Membrane Anchoring of Epstein-Barr Virus gp42 Inhibits Fusion with B Cells Even with Increased Flexibility Allowed by Engineered Spacers

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ABSTRACT We recently described the architecture of the Epstein-Barr virus (EBV) fusion-triggering complex consisting of the EBV B cell receptor human leukocyte antigen (HLA) class II and the EBV-encoded proteins gp42 and gH/gL. The architecture of this structure positioned the main body of gp42, comprising the C-type lectin domain (CTLD), away from the membrane and distant from where the membrane-bound form of gp42 might be tethered. gp42 is a type II membrane glycoprotein, with functional gp42 formed by cleavage near the gp42 amino-terminal transmembrane domain. This cleavage results in an approximately 50-amino-acid unstructured region that is responsible for binding gH/gL with nanomolar affinity. Our previous studies had shown that membrane-bound gp42 is not functional in B cell fusion. To investigate whether we could restore gp42 function by extending it from the membrane, we introduced one, two, and four structured immunoglobulin-like domains from muscle protein titin into a membrane-bound form of gp42 and tested function in binding to gHgL and HLA class II and function in fusion. We hypothesized that cleavage of gp42 generates a soluble functional form that relieves steric hindrance imposed on gHgL by membrane-bound gp42. All of the linker mutants had a dominant-negative effect on gp42 function, indicating that gp42 fusion function could not be restored simply by the addition of one to four titin domains.

IMPORTANCE Epstein-Barr virus (EBV) is associated with numerous diseases from benign mononucleosis to Burkitt’s and Hodgkin’s lymphoma, nasopharyngeal and gastric carcinoma, and lymphoproliferative disorders in patients with immune dysfunction resulting from immune suppression. Among the glycoproteins important for fusion, gp42, along with gH/gL, determines EBV tropism between epithelial and B cells. The function of gp42 is dependent on N-terminal cleavage, since membrane-bound gp42 cannot mediate fusion. We further investigated whether insertion of a linker into membrane-bound gp42 would relieve steric hindrance imposed on membrane-bound gp42 and restore fusion function. However, adding one, two, or four structured immunoglobulin-like domains to membrane gp42 did not restore fusion activity, indicating that the architecture and membrane orientation of the B cell fusion-triggering complex of EBV may be easily perturbed and that gp42 cleavage is essential for B cell fusion.

Epstein-Barr virus (EBV) (also called human herpesvirus 4 [HHV-4]) is an enveloped gammaherpesvirus and one of only eight human herpesviruses (1). The two major cell types that EBV infects are epithelial cells and B cells. EBV enters these different cell types by the concerted effort of three or four essential viral glycoproteins, depending on the cell type (1, 2). Virions produced from B cells contain gB and gHgL which mediate fusion of virions with epithelial cells (1, 2). In contrast, virions produced from epithelial cells also contain gp42 in addition to gB and gHgL (1, 2). It is gp42 that activates entry into B cells by binding to human leukocyte antigen (HLA) class II (1–4), whereas gHgL binding to integrins allows entry into epithelial cells (1, 2, 5–7). gp42 binding to gHgL also inhibits entry into epithelial cells, and thus serves as a viral tropism switch (5).

