Serp2, an Inhibitor of the Interleukin-1β-Converting Enzyme, Is Critical in the Pathobiology of Myxoma Virus

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Recently, myxoma virus was shown to encode an additional member of the serpin superfamily. The viral gene, called serp2, was cloned, and the Serp2 protein was shown to specifically bind to interleukin-1β (IL-1β)-converting enzyme (ICE), thus inhibiting the cleavage of pro-IL-1β by the protease (F. Petit, S. Bertagnoli, J. Gelfi, F. Fassy, C. Boucraut-Baralon, and A. Milon, J. Virol. 70:5860–5866, 1996). Here, we address the role of Serp2 in the development of myxomatosis, a lethal infectious disease of the European rabbit. A Serp2 mutant myxoma virus was constructed by disruption of the single-copy serp2 gene and insertion of the Escherichia coli gpt gene serving as the selectable marker. A revertant virus was obtained by replacing the E. coli gpt gene by the intact serp2 open reading frame. The Serp2 mutant virus replicated with wild-type kinetics both in rabbit fibroblasts and a rabbit CD4+ T-cell line (RL5). Moderate reduction of cell surface levels of major histocompatibility complex I was observed after infection with wild-type or Serp2− mutant myxoma virus, and both produced white pocks on the chorioallantoic membrane of the chick embryo. After the infection of European rabbits, the Serp2− mutant virus proved to be highly attenuated compared to wild-type myxoma virus, as demonstrated by the clinical course of myxomatosis and the survival rates of infected animals. Pathohistological examinations revealed that infection with wild-type myxoma virus resulted in a blockade of the inflammatory response at the vascular level. In contrast, rapid inflammatory reactions occurred upon infection with the Serp2− mutant virus. Furthermore, lymphocytes in lymph nodes derived from animals inoculated with Serp2 mutant virus were shown to rapidly undergo apoptosis. We postulate that the virulence of myxoma virus in the European rabbit can be partially attributed to an impairment of host inflammatory processes and to the prevention of apoptosis in lymphocytes. The weakening of host defense is directly linked to serp2 gene function and is likely to involve the inhibition of IL-1β-converting-enzyme-dependent pathways.

Virus survival within immunocompetent hosts requires multiple defensive strategies to evade antiviral and inflammatory responses. Poxviruses, which are among the largest animal viruses, have developed specific and efficient strategies, including interference with cytokines and growth factors, inhibition of the complement cascade, reduction of inflammation, and repression of cellular immune recognition, to effectively propagate within the infected host (3, 38, 47, 57, 60).

The genome of poxviruses is a linear large, double-stranded DNA molecule, encoding all the enzymes required for replication and transcription of its DNA, in addition to virulence factors. Whereas the essential genes are located in the central part of the genome, the genes responsible for virulence and host-range (usually not essential) mostly map near the termini (17, 65, 66). Some viral proteins help circumvent the host immune response, usually by mimicking cytokines or cytokine receptors. Poxviruses produce homologues to tumor necrosis factor (TNF) receptor (27, 56), interleukin-1β (IL-1β) receptor (2, 59), gamma interferon (IFN-γ) receptor (4, 5, 61, 69), and chemokine inhibitors (25, 55). Poxviruses are also able to block the production of some important cytokines by inhibiting the enzymes required for their processing. This is the case for the Orthopoxvirus genus members, for which SPI-2 protein (also known as CrmA) is an inhibitor of the IL-1β-converting enzyme (ICE) (33, 48, 50, 58, 64).

Previously, we described the cloning and characterization of Serp2, a new myxoma virus-encoded serpin protein closely related to CrmA (46). Myxoma virus (MV), a member of the genus Leporipoxvirus, is responsible for myxomatosis, a disease fatal to the European rabbit (Oryctolagus cuniculus). After an incubation period of a few days following infection, the primary site of intradermal inoculation evolves as a lesion characterized by tissue degeneration and necrosis. Viral dissemination leads to a generalization of the symptoms in the skin, head, and genital region, together with the development of gram-negative infections of the nasal and conjunctival mucosae (20, 37). The disease is characterized by general dysfunction of cellular immunity and multiple interruptions of the host cytokine network, with death as the most common outcome due to extreme weakness and secondary respiratory tract infections.

A number of MV proteins have been described to function as virulence factors, including M-T1 (25), M-T5 (39), M-T7 (40), M-T2 (69), SERP-1 (34, 67), M11L (45), and myxoma growth factor (45); most of these are proteins that interfere directly with effectors of the host immune system. The importance of Serp2 to myxoma virus replication in vitro and in vivo has not been investigated. We report here the characterization of Serp2 as another critical virulence factor of MV. Disruption of the Serp2 open reading frame resulted in an MV mutant that replicated normally in vitro and in vivo but was highly attenuated upon infection of European rabbits. Loss of MV virulence was correlated with marked increase of inflammatory...
and apoptotic responses in animals inoculated with Serp2− mutant virus. These in vivo findings are well explained by the results from our previous in vitro experiments suggesting that Serp2 can specifically inhibit ICE.

MATERIALS AND METHODS

Cells and viruses. The MV strains T1 and Lausanne, the Serp2− mutant virus, the Shope fibroma virus (SFV), the vaccinia virus MVA, and the recombinant vaccinia virus MVA-HIV-nef were grown in the rabbit kidney cell line RK13 maintained in Dulbecco's minimum essential medium supplemented with 10% fetal calf serum. The revertant MV-Serp2 rev was selected in HGPSRT HeLa cells in which the rabbit CD4 glycoprotein and lymphocytes, T cells, were maintained in RPMI 1640 (Gibco BRL) supplemented with 10% fetal calf serum.

