Herpes simplex virus 1 protein pUL21 alters ceramide metabolism by activating the interorganelle transport protein CERT

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Tomasz H. Benedyk* , Viv Connor † , Eve R. Caroe † , Maria Shamin † † , Dmitri I. Svergun † † , Janet E. Deane † † , Cy M. Jeffries † † , Colin M. Crump † † , and Stephen C. Graham* † †

From the 1 Department of Pathology, and 2 Cambridge Institute for Medical Research, University of Cambridge, Cambridge, UK; 3 European Molecular Biology Laboratory (EMBL) Hamburg Site, Hamburg, Germany

Herpes simplex virus (HSV)-1 dramatically alters the architecture and protein composition of cellular membranes during infection, but its effects upon membrane lipid composition remain unclear. HSV-1 pUL21 is a virus-encoded protein phosphatase adaptor that promotes dephosphorylation of multiple cellular and virus proteins, including the cellular ceramide (Cer) transport protein CERT. CERT mediates nonvesicular Cer transport from the endoplasmic reticulum to the trans-Golgi network, whereupon Cer is converted to sphingomyelin (SM) and other sphingolipids that play important roles in cellular proliferation, signaling, and membrane trafficking. Here, we use click chemistry to profile the kinetics of sphingolipid metabolism, showing that pUL21-mediated dephosphorylation activates CERT and accelerates Cer-to-SM conversion. Purified pUL21 and full-length CERT interact with submicromolar affinity, and we solve the solution structure of the pUL21 C-terminal domain in complex with the CERT Pleckstrin homology and steroidogenic acute regulatory–related lipid transfer domains using small-angle X-ray scattering. We identify a single amino acid mutation on the surface of pUL21 that disrupts CERT binding in vitro and in cultured cells. This residue is highly conserved across the genus Simplexvirus. In addition, we identify a pUL21 residue essential for binding to HSV-1 pUL16. Sphingolipid profiling demonstrates that Cer-to-SM conversion is severely diminished in the context of HSV-1 infection, a defect that is compounded when infecting with a virus encoding the mutated form of pUL21 that lacks the ability to activate CERT. However, virus replication and spread in cultured keratinocytes or epithelial cells is not significantly altered when pUL21-mediated CERT dephosphorylation is abolished. Collectively, we demonstrate that HSV-1 modifies sphingolipid metabolism via specific protein–protein interactions.

Herpes simplex virus (HSV)-1 is a human pathogen that is estimated to infect the majority of the human population, causing a lifelong latent infection (1). Latent HSV-1 resides in sensory neurons or sympathetic neurons, migrating to the periphery in periodic reactivation events throughout the lifetime of the host. In order to sustain acute (lytic) infection, HSV-1 drastically modifies infected cells (2–5). In particular, the virus extensively remodels the composition and architecture of cellular membranes to facilitate virus assembly and spread. Nascent capsids leave the nucleus via sequential envelopment and de-envelopment at the inner and outer nuclear membranes (6). These cytosolic capsids acquire a proteinaceous layer termed “ tegument ” and bud into the lumen of post-Golgi membranes that are studded with viral glycoproteins (so called “ secondary envelopment ”) (7). The resultant virus-containing vesicles are transported to cell contact sites where they release the mature virus particles to disseminate the infection (8).

HSV-1 pUL21 is a tegument protein that is conserved in all alphaherpesviruses (9). This multifunctional protein and its homologs are known to interact with multiple cellular and viral partners, including pUL16 (10, 11), pUL11 (12), gE (12), tubulin (13), and Roadblock-1 (14), and it has been implicated in a number of important processes including capsid nuclear egress (15), viral cell-to-cell spread (12, 16), and retrograde transport along axons to the neuronal cell bodies where latency is established (14, 17). Mutant HSV-1 lacking pUL21 expression exhibits a 10-fold to 100-fold replication defect and severely impaired cell-to-cell spread, both of which can be at least partially ascribed to the phosphomodulatory role of pUL21 as a protein phosphatase 1 (PP1) adaptor (18). PP1 is a highly active and abundant cellular phosphatase (19), and pUL21 recruits PP1 to promote dephosphorylation of multiple substrates, including the viral protein pUL31 that is implicated in viral nuclear egress (18, 20) and the cellular protein CERT that regulates sphingomyelin (SM) metabolism (18).

The cytoplasmic ceramide (Cer) transport protein CERT (a.k.a. Goodpasture’s antigen-binding protein, GPBP, encoded by the gene COL4A3BP) mediates the nonvesicular trafficking of Cer from the endoplasmic reticulum (ER) to the trans-Golgi network (TGN) and, in doing so, defines the rate of SM metabolism. This process is partially regulated by the virus-encoded protein pUL21, which acts as a phosphatase adaptor to promote dephosphorylation of CERT, thus accelerating Cer-to-SM conversion and altering the composition of the cellular membranes. The resulting changes in lipid metabolism can have significant implications for viral replication and host cell biology. The identification of specific protein interactions and the characterization of molecular mechanisms underlying these processes are crucial for understanding the complex interplay between virus and host during HSV-1 infection.
syntehsis (21–23). CERT contains two well-folded globular domains: an N-terminal Pleckstrin homology (PH) domain that mediates its interaction with TGN membranes by binding phosphatidylinositol 4-phosphate and a C-terminal steroidal- genic acute regulatory–related lipid transfer (START) domain that directly binds Cer to mediate its transfer (23). Crystal structures of both domains have been solved (24, 25). The “middle region” (MR) that connects the PH and START domains of CERT does not adopt a globular fold, instead containing a coiled-coil region that is likely to mediate CERT self-association (26), a “two phenylalanines in an acidic tract” (FFAT) motif that recruits CERT to the ER membranes via binding proteins VAPA and VAPB (27), and a serine-rich motif, hyperphosphorylation of which represses CERT activity (28). In cultured HeLa cells, the vast majority of CERT is in an inactive hyperphosphorylated state (CERTP), available to be mobilized via dephosphorylation to increase the rate of ER-to-TGN Cer transport in response to stimuli such as SM dephosphorylation (28).

Cer and sphingolipids like SM are essential for mammalian cell growth, and they play important roles in cell signaling, apoptosis, and membrane trafficking (29). Furthermore, sphingolipids and cholesterol participate in the formation of membrane microdomains (including “lipid rafts”) that compartmentalize the lipid and protein composition of cellular membranes (30). This compartmentalization is especially important in highly polarized cells such as neurons (31). Although sphingolipids are crucial to host cell biology, relatively little is known about the interaction between pathogens and cellular sphingolipid metabolism. The bacteria Chlamydia trachomatis is known to directly recruit CERT to bacteria-containing intracellular inclusions, thereby increasing the abundance of SM in the inclusion membrane (32), and CERT activity is necessary for the biosynthesis of double-membrane vesicles that serve as the sites of hepatitis C virus and polio virus replication (33). However, to the best of our knowledge, the binding of CERT to HSV-1 pUL21 (18) is the only known example of a direct interaction between CERT and a virus protein.