When gp42 is synthesized, it is a 223-amino-acid type II membrane protein (1, 8). A soluble form of gp42 is generated by proteolytic cleavage in the endoplasmic reticulum (ER) with cleavage occurring around amino acids 40, 41, and 42 (9). Deletion of the predicted cleavage site (residues 37 to 41) results in a membrane-bound form of gp42 that significantly abrogates B cell fusion (10), verifying the results of earlier studies indicating that soluble gp42 (sgp42) functions in B cell fusion (11, 12). Soluble gp42 mutants, containing an EBV gB signal sequence followed by gp42 N-terminal deletions up to residue 46, are functional for fusion (10). Deletions up to residue 52, however, are not well tolerated, as this affects the region within gp42 (amino acids 44 to 81) that is important for gHgL binding (10, 13). In addition to being essential for B cell fusion, gp42 inhibits epithelial cell fusion (5, 11) presumably by binding an overlapping region in gHgL that also binds the receptor for epithelial cell fusion (11, 14, 15). Mutational
studies have shown that amino acids within residues 44 to 81 of the N terminus of soluble gp42 interact with domain II (DII) of gHgL, and co-crystallization studies between gp42 and HLA-DR1 have shown that the gp42 C-type lectin domain (CTLD) (solid blue circle in Fig. 1A) interacts with HLA class II (13, 16). Crystalization studies have shown that the EBV gHgL structure is comprised of four sequential semiautonomous domains (D1, DII, DIII, and DIV) and gL forms a stable heterodimer with gH and is integral to DII folding and structure (17). The prominent KGD loop in DII has been implicated in binding residues 62 to 66 of gp42 (Fig. 1A), as well as the gHgL epithelial receptor integrin αvβ6, αvβ8, or αvβ5 (7, 15). Membrane-bound gp42 (Fig. 1B), which contains a deletion of the predicted cleavage site from residues 37 to 41 (d37-41gp42), efficiently binds gHgL but is unable to mediate fusion (10, 18).

We recently studied the assembly and architecture of the EBV B cell entry complex by electron microscopy (19). In this structure, gp42 binds to gHgL and HLA class II. It is the binding of gp42 to the B cell receptor HLA class II that is thought to trigger B cell fusion by the concerted action of gHgL and gB (1, 2). This structure, which likely represents an intermediate state of EBV entry, suggests that the B cell fusion-triggering complex may bring the two membrane bilayers of the virion and cell into proximity, orienting critical regions of the N- and C-terminal ends of gHgL to promote the activation of gB and efficient membrane fusion (19). Furthermore, we found that in this structure, gHgL also interacts with a functionally important hydrophobic pocket on gp42 (19).

In our current studies, we sought to determine whether gp42 cleavage is essential for function, allowing flexibility so that the protein components of the B cell fusion-triggering complex can interact efficiently with each other to enable the successful triggering of membrane fusion.
RESULTS

Construction of d37-41gp42 titin linker insertion mutants. To investigate whether membrane-bound gp42 sterically inhibits gHgL, shown bent and constrained in Fig. 1B, we designed a number of gp42 mutants (Fig. 1D). In the constrained conformation, interaction of gHgL with other viral glycoproteins, such as gB, or viral receptors may be prevented or reduced, resulting in a reduction in viral membrane fusion with the host cell membrane. To determine whether potential steric inhibition of membrane-bound gp42 could be relieved by increasing the distance or spacing of gp42 from the membrane, we inserted structural linkers immediately downstream of the gp42 transmembrane domain (red boxes in Fig. 1C and D). We chose to insert one, two, or four copies of the 27th immunoglobulin-like (I27) domain of the muscle protein titin. The titin Ig-like structure has been solved by both nuclear magnetic resonance (NMR) and crystallization, indicating the following: (i) The titin sequence consists mostly of repetitive motifs of tandem immunoglobulin-like (Ig-like) modules that are conformationally rigid segments interspersed with pliant hinges. (ii) The Ig-like domains consist of a beta sandwich formed by two four-stranded sheets which results in a domain with the N and C termini on opposite sides of the domain separated by 44 Å. (iii) The Ig-like domains fold independently in solution (20–23).

In addition, the naturally occurring I27 domains have previously been successfully used to study the spacing requirements for proteasome binding and the initiation of protein degradation (22). Thus, addition of one to four titin I27 domains would space membrane-bound gp42 anywhere from approximately 44 to 176 Å from the membrane, and a particular spacing may be predicted to alleviate steric hindrance and allow for functional interaction with gHgL or other viral glycoproteins and cellular receptor(s) to promote fusion.