Construction of MV-Serp2− mutant and revertant viruses. The MV serp2 gene was cloned into the Bluescript phagemid expression vector as described previously (46). DNA of this plasmid was used as template to PCR amplify the DNA fragments MV-serp2L and MV-serp2R. The following primers were used: 5′-serp2  AAG GCT GCT GCA TTT TCT CTC CCG CAG-3′; 5′-serp2  AGG CAG GGC ACC GCC CTT CCT CAC GTC CAG C-3′; 3′-serp2  5′-ClaIIATGCATGTTCAAGTGTGAGTGG-3′; 3′-serp2  5′-SacIIATGGAAACTTGAGCTGACGCTTCGCCGATGGACTGTATAACGCT-3′; and 5′-SacIIATGGAAACTTGAGCTGACGCTTCGCCGATGGACTGTATAACGCT-3′. Amplified MV-serp2L and MV-serp2R DNA was digested with XhoI and ClaI, respectively, inserted into pBluescript II SK(+) containing the Escherichia coli guanosine 5′-triphosphorylsf-0 transactivator (gpt) gene under the control of the vaccinia virus early-late promoter 7.5K (18). The resulting plasmid, called pserp2gpt, was used for transfection into MV-infected cells. MV-Serp2− mutant virus was isolated selecting for resistance against mycophenolic acid (19).

An MV-Serp2 rev revertant virus containing a wild-type SERP2 open reading frame was constructed by transfecting plasmid DNA containing the complete serp2 gene into MV-Serp2−infected RK13 cells and by using reverse ge-selection on HeLa cells (28) to isolate revertants. PCR and Southern blot analysis of viral DNA was used to confirm the disruption of the Serp2 open reading frame by insertion of the p7.5 gpt expression cassette, the absence of detectable wild-type virus in preparations of MV-Serp2− revertant virus, and the restoration of the serp2 gene in MV-Serp2 rev revertant virus. Immunoprecipitation of labeled proteins with anti-Serp2 antibodies was used to confirm the presence of a 34-kDa protein in extracts of cells infected with the wild-type or the MV-Serp2 rev revertant viruses, as well as the absence of any specific protein in extracts from MV-Serp2− mutant virus-infected cells.

Single-step growth analysis in cell culture. RBL5 cells (5 × 10⁴) infected with MV strain T1 or MV-Serp2− at a multiplicity of infection of 5 for 2 h. Unadsorbed free virus was removed, cells were washed with serum-free medium three times, growth medium was added, and cells were incubated at 37°C. Cultures were harvested at multiple points in the infection cycle, and virus was released by freeze-thawing and brief sonication. Virus titers in these lysates were determined by plaque titration on RK13 cells monolayers. Single-step growth experiments were performed in triplicate.

Apoptosis assays. To determine whether Serp2 could inhibit TNF-α-mediated apoptosis, 2 × 10⁶ HeLa cells in microtiter plates were infected with wild-type or MV-Serp2− at a multiplicity of infection of 100 in RPMI containing 10% BrdU (bromodeoxyuridine). At 15 h postinfection, the cells were rinsed twice and fresh medium supplemented with serum was added. TNF-α (10 ng/ml; Boehringer-Mannheim) and cycloheximide (40 μg/ml; Sigma) were added, and apoptosis was assessed 8 h later by cellular DNA fragmentation enzyme-linked immunosorbent assay (ELISA) (Boehringer-Mannheim) according to the recommendations of the manufacturer. The cells were harvested by the addition of lysis buffer, which leads to a release of fragmented DNA from the cytoplasm to the supernatant. After an incubation of 30 min at room temperature, followed by a centrifugation at 250 × g for 10 min, the supernatants were transferred to an ELISA plate precoated with anti-DNA antibodies, and the amount of BrdU present in each sample was determined by using anti-BrdU peroxidase conjugate solution and its substrate with a spectrophotometer at 450 nm. Mock-infected cells, in the presence of TNF-α and cycloheximide, were used as a positive control, and cells cultured for 8 h with cycloheximide in the absence of TNF-α were used as a negative control.

Antibody binding and flow cytometry analysis. RBL5 cells (2 × 10⁶) were infected with MV strain T1, MV-Serp2−, MV strain Lausanne, SFV, vaccinia virus MVA, or recombinant vaccinia virus MVA-HIV-nef at a multiplicity of infection of 5 for 2 h. Unadsorbed virus was removed, and the cells were washed with serum-free medium three times and then with medium supplemented with serum; the cultures were then harvested at 24 h postinfection. Cells were rinsed twice in RPMI supplemented with 1% serum and resuspended in 100 µl RPMI or binding buffer (137 mM NaCl, 12 mM NaHCO₃, 2.6 mM KCl, 2 mM MgCl₂, 1 mM CaCl₂, 0.5 mM Na₂-EDTA, pH 7.4) (11). CD4 monoclonal antibodies (Spring Valley Laboratories) were added at a concentration of 25 µg/ml to the cells in RPMI, and class I major histocompatibility complex (MHC) monoclonal antibodies (Spring Valley Laboratories) were added at a concentration of 25 µg/ml to the cells in binding buffer. After a 20 min incubation at 4°C, the cells were washed in RPMI before addition of fluorescein isothiocyanate-conjugated goat anti-mouse antibody. The mixture was incubated for a further 20 min at 4°C. The cells were then rinsed twice in RPMI and resuspended in phosphate-buffered saline for analysis on a fluorescence-activated cell sorter Calibur flow cytometer (Becton Dickinson). Data were acquired from 20,000 cells and analyzed with CellQuest software. Experiments using isotypic immunoglobulin G1 were used as negative controls.