Previous studies have shown that HSV-1 infection increases Cer synthesis (34) and that depletion of sphingomyelinase causes a >30-fold decrease in HSV-1 replication (35). Sphingomyelinase treatment of cultured epithelial cells has only a very modest effect upon HSV-1 entry (36), suggesting that SM is not absolutely required for virus infection. In contrast, recent studies in macrophages demonstrated that acid ceramidase activity is potently antiviral, helping sequester incoming virus particles by promoting their neutralizing association with sphingosine (Sph)-rich intraluminal vesicles within endocytic multivesicular bodies (37). Furthermore, siRNA depletion of CERT has been shown to promote secretion of usually cell-associated HSV-1 virions (38). It is therefore increasingly clear that cellular sphingolipid metabolism is important for HSV-1 biology and that further studies are required to obtain a full picture of how specific sphingolipids support and/or prevent HSV-1 infection.

We sought to define the functional consequences of the pUL21–CERT interaction for HSV-1 infection. Using click chemistry, we demonstrated that pUL21-mediated CERT dephosphorylation increases the rate of Cer to SM conversion in cultured cells. We characterized the solution structure of the pUL21 C-terminal domain in complex with the PH and START domains of CERT, identifying a specific pUL21 amino acid required for the interaction. Functional characterization of mutant HSV-1 encoding this mutated form of pUL21 confirmed that pUL21-mediated CERT dephosphorylation alters the rate of SM synthesis in infected cells. Furthermore, we observed a dramatic increase in the rate of Cer accumulation in infected cells, which was only partly counteracted by pUL21-stimulated CERT activity, but this increase in Cer abundance does not significantly alter virus replication or spread in cultured keratinocytes or epithelial cells.

Results

Human cells typically maintain lipidome homeostasis via robust feedback and control mechanisms, which respond to perturbations via local activation of specific signaling pathways (39, 40). We hypothesized that pUL21, expressed at late points of infection (3), would likely alter the rate of sphingolipid synthesis and thereby affect lipid-mediated signaling events, rather than altering the overall steady-state lipid composition of infected cells. Synthetic lipids with alkyne-containing acyl chains are efficiently processed by mammalian lipid-modifying enzymes and represent powerful tools to probe lipid metabolism (41). The impact of pUL21-directed CERT dephosphorylation on sphingolipid metabolism was therefore probed using a clickable analog of Sph, alkyne-Sph, to monitor the rate of SM biogenesis in immortalized human keratinocyte (HaCaT) cells. Exogenous Sph is efficiently incorporated into cellular metabolic pathways, being rapidly converted into Cer and then SM or hexosylceramides like glucosylceramide or galactosylceramide (42). It is also converted into phosphatidylcholine (PC) via a so-called “salvage” pathway (Fig. 1A) that directs Sph to palmitoyl-CoA, which serves as substrate for reacylation of lyso phosphatidylcholine (43).

To monitor Sph metabolism, HaCaT cells, either parental or stably expressing pUL21 (HaCaT21), were incubated with alkyne-Sph for 5 min (pulse), and the rate of alkyne-Sph incorporation into the competing metabolic pathways was monitored for 2 h (chase) by high-performance TLC (HPTLC) separation and detection of lipids conjugated to coumarin-azide via a “click” reaction (Fig. 1, B and C). Alkyne-Sph was very efficiently converted to alkene-Cer in both cell types, the levels of alkene-Cer remaining relatively stable throughout the chase, and for both cell types, synthesis of alkene-PC and alkene-SM was observed but not synthesis of alkene-hexosylceramide. HaCaT21 cells exhibited significantly reduced rates of alkene-Sph conversion, and the level of alkene-Cer is significantly lower (Fig. S1). The rate of alkene-PC synthesis was also reduced, although this reduction was not significant owing to the high interexperiment variability of the alkene-PC signal for the parental HaCaT cells. Taken
together, these results indicated that lipid metabolism is altered in HaCaT cells constitutively expressing pUL21. However, it remained unclear whether these differences reflected adaptations of cellular lipid metabolism in response to constitutive pUL21 expression rather than a direct effect upon CERT activity. Since CERT-mediated nonvesicular transport of Cer from the ER to the TGN defines the rate of Cer-to-SM conversion (21–23), pUL21-directed CERT dephosphorylation (and thus activation) should increase the rate of SM synthesis. Consistent with this hypothesis, the abundance of alkkyne-SM as a fraction of total alkkyne-Cer plus alkkyne-SM was significantly higher in HaCaT21 versus HaCaT cells at early time points (0–30 min) during the chase (Fig. 1D). By the 60 min chase time point, the rate of alkkyne-SM accumulation slows and the difference in relative abundance between cell lines was diminished, consistent with the alkkyne-SM levels in both cell types approaching equilibrium. This increase in alkkyne-SM accumulation as a fraction of alkkyne-Cer plus alkkyne-SM abundance was observed despite similar overall signal for alkkyne-SM and alkkyne-Cer in HaCaT21 cells when compared with parental cells at early time points (0–30 min), the abundance of these lipids being significantly lower in HaCaT21 cells at later time points (Fig. 1E). Reduced overall levels of SM plus Cer in HaCaT21 cells likely result from adaptation to constitutive pUL21 expression (and thus constitutive CERT hyperactivation) via stimulation of feedback pathways that reduce Sph to Cer conversion and/or increased back conversion of SM and/or Cer to Sph. Treatment of HaCaT cells with the highly specific CERT inhibitor HPA-12 (44) confirmed that the rate of Cer-to-SM conversion...
was defined by CERT activity: SM synthesis was significantly decreased in HPA-12–treated cells when compared with untreated cells, and there was concomitant accumulation of alkyne-Cer in treated cells (Figs. 1, B–D and S1). Immunoblot analysis of protein samples that were resolubilized following their precipitation during lipid extraction (Fig. 1F) confirmed that pUL21 was expressed and that dephosphorylated CERT (CERT\textsuperscript{C}) predominates in HaCaT21 cells, in contrast to the parental and HPA-12–treated HaCaT cells.