d37-41gp42 titin linker insertion mutants are expressed. Plasmids encoding each of the linker mutants were transfected into Chinese hamster ovary cells (CHO-K1) cells, and protein expression was monitored in cell lysates (Fig. 2A) and medium supernatants (Fig. 2B) 48 h posttransfection by SDS-PAGE, followed by Western blotting using a polyclonal antibody specific for gp42. As expected, wild-type (wt) gp42 produced both soluble gp42 (sgp42) and cell-associated gp42 with most of the gp42 detected in the medium supernatants. Also as expected, a previously published soluble gp42 (d36FLAGgp42) (24) produced only soluble gp42, whereas d37-41gp42 predominantly produced cell-associated gp42. The linker mutants produced cell-associated gp42 of approximately the predicted size; however, surprisingly, soluble gp42 was also produced for each of the mutants. Deletion of a larger region (residues 22 to 46) around the cleavage site and insertion of the linkers did not alter this result but decreased expression, so these clones were not analyzed further (data not shown), and we decided to focus on the first group of mutants (Fig. 1D) that had higher levels of expression.

d37-41gp42 titin linker insertion mutants are expressed on the surface regardless of the presence of gHgL or gB and are not functional in cell-cell fusion. To study whether the gp42 linker mutants were expressed on the cell surface, their expression was analyzed in the presence and absence of gHgL and gB by cell-based enzyme-linked immunosorbent assay (cELISA). We cotransfected CHO-K1 cells with plasmids encoding vector, wt gp42, d37-41gp42, d37-41gp42 (I27-1), d37-41gp42 (I27-1,2) and d31-44gp42 (I27-1,2,3,4) with or without gHgL and gB. We found that the presence or absence of gHgL and gB did not make a significant difference in the surface expression of any of the membrane-bound mutants which were expressed at levels similar to that of wt gp42 (set at 100%) (Fig. 3). For a control, the soluble
FLAGgp42 was expressed on the surface only in the presence of gHgL. In addition, we confirmed that the linker mutants were expressed on the surface of the cell by biotinylating the cell surface and performing immunoprecipitation with antibody to gp42 (F-2-1) (data not shown).

However, when we tested the ability of the mutants to mediate fusion of Daudi B cells expressing T7 polymerase in the presence of gHgL, gB, and a T7 polymerase-driven luciferase reporter, we found that d37-41gp42, d37-41gp42 (I27-1), d37-41gp42 (I27-1,2) and d31-44gp42 (I27-1,2,3,4) were all significantly reduced in fusion (Fig. 3). As expected from previous studies, d37-41gp42 fusion function was greatly reduced compared to wt gp42 (10). d37-41gp42 (I27-1), d37-41gp42 (I27-1,2), and d31-44gp42 (I27-1,2,3,4) mediated fusion at levels less than 20% of wt gp42. Adding one to four I27 domains to the N terminus of sgp42 (d36FLAGgp42) did not significantly alter fusion compared to wild-type sgp42 (data not shown), indicating that the addition of linkers alone does not make gp42 nonfunctional.

d37-41gp42 titin linker insertion mutants bind to exogenously expressed gHgL and HLA class II. To determine whether the reduction in fusion was due to altered or decreased binding to gHgL or HLA class II, CHO-K1 cells were transiently transfected with 2 μg of the gp42 linker mutants either in the presence or absence of gHgL and gB (0.5 μg each) and T7 luciferase (0.8 μg). Eighteen hours posttransfection, 40,000 cells were transferred to each well of a black 96-well plate and overlaid with 40,000 Daudi B cells for fusion assay. Eighteen hours after overlay, cells were washed with phosphate-buffered saline (PBS) and lysed for 10 min with 50 μl of passive lysis buffer (Promega) per well. Luciferase activity was measured with a PerkinElmer Victor plate reader immediately after addition of 50 μl/well of luciferase reagent (Promega). Eighteen hours posttransfection, 80,000 cells were transferred to each well of a clear 96-well plate, and eighteen hours later, surface expression was determined by cELISA using anti-gp42 (3H3), fixation, secondary biotinylated anti-mouse IgG antibody (Sigma), tertiary streptavidin-horseradish peroxidase (GE Healthcare) and TMB (3,3′,5,5′-tetramethylbenzidine) after TMB one-component substrate (BioFX). Absorbance readings were taken at 380 nm using a PerkinElmer Victor plate reader. Binding was normalized to wild-type gp42 binding levels, which were set at 100%. Data shown are representative of the results from three independent experiments.