Infection of rabbits with MV-Serp2− mutant virus. Eight-week-old male New Zealand White rabbits (Oryctolagus cuniculus) were obtained from a local supplier and housed in biocontainment facilities under the guidelines of the European Community Council on Animal Care. Infections were performed intradermally in the right ear with 5 × 10⁶ PFU of virus per animal. Rabbits were bled from the ear vein by retroorbital punction at 6, 10, and 16 weeks after infection. At 4, 8, and 11 days postinoculation, two animals from each group were sacrificed. Two mock-infected rabbits were sacrificed and used as controls.

Histological examination. All animals were subjected to a complete postmortem examination. Tissue material from the injection site (ear; primary site) and ocular conjunctiva, parotid lymph nodes, spleen, lungs, and tests were taken and stored in 10% neutral formalin for further analysis. After fixation, tissues were processed routinely into paraffin blocks, sectioned at 4 μm, and stained with hematoxylin and eosin for microscopic examination. Histologic lesions were assessed and graded as follows: +, minimal; ±, light; +, +, moderate; +, +, +, marked; and +, +, +, severe. The TUNEL method was used to assess apoptosis of lymphocytes in parotid lymph nodes and spleen. For this reaction, two thymuses of young mice treated with corticoids were used as positive controls. In conventional histologic sections, apoptosis was detected with a standard avidin-biotinylated horseradish peroxidase-conjugated antibody complex as described previously (39). Next, a solution of fluorescein isothiocyanate-conjugated goat anti-mouse IgG1 was applied (1:50 dilution) to all sections. After rinsing twice in phosphate-buffered saline, any remaining unbound antibody was blocked by incubating the slides with 1% bovine serum albumin in phosphate-buffered saline. To determine the number of apoptosis-positive cells, slides were counterstained with hematoxylin and eosin for microscopic examination. Histologic lesions were assessed, and a grading scheme based on the number of apoptosis-positive cells was used: 0, zero; 1, 1 to 10 apoptosis-positive cells; 2, 11 to 50 apoptosis-positive cells; 3, 51 to 100 apoptosis-positive cells; 4, 101 to 200 apoptosis-positive cells; 5, 201 to 500 apoptosis-positive cells; 6, 501 to 1,000 apoptosis-positive cells; 7, more than 1,000 apoptosis-positive cells.

Immunostaining of histologic sections. Viral antigens in paraffin-embedded sections of ear (primary site) and parotid lymph node were reactivated with 0.1% trypsin in phosphate-buffered saline (pH 7.6) for 30 min at 37°C. After neutralization of endogenous peroxidase and incubation with the secondary primary antibody (rabbit hyperimmune serum anti-MV), the samples were incubated in goat preimmune serum. This step was performed in order to minimize the background generated by the secondary antibody (biotinylated goat anti-rabbit immunoglobulin G) provided by Dako (kit K492). Horseradish peroxidase-conjugated streptavidin was added and visualized with 3,3′-diaminobenzidine tetrahydrochloride (DAB), which formed a brown precipitate. Nuclei were counterstained with Mayer’s hematoxylin.

RESULTS

Construction and characterization of MV-Serp2− mutant and revertant. The Serp2 open reading frame is present as a single copy gene in the MV genome (46) located within the EcoRI-Fl restriction fragment. To construct MV mutant virus, we disrupted the Serp2 open reading frame by targeting insertion of the EcoR1 marker gene under the control of the vaccinia virus p7.5 promoter by homologous recombination precisely to a site within the Serp2 coding sequence, resulting in a 30-bp deletion in the coding sequence. Using a standard dominant selection method (12, 19, 68), we isolated an MV-Serp2− mutant virus able to replicate in the presence of mycophenolic acid. As a control we constructed a revertant virus from MV-Serp2− in which the complete Serp2 open reading frame was restored and which was referred to as MV-Serp2 rev.

To ensure correct disruption of the serp2 gene and its authentic restoration in MV-Serp2 rev, viral DNA was analyzed by PCR with the specific primers 5′serp2-1 and 3′serp2-2. Agarose gel electrophoresis of DNA amplified from wild-type and MV-Serp2 rev-specific template DNA revealed fragments corresponding in size (1,130 bp) to the expected molecular weight of the complete Serp2 coding sequence (data not shown). In contrast, a DNA fragment of 2,850 bp could be amplified from MV-Serp2− DNA, which indicated the presence of additional genomic sequence due...
to the integration of the gpt expression cassette. Furthermore, this analysis confirmed the genetic purity of MV-Serp2- mutant virus as shown by the failure to amplify a detectable DNA band corresponding in size to the wild-type Serp2 coding sequence. Additionally, Southern blot analysis of viral DNA revealed characteristic restriction patterns for the wild-type, MV-Serp2- and MV-Serp2 rev genomes (data not shown). Since only 30 bp of the serp2 open reading frame were missing, we wanted to make sure that the protein could not be produced, even as a truncated form. Immunoprecipitation with a specific anti-Serp2 antiserum (46) revealed a polypeptide of 34 kDa, corresponding to Serp2, in extracts from RK13 cells infected by either the wild-type or the MV-Serp2 rev virus, but no specific band could be visualized on extracts from cells infected with the MV-Serp2- mutant (data not shown).