pUL21 has multiple functions during infection, promoting the PPI-mediated dephosphorylation of not just CERT but of multiple different cellular and viral proteins (18). It was therefore essential to identify a point mutant of pUL21 with impaired binding to CERT, but not to other cellular or viral partners, to dissect the functional significance of increased CERT activity during infection. Defining the interaction surfaces of transient protein complexes presents a considerable challenge (45), made more difficult in the case of pUL21 and CERT by their containing multiple domains and regions of intrinsic disorder (18). The minimal region of CERT required for pUL21 binding was thus probed using pull-down experiments where pUL21-glutathione-S-transferase (GST) purified following bacterial expression (bait) was used to capture CERT or truncations thereof (preys) expressed by \textit{in vitro} coupled transscription/translation in wheat germ extract (Fig. 2A). The longer cytoplasmic isoform of CERT (CERT\textsubscript{L}), where expression of exon 11 extends the amino-terminal region of the START domain by 26 amino acids (START\textsubscript{L}), was used since both CERT and CERT\textsubscript{L} have been shown to possess Cer transfer activity (23), and a contribution of these additional 26 amino acids to pUL21 binding could not be excluded. Full-length CERT\textsubscript{L} was efficiently captured by pUL21-GST. The PH and START\textsubscript{L} domains were also captured by pUL21-GST when expressed individually, albeit less efficiently, suggesting that the CERT\textsubscript{L} possesses multiple pUL21-binding motifs. The CERT MR was not captured by pUL21-GST and is thus likely to be dispensable for binding. A truncated CERT\textsubscript{L} construct that retained the pUL21-binding PH and START\textsubscript{L} domains, but lacked the majority of the highly flexible MR, was thus designed for use in subsequent biochemical and structural studies (miniCERT\textsubscript{L}) (Fig. 2B). The extent of the MR retained to link the two domains was informed by the crystal structure of the CERT PH–START domain complex (46), defining a minimal distance of at least 55 Å between the PH domain C terminus and the START domain N terminus, and the desire to exclude the predicted coiled-coil region that may cause oligomerization (26). MR residues 132 to 350 were thus excluded from miniCERT\textsubscript{L}. The first 19 amino acids of the PH domain were also omitted from miniCERT\textsubscript{L} as this region is predicted to be highly disordered and was excluded in previous structural studies (24, 47).

Previous studies identified that the MR mediates CERT trimerization (26, 48). Size-exclusion chromatography with inline multiangle light scattering (SEC–MALS) (Fig. 2C) confirmed that hyperphosphorylated StrepII-tagged CERT\textsubscript{L}, purified following expression in mammalian (Freestyle 293F) cells, has an observed mass (184.3 kDa) approaching that expected for a trimer (215.7 kDa). Similarly, a constitutively hypophosphorylated form of CERT\textsubscript{L} where S132 is mutated to alanine, and thus cannot become phosphorylated to initiate the serine-rich motif phosphorylation cascade (28), had an observed mass (199.2 kDa) approaching that expected for a trimer. In contrast, H\textsubscript{6}-miniCERT\textsubscript{L} purified following bacterial expression was predominantly monomeric and monodisperse (observed mass of 43.9 kDa, expected monomeric mass of 44.9 kDa), consistent with trimerization of CERT\textsubscript{L} being driven by the MR and not being dependent upon CERT\textsubscript{L} phosphorylation.

Isothermal titration calorimetry (ITC) demonstrated that pUL21-H\textsubscript{6} (18) binds StrepII-CERT\textsubscript{L} with approximately micromolar affinity (Fig. 2, D and E, Tables 1 and S1), the observed affinity not differing significantly between the hyperphosphorylated (CERT\textsubscript{L}\textsuperscript{P}) and hypophosphorylated (CERT\textsubscript{L}\textsuperscript{S132A}) forms of the protein. While StrepII-CERT\textsubscript{L}\textsuperscript{S132A} forms a 1:1 complex with pUL21, the observed binding stoichiometry (N) was consistently lower for StrepII-CERT\textsubscript{L}\textsuperscript{P} (0.74), consistent with a proportion of the pUL21 binding sites on CERT\textsubscript{L} being sterically occluded when the protein is hyperphosphorylated. H\textsubscript{4}-miniCERT\textsubscript{L} and pUL21-H\textsubscript{4} form an equimolar complex, binding with micromolar affinity similar to the full-length protein (Fig. 2F and Table 1). However, the thermodynamics of binding differ significantly; whereas pUL21 binding to CERT\textsubscript{L} is endothermic and entropically driven, binding to miniCERT\textsubscript{L} is exothermic and enthalpically driven with minimal change in overall entropy (Table 1). This is consistent with CERT\textsubscript{L} undergoing significant conformational rearrangement upon binding to pUL21, whereas the conformational changes to miniCERT\textsubscript{L} (if any) upon pUL21 binding are likely to be much more modest.

Small-angle X-ray scattering (SAXS) in batch mode was used to probe the solution structure of H\textsubscript{4}-miniCERT\textsubscript{L} (Fig. 2G and Table S2). The frequency distribution of real-space distances (p(r) profile) is largely symmetric (Fig. 2H), and the dimensionless Kratky plot (Fig. 2I) has a bell-shaped peak at sRg = V3, consistent with miniCERT\textsubscript{L} having a compact globular conformation in solution. \textit{Ab initio} modeling of the H\textsubscript{4}-miniCERT\textsubscript{L} scattering profile using GASBOR (49) yields a dummy-residue model (Fig. 2, J and K) that closely resembles the crystal structure of CERT START and PH domain complex (46). A pseudoatomic model of the H\textsubscript{4}-miniCERT\textsubscript{L} solution structure, generated using the crystal structures of the CERT PH and START domains determined in isolation (24, 25) as rigid bodies, confirms the similarity of the H\textsubscript{6}-miniCERT\textsubscript{L} solution model to the crystal structure of the CERT PH–START domain complex (Fig. 2, L and M). Collectively, the MALS, ITC, and SAXS analyses demonstrate that miniCERT\textsubscript{L} forms a compact monomeric protein with high affinity for pUL21, representing an excellent tool for structural studies.

