Insulin Degrading Enzyme Induces a Conformational Change in Varicella-Zoster Virus gE, and Enhances Virus Infectivity and Stability

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Abstract

Varicella-zoster virus (VZV) glycoprotein E (gE) is essential for virus infectivity and binds to a cellular receptor, insulin-degrading enzyme (IDE), through its unique amino terminal extracellular domain. Previous work has shown IDE plays an important role in VZV infection and virus cell-to-cell spread, which is the sole route for VZV spread in vitro. Here we report that a recombinant soluble IDE (rIDE) enhances VZV infectivity at an early step of infection associated with an increase in virus internalization, and increases cell-to-cell spread. VZV mutants lacking the IDE binding domain of gE were impaired for syncytia formation and membrane fusion. Pre-treatment of cell-free VZV with rIDE markedly enhanced the stability of the virus over a range of conditions. rIDE interacted with gE to elicit a conformational change in gE and rendered it more susceptible to proteolysis. Co-incubation of rIDE with gE modified the size of gE. We propose that the conformational change in gE elicited by IDE enhances infectivity and stability of the virus and leads to increased fusogenicity during VZV infection. The ability of rIDE to enhance infectivity of cell-free VZV over a wide range of incubation times and temperatures suggests that rIDE may be useful for increasing the stability of varicella or zoster vaccines.

Introduction

Varicella-zoster virus (VZV), a member of the alpha-herpesvirus family, is the etiologic agent of chickenpox and shingles. In humans, cell-free virions are released from skin lesions and are transmitted to epithelial cells in the respiratory tract of susceptible hosts [1]. In cell culture, however, no cell-free infectious viruses are spontaneously released, and infection is exclusively by cell-to-cell spread of virus. While cell-free virus can be obtained by sonication of infected cells, the lack of high titer cell-free virus has hindered the progress of studies to define the mechanism by which VZV enters into target cells.

Previous studies have identified cellular molecules that are important for entry of VZV into cells. Cation-independent mannose 6-phosphate receptor (MPR) has been proposed to facilitate an early step of VZV infection [2]. Previous studies from our laboratory showed that insulin-degrading enzyme (IDE), a member of the zinc metalloproteinase family, is a putative cellular receptor for VZV [3]. Down-regulation of IDE by specific siRNA, inhibition of IDE activity with bacitracin, or blocking IDE with antibody inhibited VZV infection and impaired cell-to-cell spread of the virus. Over-expression of human IDE by transfection into cell lines resulted in increased entry of both cell-free and cell-associated virus. VZV glycoprotein E (gE), which is essential for virus infectivity [4,5], interacts with IDE through a binding domain located at the amino terminus of the ectodomain of gE, that is not conserved in other human herpesviruses [3,6,7,8]. VZV deleted for the IDE binding domain in gE is impaired for infectivity of cell-free virus [5] and shows reduced cell-to-cell spread of virus both in vitro and in human skin xenografts in SCID mice [5,8]. Here, we show that the interaction of IDE with gE is important for VZV-induced syncytia formation and fusogenicity, and that recombinant soluble IDE (rIDE) modifies gE, induces a conformational change in gE, enhances VZV infectivity, and stabilizes cell-free virus.

Results

rIDE augments cell-free VZV infectivity at an early stage of infection and enhances stability of cell-free virus

The open reading frame of human IDE contains two ATGs near the amino terminus that could serve as translation initiation codons. Previous studies with cloned IDE cDNA showed that the second ATG encoding amino acid 42, which better matches a Kozak consensus sequence, serves as the canonical start site for translation [9,10]. Recombinant baculovirus was constructed to express human IDE with a hemaglutinin (HA) tag inserted after the second methionine (amino acid 42) of IDE driven by polyhedrin promoter [3,11] (Fig. 1A). rIDE was expressed as a
110 kD protein (Fig. 1B), although gel filtration showed oligomerization of the protein as has been reported previously [12]. Incubation of rIDE with radiolabeled insulin resulted in a similar profile of degradation products as seen with endogenous IDE from rat liver [13] or another form of recombinant IDE [14] (Fig. 1C). rIDE had insulin degrading activity similar to recombinant 6HisFlag-IDE (Fig. 1D).

Previously we showed that IDE interacts with the extracellular domain of VZV gE, that a gE mutant lacking amino acids 32–71 cannot bind IDE, and that a gE mutant lacking amino acids 163–208 is unable to bind to VZV gI and shows enhanced binding to IDE [6]. Here we found that rIDE also formed a complex with the extracellular domain of gE, that rIDE did not interact with gE lacking the IDE binding domain, and that rIDE interacted to a greater extent with the mutant gE that does not bind gI (Fig. 1E).

