In Vivo Role of Complement-interacting Domains of Herpes Simplex Virus Type 1 Glycoprotein gC

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Summary

Immune evasion is critical for survival of viruses that establish persistent or recurrent infections. However, at the molecular level, little is known about how viruses evade immune attack in vivo. Herpes simplex virus (HSV)-1 glycoprotein gC has two domains that are involved in modulating complement activation; one binds C3, and the other is required for blocking C5 and properdin (P) binding to C3. To evaluate the importance of these regions in vivo, HSV-1 gC mutant viruses were constructed that lacked one or both gC domains and studied in a murine model of infection. Each gC region of complement regulation contributed to virulence; however, the C3 binding domain was far more important, as virus lacking this domain was much less virulent than virus lacking the C5/P inhibitory domain and was as attenuated as virus lacking both domains. Studies in C3 knockout mice and mice reconstituted with C3 confirmed that the gC domains are inhibitors of complement activation, accounting for a 50-fold difference in virulence between mutant and wild-type viruses. We conclude that the C3 binding domain on gC is a major contributor to immune evasion and that this site explains at a molecular level why wild-type virus resists complement attack.

Key words: immunology • pathogenesis • innate immunity • disease • C3

Herpes simplex virus has developed many strategies for survival in the human host. One of the most successful is its ability to establish latent infection in neurons without expressing viral proteins that serve as targets for immune attack (1, 2). However, during latency, virus cannot spread person to person; therefore, the virus must reactivate to successfully infect another host. HSV has the ability to replicate for days after reactivation in the immune host, which may be related to viral immune evasion strategies. HSV-1 immune evasion molecules block activities of antibodies (3–7) and complement (8–10) and prevent antigen presentation by the MHC class I complex (11, 12). Molecules that perform similar functions have been identified in adenoviruses, vaccinia virus, and other members of the herpes virus family (13–18).

HSV-1 glycoprotein gC is among the best characterized of the viral immune evasion molecules. gC binds complement components C3, C3b, iC3b, and C3c, accelerates the decay of the alternative pathway C3 convertase, and blocks the interaction of C5 and properdin (P) with C3 (9, 19, 20). The C5 and P blocking domain is near the NH₂ terminus, whereas the C3 binding domain is in the central region of the molecule (9, 21). gC inhibits complement activation (9, 20), protects cell free virus from complement-mediated neutralization (10, 22, 23), inhibits complement-mediated lysis of HSV-infected cells (24, 25), and protects the virus against complement attack in vivo (26).

Vaccines incorporating molecules required for virus entry, including glycoproteins gB and gD, failed to prevent infection and had little effect on recurrence rates (27–29). A better understanding of HSV immune evasion strategies may be required to design more effective vaccines. With this goal in mind, we evaluated virulence of gC mutant viruses deleted in the C3 binding domain, the C5/P blocking domain, or both to define the contributions of each domain to immune evasion. We chose to study gC–complement interactions in mice because C3 knockout mice are available to establish the importance of C3. We selected the murine flank model, as we previously showed that complement significantly contributes to defense against HSV-1 in this model (24). We used C3 reconstitution (30) to confirm that gC is highly effective in protecting wild-type virus from complement attack and to define an impressive role for the C3 binding domain in pathogenesis.

Materials and Methods

Construction of gC Mutant Viruses. HSV-1 strain NS, a low passage clinical isolate, was the parental strain for the mutant viruses.

*Abbreviation used in this paper: P, properdin.
NS-gC null virus has the entire gC protein coding sequence replaced by a lacZ expression cassette under the control of the HSV-1 infected cell protein 6 early promoter, as previously described (23). NS-gCΔC5/P virus has a deletion of the gC domain involved in blocking C5 and P binding to C3 (9, 21). This deletion was constructed from the N S BamH I “I” fragment (reference 31; nucleotides 91,602–98,254; sequence available from EM BL/GenBank/DBJ under accession numbers X14122, D00317, D00374, S40493, and X21238) by cutting with Nsi I and EcoR I and cloning this fragment (nucleotides 94,911–96,751) into pGEM 7Z (Promega Corp.) to create plasmid pLW1. An N hel–EcoR I fragment (nucleotides 96,276–96,751) was excised from pLW1 I and cut with Apal to delete 91 gC amino acids (amino acids 33–123; nucleotides 96,407–96,679). The N hel–EcoR I fragment deleted of gC amino acids 33–123 was inserted back into pLW1 I, and then the N si1–EcoR I fragment from pLW1 I was cloned back into the BamH I I fragment to create a flanking sequence vector with a deletion of the gC C5/P domain.

gCΔC3 has a deletion of 92 amino acids (gC amino acids 275–367), prepared by cutting plasmids B29 and H71 (32) with BglII, reliating the 5′ portion of B29 with the 3′ portion of H71, and then cloning this fragment into the EcoR I–EcoRV site of the BamH I I fragment. The construction adds a 12-mer BglII linker after gC amino acid 275 and creates a flanking sequence vector that has a deletion of two gC C3 binding regions, resulting in a net loss of 88 amino acids.