Structural characterization \textit{via} X-ray crystallography and SAXS has revealed pUL21 to comprise two domains joined by a highly flexible linker (Fig. 3A) (18, 50, 51), and immunoprecipitation experiments mapped CERT binding to the pUL21 C-terminal domain (18). The C-terminal domain of pUL21 (pUL21C), spanning amino acids 275 to 535, was
Figure 2. pUL21 binds to the PH and STARTL domains of CERTL and to a monomeric form of CERTL, comprising just the PH and STARTL domains (miniCERTL). A, minimal binding elements of CERTL were determined via pull down (PD) experiment using immobilized purified pUL21-GST or GST alone to capture myc-tagged full-length CERTL or truncations thereof expressed via in vitro transcription/translation. Captured proteins were subjected to SDS-PAGE and immunoblotting using an anti-myc antibody. Ponceau S (PonS) staining of the immunoblot membrane before blocking shows equal and efficient capture of the bait proteins across the tested conditions. B, schematic representation of miniCERTL. Dotted lines indicate the regions of full-length CERTL that were omitted. C, SEC–MALS elution profiles (normalized differential refractive index, dRI) of StrepII-CERTL (black dotted), StrepII-CERTL, S132A (gray solid), and H₆-miniCERTL (green solid). Weight-averaged molecular masses (colored solid lines) are shown across the elution peaks. The expected molecular masses for trimeric StrepII-CERTL and StrepII-CERTL, S132A, and for monomeric H₆-miniCERTL, are shown as dotted horizontal lines. Inset shows Coomassie-stained SDS-PAGE of the samples used for SEC–MALS (left), plus SDS-PAGE supplemented with PhosTag reagent (right) of the StrepII-CERTL proteins. D–F, representative ITC titration curve of pUL21-H₆ binding to (D) StrepII-CERTL, (E) StrepII-CERTL, S132A, and (F) H₆-miniCERTL. Insets show normalized binding curves with integrated changes in enthalpy (ΔH) as a function of molar ratio. The affinity (Kᵰ), ΔH, and stoichiometry (N) for the presented titrations are displayed below. G, SAXS profile measured for H₆-miniCERTL. The reciprocal-space fit of the p(r) profile to the SAXS data is shown as a yellow line. χ², fit quality; p, Correlation Map (CorMap) probability of systematic deviations between the model fit and the scattering data. H, fit to the SAXS data of the pseudoatomic model of H₆-miniCERTL obtained using CORAL, M, CORAL pseudoatomic model of H₆-miniCERTL (green ribbons) superimposed onto the crystal structure of the complex between the CERT PH and START domains (cyan ribbons, PDB ID: 5JJD) by aligning the START/STARTL domains. Superposition is shown in orthogonal orientations where the linker

**HSV-1 pUL21 modulates cellular ceramide transport**
**HSV-1 pUL21 modulates cellular CERTL ceramide transport**

**Table 1** Thermodynamic properties of the pUL21–CERTL interaction

| Syringe | Cell   | $K_D$ (nM) | $\Delta H$ (kcal/mol) | $\Delta G$ (kcal/mol) | $-\Delta S$ (kcal/mol) | $N$ (stoichiometry) | $n$ (replicates) |
|---------|--------|------------|------------------------|-----------------------|------------------------|---------------------|-----------------|
| CERTLpUL21 | pUL21 | 772.7 ± 274.0 | 3.89 ± 1.29 | $-8.37$ ± 0.20 | $-12.27$ ± 1.07 | 0.74 ± 0.02 | 3 |
| CERTLpUL21 | pUL21 | 642.0 ± 257.4 | 2.385 ± 0.29 | $-8.47$ ± 0.25 | $-10.85$ ± 0.06 | 0.97 ± 0.04 | 2 |
| miniCERTL | pUL21 | 1042.0 ± 537.7 | $-8.65$ ± 0.63 | $-8.21$ ± 0.24 | $-0.44$ ± 0.78 | 0.89 ± 0.06 | 6 |
| miniCERTL | pUL21C | 4325.0 ± 1661.7 | $-3.84$ ± 0.02 | $-7.35$ ± 0.23 | $-3.51$ ± 0.25 | 0.97 ± 0.04 | 2 |
| miniCERTL | pUL21C | 8086.7 ± 932.8 | $-7.20$ ± 0.05 | $-6.95$ ± 0.06 | 0.26 ± 0.02 | 1.16 ± 0.13 | 3 |

As quantitated by ITC. Experiments were performed $n$ (replicates) times and mean ± SD values are shown. Data for individual titrations are presented as supporting information (Table S1).

expressed with a N-terminal hexahistidine tag and purified following bacterial expression. In contrast to a previous report (51), we did not observe copurification of nucleic acid with H$_6$-pUL21C, the purified protein having an absorbance ratio of ~0.6 at 260/280 nm. SEC–MALS and SAXS analysis (Fig. S2) confirmed that H$_6$-pUL21C is monomeric and monodisperse, adopting a compact structure in solution that matches the previously determined crystal structure ($\chi^2 = 0.99$, CorMap $p = 0.133$) (51). SEC analysis of a preformed complex between purified H$_6$-miniCERTL and a 1.3-fold molar excess of H$_6$-pUL21C confirmed that the two proteins form a stable complex (Fig. 3A), suitable for structural studies. ITC demonstrated that H$_6$-pUL21C forms an equimolar complex with H$_6$-miniCERTL with a dissociation constant ($K_D$) of 4.3 ± 1.2 μM (Fig. 3C, Tables 1 and S1). The approximately fourfold reduction in miniCERTL binding by pUL21C versus full-length pUL21 is consistent with previous immunoprecipitation results, where transfected pUL21C-GFP captured endogenous CERT slightly less efficiently than did pUL21-GFP in human embryonic kidney 293T (HEK293T) cells (18). These ITC experiments confirm that the C-terminal domain of pUL21 is the major determinant of CERT binding. Attempts to predict the structure of pUL21C in complex with miniCERTL using AlphaFold-Multimer (52) did not yield high confidence models, presumably because of a paucity of coevolutionary signal between these viral and human proteins, and structural characterization of this complex thus required an experimental approach.

To probe the structural basis of the CERT recruitment by pUL21, a preformed complex of H$_6$-miniCERTL and H$_6$-pUL21C was subjected to SEC with inline SAXS measurement (SEC–SAXS, Fig. 3D). SAXS data were processed by averaging frames with a consistent calculated radius of gyration ($R_g$) and then subtracting averaged buffer frames to yield the H$_6$-miniCERTL–H$_6$-pUL21C complex scattering profile (Fig. 3E). The probable frequency of real-space distances ($p(r)$ profile) of the complex is moderately asymmetric (Fig. 3F), in contrast to the highly symmetric $p(r)$ profiles of H$_6$-pUL21C (Fig. S2D) or H$_6$-miniCERT (Fig. 2H) alone, suggesting a less spherical particle, and the dimensions of the complex ($R_g = 3.5$ nm, $D_{max} = 13.6$ nm) are substantially larger than for H$_6$-pUL21C ($R_g = 2.2$ nm, $D_{max} = 8.5$ nm) or H$_6$-miniCERT ($R_g = 2.7$ nm, $D_{max} = 9.1$ nm). The peak of the dimensions

Kraty plot is slightly higher, with its peak away from $sr_g = \sqrt{3}$ (Fig. 3G), suggesting some flexibility in the system granted either by modest dissociation of the complex or some mobility of the H$_6$-miniCERTL domains with respect to each other and H$_6$-pUL21C. *Ab initio* modeling using GASBOR indicated an elongated molecule (Fig. 3H). While initial pseudoatomic models of the complex generated using a fixed conformation of H$_6$-miniCERTL did not fit the SAXS profile acceptably, allowing the PH and STARTL domains freedom to move with respect to each other and to H$_6$-pUL21C yielded three pseudoatomic models with high-quality fits to the SAXS profile (Figs. 3, I and J and S3). In all three top models, miniCERTL binds "end-on" to the pUL21 molecule, forming an ellipsoidal “rugby ball”–like particle. In two of the top three models, miniCERTL binds the “left wing” of the dragonfly-shaped pUL21C domain (51), whereas in the other, it binds the "right wing" (Figs. 3, I and J and S3). All these models have a similar overall shape, and thus all explain the SAXS scattering data well, but the relative orientations of H$_6$-miniCERTL and H$_6$-pUL21C differ. All three top models were thus used to design specific pUL21C mutations that might disrupt (mini) CERT binding.