Previously, we reported that purified endogenous IDE protein extracted from liver blocked VZV infection, while rIDE from cloned cDNA expressed in baculovirus-infected cells enhanced VZV infectivity [3]. The difference in the activity of these two proteins could be due to the difference in their amino termini. rIDE begins at a canonical initiation codon at amino acid 42. Purified rIDE showed a single band of 110 kD while native IDE purified from liver is actually a complex mixture of IDE forms. To further investigate the effect of rIDE on VZV infectivity, we incubated cell-free VZV ROka-lacZ with rIDE or control proteins for 15 min at 37°C before infecting human melanoma cells with the virus. Three days later, the cells were fixed and treated with X-gal and blue foci were counted. rIDE significantly enhanced VZV infectivity compared with control protein or SPGC buffer alone (P < 0.001 for rIDE vs. control protein, Student t test, Fig. 2A). Enhancement of infectivity was not observed when rIDE was added at 4 hr post-infection which bypasses the entry step (P < 0.0001 for rIDE added at 15 min before infection vs. rIDE added 4 hr after infection by Student t test, Fig. 2A). Denaturation of rIDE with 12.5 mM citric acid at pH 2.3 followed by neutralization with 0.5 M HEPES buffer at pH 9.0 to a final pH 7.0, inactivated the enhancing activity of IDE. These results indicate that the native form of rIDE is required to enhance infectivity and that rIDE functions at an early stage of infection.

Incubation of rIDE with cell-free VZV at 37°C for 15 min before adding the virus to cells enhanced infectivity in a dose-dependent manner (P = 0.0002 by ANOVA) (Fig. 2B).

Since VZV is highly cell-associated in vitro, and cell-free virus produced by sonication is very labile and its infectivity declines...
quickly over time, we tested whether incubation of cell-free VZV with rIDE enhances stability of the virus. rIDE increased virus infectivity when incubated with cell-free ROka-lacZ for 1 or 3 hr at 37°C compared with buffer or with filtrate passing through the rIDE purification (Fig. 2C). rIDE also enhanced cell-free VZV infectivity and/or stability after incubation for 18 hrs at 25°C prior to infecting cells (P = 0.002 for rIDE vs. buffer or controls, Student t test) (Fig. 2A last two columns and Fig. 2D). Enhancement of virus stability by incubating rIDE with VZV was observed over a range of incubation periods at room temperature (Fig. 3A, P = 0.0002 for rIDE vs. BSA control for combined data from 11 experiments (each with ≥2 replicates). Incubation of rIDE and cell-free virus at 4°C for 60 min before infection of cells also augmented VZV infectivity (Fig. 3B). These results indicate that rIDE enhances both infectivity and stability of cell-free VZV over a wide range of incubation times and incubation temperatures.

To further determine if the effect of rIDE on VZV infectivity is at an early stage of virus infection, we performed quantitative PCR for VZV DNA to determine the copy number of viral genomes inside the cells 90 min after infection. Incubation of rIDE with virus prior to infection resulted in increased numbers of VZV genomes inside the cells, compared with virus incubated with a control protein, indicating that rIDE increases virus internalization (P<0.0001 for rIDE vs. P7.5, Fig. 4A). Addition of rIDE at 2.5 hr after infection with VZV resulted in an increase in the size of...
plaques (P<0.0001 for both rIDE vs. BSA control and rIDE vs. Buffer, Fig. 4B), suggesting that rIDE not only augments infectivity of cell-free virus at entry but also promotes infectivity through enhanced cell-to-cell spread of virus. This is consistent with our earlier observation that IDE is important for infection with both cell-free and cell-associated VZV [3].
Since the prior experiments were all performed using recombinant-derived VZV, we also tested the effect of rIDE on non-recombinant viruses. Cell-free zoster vaccine virus, which contains various stabilizers, was reconstituted according to the manufacturer’s instruction, and incubated with rIDE or control proteins for 30 min at 37°C. rIDE increased the infectivity of zoster vaccine virus (P < 0.0001 for rIDE vs. buffer, bovine serum albumen (BSA), or P7.5 control protein by Student t test; Fig. 4C).