FIG 3 The gp42 linker mutants are expressed independently of gHgL expression and are not functional in B cell fusion. CHO-K1 cells were transiently transfected with 2 μg of the gp42 linker mutants either in the presence or absence of gHgL and gB (0.5 μg each) and T7 luciferase (0.8 μg). Eighteen hours posttransfection, 40,000 cells were transferred to each well of a black 96-well plate and overlaid with 40,000 Daudi B cells for fusion assay. Eighteen hours after overlay, cells were washed with phosphate-buffered saline (PBS) and lysed for 10 min with 50 μl of passive lysis buffer (Promega) per well. Luciferase activity was measured with a PerkinElmer Victor plate reader immediately after addition of 50 μl/well of luciferase reagent (Promega). Eighteen hours posttransfection, 80,000 cells were transferred to each well of a clear 96-well plate, and eighteen hours later, surface expression was determined by cELISA using anti-gp42 (3H3), fixation, secondary biotinylated anti-mouse IgG antibody (Sigma), tertiary streptavidin-horseradish peroxidase (GE Healthcare) and TMB (3,3′,5,5′-tetramethylbenzidine) after TMB one-component substrate (BioFX). Absorbance readings were taken at 380 nm using a PerkinElmer Victor plate reader. Binding was normalized to wild-type gp42 binding levels, which were set at 100%. Data shown are representative of the results from three independent experiments.
bars) show slight competition with increasing wild-type gp42 (0.01, 0.1, and 1.0 μg/H9262). Lanes 6 to 8 (white bars) show that with the highest concentration of d37-41gp42 (1.0 μg/H9262) [lane 8], there is approximately 25% reduction in fusion. In contrast, lanes 9 to 11, 12 to 14, and 15 to 17 (white bars) show that with the highest concentration (1.0 μg) of d37-41gp42 (I27-1), d37-41gp42 (I27-1,2), and d31-44gp42 (I27-1,2,3,4), respectively, there is approximately 90% reduction in fusion, suggesting that the gp42 linker mutants are better at blocking fusion than the d37-41gp42 mutant and that this effect is dose dependent.

To determine whether the dominant-negative effect was alleviated by adding sgp42, we transfected CHO-K1 cells with d36FLAGgp42 and isolated soluble gp42 from supernatants 48 h posttransfection. Either 2 or 20 μl of supernatant (gray bars and black bars, respectively, in Fig. 5) was added to duplicate and triplicate plates of the fusion assay described above (with the exception of lane 1). We found that increasing the amount of sgp42 increased the level of fusion in a dose-dependent manner when no gp42 was transfected in the plate (lane 18, compare white, gray, and black bars). However, fusion did not increase with addition of sgp42 when any of the membrane-bound gp42s were transfected first (lanes 6 to 17 [compare white, gray, and black bars]). This confirms that d37-41gp42 and the linker mutants work in a dominant-negative manner, and once gHgL is bound by membrane-bound gp42, sgp42 cannot compete for binding.

DISCUSSION

We recently reported that deletion of the gp42 N-terminal cleavage site blocks gp42 function in fusion (10), indicating that when bound to a membrane by the transmembrane domain, gp42 does not function in fusion, despite binding to gHgL. This is in contrast with gD, the functional homolog of gp42 found in herpes simplex virus (HSV) that functions both as a soluble form and a membrane-bound form, although at reduced levels when soluble (26). Presumably gp42 is cleaved in the ER and is membrane bound by virtue of being tethered to gHgL, which contains a transmembrane domain.