Table 1

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As quantitated by ITC. Experiments were performed $n$ (replicates) times and mean ± SD values are shown. Data for individual titrations are presented as supporting information (Table S1).

regions or termini that were modeled by CORAL are depicted as yellow spheres. DP, differential power; GST, glutathione-S-transferase; ITC, isothermal titration calorimetry; PDB, Protein Data Bank; PH, Pleckstrin homology; SAXS, small-angle X-ray scattering; SEC–MALS, size-exclusion chromatography with inline multilaminate light scattering; START, steroidogenic acute regulatory–related lipid transfer.
Figure 3. Solution structure of the H6-pUL21C–H6-miniCERTL heterodimer. A, schematic of pUL21, with N- and C-terminal domains joined by a highly dynamic linker (18). Crystal structures of domains (N, PDB ID: 4U4H (50); C, PDB ID: 5ED7 (51)) are shown. B, Superdex 75 10/300 SEC elution profile of the H6-pUL21C–H6-miniCERTL complex, preformed in the presence of 1.32-fold molar excess of H6-pUL21C. Dotted lines indicate fractions that were collected and subjected to SDS-PAGE analysis with Coomassie staining, revealing coelution of the two proteins. C, representative ITC titration curve of H6-pUL21C binding to H6-miniCERTL. Inset shows normalized binding curve with integrated changes in enthalpy ($\Delta H$) as a function of molar ratio. The ranges of the vertical axes are identical to Figure 2. The affinity ($K_D$), $\Delta H$, and stoichiometry (N) for the presented titration are displayed below. D, SEC elution profile (partial integrated scattered X-ray intensity versus data frame number) obtained during SEC–SAXS analysis of H6-pUL21C–H6-miniCERTL complex. Dashed line indicates the calculated radius of gyration ($R_g$) across the frames averaged for structural analyses. E, averaged SAXS profile of the H6-pUL21C–H6-miniCERTL complex. The reciprocal-space fit of the $p(r)$ profile to the SAXS data is shown as a yellow line. $\chi^2$, fit quality; $p$, Correlation Map (CorMap) probability of systematic deviations between the model fit and the scattering data (92). Inset displays the Guinier plot ($sR_g < 1.3$), which is linear as expected for an aggregate- and repulsion-free system. F, real-space distance distribution function, $p(r)$, calculated from the SAXS profile. G, dimensionless Kratky plot of the SAXS data. Gray dotted lines indicate the expected maximum of the plot for a compact protein ($sR_g = \sqrt{3}$, $(sR_g)^2I(s)/I(0) = 3e^{-1}$). H, ab initio modeling of the H6-miniCERTL–H6-pUL21C complex using GASBOR. Fit of the calculated scattering (yellow) to the SAXS profile is shown, as is a representative dummy-residue model (blue spheres) with modeled water beads of the hydration shell (white spheres). I and J, pseudoatomic models of the H6-miniCERTL–H6-pUL21C complex generated using CORAL. The fits of the computed scattering (yellow) to the H6-pUL21C–H6-miniCERTL SAXS profile (blue) are shown. High-quality fits are obtained with models where miniCERTL binds either the left or right “wings” of the dragonfly-like pUL21C domain (51). pUL21C is shown as a violet ribbon with “wings” and termini labeled, and miniCERTL, as a green molecular surface (PH, light green; STARTL, dark green). For clarity, regions absent from the crystal structures that were modeled by CORAL are not displayed. Residues from the left and right “wings” of pUL21C that were selected for further investigation are shown as sticks with red (I) or pink (J) carbon atoms, respectively. An additional pseudoatomic model plus its fit to the H6-pUL21C–H6-miniCERTL SAXS profile is shown in Figure S3, where the fits presented in (I) and (J) are also reproduced for ease of comparison. DP, differential power; ITC, isothermal titration calorimetry; PDB, Protein Data Bank; PH, Pleckstrin homology; SAXS, small-angle X-ray scattering; SEC, size-exclusion chromatography; START, steroidogenic acute regulatory–related lipid transfer.
Figure 4. Identification of pUL21 point mutants with disrupted binding to CERT or pUL16. A and B, immunoblot following immunoprecipitation from HEK293T cells transfected with plasmids encoding GFP-tagged pUL21, either WT or with amino acid substitutions at the putative CERT binding interface on the left (A) or right (B) wings of pUL21C, or encoding GFP alone. At 24 h post-transfection, cells were infected with ΔpUL21 HSV-1 (MOI = 5), and at 16 h postinfection, cells were lysed, tagged proteins were captured using GFP affinity resin, and the bound proteins were subjected to SDS-PAGE and immunoblotting using the antibodies listed. Ponceau S (PonS) staining of the nitrocellulose membrane before blocking is shown to confirm efficient capture of GFP-tagged proteins. GAPDH (A) and tubulin (B) are used as loading controls. C, Superdex 75 16/600 SEC elution profile of pUL21V382E-H6 following bacterial expression and affinity capture. Indicated peak was pooled and used for subsequent experiments. D, differential scanning fluorimetry of WT (purple) and V382E (red) pUL21-H6. Representative curves are shown, and melting temperature (Tm) is mean ± SD of three technical replicates. Inset shows Coomassie-stained SDS-PAGE of the purified proteins. E, representative ITC titration curve of pUL21V382E-H6 binding to H6-miniCERT. Inset shows normalized binding curve with integrated changes in enthalpy (ΔH) as a function of molar ratio. The ranges of the vertical axes are identical to Figure 2F. The affinity (Kd), ΔH, and stoichiometry (N) for the presented titration are displayed below. F, in vitro dephosphorylation assay using all-purified proteins. About 0.5 μM CERT was incubated with varying concentrations of pUL21-H6 WT or V382E (twofold dilution from 4 μM to 62.5 nM) in the presence of 25 nM GST-PP1 for 30 min at 30°C. Proteins were resolved using SDS-PAGE supplemented with PhosTag reagent to enhance separation of CERT that is hyperphosphorylated (CERTP) or hypophosphorylated (CERTO), and protein bands were visualized using Coomassie. Images are representative of two independent experiments. G, per-residue predicted Local Distance Difference Test (pLDDT) scores (top) and predicted aligned error (PAE) matrix (bottom) for predicted pUL16–pUL21C complex. DP, differential power; HEK293T, human embryonic kidney 293T cell line; HSV-1, herpes simplex virus 1; ITC, isothermal titration calorimetry; MOI, multiplicity of infection; PP1, protein phosphatase 1; SEC, size-exclusion chromatography.
CERT binding while maintaining the ability of pUL21 to coprecipitate its other known binding partners, PP1 (18) and pUL16 (10). The presence of a valine residue at this position is absolutely conserved in HSV-1 and HSV-2 and is highly conserved across members of the simplexvirus genus (Fig. S4). H$_6$-pUL21V$_{382E}$ was purified following bacterial expression (Fig. 4C). Differential scanning fluorimetry (a.k.a. Thermofluor) showed pUL21V$_{382E}$ to be well folded as its thermal stability is similar to WT H$_6$-pUL21 (Fig. 4D). ITC analysis demonstrated that H$_6$-pUL21V$_{382E}$ has approximately eightfold reduced binding affinity for H$_6$-miniCERT$_{\text{C}}$ when compared with WT pUL21 (Fig. 4E, Tables 1 and S1). The effect of the V382E substitution upon the ability of pUL21 to promote CERT dephosphorylation, converting CERT$_{\text{P}}$ to CERT$_{\text{O}}$, was probed using an in vitro dephosphorylation assay with all-purified reagents (Fig. 4, F and G). The dose-dependent acceleration of GST-PP1-mediated CERT dephosphorylation is significantly greater for WT pUL21 than pUL21V$_{382E}$ (p = 0.040; two-way ANOVA with Sidak’s multiple comparison test), consistent with the GST-PP1–pUL21 complex having lower affinity for the substrate CERT$_{\text{P}}$ (EC$_{50}$[pUL21] = 0.491 ± 0.123 μM, EC$_{50}$[pUL21V$_{382E}$] = 3.329 ± 0.7874 μM; three parameter dose–response curve, n = 2 independent experiments).