rIDE modifies VZV gE and induces a conformational change in gE

IDE is a metalloproteinase. While IDE interacts with a variety of substrates, it cleaves only a subset of its substrates such as insulin and β-amyloid protein at perceptible rates [15,16]. Previously, we incubated the extracellular domain of gE (gEt) with IDE protein extracted from liver and did not detect cleavage or degradation of gEt using a monoclonal antibody [3]. To determine if rIDE modifies gE, we incubated biotin-labeled gEt protein with rIDE or control proteins (P7.5-Fc which encodes vaccinia virus P7.5 protein or BZLF2-Fc which encodes Epstein-Barr virus BZLF2 protein) at 37°C, 22°C, or 4°C for 30 min. The proteins were then boiled in sample buffer, separated on an SDS-PAGE gel, and gEt was stained with streptavidin conjugated-horse radish peroxidase. Incubation of gEt with rIDE, but not with the control proteins, resulted in a slightly smaller sized band than full length gEt, indicating that rIDE modifies gEt (Fig. 5A). While it was somewhat surprising that rIDE modified gE after incubation at 4°C, this is consistent with our observations that rIDE enhanced VZV infectivity at 4°C (Fig. 3B).

Many viral glycoproteins undergo conformational changes after binding their receptor or co-receptor which results in the exposure of fusion peptides embedded in the viral glycoproteins that initiate a series of events leading to fusion between the viral envelope and the cell membrane [17,18]. Since VZV gE has an important role in syncytia formation and membrane fusion [19], we postulated that gE might undergo a conformational change after binding to rIDE. Limited exposure of ligand-receptor complexes to proteinase has been used to detect receptor-induced conformational changes in several proteins, including avian sarcoma/leukosis virus (ALV) envelope protein complexed with its receptor Tva and spike protein of mouse hepatitis virus complexed with soluble receptor, CEA-CAM1a [20,21,22]. The extracellular domain of gE was labeled with

Figure 5. rIDE modifies gEt and induces a conformational change in gEt. (A) Biotin labeled gEt-Fc protein (arrow) was incubated with IDE at the indicated temperature for 30 min. After electrophoresis and transfer onto a nitrocellulose membrane, proteins were visualized with streptavidin-conjugated-horse radish peroxidase. (B) Biotin-labeled gEt-His protein (arrow) was incubated with buffer, control protein B2LF2 or P7.5, rIDE (produced in baculovirus) at 37°C for 30 min, followed by urea for 18 hr and thermolysin for 30 sec. The proteins were separated by electrophoresis, transferred to a nitrocellulose membrane and proteins were detected using streptavidin-conjugated horse radish peroxidase. (C) Biotin-labeled gEt-His protein (arrow) was incubated with rIDE or control proteins at 37°C for 30 min, followed by incubation with 4 μg/ml of endoproteinase Asp-N (Roche Applied Science, Indianapolis, IN) at 37°C for 45 sec. The digestion was then terminated by adding 0.5 M EDTA and samples were boiled in SDS-PAGE gel loading buffer with 2.5% 2-mercaptoethanol and separated by electrophoresis. Protein fragments were detected by streptavidin conjugated-horse radish peroxidase. (D) Binding of catalytically inactive IDE mutant protein IDE-E111Q to gE fails to induce a conformational change in gE. Biotin-labeled gEt-His protein (arrow) was incubated with rIDE (produced by baculovirus), IDE-E111Q (produced in bacteria), or IDE-w. t. (produced in bacteria), or negative control proteins as indicated at 37°C for 30 min, followed by thermolysin and processed as described in panel B. (E) Coomassie Blue stained SDS-PAGE gel showing the amount of IDE-E111Q and IDE-w.t. proteins used for pulse proteolysis assay in panel D. doi:10.1371/journal.pone.0011327.g005
target cells with ROka68D32-71-infected cells resulted in reduced IDE binding domain (amino acids 32 to 71) of gE. Incubation of VZV (ROka) or a mutant virus, ROka68D32-71 that lacks the impaired for membrane fusion and syncytia formation. A VZV mutant lacking the IDE binding domain of gE is reduced in the intensity of the 70 kDa gEt band (Fig. 5B and 5D). Similar results were also observed with endoproteinase Asp-N digestion (Fig. 5C). Therefore, rIDE triggered a conformational change in VZV gEt that confers an increased susceptibility to proteinase. Repeated attempts to determine a cleavage site on gEt by N-terminal protein sequencing failed to generate any sequences.

Glutamic acid 111 in the zinc-binding site of human IDE is a major catalytic residue for its insulin degrading function [23]. Mutation of glutamic acid 111 to glutamine in IDE results in a mutant protein, IDE-E111Q, which is catalytically inactive for insulin degradation. Although IDE-E111Q formed a complex with gEt (Fig. 6), it failed to elicit a conformational change in gEt as compared with rIDE or wild-type IDE in the thermolysin assay (Fig. 5D, E). These results suggest that the catalytic activity of IDE for insulin degradation might be important for inducing a conformational change in gEt.