To determine whether membrane-bound gp42 could be made functional by increasing separation from the membrane, the naturally occurring structured immunoglobulin-like domain I27 from the multidomain muscle protein titin was inserted between the gp42 transmembrane domain and the gHgL binding region of gp42. While I27 is rigid in nature, the N and C termini that connect each I27 domain have some flexibility (20–23). We predicted that the addition of one, two, and four I27 domains would approximately match, double, or quadruple the wild-type gp42 spacing from the membrane and thus potentially relieve any steric hindrance imposed on gHgL by being membrane bound. Our results show that while all the gp42 linker mutants are expressed on the surface, cleavage occurs in the absence of the canonical cleavage site. Because the cleavage products were identical in size, it is likely that the cleavage occurs after the linker additions. We tried to alleviate this problem by making another series of insertion mutants with a larger deletion of the region that is cleaved in gp42, but these mutants also had some cleavage and were poorly expressed. Of the mutants we studied in detail, each of the soluble gp42 variants produced were functional in fusion when added as an overlay to gHgL/gB-expressing cells (data not shown). However, cotransfection of the linker mutants together with wild-type gH/gL and gB resulted in significantly reduced fu-
sion compared to wild-type gp42 or sgp42. These results suggest that the membrane-bound linker mutants bind gHgL prior to cleavage and that this binding is not readily replaced by functional, soluble gp42, even though it is produced by the mutants. This finding is reinforced by the observation that the titin linker mutants effectively act in a dominant-negative manner, even more efficiently than the d37–41gp42 cleavage mutant, perhaps due to their wild-type-like levels of binding gHgL as well as HLA. Overall, our results suggest that the structured titin linkers that match, double, or quadruple the presumed wild-type gp42 spacing from the membrane do not alleviate the steric hindrance imposed on gHgL by membrane-bound gp42 to sufficiently promote fusion similar to that induced by wild-type gp42 or sgp42. These results indicate that the architecture of the B cell fusion-triggering complex (19) has additional, specific requirements for membrane fusion to proceed. Tethering gp42 to the membrane through its N terminus may generate a steric block that could interfere with membrane fusion by blocking the recruitment of gB or potentially altering other gHgL interactions that promote gB activation.

MATERIALS AND METHODS

Cells and antibodies. Chinese hamster ovary cells (CHO-K1) were grown in 75-cm² cell culture flasks (Corning) in Ham’s F-12 medium (BioWhittaker) supplemented with 10% fetal bovine serum (HyClone) and 1% penicillin-streptomycin (BioWhittaker). Trypsin-Versene (BioWhittaker) was used to detach adherent cells. Polyclonal anti-gp42 antibody serum (PB114) was used as previously described (27). Monoclonal anti-gp42 antibody (3H3) (catalog no. ab24265; Abcam) and anti-FLAG-M2 (catalog no. F1804; Sigma-Aldrich Chemical Company) were used to detect HLA class II and gp42, respectively. Mono-
clonal anti-FLAG M2 antibody (F1804) and polyclonal anti-FLAG antibody (F7425) were obtained from Sigma-Aldrich Chemical Company.

**Plasmids.** The 27th immunoglobulin-like (I27) domain of muscle protein titin (schematically represented in Fig. 1D) was PCR amplified with sequence-specific primers containing PstI-modified ends and cloned so that one, two, and four copies of I27 (22) were obtained. The I27 domains were unidirectionally cloned into a unique PstI site (residue 29) of d37-41gp42 in the pCAGGS plasmid. The linkers were cloned just downstream of the transmembrane domain which ends at residue 22 and upstream of the gHgL binding domain which begins at residue 44. Other functional regions of gp42 were not disturbed. The ligated products were transformed into competent Escherichia coli DH5α and selected on plates containing ampicillin. DNA was isolated from overnight cultures using the Qiagen miniprep kit, digested to confirm the presence of the insert and in both directions by the Northwestern Genomic Core Facility to confirm the resulting sequences. The resulting linker insertion sequences are shown in Table 1. Large-scale DNA preparations were isolated using Qiagen Endo-Free plasmid maxiprep kit and used in subsequent experiments. EBV gl in pCAGGS was previously described (28).

**Transfection and Western blotting.** CHO-K1 cells were transfected in Opti-MEM (Gibco) medium using Lipofectamine 2000 (Invitrogen) according to the manufacturer’s instructions. Briefly, 24 h after plating the cells in a six-well plate, various combinations of expression vectors were transfected with Lipofectamine in Opti-MEM overnight. The medium was changed 16 h posttransfection, and 1 h later, the cells were detached with 0.05% trypsin-EDTA. The cells were washed, fixed and incubated with biotinylated goat anti-mouse IgG (Sigma) for 30 min at room temperature, followed by incubation with streptavidin-horseradish peroxidase (HRP) (GE Healthcare) for 30 min at room temperature and with 3,3′,5,5′-tetramethylbenzidine (TMB) one-component HRP substrate (BioFX). Absorbance readings were taken at 380 nm using a Wallac-Victor plate reader (PerkinElmer).