While mutation of the pUL21C right wing did not prevent CERT binding, one substitution (L490E) abolished binding to pUL16 (Fig. 4B). pUL21L$_{490E}$ retains the ability to bind CERT and PP1, suggesting that the mutant is stable and maintains a correct overall fold. A model of pUL21 in complex with pUL21C was generated using AlphaFold-Multimer (52), revealing that pUL16 is predicted to bind the right wing of pUL21C, with helix α9 spanning residues 486 to 497 being central to this interaction (Fig. 4H). The structures of pUL16 and pUL21C, and their relative orientations in the complex, were predicted with high overall confidence (Fig. 4I). pUL16 is predicted to comprise two domains, an N-terminal domain (residues 35–172) that is joined by a short linker to a C-terminal domain (residues 179–371) (Fig. S5A). A structural homology search using DALI (53) revealed that the pUL16 C-terminal domain shares greatest structural similarity to the N-terminal domain of pUL21 (Fig. 5B and Table S3), whereas the pUL16 C-terminal domain is similar in fold to a variety of bacterial and eukaryotic proteins that have diverse annotated functions ranging from phosphatase activity to lipid binding (Fig. 5, C and D and Table S3). Residues 1 to 34 of pUL16 were predicted with low confidence and are likely to be intrinsically disordered. Residue L490 of pUL21C is predicted to bind a hydrophobic pocket formed by L91, R362, and residues 73 to 78 of pUL16 (Fig. 4H). Loss of pUL16 binding by pUL21C L$_{490E}$ is consistent with the introduction of a charged glutamic acid side chain, which could still form polar interactions with these residues. pUL21 E409 is not predicted to lie at the pUL16 binding interface, consistent with mutation of this residue not affecting the pUL21C–pUL16 interaction.

To probe the ability of the pUL21 mutant with reduced CERT binding (pUL21V$_{382E}$) to stimulate CERT dephosphorylation during infection, a mutant strain of HSV-1 encoding pUL21V$_{382E}$ was generated using two-step Red recombination (54). Dephosphorylated CERT (CERT$_{\text{O}}$) is significantly less abundant in HaCaT cells infected with HSV-1 expressing pUL21V$_{382E}$ or lacking pUL21 expression (ΔpUL21) when compared with WT HSV-1-infected cells (Fig. 5, A and B). In addition to CERT, pUL21 expression reduces the phosphorylation of multiple substrates of the viral kinase pUS3 in HSV-1–infected cells (18), the phosphorylated forms of these substrates being detectable using an antibody that recognizes phosphorylated substrates of the cellular kinase Akt (55). While infection with ΔpUL21 HSV-1 causes a dramatic increase in the abundance of multiple phosphorylated pUS3 substrates, the abundance of these phosphoforms is indistinguishable between WT and pUL21V$_{382E}$ HSV-1 (Fig. 5A). This confirms that the V382E substitution specifically disrupts CERT dephosphorylation, rather than generally inhibiting the ability of pUL21V$_{382E}$ to recruit PP1 to substrates. Similar changes in CERT dephosphorylation, but not in the dephosphorylation of other pUS3 substrates, are observed when Vero cells are infected with pUL21V$_{382E}$ HSV-1 (Fig. 5c, A and B). Immunocytochemistry confirms that both WT and V382E pUL21 have the similar subcellular localization, being observed predominantly at the nuclear rim of infected Vero cells (Fig. 5C).

Metabolic labeling was used to monitor the impact of pUL21-mediated CERT dephosphorylation on sphingolipid biogenesis during infection. A pulse-chase experiment was performed where HaCaT cells infected with WT or pUL21V$_{382E}$ HSV-1, or mock infected, were incubated for 5 min with alkyne-Sph at 14 h postinfection (hpi), and its metabolites were monitored for 2 h. Cer accumulates, and the rate of SM synthesis is significantly decreased in cells infected with WT and pUL21V$_{382E}$ HSV-1 when compared with uninfected cells (Fig. 5D). Although a substantial decrease in the rate of SM synthesis is seen for both WT and mutant HSV-1 infection, the defect is significantly larger in cells infected with HSV-1 pUL21V$_{382E}$ (Fig. 5E). The overall abundance of SM plus Cer is similar between infected and uninfected cells, suggesting that Cer to SM conversion is specifically impaired rather than influx of alkyne-Sph into the SM biogenesis pathway being defective (Fig. 5F). Taken together, this suggests that pUL21-mediated activation of CERT accelerates Cer to SM conversion during infection, albeit from a much lower base than in uninfected cells.