A VZV mutant lacking the IDE binding domain of gE is impaired for membrane fusion and syncytia formation

To determine if the interaction of gE with IDE is important for membrane fusion, we used a reporter system in which VZV permissive HeLa cells were transfected with a plasmid encoding β-galactosidase under the T7 promoter (pG1N-T7-β-gal) and served as target cells. One day after transfection, the target cells were incubated for 20 hr with melanoma cells stably expressing T7 polymerase that contained equal titers (10⁷ PFU/ml) of control VZV (ROka) or a mutant virus, ROka68D32-71 that lacks the IDE binding domain (amino acids 32 to 71) of gE. Incubation of target cells with ROka68D32-71-infected cells resulted in reduced membrane fusion activity as measured by lower levels of β-galactosidase compared with ROka-infected cells (P = 0.0001 for ROka vs. ROka68D32-71 by Student t test; Fig. 7A). Similar results were observed with ROka68D32-71-GFP (which also expresses GFP) and ROka-GFP (Fig. 7B). Transfection of HeLa cells with vector control plasmid, in place of the reporter plasmid, followed by infection with VZV ROka resulted in a background level of fusion activity. Therefore, VZV mutants lacking the IDE binding domain of gE have reduced membrane fusogenicity.

To further examine the function of IDE in VZV infection, we analyzed syncytia formation in cells infected with ROka68D32-71 and ROka68D32-71-GFP. While the truncated gE in ROka68D32-71 virus is impaired for binding to IDE in a pull-down assay, the mutant virus is not impaired for maturation and egress to the cell surface [5]. Both ROka68D32-71 and ROka68D32-71-GFP showed smaller syncytia with fewer nuclei in the syncytia compared with their parental viruses, ROka or ROka-GFP (Fig. 8A and B). Cells infected with ROka68D32-71-GFP had significantly fewer nuclei per syncytium than cells infected with ROka-GFP when the total number of nuclei in syncytia were quantified using 3 dimensional reconstruction of sequential Z-sections of cells with confocal microscopy (Fig. 8C, p<0.001).

A VZV mutant lacking the IDE binding domain of gE accumulates on the cell surface or the cell-cell junction

To rule out the possibility that the reduced level of syncytia formation of the VZV mutant might be due to defective maturation and transport to cell surface and/or to cell-cell junctions, we performed immuno-electron microscopy studies. Melanoma cells infected with ROka68D32-71 showed large numbers of enveloped virions accumulating at cell-cell junctions (Fig. 9A) and on the cell surface (Fig. 9B). In 53 randomly selected immune-electron micrographs, there were a total of 873 gE-positive ROka68D32-71 virions on the cell surface and/or in cell junctions, compared with a total 140 gE-positive ROka virions in 41 pictures (Fig. 9C, P<0.0001 for ROka68D32-71 vs. ROka), indicating that the impairment in syncytia formation and cell-to-cell spread of the IDE binding domain mutant virus is not due to defective maturation with reduced transport to the cell surface or to cell-cell junctions. A previous study showed that VZV gE is myristylated [24]. Myristylation of proteins is important for their association with membranes. To rule out the possibility that deletion of the IDE binding domain of gE does not adversely affect gE myristylation, we labeled ROka or ROka68D32-71-infected cells with 3H-myristic acid and immunoprecipitated gE. gE from both parental and mutant VZV showed similar levels of myristylation, indicating that differences in myristylation do not explain the phenotype observed with the mutant virus (Fig. 10). Therefore, the IDE binding domain of gE is important for membrane fusogenicity, syncytia formation, and for cell-to-cell spread of the virus.

**Discussion**

We have shown that soluble rIDE enhances VZV infectivity and cell-to-cell spread in human melanoma cells at an early step of virus infection. rIDE interacted with gE to elicit a conformational change in gE and modified the size of gE. In addition, a VZV mutant virus lacking the IDE binding domain of gE was impaired for syncytia formation and membrane fusion, suggesting that IDE enhances infectivity and stability of the virus through eliciting a conformational change in gE and modulating fusogenicity during VZV infection.