**TABLE 1 d37-41gp42 titin linker insertion mutants**

| Mutant gp42       | Length of insert (no of amino acids) | Sequence of linker insert       |
|-------------------|--------------------------------------|---------------------------------|
| d37-41(I27-1)     | 92                                   | AILEVEKPLYGVLFVGATEHFEIELSEPVDHGQWKLGQPLTASPDAEIEDGKHHILILHNAQLGTMGEVSFAQ AANKSAANLKKVEKPR |
| d37-41(I27-1,2)   | 184                                  | AILEVEKPLYGVLFVGATEHFEIELSEPVDHGQWKLGQPLTASPDAEIEDGKHHILILHNAQLGTMGEVSFAQ AANKSAANLKKVEKPR |
| d37-41(I27-1,2,3,4)| 368                                  | AILEVEKPLYGVLFVGATEHFEIELSEPVDHGQWKLGQPLTASPDAEIEDGKHHILILHNAQLGTMGEVSFAQ AANKSAANLKKVEKPR |

*a* Titin immunoglobulin-like 127 domains were PCR amplified from clones kindly provided by Andreas Matouschek (21). Following PCR, bands representing one, two, and four titin domains were digested with PstI and ligated into a unique PstI in d37-41gp42.

**Cell-cell fusion assay.** CHO-K1 cells were transiently transfected as described above. The medium was changed 16 h posttransfection, the cells were detached with Versene, and 37,500 cells per well in triplicate were transferred to duplicate 96-well plates. One plate was used for cELISA (described above), and the other plate was overlaid with equal numbers of CHO-K1 target cells transfected with T7 polymerase luciferase reporter plasmid and relevant test plasmids and Daudi B cells expressing T7 polymerase. The total volume was adjusted to 150 μl with complete Ham’s F-12 medium. Eighteen to 20 h after the cells were laid over the plate, the cells were washed with PBS and lysed for 10 min with 50-μl passive lysis buffer (Promega) per well. Luciferase activity was measured with a PerkinElmer Victor plate reader immediately after the addition of 50 μl/l well of luciferase reagent (Promega).

**HLA class II and gH/gL binding.** cELISA was used to determine soluble gH (sgH) binding or soluble HLA class II (sDQ2-α1l) (sDQ2-α1l). Briefly, CHO-K1 cells were cotransfected in a six-well dish with each of the FLAG-tagged mutants and wild-type gp42, gB, or wild-type gH. The medium was changed 16 h posttransfection, and 1 h later, the cells were detached with Versene, counted using a Beckman Coulter Z1 particle counter, 37,500 cells were transfected to each well of a 96-well plate, and the total volume was adjusted to 150 μl with complete Ham’s F-12 medium. Twenty-four hours later, the cells were washed once with PBS. To determine whether the significantly reduced fusion was due to altered or decreased binding to either gHgL or HLA class II, we transfected CHO-K1 cells with a previously purified sDQ2-α1l (HLA class II) that was kindly provided by Elizabeth Mellins at Stanford University and used to monitor HLA class II binding (19), and a previously described published soluble gHgL (sgHgL) collected from culture supernatant was used to monitor gH/gl binding (25). CHO-K1 cells were transfected with the vector control, wt gp42, d37-41gp42, d37-41gp42 (I27-1), d37-41gp42 (I27-1,2), d37-41gp42 (I27-1,2,3,4), and 24 h later, they were overlaid with sgHgL or HLA class II for 1 h at 4°C. Bound protein was then determined by cELISA either using an anti-HLA class II DQ antibody (1a3) (catalog no. ab24265; Abcam) and anti-FLAG-M2 (catalog no. F1804; Sigma) which recognized the epitopetagged gH/gl. The cells were washed, fixed, and incubated with biotinylated goat anti-mouse IgG (Sigma) for 30 min at room temperature, followed by incubation with streptavidin-horseradish peroxidase (HRP) (GE Healthcare) for 30 min at room temperature and with 3,3′,5,5′-tetramethylbenzidine (TMB) one-component HRP substrate (BioFX).
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