Comparison of pocks formed on the chorioallantoic membrane. The chorioallantoic membrane (CAM) of chick embryos has been widely used to study the acute inflammatory response to poxvirus infection. In that system the wild-type cowpox virus produces hemorrhages, whereas a mutant lacking the cmt4 gene produces white pocks characterized by an influx of inflammatory cells (22, 48). To compare the lesions produced on CAM, 9-day-old embryonated eggs were infected with 10⁶ PFU of wild-type MV or MV-Serp2- mutant. After 5 days of incubation at 33°C, an examination of the CAM revealed small white pocks (ca. 1 mm in diameter) which were present on CAM infected with either virus. No hemorrhage was detected. Furthermore, the pocks formed by both viruses did not differ in number or other morphology.

MV-Serp2- mutant efficiently replicates in rabbit kidney cell and CD4+ T-cell lines. No defects in the ability of MV-Serp2- to replicate in cultured rabbit RK13 fibroblasts in vitro were noted in a single-step growth curve analysis. Similar results were also found in a cultured rabbit RL5 CD4+ T-cell line (data not shown).

MV-Serp2- mutant fails to induce apoptosis upon cell culture infection. Since we have demonstrated previously that Serp2 exhibits ICE inhibiting activity and that the ICE family, recently named the caspases, plays a central role in the execution of apoptosis, we were interested in investigating apoptotic cell death after infection with MV-Serp2- mutant virus.

Since MV-Serp2- could replicate with wild-type kinetics in RL5, these cells were not expected to be a good model for investigating an antiapoptotic role for Serp2. Indeed, when using DNA fragmentation ELISA, we found no apoptosis in RL5 cells infected with either wild-type or MV-Serp2- MV.

MV-Serp2- mutant virus inhibits TNF-α-mediated apoptosis. In a similar way we also studied HeLa cells, which are susceptible to TNF-α-mediated apoptosis. We measured the cellular DNA fragmentation (ELISA test) after infection with wild-type MV or MV-Serp2- and after treatment with TNF-α and cycloheximide. Mock-infected cells, in the absence of TNF-α, showed no sign of apoptosis. Incubation with TNF-α and cycloheximide induced apoptosis, as measured 8 h after treatment. There was no significative apoptosis in cells infected with wild-type MV. The results were the same with cells infected with MV-Serp2- indicating that Serp2 is not necessary for the inhibition of TNF-induced apoptosis in HeLa cells (data not shown).

Effect of Serp2 on the level of cell surface antigens. Since it has been shown that there is a decrease of the class I MHC surface expression at 24 h after infection with MV strain Lau sanne (11), we investigated whether disruption of the Serp2 open reading frame could influence the expression of these antigens at the cell surface. Having checked that mock-infected RL5 cells showed strongly positive surface staining with mono-}

clonal antibody against the class I MHC, we saw no striking difference in the downregulation of MHC I on the surface of RL5 cells infected with wild-type MV strain T1 or with MV-Serp2- mutant virus (data not shown). These results suggest that Serp2 is not implicated in the downregulation of MHC I antigens on the surface of T-lymphocyte cells.

It has also been published that upon infection with MV, the level of CD4 surface antigens was markedly reduced (8). In a similar way, we also investigated the influence of serp2 disruption on the described phenotype. Mock-infected RL5 cells showed a strong positive surface staining with anti-CD4 antibodies. After infection with wild-type MV strain T1 or Lau sanne, there was a slight decrease in surface CD4 levels; the same observation could be made after infection of RL5 cells with MV-Serp2- mutant. Infection of RL5 cells with vaccinia virus expressing HIV-nef led to a severe downregulation of CD4 molecules and served as a positive control (data not shown). These results suggest that Serp2 has no impact on the downregulation of CD4 antigens on the surface of RL5 cells.

MV-Serp2 is an important virulence factor in the European rabbit. European rabbits infected with wild-type MV (strain T1 or Lau sanne) rapidly develop a routinely 100% fatal disease known as myxomatosis. As well as being able to activate CPP32, an important cell death protease, ICE is known to liberate bioactive IL-1β, a proinflammatory lymphokine that can alert neighboring cells of the immune system. This signaling may allow inflammatory cells to activate and accumulate at

FIG. 1. Clinical course of rabbits at 10 days postinoculation. Rabbits infected with the wild-type MV (A and B) show secondary skin lesions turning necrotic on the head, the back, and the legs and edema of the testicles (A) and show primary and secondary myxomas on the ears, leading to abnormal posture, and on the nose and around the eyes, and blepharocconjunctivitis (B). Rabbits infected with the MV-Serp2- mutant (C and D) show no secondary skin lesions and little edema of the genital region (C); they also show marked diffuse inflammation of the ears and blepharocconjunctivitis (D).
a site where cell suicide is being used as an antiviral defense (70). Therefore, the putative anti-ICE activity of Serp2 tempted us to determine the effects of a serp2 disruption on MV virulence in vivo. Three groups of animals were infected with wild-type MV \((n = 4)\), MV-Serp2\(^{-}\) mutant virus \((n = 10)\), or MV-Serp2 \(rev\) \((n = 4)\). We observed a marked reduction in virulence in rabbits infected with MV-Serp2\(^{-}\) compared to rabbits infected with wild-type MV or MV-Serp2 \(rev\), respectively (Table 1). On day 4 postinfection, the inoculation site was a diffuse inflammation in rabbits infected with MV-Serp2\(^{-}\), larger and less circumscribed than in rabbits infected with wild-type MV or MV-Serp2 \(rev\), in which a red and soft nodule, known as the primary myxoma, was present. At day 7 postinfection, when rabbits inoculated with either wild-type MV or MV-Serp2 \(rev\) demonstrated classical symptoms of myxomatosis and were prostrated, rabbits infected with MV-Serp2\(^{-}\) had no skin lesions, such as secondary myxomas, on the head, body, or legs. They behaved normally and presented only a mild conjunctivitis. The ears were thickened and red. By day 11 postinfection, all eight rabbits infected with wild-type MV or MV-Serp2 \(rev\) had to be sacrificed, whereas in MV-Serp2\(^{-}\)-infected rabbits, clinical signs of gram-negative infections in the conjunctiva and respiratory tract had developed, although the usual state of health remained unchanged (Fig. 1). The respiratory symptoms worsened between days 10 and 15, but the usual state of health remained unchanged (Fig. 1). The rabbits sacrificed because of increased severity of the symptoms (edema of the testicles, edema of the nasal turbinals, and ataxia) showed systemic lymphodepletions of lymphocytes. The latter finding could well be the consequence of vigorous cell death triggered by the viral infection even if this process cannot be determined on conventional histologic sections.