![Figure 6. IDE mutant protein, IDE-E111Q binds to gEt. gEt but not control protein p7.5, pulled down both IDE-E111Q protein and wild-type IDE (arrow). doi:10.1371/journal.pone.0011327.g006](image-url)
Previous studies using fibroblasts and melanoma cells suggested that IDE functions as a cellular receptor for gE enhancing VZV infectivity and cell-to-cell spread in these two cell types [3,5]. However, a recent study found that although a VZV mutant with a deletion in the IDE binding domain of gE showed reduced cell-to-cell spread of virus in melanoma cells and impaired infectivity of
skin xenografts in vivo, it was not defective in infecting T cells [8], suggesting that IDE may facilitate virus entry in a cell-type dependent manner, similar to gp42 of Epstein-Barr virus [25]. For the closely related herpes simplex virus (HSV), cellular molecules have also been shown to preferentially mediate virus entry in different cell types; nectin-1 is important for entry of HSV in neurons, while HVEM is used for entry in lymphocytes [26].

VZV gE binds to IDE through its amino-terminal region, although the overall conformation of gE is important for its interaction with IDE [6,8]. The observation that gE produced by the baculovirus expression system is smaller in size than its counterpart produced from mammalian cells (Li et al., unpublished data) suggests that it is less glycosylated in insect cells. gE produced by baculovirus forms a complex with IDE [3], which implies that glycosylation of gE may not be important for interaction with IDE. Carpenter et. al. reported that IDE binds to the mature 98-kDa form of VZV gE under low salt conditions, but IDE binds only to the 73 kDa form of gE at high salt conditions [27]. They showed that tunicamycin blocks glycosylation of gE in the endoplasmic reticulum resulting in only production of the 73 kDa form of gE, and that in the presence of tunicamycin, gE was blocked in its transit out of the endoplasmic reticulum. Therefore they concluded that IDE must interact only with the 73 kDa form of gE in the cytosol and that this binding does not represent a receptor/ligand interaction. Interestingly, a previous study by these authors showed that the smaller form of gE is present on cell surface, recycles through the endocytotic pathway, and is incorporated into virions [28]. Studies from other groups also showed that a mixture of forms of gE ranging from ~50–100 kDa are present in purified virions [29,30,31]. Therefore, the 73 kDa form of gE is present in virions where it may interact with IDE.

Soluble receptor-mediated enhancement of virus infectivity has been reported for several viruses including HIV, avian leukemia/sarcoma virus, mouse coronavirus, [32,33,34,35] and the closely related herpes simplex virus (HSV). Nectin-1 and 3-O-sulfated heparan sulfate function as receptors for HSV [36,37]. Soluble 3-O-sulfated heparin sulfate triggers HSV entry into non-permissive cells and augments viral glycoprotein-mediated membrane fusion [38]. A soluble truncated form of nectin-1 lacking the transmembrane domain confers HSV susceptibility to non-permissive cells [39]. Further studies showed that the soluble V domain of nectin-1, [amino acid residues 1–123), promotes HSV entry. Interestingly, in this study neither free nor gD-bound sNec1123 was bound to the cell surface, suggesting that the interaction of soluble receptor with its gD ligand facilitates virus penetration by activating viral fusion

Figure 9. Accumulation of a VZV mutant deleted for the IDE binding domain of gE at the cell surface and at the cell-cell junction. Melanoma cells were infected with VZV ROka or ROka68D32-71, fixed, and incubated with monoclonal antibody to VZV gE followed by FluoroNanogold-conjugated anti-mouse antibody and visualized by transmission electron microscopy. (A) Representative virions at cell-cell junctions are indicated with arrows. Magnification 12000×. The insets in panel A provide higher-power images of selected virions. (B) Representative virions on the cell surface are indicated with arrows. Magnification 5000×. (C) Quantification of gE-positive virions on the cell surface and at cell-cell junctions. Multiple cells from the experiment in panel A were coded and observed under a Hitachi H7500 transmission electron microscope. gE positive viral particles at cell surface and/or junctions were counted.
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machinery rather than directly enhancing virus attachment to cells [40]. While rIDE increased VZV infectivity at an early stage of infection, treatment of radiolabeled VZV with rIDE did not enhance virus binding (Fig. 11). Thus, soluble rIDE appears to promote VZV infectivity by increasing the efficiency of entry of virus into cells, rather than enhancing attachment. This conclusion is further supported by the observation that rIDE increases internalization of VZV DNA within 90 min of initiating infection.

We found that rIDE induces a conformational change in VZV gE. Numerous studies have established that receptor binding-induced conformational change in viral glycoproteins is critical in initiating virus-mediated membrane fusion. The conformational change results in exposed viral fusion peptides and provides energy for the fusion process through refolding [18,41,42]. For larger viruses such as HSV, the fusion machinery consists of more than one glycoprotein working in concert [43,44]. Since VZV gE is not fusogenic when expressed alone, but does act as a fusogen in concert with other glycoproteins [45], it is possible that the conformational changes elicited by rIDE may either directly enhance fusogenicity of gE, or allow recruitment of glycoproteins gB and gH, to form a fusion complex.