A flanking sequence vector to construct gCΔC5/P.C3 was prepared by replacing the BamH I I EcoR I–EcoRV fragment in the gCΔC5/P flanking sequence vector with the EcoR I–EcoRV fragment of the gCΔC3 flanking sequence vector. Recombinant viruses were prepared in Vero cells by cotransfection using NS-gCnull virus DNA and the various flanking sequence vectors. Recombinant viruses were identified by anti-gC immunoperoxidase staining of infected cells (21). Virus pools were prepared after three rounds of plaque purification.

Sucrose Gradient Purification of Virus. Vero cells were infected at an M O I of 5 for 20 h at 37°C, and then virus from supernatant fluids was purified on a 5%–65% sucrose gradient (23). The viable virus band was collected, dialyzed against PBS at 4°C, and stored at −70°C. Virus titers were determined by plaque assay on Vero cells.

One-Step Growth Curves. Vero cells were inoculated at an M O I of 5, and at 1, 4, 8, 12, 20, and 24 h after infection, cells plus supernatant fluids were harvested, cells were lysed by sonication, and viral titers were determined by plaque assay.

Southern Blot Analysis. DNA was digested with BamH I and HindIII, electrophoresed in 1.2% agarose gels, transferred to Immobilon-S membranes (Millipore Corp.), and U V cross-linked. A biotin-labeled gC-1 fragment spanning the entire protein coding sequence (nucleotides 96,276–97,945) was used as probe, and bands were detected by chemiluminescence (New England Biolabs) (6).

Western Blot Analysis. Purified virus was run on 10% SDS-PAGE, transferred to Immobilon-P membranes (Millipore Corp.), and visualized using polyclonal rabbit anti-gC antibody and horseradish peroxidase–conjugated goat anti–rabbit IgG (Amer sham Pharmacia Biotech) (23).

A nontybody-independent complement neutralization. N eutralization assays were performed by incubating 104–106 PFU of purified virus with H SV-1 and -2–seronegative human serum as the source of complement for 1 h at 37°C. As a control, complement was inactivated by heat treatment at 56°C for 30 min. Virus titers were then determined by plaque assay on Vero cells. Complement neutralization was calculated as the difference in titers between virus incubated with active or heat-inactivated serum (23).

C3b Rosetting Assays. Vero cells were infected at an M O I of 2 for 20 h, and then cells were removed using cell dissociation buffer (GIBCO BR.), incubated with C3b-coated erythrocytes for 1 h at 37°C, and viewed by microscopy for rosettes. Cells with four or more bound erythrocytes were considered positive (23).

C3 Purification. Human C3 was purified according to Hammer et al. (33) with modifications. 20 parts human plasma were treated with 1 part inhibitor containing 1 M KH4PO4, 0.2 M Na2EDTA, 0.2 M benzamidine, and 1 mM PMSC. C3 was precipitated from plasma first with 4.5% polyethylene glycol (PEG) and then with 12% PEG at 0°C. The pellet was dissolved in buffer A (20 mM NaH4PO4, pH 7.4) and loaded onto a DEAE 40HR column (5 × 6.3 cm; Whaters) equilibrated with buffer A. The bound proteins were eluted using a 0–0.5 M NaCl linear salt gradient. Fractions containing C3 were detected by SDS-PAGE and immunodiffusion, dialyzed against buffer A, loaded onto an M ono Q HR 10/10 column (Amersham Pharmacia Biotech), and eluted with a linear salt gradient to 0.5 M NaCl. Homogeneous C3 fractions were identified by SDS-PAGE and immunodiffusion, pooled, and dialyzed against PBS (10 mM NaH4PO4 and 145 mM NaCl, pH 7.4). The purified C3 contained 80% native C3 and 20% C3H (H O), as determined by analyzing the protein sample on a M ono S column (Amersham Pharmacia Biotech) (34).