In view of the significant difference in the resulting pathologic pattern became even more evident. In primary sites, the lesions associated with the wild-type virus were a marked perivascular dermatitis with diffuse edema, focal interstitial hemorrhages, accumulation of heterophils, and a well-developed myxoma (i.e., activated fibroblasts and interstitial mucinosis). In contrast, the lesions associated with the MV-Serp2\(^{-}\) mutant presented two striking differences: they were much less intense, and the sequential inflammatory cellular reactions progressed more rapidly, as attested by the presence of infiltrates of mononuclear cells (histiocytes and lymphocytes). These observations are also valid for the conjunctival lesions. The parotid lymph nodes of rabbits inoculated with the wild-type MV showed a marked lymphadenitis with secondary myxomas. The parotid lymph nodes of rabbits inoculated with MV-Serp2\(^{-}\) mutant exhibited a severe lymphodepletion and no secondary myxoma was observed.

### Table 1. Pathogenicity of MV-Serp2\(^{-}\) in European rabbits

<table>
<thead>
<tr>
<th>Day</th>
<th>Wild-type MV and MV-Serp2 (rev)(^{a})</th>
<th>MV-Serp2(^{-}) mutant(^{b})</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>Inoculation of four rabbits with wild-type MV strain T1 and four rabbits with MV-Serp2 (rev) intradermally with 5,000 PFU</td>
<td>Inoculation of 10 rabbits intradermally with 5,000 PFU</td>
</tr>
<tr>
<td>4</td>
<td>Primary lesions at inoculation sites: raised, soft, red, ca. 2 cm</td>
<td>Primary lesions at inoculation sites: red, diffuse, ca. 3 cm</td>
</tr>
<tr>
<td>7</td>
<td>Emaciated; prostrated; gram-negative bacterial infections of nasal and conjunctival mucosa; multiple secondary lesions on face and ears; edema of the testicles</td>
<td>Normal fatness; regular activity; mild conjunctivitis; no gram-negative bacterial infections detected; diffuse inflammation of the ears</td>
</tr>
<tr>
<td>11</td>
<td>Emaciated; ataxia; dyspnea; severe gram-negative bacterial infections in conjunctiva and respiratory tract; secondary lesions turning necrotic; all rabbits sacrificed because of increased severity of the symptoms</td>
<td>Regular activity; purulent blepharoconjunctivitis and nasal discharge; marked inflammation of the ears; no secondary myxomas (6/10) or minute myxomas on the back and nose (4/10)</td>
</tr>
<tr>
<td>15</td>
<td></td>
<td>Gram-negative bacterial infections of nasal and conjunctival mucosa; no secondary or minute myxomas; one rabbit sacrificed at day 14; one rabbit sacrificed at day 15</td>
</tr>
<tr>
<td>30</td>
<td></td>
<td>Full regression of the symptoms, except for one rabbit sacrificed at day 25 due to respiratory bacterial infection</td>
</tr>
</tbody>
</table>

\(^{a}\) Rabbit survival: 0 of 4 and 0 of 4.

\(^{b}\) Rabbit survival: 7 of 10.
At day 11, the same major differences were noted, and some features must be pointed out: (i) a severe infiltration by heterophils persisted at the primary sites of infection with the wild-type virus, whereas for the MV-Serp2 mutant, mononuclear cells were predominant (Fig. 2); (ii) well-developed secondary myxomas were seen only in secondary sites of rabbits inoculated with wild-type MV; and (iii) a severe lymphodepletion was present only in the parotid lymph node of MV-Serp2 mutant-inoculated rabbits.

From these observations we can conclude that the inflammatory reaction is far less hemorrhagic and more rapidly progressive in lesions induced by the MV-Serp2 mutant. This virus leads to a progressive, severe lymphodepletion in the parotid lymph node. No secondary myxoma was seen in the secondary sites.

