an enhanced sensitivity to T cell control.

B cell–deficient (μMT−/−) mice were employed to identify the relative role of T cell subsets (Fig. 5 B). Due to the lack of specific antibodies, MCMV spreads rapidly in μMT−/− mice, and detection of infectious virus is facilitated (35). 8-wk-old μMT−/− mice were depleted of only CD8+ T lymphocytes, depleted of both CD4+ and CD8+ T cell subsets, or left undepleted. Virus titers were determined 10 d after infection. The growth restriction of the m152 deletion mutant was notable in particular in the lungs of nondepleted mice, resulting in titer differences ranging from 2 to 3 log10 (Fig. 5 B, left panels). After depletion of CD8+ T lymphocytes or of both T cell subsets, mutant and revertant virus reached comparably virus titers (Fig. 5 B, center and right panels). These data demonstrate that CD8+ T cells are the relevant cell subset responsible for the replication inhibition associated with the m152 gene deficiency. Furthermore, the attenuating effect is also seen in adult mice. Although the differences are not significant, in BALB/c μMT−/− mice, the m152 deletion mutant reached slightly lower titers in the spleen and lungs as compared with wild-type and revertant virus (Fig. 5 C, top and bottom panels). These data suggest that the attenuation is also caused by an enhanced sensitivity to T cell control in the absence of B cell–deficient mice, which are known to have an imbalanced T cell population (41).

Figure 5. Attenuation of the m152 deletion mutant is T cell dependent. (A) 4-d-old BALB/c and C57BL/6 mice were depleted of CD4+ and CD8+ T lymphocytes or were left untreated, and they were then infected with 1,000 PFU i.p. of the ΔmC95.24 (○) or rMC96.27 viruses (●). 10 d after infection, virus titers were determined. Titers of individual mice (circles) and median values (horizontal bars) are shown. There was a significant difference in virus titers between ΔmC95.24 and rMC96.27 in both mouse strains (P < 0.005; left panels). Depletion of T cells abrogated the difference (right panels). (B) 8-wk-old B cell–deficient mice (μMT−/−, BALB/c background) were depleted of CD8+ T cells or both CD4+ and CD8+ T lymphocytes, or they were left untreated. Mice were infected with 2 × 103 PFU i.p. of the ΔmC95.24 and rMC96.27 viruses, and virus titers were determined 10 d after infection. Titers in individual animals and median values (horizontal bars) are shown. The differences in virus titers between the groups of nondepleted mice infected with ΔmC95.24 and rMC96.27 were significant (P < 0.005) for titers in lungs and spleens (left panels). Depletion of both T cell subsets abrogated the differences between the two recombinants in both organs tested (right panels). Depletion of only CD8+ T cells reduced but did not abolish the differences between the two viruses (P < 0.05; center panels). DL, detection limit.
lower titers than the revertant, even after depletion of T cell subsets.

No Phenotype of the m152 Deletion Mutant in MHC Class I-Deficient and CD8− T Cell-Deficient Mice. The m152 gene function affects antigen presentation in the MHC class I pathway. Therefore, in mice in which this presentation pathway is defective, the specific defect of the virus should be phenotypically complemented. To test this, we used MHC class I-deficient C57BL/6 and mice deficient for CD8− T lymphocytes due to the deletion of the CD8 gene (C57BL/6 CD8−/− mice). 4-d-old mice were infected with 1,000 PFU of either the m152 deletion mutant or the revertant virus. In contrast to the situation in immunocompetent mice, no difference in the titers between the two viruses was found in three replicate experiments performed in β2m−/− and CD8−/− mice. One representative experiment is shown in Fig. 6. Essentially, the same message was obtained in adult CD8−/− mice infected with the m152 deletion mutant or the revertant virus. However, adult mice of the C57BL/6 strain cleared both viruses so efficiently that the titers in tissues were below the threshold levels when assayed 10 d after infection. Therefore, to enhance the virus replication and get to a measurable virus load in tissues, we had to deplete NK cells in vivo (data not shown). Altogether, these experiments show that attenuation of the m152 deletion mutant is directly linked to functions required for antigen presentation and recognition in the MHC class I pathway.