M urine F lank M odel. The flank model and the C3 knockout mice derived from C57BL/6 mice have been described previously (26, 35). In brief, purified virus was scratched onto the flank that was shaved and chemically denuded. Disease at the inoculation site was scored for disease at the inoculation site was scored.
Results

Characterization of gC Mutant Viruses. Features of wild-type gC and three gC mutants are shown in Fig. 1. Wild-type gC (Fig. 1, top) has four C3 binding regions (light gray), each of which is required to bind C3 (32). The gC domain that interferes with C5 and P binding to C3 is located at the NH2-terminal region between residues 33 and 123 (9, 21). gCΔC5/P mutant (Fig. 1, second from top) contains a deletion of gC amino acids 33–123. gCΔC3 mutant (Fig. 1, third from top) has a deletion of amino acids 275–367 that includes C3 binding regions II and III. This deletion removes a disulfide-bonded cysteine pair (37) to minimize the effect of the deletion on gC conformation. gCΔC5/P, C3 mutant (Fig. 1, bottom) has both domains deleted that interact with complement.

The mutant gC proteins were recombined into virus, and the size of the gC DNA was evaluated by Southern blot analysis. HSV-1 was cut with BamH1 and HincII to generate two gC fragments of different sizes consistent with the deletion mutations introduced. Right panel: DNA from NS and gC mutant virus N S-gCΔC5/P, C3 and N S-gCΔC3; the right blot shows the smaller upper band for gC mutant N S-gCΔC5/P. Bands were of the expected size for each of the gC mutant viruses. Western blots were performed on purified virus to demonstrate that mutant gC glycoproteins are incorporated into the virion and are of the expected size (Fig. 2 B). The double mutant virus N S-gCΔC5/P, C3 makes the smallest of the three mutant proteins. N S-gCΔC5/P has a deletion of 91 amino acids and on SDS-PAGE runs smaller than N S-gCΔC3, which has a deletion of 88 amino acids. However, the difference in migration of the two glycoproteins was greater than expected based on amino acid size. A likely explanation is that five N-linked glycosylation sites and numerous predicted O-linked glycosylation sites lie within the deleted domain of N S-gCΔC5/P, whereas one N-linked and few O-linked glycosylation sites lie within the deleted fragment of N S-gCΔC3.

The kinetics of virus replication and peak titers achieved for each gC mutant virus were similar to those of wild-type virus (Fig. 3). C3b binding was assessed by a rosetting assay. Wild-type virus, N S, and N S-gCΔC5/P (C5/P) form rosettes, whereas N S-gCΔC3 (C3), N S-gCΔC5/P, C3 (C5/P, C3), and N S-gCnull (gCnull) viruses do not form rosettes.

Antibody-independent complement neutralization of wild-type virus, gE null mutant virus, and a panel of gC mutant viruses 101–106 PFU of purified virus was incubated with HSV-1 and 2 antibody-negative human serum as a source of complement or heat-inactivated serum as control. After a 1-h incubation at 37°C, virus titers were determined by plaque assay. Results are expressed as the differences in titers (log10) ± SEM comparing virus incubated with inactivated complement or active complement. N S, wild-type HSV-1; gE null, gE deletion virus N S-gE null (6) derived from N S; C5/P, N S-gCΔC5/P; C3, N S-gCΔC3; C5/P, C3, N S-gCΔC5/P, C3; gC null, N S-gC null. Experiments were performed three to five times, as indicated by the numbers above the bars.
Figure 5. Lesion scores in mice infected with wild-type or gC mutant viruses. The left panels (A, C, and E) show disease scores ± SEM at the inoculation site, and the right panels (B, D, and F) show disease scores ± SEM at the zosteriform spread site. A and B demonstrate disease scores in complement-intact mice (normal mice), C and D in C3 knockout mice, and E and F in C3 knockout mice reconstituted with human C3. WT, virus NS; C5, NS-gCΔC5/P; C3, NS-gCΔC3; C5,C3, double mutant virus NS-gCΔC5/P,C3. The numbers of animals evaluated in each group are indicated in the legends at right.
virus when evaluated by single-step growth curves (result not shown). gC protein expression at the infected cell surface was confirmed for each mutant virus by immunoperoxidase staining, and the ability of gC to bind C3b was determined by rosetting assays. Mutant viruses NS-gCΔC3, N S-gCΔC5/P, C3, and N S-gCnull lack the C3 binding domain and fail to form C3b rosettes, whereas NS and N S-gCΔC5/P mutant viruses have the C3 binding domain and form rosettes (Fig. 3). The results suggest that the gC mutations have not drastically altered protein conformation, as mutant virus NS-gC forms rosettes (Fig. 3). The results suggest that the gC mutations have not drastically altered protein conformation, as mutant virus NS-gCΔC3 virus caused significantly less disease than N S-gCΔC5/P virus (P < 0.001 for days 6–8 at the inoculation and zosteriform sites). We conclude that both gC domains have a significant effect on disease severity; however, the C3 domain is more important because the mutant virus lacking this domain is significantly less virulent than the C5/P mutant virus and is impaired as the double mutant virus.