Having confirmed that pUL21V$_{382E}$ HSV-1 specifically lacks the ability to stimulate CERT dephosphorylation, and that HSV-1 encoding pUL21V$_{382E}$ has a reduced rate of Cer to SM conversion, the impact of this deficit upon virus replication and spread in cultured cells was assessed. WT and pUL21V$_{382E}$ HSV-1 form similar-sized plaques on HaCaT and Vero cells (Fig. 6A), suggesting that CERT dephosphorylation is dispensable for efficient viral cell-to-cell spread. A single-step...
growth curve, where Vero and HaCaT cells are infected at a high multiplicity of infection (MOI) and the production of infectious progeny is monitored over time, was used to compare the replication of WT and pUL21V382E HSV-1 (Fig. 6B). Both viruses produce similar abundance of infectious progeny by 24 h postinfection. In two biologically independent experiments performed for each cell type, the kinetics of virus replication appeared to be accelerated, with higher titers of pUL21V382E HSV-1 being observed between 6 and 12 h postinfection, but the difference in growth rate was not statistically significant for either cell line (two-way ANOVA with Sidak’s multiple comparisons test).

While HSV-1 preferentially remains cell attached, spreading via direct cell–cell contacts, the cell-free secretion of virions from infected cells is altered when CERT is depleted or overexpressed (38). Cell-free release of pUL21V382E HSV-1 was determined by quantitating the amount of infectivity secreted into the medium as a percentage of overall infectivity (secreted plus cell associated), normalized for each independent replicate to the secretion of a WT virus in control experiments performed at the same time. HaCaT cells infected for 12 h with various MOI of pUL21V382E HSV-1 did not release significantly more or less infectivity when compared with WT HSV-1 infection (Fig. 6C). Similarly, the cell-free release of infectivity at 12 hpi did not differ between Vero cells infected with pUL21V382E or WT HSV-1 (Fig. S6C).

Discussion

HSV-1 extensively remodels the proteome of infected cells, altering both the abundance (3) and post-translational
modulation status (2, 18, 56) of multiple cellular proteins. We know much less about HSV-1-mediated changes to the cellular lipidome, although previous studies have identified that HSV-1 infection changes phosphoinositide levels (57) and increases the rate of de novo phospholipid synthesis (58). By combining biochemical and structural studies with cell-based models of infection, we show here that pUL21 accelerates the conversion of Cer to SM in infected cells by promoting dephosphorylation and activation of the Cer transport protein CERT (Fig. 5). This is the first observation of a viral protein directly binding CERT and altering its activity. While pUL21-mediated CERT activation causes an apparent modest acceleration in the rate of virus replication, this effect is not statistically significant and CERT activation does not contribute to replication or cell-to-cell spread of HSV-1 in cultured keratinocytes or epithelial cells (Fig. 6).

Our previous work showed that pUL21 is a phosphatase adaptor with multiple different targets (18). Mutation of the pUL21 TROPPO motif, required for PP1 binding and thus stimulation of dephosphorylation, dramatically reduced the replication and spread of HSV-1. While the TROPPO motif was identified via its conservation across alphaherpesvirus sequences, the molecular basis of CERT recruitment remained unknown. Our previous immunoprecipitation experiments demonstrated that the N-terminal domain of pUL21 was dispensable for CERT binding (18). Biochemical mapping and SAXS structural characterization presented here have now identified a specific mutation, V382E, that decreases the affinity of pUL21 for miniCERT1 by approximately eightfold (Fig. 4E). While this reduction in affinity is moderate when assayed in dilute biochemical solution, it is sufficient to disrupt coprecipitation of CERT from transfected cells (Fig. 4A). The V382E substitution abolishes pUL21-mediated dephosphorylation of CERT during infection (Fig. 5, A and B) and significantly decreases the rate of Cer to SM conversion (Fig. 5E). These results are consistent with other binding partners competing to bind CERT and/or pUL21 within the context of infected cells, these competing interactions amplifying the effect of reduced pUL21–CERT binding affinities. The V382E mutation does not prevent pUL21 binding to pUL16 (Fig. 4A) nor the ability of pUL21 to stimulate dephosphorylation of other targets (Fig. 5A), confirming that pUL21 binds CERT via a molecular surface that is distinct from the binding site(s) of other targets. The WT levels of virus replication and cell-to-cell spread observed for pUL21V382E HSV-1 are consistent with our previous in vitro evolution studies of HSV-1 mutants where the pUL21 TROPPO motif was mutated (18). Adaptation of the virus to the loss of pUL21 PP1 binding, via suppressor mutations that reduce the activity of the kinase pUS3, restored virus replication and spread without restoring enhanced CERT dephosphorylation (18). We therefore conclude CERT is not a critical substrate of pUL21 for virus replication and spread in cultured fibroblasts and keratinocytes. The C-terminal domain of pUL21 is highly conserved across HSV-1 and HSV-2 (>93% and 80–84% identity, respectively). V382 is absolutely conserved across these species, and a small hydrophobic residue (valine, leucine, or alanine) at this position is very highly prevalent across the genus Simplexvirus (Fig. S4A). The C-terminal domain of pUL21 is much less well conserved when considering the entire subfamily Alphaherpesvirinae (>20% sequence identity across the subfamily), with the amino acid residue at position 382 being correspondingly variable (Fig. S4B). This is consistent with our previous observation that the pUL21 homolog ORF38 from varicella-zoster virus (genus: Varicellaviruses) does not bind CERT (18). It is therefore likely that CERT binding and activation is a conserved feature of HSV-1, HSV-2, and other simplexviruses but does not extend to other genera of alphaherpesvirus. Given the lack of phenotype observed for viruses encoding the CERT-binding pUL21V382E mutant in cultured cells, it is possible that this interaction plays a role in immune regulation or transmission fitness (59). Alternatively,
HSV-1 pUL21 modulates cellular ceramide transport