In addition to the receptor-binding mediated conformational change of viral glycoprotein, sequential conformational changes induced by proteolysis are required during the entry process of several viruses. Proteases cleave viral fusion proteins, including glycoproteins, into mature, fusogenic forms and enhance virus binding and infectivity [46,47,48,49,50,51,52,53]. Our finding that rIDE, a metalloproteinase, triggers a conformational change in gE and modifies the size of gE suggests that a proteolytic activity of IDE might be important for VZV infectivity. This is further supported by the fact that the IDE mutant (IDE-E111Q) which is impaired for degradation of insulin, fails to induce a conformational change in gE. Since gE is much larger than insulin and gE lacks any detectable amino acid homology with other substrates of IDE, the interacting amino acid residues of IDE with insulin and gE are likely to be different. Therefore, there may be a proteolytic motif of IDE for gE, which is different from that for degrading insulin.

HSV enters cells through different pathways involving either the cell surface membrane or intracellular vesicles such as endosomes in a cell-type dependent manner [26,54,55]. Previous work by Hambleton and colleagues suggested VZV enters human fibroblasts through an endocytosis pathway [56]. Although we have shown that rIDE enhances VZV internalization, it is not clear whether this occurs at the cell surface or in endosomes. In most cell types, endogenous IDE localizes primarily in endosomes and other subcellular compartments with about 10% on the cell surface or in extracellular spaces. It is interesting to note that upon receptor binding, activation of coronavirus glycoprotein by extracellular proteases facilitates virus entry through the cell surface plasma membrane, while in the absence of extracellular proteases the virus enters the cell using the endosomal pathway [22].

We found that rIDE increased the stability of cell-free VZV. There are two possible explanations for how rIDE might enhance
VZV infectivity assay

Cell-free VZV virus encoding β-galactosidase or GFP was incubated with rIDE or other proteins and added to MeWo cells in SPGC buffer at 33°C for 90 min before changing to normal growth medium. Infected foci were scored at day 3–5 post-infection by staining with 5-bromo-4-chloro-3-indolyl β-D-galactopyranoside (X-gal).

Cell-free vaccine virus was reconstituted in sterile water (as recommended by the manufacturer) and added to MeWo cells as described above. Four days after infection, the cells were fixed and stained with anti-gE monoclonal antibody (Chemicon, Temecula, CA) followed by anti-mouse-Alexa 488 secondary antibody (Invitrogen, Carlsbad, CA) and infected foci were visualized by fluorescence microscopy.

Construction of recombinant IDE and IDE mutants

Construction of recombinant baculovirus expressing HA-tagged human IDE (rIDE) has been reported elsewhere [3] and was used to infect SF9 or H5 insect cells. Three days post-infection the cells were collected, treated with lysis buffer (25 mM Tris-HCl pH 7.4, 5 mM EDTA, 15 mM NaCl, and 0.1% NP40), and the lysate was centrifuged at 2,450 g for 30 min at 4°C. HA-tagged rIDE was purified from the cleared supernatant through an anti-HA conjugated affinity column (Sigma, St. Louis, MO) at 4°C. After extensive washing with PBS, rIDE was eluted with HA peptide (200 μg/ml in PBS). Excess HA peptide was removed using a Centriprep column with a 10 kDa molecular weight cut-off (Millipore, Billerica, MA). For some experiments HA-IDE was further purified through a gel filtration column in 25 mM Tris, pH 8.0, 150 mM NaCl buffer.

Expression and purification of polyhistidine-tagged human wild-type (IDE-w.t.) and mutant (IDE-E111Q) IDE proteins in E. coli were performed as previously described [12] with minor modifications. Briefly, plasmids pProEX-IDEwt and pProEX-IDE-E111Q [12] were propagated in E. coli Rosetta (DE3) cells and protein expression was induced with isopropyl-1-thiogalactoside at a final concentration of 200 μM for 16 hrs. The cells were then lysed in RIPA buffer (10 mM Tris-HCl, pH 8.0, 100 mM NaCl, 1 mM EDTA, 0.1% NP40, 0.5% deoxycholic acid, 0.5% SDS), and polyhistidine-tagged IDE was purified through a Talon metal affinity column (Clontech, Mountain View, CA), eluted with 250 mM imidazole, and dialyzed overnight in PBS at 4°C.