Increased Susceptibility of the m152 Deletion Mutant to Naive T Lymphocytes. Adoptive cell transfer into immunodepleted recipients was used to determine the sensitivity of MCMV N2 virus-primed as well as to naive lymphocytes. Lethal MCMV infection in gamma-irradiated BALB/c mice is therapeutically prevented by adoptive transfer of as few as 105 MCMV-primed CD8+ T cells, whereas the same number of naive lymphocytes or primed CD4+ T cells is ineffective (37, 41, 42). As the product of the m152 gene down-regulates presentation of viral antigens in the MHC class I pathway, we expected to see an increased sensitivity of the mutant to primed T cells and perhaps also a more effective priming of T lymphocytes.

To test this, 2 × 105 lymphocytes derived from BALB/c mice, either MCMV primed or naive, were intravenously transferred into syngeneic gamma-irradiated recipients 12 h after infection with wild-type MCMV, the m152 deletion mutant, or the revertant virus strain. Adoptive T cell control of MCMV is a selective function of CD8+ T cells but not of CD4+ T cells and is more effective in spleen and liver than in the lungs (37, 42). Accordingly, the replication of the m152 deletion mutant is more efficiently controlled in these organs than the revertant virus.

Small numbers (~105) of naive T lymphocytes fail to protect mice against MCMV infection (41). This was reproduced for mice infected with the revertant virus; however, the number of 2 × 105 naive lymphocytes already decreased the titers of the m152 deletion mutant (Fig. 7). This is a function of T lymphocytes, as depletion of the CD8+ T cell subset eliminated this activity (data not shown). Transfer of graded numbers of naive cells into gamma-irradiated mice showed that the number of naive T cells had to be increased by 100-fold to achieve an effect on wild-type MCMV comparable to the effect on revertant MCMV.

![Figure 6](image1)

**Figure 6.** No growth difference of m152 deletion and revertant viruses in β2m−/− and CD8−/− mice. Normal C57BL/6, β2m−/−, and CD8−/− mice (all 4 d old) were inoculated with 1,000 PFU i.p. of the ΔMC 95.24 (open bars) or MC 96.27 (shaded bars) recombinant viruses. Shown are virus titers in lungs 10 d after infection. Data represent the mean value of five mice. There was a significant difference in virus titers between ΔMC 95.24 and MC 96.27 viruses (P < 0.005). The titer difference between the two viruses in β2m−/− and CD8−/− mice is not significant.

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![Figure 7](image2)

**Figure 7.** Susceptibility of the m152 deletion mutant to MCMV-primed and naive T lymphocytes. 8-wk-old BALB/c gamma-irradiated mice were injected with 105 PFU of ΔMC 95.24 or MC 96.27 virus. 2 × 105 T lymphocytes were obtained from latently infected or uninfected BALB/c mice, and cells were transferred intravenously into recipients immediately after infection. Mice that did not receive cell transfer were used as negative controls. Shown are titers in individual recipients, measured 13 d after transfer and infection. Horizontal bars indicate the median values. DL, detection limit.
fulfilled by the cation in immunocompetent animals. This prediction was able for virus growth in fibroblasts but should restrict replication of class I molecules (16). If this was the major function of the gp40, which arrests the export of nascent mouse but not human MHC class I molecules (16). If this was the major function of the protein, then the deletion of the gene should be dispensable for virus growth in fibroblasts but should restrict replication in immunocompetent animals. This prediction was fulfilled by the m152 deletion mutant virus. Virus growth in vivo but not in fibroblasts was affected by the mutation. Furthermore, the MHC class I complex transport and the capacity to present viral peptides to CD8 T lymphocytes was restored.