Histologic assessment of lymphoid apoptosis. The TUNEL method was used to assess apoptosis of lymphocytes in the parotid lymph node and spleen of 12 inoculated rabbits and 2 control animals. The main results for the parotid lymph node are summarized in Table 3. The two controls showed minimal apoptosis (25 to 50 apoptotic bodies for each microscopic field at a ×400 magnification), mainly located in the germinal centers. For rabbits inoculated with wild-type MV, apoptosis of lymphocytes was at the same level and showed the same localization as controls, whatever the day of the experiment (Fig. 3A). The foci of apoptotic heterophils were seen; they could be easily distinguished from apoptotic lymphocytes by cytologic criteria. The parotid lymph nodes of rabbits inoculated with the MV-Serp2 mutant underwent apoptosis, with major differences from the controls: at day 4 postinoculation, a focal extensive apoptosis was initiated in an area of the lymph node (Fig. 3B and C). The grade of the two rabbits was different (light versus severe). Foci of apoptotic heterophils were also seen. At day 8 postinoculation, the parotid lymph node of both rabbits showed a severe focal extensive apoptosis. At day 11, large areas of the parotid lymph node contained only remnants of apoptotic bodies for each animal (Fig. 3D). Traces of localized extensive foci persisted. In intact remaining lymphoid tissue, a light increase in the number of apoptotic lymphocytes was noticed.

From these results we conclude that lymphodepletion in the lymph node draining the primary site is imputable to apoptosis of lymphocytes. Apoptosis of lymphocytes in the spleen was the same in rabbits injected either with the wild-type MV or with the MV-Serp2 mutant and was identical to the controls.

Virus load in tissues. In order to make sure that the striking phenotypic differences between the wild-type and the MV-Serp2 mutant could not be attributed to an impairment of the latter virus to replicate in vivo, we quantified the level of viral replication at the inoculation site and the parotid lymph node. Standard immunohistochemical studies were carried out, and the levels of virus replication in tissues stained with anti-MV antibodies were quantified by counting the number of infected cells per square millimeter.

At days 4, 8, and 11 postinoculation, there was the same level of virus replication at the injection site of either the wild-type MV or with the MV-Serp2 mutant (not shown); comparable results were

### TABLE 2. Histologic observations of lesions from rabbits infected with the wild-type MV or the MV-Serp2 mutant

<table>
<thead>
<tr>
<th>Site</th>
<th>Wild type</th>
<th>MV-Serp2</th>
<th>Wild type</th>
<th>MV-Serp2</th>
<th>Wild type</th>
<th>MV-Serp2</th>
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<tbody>
<tr>
<td>Primary&lt;sup&gt;b&lt;/sup&gt;</td>
<td></td>
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<tr>
<td>Perivascular dermatitis with:</td>
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<tr>
<td>Epidermal spongiosis and vesicles</td>
<td>+ +</td>
<td>f</td>
<td>+</td>
<td>++++</td>
<td>d</td>
<td>+</td>
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<tr>
<td>Edema</td>
<td>+</td>
<td>f</td>
<td>+</td>
<td>++++</td>
<td>d</td>
<td>+</td>
</tr>
<tr>
<td>Focal interstitial microhemorrhage</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
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<tr>
<td>Focal thrombosis</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
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<tr>
<td>Inflammatory infiltrates of:</td>
<td></td>
<td></td>
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<td></td>
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<tr>
<td>Heterophils (perivascular and interstitial)</td>
<td>+</td>
<td>s</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Histiocytes, lymphocytes (interstitial)</td>
<td>+</td>
<td></td>
<td>+</td>
<td>+</td>
<td>s</td>
<td>+</td>
</tr>
<tr>
<td>Activated fibroblasts</td>
<td>+</td>
<td></td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Intestinal mucinosis</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
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<tr>
<td>Secondary&lt;sup&gt;c&lt;/sup&gt;</td>
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<tr>
<td>Superficial perivascular conjunctivitis with:</td>
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<tr>
<td>Inflammatory infiltrates of:</td>
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<tr>
<td>Heterophils</td>
<td>+</td>
<td>f</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Mononuclear cells (histiocytes, lymphocytes)</td>
<td>+</td>
<td>s</td>
<td>+</td>
<td>+</td>
<td>s</td>
<td>+</td>
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<tr>
<td>Focal lymphoid hyperplasia</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
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<tr>
<td>Focal lymphoid depletion</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Activated fibroblasts</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
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<tr>
<td>Intestinal mucinosis</td>
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<td>+</td>
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<td>Lymphadenitis with:</td>
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<tr>
<td>Lymphodepletion</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>(m)</td>
<td>+</td>
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<tr>
<td>Focal microhemorrhage</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
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<td>+</td>
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<tr>
<td>Focal histiocytes</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>(e)</td>
<td>+</td>
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<tr>
<td>Infiltration by heterophils</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>(e)</td>
<td>+</td>
</tr>
<tr>
<td>Activated fibroblasts</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
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<td>Intestinal mucinosis</td>
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<td>+</td>
<td>+</td>
<td>+</td>
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</tr>
</tbody>
</table>

* Lesion intensity: +, minimal; ++, light; ++++, moderate; ++++, marked; ++++++, severe. Lesion topography: s, scattered; f, focal; m, multifocal; e, extensive; d, diffuse. p.i., postinoculation.
* Samples were taken from lesions at the inoculation site.
* Samples were taken from lesions on the ocular conjunctiva and parotid nodes.
obtained in the parotid lymph node (Fig. 4). Eight days after infection with the wild-type MV, the cells in the lymph node (mostly fibroblasts and histiocytes, which are characteristic of the secondary myxomas) were positively stained with the anti-MV antibodies (Fig. 4A). With the MV-Serp2 mutant, the cell populations were different (mostly mononuclear cells, which from morphologic criteria could be defined as lymphocytes), but the viral load, according to our semiquantitative measure, was on the same order of magnitude (Fig. 4B). From these results we conclude that the MV-Serp2 mutant is able to replicate in vivo at a level comparable to that of the wild-type MV.