Purification and biotinylation of gEt protein

Insect SF9 cells were infected with a recombinant baculovirus expressing the extracellular domain of VZV gE with a C-terminal (histidine)6 tag [64]. Three days after infection, tissue culture supernatant was harvested and gEt protein was purified through a Talon metal affinity column (Clontech, Mountain View, CA).
After extensive washing with 5 mM imidazole diluted in PBS and PBS buffer alone, 0.5 mg/ml of EZ-Link Sulfo-NHS-Biotin in PBS (pH 8.0) (Pierce, Rockford, IL) was added to the resin and incubated for 30 min at 25°C followed by three washes with cold PBS. The biotinylated protein was then eluted with 250 mM imidazole and dialyzed overnight in PBS.

**ELISA assay to detect gE-Fc fusion proteins and rIDE interactions**

gE-Fc and control (vaccinia P7.5 and EBV BZLF2) Fc fusion proteins were described previously [3]. Serial dilutions of Fc fusion proteins were coated onto an ELISA plate at 4°C overnight. After washing with PBS containing 0.05% Tween-20, anti-human Fc-horseradish peroxidase was added (Pierce, Rockford, IL). The amount of Fc fusion protein, designated as specific Fc units, was measured spectrophotometrically at OD450 nm after adding substrate-chromogen TMB (Dakocytomation, Carpinteria, CA).

rIDE was coated onto an ELISA plate at 500 ng/well at 4°C overnight. After washing with PBS containing 0.05% Tween-20 and blocking with 5% BSA, Fc fusion proteins were added at the same specific units (see above) at room temperature for 45 min to allow binding, followed by addition of anti-human Fc-horse radish peroxidase and TMB substrate.

**Insulin degradation assay**

rIDE was purified from baculovirus-infected cells using an anti-HA affinity column and eluted with HA peptide [3]. IDE from rat liver was purified by multiple chromatographic steps as described previously [13]. Recombinant human IDE containing hexahistidine and FLAG epitopes fused to the amino terminus of IDE was expressed in HEK-293T cells and purified using cobalt affinity chromatography [14]. Protein concentrations were determined using the BCA protein assay kit (Pierce, Rockford, IL). For degradation assays, samples of IDE were incubated with 125I-labeled insulin for 15 min at 37°C. IDE preparations were each diluted to attain an equal level of insulin degradation. Samples were resolved by reverse-phase chromatography using a C8 column as described previously [65]. Fractions were collected and the radioactivity was counted. The degree of insulin degradation was defined as the proportion of remaining intact insulin to total radioactivity. The pmoles of insulin degraded in each sample was calculated from the specific activity of the 125I-labeled insulin.

**Myristylation assay of gE**

MeWo cells were infected with cell-associated VZV ROka or ROka68D32-71 for 24 hr, labeled with 3H-9,10 myristic acid (167 µCi/ml) (Perkin-Elmer, Waltham, MA) for 18 hr, and lysed in RIPA buffer. VZV gE was immunoprecipitated with gE monoclonal antibody (Chemicon, Temecula, CA) and protein A-Sepharose beads (Sigma-Aldrich, St. Louis, MO), and separated on a 4–20% SDS-PAGE gel. The gel was incubated with [3H]-ENHANCE (Perkin-Elmer, Waltham, MA) according to manufacturer’s instructions before autoradiography.

**Immuno-electron microscopy**

Human melanoma cells were seeded on Theranox coverslips (Nalge Nunc International, Rochester, NY) and infected with cell-associated VZV ROka or parental virus (ROka). When similar CPE was noted, cells were fixed with 2% paraformaldehyde (Electron Microscopy Sciences, Hatfield, PA) on ice for 20 min. The coverslips were then incubated with 5% milk, and stained with anti-gE antibody (Millipore, Billerica, MA) and FluoroNanogold conjugated-anti-mouse Fab’-AlexaFluor488 (Nanoprobes, Yaphank, NY) followed by a final fixation with 4% paraformaldehyde, 2.5% glutaraldehyde in 0.1 M sodium cacodylate buffer overnight at 4°C. Samples were washed three times with distilled water followed by 0.01 M sodium citrate and silver enhancement for 4 min with HQ, silver reagents (Nanoprobes, Yaphank, NY). Samples were post-fixed with 1.0% osmium tetroxide/0.8% potassium ferrocyanide in 0.1 M sodium cacodylate, dehydrated with a graded ethanol series, and embedded in Spurr’s resin. Thin sections were cut with an RMC MT-7000 ultramicrotome (Ventana, Tucson, AZ), and stained with 1% uranyl acetate prior to viewing at 80 kV on a Hitachi H-7500 transmission electron microscope (Hitachi, Tokyo, Japan). Digital images were acquired with a Hamamatsu XR-100 bottom mount digital camera system (Advanced Microscopy Techniques, Danvers, MA) and processed using Adobe Photoshop v. 7 (Adobe Systems Inc, San Jose, CA).