Experiments were performed in C3 knockout mice to determine if interactions between gC and complement account for the lowered virulence of gC mutant viruses. Virulence of gC mutant viruses should be similar to wild-type virus in C3 knockout mice, as C5, P, or C3 cannot become activated. Fig. 5, C and D show that virulence for each of the mutant viruses is comparable to wild-type virus in C3 knockout mice.

As further evidence for the importance of the interaction between gC and complement, C3 knockout mice were reconstituted with C3. Total hemolytic complement activity was undetectable in serum from C3 knockout mice; however, after injection of human C3, total hemolytic complement activity was restored to levels detected in complement-intact animals (result not shown). Fig. 5, E and F show that comparison N S-gCΔC3 or double mutant virus N S-gCΔC5/P, C3 with N S (P < 0.001 for days 5–8 at the inoculation and zosteriform sites). Notably, N S-gCΔC3 virus caused significantly less disease than N S-gCΔC5/P virus (P < 0.001 for days 6–8 at the inoculation and zosteriform sites). We conclude that both gC domains have a significant effect on disease severity; however, the C3 domain is more important because the mutant virus lacking this domain is significantly less virulent than the C5/P mutant virus and is impaired as the double mutant virus.

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the gC double mutant virus NS-gCΔC5/P,C3 is markedly less virulent in C3 knockout mice reconstituted with C3, whereas the virulence of wild-type virus is unchanged (comparing NS and gC double mutant virus inoculation site lesions, \( P < 0.05 \) on day 3, \( P < 0.01 \) on day 4, and \( P < 0.001 \) on days 5–8, whereas for zosteriform lesions, \( P < 0.001 \) on days 5–8).

Fig. 6 shows photographs of flank lesions on day 6 after infection. The left panels show NS, and the right panels show NS-gCΔC5/P,C3, demonstrating that lesions caused by NS are comparable in the presence or absence of complement, whereas lesions produced by the gC mutant virus are greatly affected by complement. These experiments illustrate the important role of gC–complement interactions in pathogenesis.

Defining the Magnitude of the gC–Complement Effect on Virulence. Experiments were performed using NS and gC double mutant virus NS-gCΔC5/P,C3 to compare disease scores over a range of inocula. In Fig. 7, the disease scores are expressed as the average cumulative disease scores from days 3–8 after infection (26). In complement-intact mice, infection with \( 5 \times 10^6 \) PFU of the gC double mutant virus produced a disease score of 19.3 ± 1.9 at the inoculation site (Fig. 7A). Producing a comparable disease score required 50-fold less wild-type virus. Infection with \( 5 \times 10^5 \) PFU of gC double mutant virus resulted in a disease score of 8.9 ± 0.9, which also correlated with the disease caused by 50-fold less wild-type virus. Similarly, ∼50-fold less NS than gC mutant virus was required to produce comparable zosteriform disease (Fig. 7B). In marked contrast, in C3 knockout mice, no significant differences were noted comparing wild-type and gC mutant viruses (Fig. 7C and D). We conclude that domains on gC that interact with comple-
ment account for a 50-fold effect on virulence in complement-intact animals, whereas in C3 knockout animals, the domains have no effect. Notably, NS disease scores are virtually identical in complement-intact and C3 knockout mice (comparing the scores for wild-type virus in Fig. 7, A with C or B with D), which indicates that gC complement-interacting domains are remarkably effective in protecting wild-type virus from complement attack.

Discussion

We have previously defined a role for gC in binding C3 (8, 9, 19), accelerating the decay of the alternative pathway C3 convertase (20) and interfering with the interaction of C5 and P with C3b (9, 21). We showed that gC null virus is highly susceptible to complement neutralization even in the absence of antibodies and that gC null virus is less virulent in mice and guinea pigs (23, 26). We now address the contribution to pathogenesis of two gC domains that interact with complement by constructing gC mutant viruses lacking the C5/P blocking domain, the C3 binding domain, or both domains.