DISCUSSION

In this study we report data to elucidate the functions of the MV Serp2 protein, which is 35% identical to the cowpox virus CrmA protein. CrmA has been characterized as viral inhibitor of ICE (caspase 1), which can prevent the onset of both

TABLE 3. Histologic assessment by the TUNEL method of lymphoid apoptosis in the parotid lymph nodes of rabbits infected with wild-type MV or MV-Serp2 mutant

<table>
<thead>
<tr>
<th>Day</th>
<th>Wild-type</th>
<th>MV-Serp2−</th>
<th>Histologic assessment (lesion intensity)</th>
</tr>
</thead>
<tbody>
<tr>
<td>4</td>
<td>A few apoptotic cells in germinal centers (as in controls). Score: homogeneous (+)</td>
<td>Focal extensive apoptosis. Score: variable (+/+++++).</td>
<td></td>
</tr>
<tr>
<td>8</td>
<td>A few apoptotic lymphocytes in germinal centers (as in controls). Score: homogeneous (+)</td>
<td>Focal extensive lymphoid apoptosis. Score: homogeneous (+++++).</td>
<td></td>
</tr>
</tbody>
</table>

* Lesion intensity: +, minimal; ++, light; ++++, moderate; +++++, marked; ++++++, severe.
flammation and apoptosis by reducing the levels of the active proinflammatory lymphokines IL-1β and IL-18 and by preferentially inhibiting the proximal components of the ICE/CED-3 protease cascade in the cell death process (33, 48, 50, 58, 64). CrmA also has a high affinity for caspase 8 or FLICE (72) and Granzyme B (49). More importantly, Serp2 was also previously shown to bind to human ICE and prevent it from processing the pro-IL-1β into its bioactive form (46). Therefore, Serp2

FIG. 3. Parotid lymph node (TUNEL method). The panels show samples at 8 days postinoculation with the wild-type MV, with minimal lymphocytic apoptosis (A), and infection with the MV-Serp2− mutant at 4 days postinoculation, with focal extensive lymphocytic apoptosis in cortical areas (arrows) (B); at 4 days postinoculation, with focal extensive lymphocytic apoptosis at a higher magnification (C); and at 11 days postinoculation, with remnants of apoptotic foci in a lymphodepleted parenchyma (arrow) (D). Magnification: ×100 (A); ×40 (B); ×100 (C); ×40 (D).

FIG. 4. Parotid lymph node (immunohistochemistry) from rabbits at 8 days postinoculation (×400). Infected cells are visualized with DAB (arrow). (A) Infection with the wild-type MV. The cells present in the lymph node are mostly histiocytes and activated fibroblasts (large nucleus, double arrows). These and other cells are infected. (B) Infection with the MV-Serp2− mutant. Most cells are infected, and mononuclear cells are predominant.
was another candidate poxvirus-encoded factor to counteract host defenses and to potentially contribute to the pathology associated with MV infection.

To assess the role of Serp2 in the pathogenesis of myxomatosis, we successfully engineered a fully replication-competent MV mutant deleted in the Serp2 open reading frame. Revealing a wild-type-like phenotype when grown in vitro, the mutant virus was found to be highly attenuated upon in vivo infection in rabbits. The clinical course of infection with MV-Serp2 mutant was characterized by the development of mild and more-diffuse primary myxoma lesions at the site of inoculation, the absence or benignity of rare secondary lesions and, most impressively, by the high rate of complete recovery seen in the infected animals.

Detailed histologic examination led to three major findings. First, the inflammatory response upon inoculation with wild-type MV seemed to be arrested at the vascular level (with heterophils predominant even at a late time), whereas with the MV-Serp2 mutant the inflammation proceeded to the cellular phase (with lymphocytes and histiocytes infiltrating the lesion). In this latter case the overall reaction was less intense and less hemorrhagic than when Serp2 was present. The inhibition of IL-1β processing by Serp2 is in accordance with these observations. IL-1 is a potent proinflammatory cytokine that can affect the function of many physiologic systems (for a review, see reference 15). Among the pleiotropic effects of IL-1β, including thrombosis and inflammation, is an increased expression of the immunoglobulin superfamily molecules ICAM-1 and VCAM-1, which bind to integrins on lymphocytes and monocytes (14, 24). By blocking IL-1β processing, Serp2 would diminish the ability of these cells to migrate into the inflamed tissue. Moreover, ICE is involved in the processing of pro-IL-18 into its active form (23, 26). IL-18, also known as IFN-γ-inducing factor, stimulates the production of IFN-γ by lymphocytes (42, 44). As for IL-1β, the effects of IFN-γ are numerous, including the chemotraction of macrophages (10, 43, 71). This phenomenon was also occurs after infection with a mutant lacking a secreted homolog of the IFN-γ receptor; Mossman et al. (40) reported that infection with the IFN-γ receptor-deficient MV resulted in an inversion of the heterophil/mononuclear cell ratio at the secondary sites. Our histologic findings, although not definitely proving which cytokine defect is involved, are consistent with the hypothesis that the presence of Serp2 leads to a reduced amount of both IL-1β and IFN-γ at the infection sites.