The revertant virus regained wild-type properties in vivo and fulfilled the second requirement by proving the causal linkage between targeted deletion and biological phenotype. As with HCMV infection in humans, the primary infection of mice even with wild-type MCMV is usually asymptomatic. Newborn mice and mice that are a few days old are much more sensitive than adult mice to tissue culture-grown virus, due to the immaturity of the NK cell response (44). In neonates, the infection with 10^4 PFU causes a high percentage of mortality and running in survivors. The attenuating effect of the m152 gene deletion resulted in a higher number of survivors and an earlier cessation of running.

The third requirement was also fulfilled: loss of the phenotypic difference between deletion mutant and revertant virus in the absence of the host immune function affected by the viral gene product. gp40 blocks the export of nascent MHC class I molecules already loaded with viral peptides. The predictable consequence is the inhibition of CD8 T cell priming and CD8 T cell effector function. Loss of the m152 gene should lead to an increased sensitivity of the virus to lymphocytes. Indeed, the virus mutant grew to smaller titers in the various tissues tested. This attenuation did reflect a more stringent control of the deletion mutant by T cell functions, as elimination of T cells resulted in comparable tissue titers of mutant and revertant virus. Furthermore, the attenuating effect of the m152 deletion mutant was absent in C57BL/6 mice that failed to form the functional MHC class I molecules due to the lack of β2-microglobulin expression and also in mice that have a defect in the maturation of MHC class I-restricted CD8^+ T cells due to the deletion of the CD8 gene. Altogether, this study proves for the first time that in their natural host, herpesviruses benefit from functions that inhibit antigen presentation in the MHC class I pathway in vivo.

It remains open whether the observed function is the only function of the m152 gene product in vivo. MHC class I molecules activate CD8^+ T cells and, at the same time, inhibit NK cells (45, 46). Accordingly, a prediction of the transport block of MHC class I molecules due to m152 gene expression is the susceptibility of MCMV-infected cells for NK cell-mediated destruction in vivo. A deletion of the m152 gene and the restoration of MHC class I molecule transport should result in an enhanced resistance of infected cells to NK cell control in vivo. Our data do not support this assumption. Preliminary studies suggest that the lack of the m152 gene certainly does not make the virus more resistant to control by NK cells (Krmpotic, A., B. Polic, and S. Jonjic, unpublished data). Both HCMV and MCMV genes code for glycoproteins that show homology to MHC class I molecules, UL18 in HCMV (47) and m144 in MCMV (27, 48). It has been hypothesized that these viral MHC class I homologues are capable of engaging NK cell inhibitory receptors to protect cells from lysis due to the downregulation of MHC class I expression. Attenuation of MCMV harboring a deletion in the m144 gene has been explained by enhanced control by NK cells in vivo (48). However, a more recent study on UL18 functions failed to confirm the inhibitory function of viral MHC class I homologues on NK cells (49). Therefore, the potential interaction of m152 with m144 needs to be addressed.

Another explanation is that the remaining functions of the genes m04 and m06 fully complement the expected NK cell effect of m152. The genes m04 and m06 have an effect on MHC class I molecules. Both genes are expressed later than m152 during the MCMV replication cycle, and both
genes encode glycoproteins that bind tightly to MHC class I molecules. gp34, encoded by the m04 gene, forms a complex with MHC class I molecules that can be detected on the surfaces of infected cells, but the functional consequence is not yet clear and remains to be tested. We are in the process of constructing double and triple deletion mutants to determine the individual contribution of each of the MHC class I–reactive genes and MHC class I homologues in immune evasion. To this end, we have recently pioneered the cloning of infectious herpesvirus genomes and have developed targeted and random mutagenesis techniques (50, 51).

Our results show for the first time that genes that inhibit antigen presentation in the MHC class I pathway provide a significant growth advantage for CMV during primary infection. What is the potential benefit for the virus? The conditions of primary infection define the load of latent viral genomes and the risk of recurrence of the CMV infection (39). Accordingly, we predict that the m152 gene allows a higher number of MCMV genomes to establish a latent infection, thereby enhancing the chance for reactivation and transmission to the next host and thus escaping extinction.

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