**Real-time PCR to detect VZV internalization**

VZV internalization was assayed by modifying a previous protocol used to determine internalization of Kaposi’s sarcoma-associated herpesvirus [66]. Cell-free VZV preparations, filtered through a 5.0 µM non-pyrogenic filter (Pall Corporation, Cornelius, OR), were added to MeWo cells seeded in 6-well plates in SPGC buffer on ice for 60 min to allow binding. Cell entry was initiated by raising the incubation temperature to 37°C for the indicated time. Extracellular virus was removed by treating the cells with 300 µg/ml heparin in PBS on ice for 20 min, washing with PBS, inactivating virus with low pH Na-Citrate buffer (40 mM sodium citrate, 10 mM KCl; 135 mM NaCl, pH 3.6) at room temperature for 2 min [67], and washing with PBS. Extracellular VZV viral DNA was removed by incubating the cells with 100 µg/ml proteinase K in HBSS buffer (Invitrogen, Carlsbad, CA) with 1 mM CaCl₂ at room temperature for 5 min, washing with PBS, and adding 4 units of DNase I per sample at room temperature for 10 min (Roche Applied Science, Indianapolis, IN). Intracellular VZV viral DNA was then extracted together with cellular genomic DNA using a DNeasy blood & tissue kit (Qiagen, Valencia, CA) following the manufacturer’s protocol. The copy number of intracellular VZV genomic DNA was determined by real-time PCR using primers and a probe targeting VZV ORF62 as described previously [68]. Serial dilutions of plasmid pCMV62 were used to generate a standard curve, and copy numbers of VZV genomic DNA were normalized using the copy number of human RnaseP DNA amplified from the same samples. The effectiveness of heparin treatment and acid inactivation to remove un-internalized virus, and proteinase K followed by DNase I digestion to remove extracellular viral DNA was verified by quantitative PCR with DNA extracted from the virus-bound cells (virus binding on ice for 60 minutes without shifting to 37°C).

**Limited proteolysis to detect conformational changes in ligand-receptor interactions**

Protease treatment ([20] of gE) was performed with the following modifications. Biotinylated gEt protein was incubated with rIDE or control proteins at room temperature or 37°C for 30 min followed by adding equal volume of 6 M urea for 18 hr at room temperature. Thermolysin diluted in buffer (2.5 M NaCl, 10 mM CaCl₂) was added at 0.2 µg/ml at room temperature for 30 sec. The digestion was then terminated by adding 0.5 M EDTA and samples were boiled in SDS-PAGE gel loading buffer with 2.5% 2-mercaptoethanol and separated by electrophoresis. Protein fragments were detected by streptavidin conjugated-horse radish peroxidase.
Membrane fusion assay

A membrane fusion assay using a β-galactosidase reporter was adapted from Feng et al. [69]. The pG1N-T7-β-gal plasmid encodes β-galactosidase under the T7 promoter and was a gift from Ed Berger (niaid, NIH, Bethesda, MD) [70]. Plasmid pAR3126 encodes T7 polymerase driven by an SV40 early promoter [71] and was kindly supplied by William Studier (Brookhaven National Laboratory, Upton, New York). McWo cells stably expressing T7 polymerase were generated by cotransfecting pAR3126 and PCIt-Neo (Promega, Madison, WI) followed by neomycin selection at 1.0 mg/ml. VZV ROka or ROka68 was propagated and titrated to equivalent PFU (plaque forming units) in the MeWo cells expressing T7 polymerase and served as effector cells. VZV permissive HeLa cells were transfected with pG1N-T7-β-gal for 24 hr and served as target cells. Effector cells (VZV-infected MeWo cells expressing T7 polymerase) were mixed with target cells (HeLa cells with the β-galactosidase gene driven by the T7 promoter) for 20-24 hrs at 37°C. Cells were then lysed in PBS containing 0.1% NP40, and β-galactosidase expression resulting from fusion of effector with target cells was measured by incubating the lysate with chloroform red β-D-galactopyranoside (Roche Applied Science, Indianapolis, IN) and performing spectrophotometry at OD570 nm.

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Author Contributions

Conceived and designed the experiments: QL, JIC. Performed the experiments: QL, MA KW DS FH ERF RB. Analyzed the data: QL, MA KW JIC. Wrote the paper: QL, JIC.

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