The murine studies indicate that each of the single mutant viruses is significantly less virulent than wild-type virus; however, the C3 binding domain mutant virus is clearly the more important, as this mutant virus was as attenuated as the gC double mutant virus. Fig. 8 A presents a model of complement inhibition mediated by gC domains. In vivo, NS-gCΔC5/P mutant virus is only moderately less virulent than wild-type virus, suggesting that gC affects more than C5 and P binding to C3b, such as accelerating the decay of the alternative pathway C3 convertase, C3bBb (20), or perhaps other C3-mediated functions yet to be determined (Fig. 8 B). The deletion of the C3 binding domain eliminates gC binding to C3b (Fig. 8 C), rendering the C5/P domain ineffective because it lacks proximity to C3b and cannot hinder C5 and P binding. Therefore, a C3 binding domain mutant virus should be as impaired as a double mutant virus (Fig. 8 D). The model explains our in vivo results, but it fails to explain why the C3 binding mutant virus was not as impaired as the double mutant virus in neutralization assays performed in vitro. A possible explanation is that the main effect of gC in vivo may be on a function other than neutralization, such as preventing complement lysis of infected cells (24).

Our results definitively demonstrate that gC is a virulence factor because it regulates complement activation. This conclusion is based on observations that gC mutant viruses are significantly less virulent than wild-type virus in complement-intact mice, whereas virulence is comparable to that of wild-type virus in C3 knockout mice. Proof for the requirement for C3 was shown by C3 reconstitution experiments in which C3 reduced virulence of the gC double mutant virus but had no effect on wild-type virus. The magnitude of the gC effect was addressed by evaluating lesion scores using inoculation titers ranging from $5 \times 10^5$ to

![Figure 8](https://example.com/figure8.png)
5 \times 10^2 \text{ PFU}. In complement-intact mice, 50-fold more gC double mutant virus was required to cause disease comparable to that caused by wild-type virus.

In vitro studies indicate that gC is also involved in the initial stage of virus attachment to cells by binding to heparan sulfate (38, 39). In the absence of gC, gB can mediate this effect (38), but a role for gC in attachment has not been established in vivo. Our results indicate that the domains deleted from the gC mutant viruses do not play an important role in attachment, because in C3 knockout mice, the mutant viruses are as virulent as wild-type virus. Of particular interest is the C5/P deletion mutant virus. This mutant virus lacks gC amino acids 33–123, identified as an important domain for gC-mediated attachment to heparan sulfate in vitro (40). Yet this virus shows no difference in virulence from wild-type virus in C3 knockout mice, suggesting that this domain is not important for virus attachment in vivo.

Disease at the inoculation site develops by day 3 after infection, and zosteriform disease generally appears on day 5 (6, 26). We scored inoculation site and zosteriform disease separately so that we could better evaluate early stages of infection. Interestingly, disease scores at the inoculation site of the C3 single and double mutant viruses did not differ from wild-type virus until day 5. This result suggests that the complement effect on gC mutant virus cannot be detected until rather late after infection. Several explanations are possible. If complement lysed infected cells rather than inhibiting cell free virus, it may take several days until enough cells are infected to create an observable effect. Another possible explanation is that complement concentrations may be too low at the inoculation site early in infection, whereas at later time points, virus-induced tissue injury leads to an increase in vascular permeability and higher levels of complement proteins. Additional considerations include that complement may be mediating its effects by stimulating chemotaxis or other proinflammatory events that may take several days to develop, or that antibodies may be required to augment the effects of complement. This last explanation seems unlikely in view of in vitro results that demonstrate that antibodies are not required for complement-mediated neutralization or lysis of infected cells (23, 24).

We have previously described a humoral immune evasion domain on glycoprotein gE that inhibits IgG Fc-mediated functions (4–6). We showed that gE blocks complement-enhanced antibody neutralization, reducing the effectiveness of antibody 100-fold in vitro and offering highly significant protection to the virus or virus-infected cell against antibodies in vivo (6). The results of this study indicate that complement has little effect on wild-type virus, as disease scores of wild-type virus are virtually identical in C3 knockout and complement-intact mice. Immune evasion mediated by gC and gE may underlie the long-held view that humoral immunity is relatively unimportant in HSV-1 pathogenesis (41). These immune evasion molecules render the Fc domain of antibody and the complement cascade far less effective in host defense. These observations have important implications for vaccines and chemotherapy. They explain why antibodies and complement are poorly protective against HSV infection and provide potential targets for therapies to block HSV-1 immune evasion and render the virus more susceptible to innate and acquired immunity.

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