Our second important histologic observation was that, in the absence of Serp2, undergo apoptosis which can be quantified in the parotid lymph nodes. According to the spatial distribution of the apoptotic foci in the lymph nodes, it is reasonable to assume that apoptosis occurred in the lymph node itself, as well as in lymphocytes drained from the inoculation site. ICE has been widely recognized as an important mediator of the apoptotic process (reviewed in references 7 and 41). The impairing of ICE by Serp2 would be in accordance with an inhibition of the apoptotic process by the wild-type virus, but not by the MV-Serp2 mutant virus. However, in our in vitro experiments, the defect in Serp2 production did not lead to apoptosis of RL5 T-helper lymphocytes. This cell line has been widely used to study other MV genes. MV lacking the TNF receptor homologue (53), the M11L gene (35), and the host-range superfamily member M-T5 (39) have all been shown to induce apoptosis in this cell line. The reason why all these mutants display the same phenotype in RL5 cells is not clear. M-T5 might counteract the shutoff of protein synthesis after MV infection (39), but M-T2 does probably not block apoptosis via its TNF binding domain (53). M11L is a transmembrane protein involved in the repression of inflammation, but its target is unknown (43). On the other hand, the cell-type dependence of apoptosis inhibition by viral products has been widely reported (6, 13, 21, 51). In that context, it is clear that RL5 cells are not the best model for checking Serp2 inhibition of ICE-mediated apoptosis.

There are several reports that either cowpox CrmA (62) or its homologue vaccinia virus SPI-2 protein (16, 32) can inhibit Fas-mediated apoptosis. However, in our experiments Serp2 was not required for the inhibition of TNF-induced apoptosis in HeLa cells. Several hypotheses underlie our findings. The first possibility is that, although reported to be unable to bind to human TNF-α (52), the MV TNF receptor homologue was responsible for this inhibition; another possibility is that there exists an as-yet-unidentified antiapoptotic factor encoded by MV, whose role would be to prevent TNF-induced apoptosis. It has recently been shown that the MCI59 protein of Molluscum contagiosum, another member of the poxvirus family, could inhibit Fas- and TNFRI-induced apoptosis through its death effector domain (9). It is thus possible that MV encodes one or more proteins that would prevent ICE activation in cell culture.

Since it has been reported that MV could downregulate the expression of CD4 (8) and MHC I molecules (11) on the surface of infected cells, we checked whether Serp2 could account for this phenotype. In fact, we could see no difference in the relative decrease of either CD4 or MHC I antigens of RL5 cells infected with the wild-type or the MV-Serp2 mutant. The effect of MV on the expression of these surface molecules was light in both cases compared to the more drastic effect observed upon the vaccinia virus-mediated expression of the human immunodeficiency virus type 1 nef gene (1, 54). It was particularly relevant to check for a possible implication of Serp2 in MHC I downregulation, since this would have interfered with the apoptosis process. Any decrease in MHC I expression at the surface of infected cells would result in the inability of the cytotoxic T lymphocytes to bind to these cells and to recognize them as targets. Since Serp2 is not interfering with MHC I expression, the marked difference in the lymphocyte apoptosis observed in vivo cannot be attributed to a modification of the immune effectors on the membrane of the infected cells. More likely, the difference is due to an antiapoptotic effect of Serp2.

The last histologic finding was that in the secondary sites (ocular conjunctiva and lymph nodes) there were no so-called secondary myxomas upon infection with the MV-Serp2 mutant, contrasting with the wild-type MV. The molecular mechanism by which MV induces the formation of myxomas is unclear, but it should not be attributed only to the presence of the virus in the tissue. Indeed, we were able to show that the MV-Serp2 mutant can replicate as efficiently as the wild-type virus at the inoculation site and in the parotid lymph node. The comparable level of viral loads between both viruses indicates that the clinical and histologic differences observed cannot be attributed simply to a more poorly growing mutant virus. We were able to show that the lymphocytes in the lymph node of rabbits infected with the MV-Serp2 mutant are indeed infected.

Our conclusions from these observations are that in the MV, Serp2 is able to block both the processing of the inflammatory reaction at an early stage and apoptosis of the lymphocytes. It seems relevant to associate these phenomena with the inhibition of caspase 1 and/or another aspartic acid-specific protease.

It was previously reported that inactivation of the crmA gene of cowpox virus or its equivalent SPI-2 gene of rabbitpox virus resulted in an attenuation of the clinical course, as measured
by the body weight of mice after intranasal infection (63). However, the cowpox and rabbitpox viruses differed in the inflammatory response. An acute inflammatory response was described as associated with disruption of the SPI-2 gene in the rabbitpox virus, whereas disruption of the open reading frame on the cowpox virus seemingly resulted in a decreased influx of inflammatory cells. Using the same murine intranasal model, other authors have reported that inactivation of the SPI-2 gene of the vaccinia virus had no effect on virus virulence (31). These contradictory results concerning three members of the Orthopoxvirus genus may be attributed to either true differences in the viruses or to mutations acquired elsewhere in the genomes, since neither cowpox nor rabbitpox revertant viruses were produced (63). However, in each case the mice had been inoculated intranasally. Here, we have been using the intradermal route, which seems more relevant since rabbits get contaminated through fleas or mosquito bites (20). It is not surprising that its extra target(s) might be different from those of CrmA or SPI-2.

(46). It is not surprising that its extra target(s) might be different from those of CrmA or SPI-2. Unlike CrmA and SPI-2, which are only expressed early in the course of a natural infection.

Inhibition of ICE and/or other caspases by Serp2 will be informative regarding the relative importance of inflammation and apoptosis for eliminating the virus.

ACKNOWLEDGMENTS

We are grateful to G. McFadden and G. L. Smith for kindly providing us with RL5 cells and HGPRT−HeLa cells, respectively. We also thank V. Lourec for monitoring the rabbits.

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REFERENCES


MYXOMA VIRUS Serp2 DELETION MUTANT