the interaction may enhance virus replication in other cell types. Specifically, we note that HSV-1 is a neurotropic virus and that sphingolipids like SM are highly enriched in neurons, where they are critical for correct neuronal development and function (31). Studying whether pUL21-mediated CERT dephosphorylation affects virus replication in neurons is an important question for future investigation. Studies of cellular lipid metabolism are complicated by robust cellular feedback mechanisms (60) and challenges in accurately detecting changes in lipid abundance (61). Here, we successfully employed lipid labeling, click chemistry, and HPTLC to monitor the kinetics of Sph metabolism, revealing that HSV-1 pUL21 significantly increases the rate of CERT-mediated Cer to SM conversion both outside (Fig. 1) and within (Fig. 5) the context of infection. Furthermore, we observe a highly significant increase in the abundance of labeled Cer when cells are infected with HSV-1, this increase being greater when pUL21-mediated CERT activation has been abolished. The dramatic change in the rate of Cer to SM conversion is most likely explained by the known propensity of HSV-1 to promote dispersal of the Golgi and TGN (4, 62). Such dispersal would alter ER–TGN contact sites, where CERT-mediated transport enables Cer to SM conversion by TGN-resident SM synthase (63). Side-by-side kinetic comparison of lipid metabolic enzymes including Cer synthase confirms that the presence of alkyl Cer groups within acyl tails does not significantly alter lipid catalysis when compared with radiolabeled or natural substrates (41). While the CERT START domain has been demonstrated to bind Cer containing an alkyl group plus a photoactivatable diazirine group (64, 65), we are unaware of any comparisons between CERT lipid transfer activity with natural or radiolabeled Cer versus alkyl-Cer. It is possible that the presence of an alkyl group may alter the affinity of CERT for Cer, and hence that pUL21-mediated CERT activation may cause a greater or lesser change to the rate of unlabeled Cer to SM conversion. Furthermore, our experimental strategy to probe CERT activity exploited the uptake of exogenous alkyl-Sph and its conversion to alkyl-Cer via the salvage pathway mediated by Cer synthase. This contrasts to the de novo synthesis pathway, where Cer is generated from serine and palmitoyl-coA via intermediates including dihydroceramide (66). While both the de novo and salvage Cer synthesis pathways are known to occur within cells, and the efficient conversion to alkyl-Cer and alkyl-SM clearly indicated efficient salvage of alkyl-Sph in our experiments (Figs. 2, B and C and 5D), it is possible that the effect of pUL21-mediated CERT activation could differ in magnitude for Cer synthesized via the de novo pathway. Future experiments could use radiolabeled serine to study how pUL21-mediated CERT activity affects the metabolism of Cer produced via the de novo pathway (67).

HSV-1 utilizes protein kinase D–mediated trafficking from the TGN to the plasma membrane (68). Disruption of protein kinase D–mediated trafficking has been shown to alter virus secretion, with siRNA-mediated depletion of CERT increasing the secretion of extracellular (cell-free) HSV-1 particles and CERT overexpression reducing virus secretion (38). We do not observe changes in the secretion of pUL21V382E HSV-1, which lacks the ability to dephosphorylate CERT and thus activate CERT-mediated Cer transport. While these results may appear at first glance to be contradictory, it should be noted that in our experiments the cells contained WT levels of CERT, and cells infected with either ΔpUL21 or pUL21V382E HSV-1 have similar levels of active (CERTP) and inactive (CERTO) protein compared with uninfected controls (Fig. 5A). In contrast, siRNA depletion would lead to a complete absence of CERT and of CERT-mediated lipid transfer. Furthermore, the previous study demonstrated that treatment of infected cells with HPA-12 did not stimulate virus particle secretion (38). This insensitivity of HSV-1 secretion to CERT pharmacological inhibition, combined with our observation that pUL21-mediated CERT activation does not alter HSV-1 secretion, strongly suggests that CERT catalytic activity is not directly linked to the regulation of virus secretion.

Our observation of labeled Cer accumulation during HSV-1 infection (Fig. 5) is consistent with an earlier report that HSV-1 infection causes an approximately twofold increase in Cer abundance within BHK-21 cells (34). Cer has many distinctive physical properties that set it apart from other membrane lipids: it has negative intrinsic curvature, it increases the order of phospholipids in membranes, and it makes biological membranes more permeable to even large solutes such as proteins (69). Nascent HSV-1 particles must traverse multiple biological membranes during virus assembly, most notably during capsid egress from the nucleus. Being too large to exit via nuclear pores, capsids leave by budding into and then out of the perinuclear space, the former step being catalyzed by the herpesvirus nuclear egress complex (NEC) (70). A recent study identified that the NEC induces lipid ordering to generate the negative curvature required for capsid budding (71). It is tempting to speculate that pUL21 regulates nuclear capsid egress by modulating Cer abundance, an effect that would be distinct from the previously demonstrated role of pUL21 in promoting nuclear egress via regulating the phosphorylation of NEC components (18, 20). In addition to potentially promoting virus nuclear egress, accumulation of Cer can lead to caspase 3 activation and apoptosis via increased mitochondrial outer membrane permeabilization (72). HSV-1 is known to encode proteins that defend against Cer-induced apoptosis (73), and it is possible that pUL21-stimulated acceleration of Cer to SM conversion serves to limit the proapoptotic activity of Cer, although confirmation of this requires further study.

In the process of probing the molecular basis of pUL21 binding to CERT, we serendipitously discovered a single amino acid substitution (L490E) that prevents pUL21 binding to HSV-1 pUL16 (Fig. 4B). Both pUL16 and pUL21 are known to facilitate efficient release of virus capsids from the nuclei of infected cells (74), and a recent study suggests that the pUL16–pUL21 interaction may help regulate capsid maturation and stability (75). Our mutagenesis is consistent with previous biochemical experiments that had mapped the binding of pUL16 to the C-terminal region of pUL21 (residues 268–535) and shown that the N-terminal 40 amino acids of pUL16 were dispensable for this interaction (10). Recent advances in deep
learning structure prediction enable the accurate prediction of protein structures and complexes, even in challenging cases such as multidomain viral proteins with no known sequence homologs (76). We therefore generated a model of the pUL16–pUL21C interaction using AlphaFold-Multimer (52). pUL21 residue L490 lies at the center of the predicted interaction interface, consistent with its substitution to glutamate inhibiting pUL16 binding (Fig. 4, B and H). The structure of pUL16 has not been solved experimentally, and inspection of the pUL16 structural model provides two interesting observations. First, the closest structural homolog of the predicted pUL16 C-terminal domain is the N-terminal domain of pUL21 (Fig. 5B and Table S3). This structural correspondence (DALI Z-score 5.0; 3.6 Å rmsd across 121 Cα atoms) is striking given the low sequence identity (7%) shared by the domains, although it is known that viral proteins can share similar folds despite having unidentifiable sequence similarly (77). While identifiable pUL21 homologs are restricted to alphaherpesviruses (9), homologs of pUL16 can be found across Herpesviridae (78). The similarity of the predicted pUL16 C-terminal domain structure to that experimentally observed for the pUL1N-terminal domain suggests that the two proteins are distant homologs, having arisen via a gene duplication event in an ancestral (alpha)herpesvirus. The second observation is that several proteins with structural similarity to the predicted pUL16 N-terminal domain have been associated with the transport of lipids or other bulky hydrophobic molecules (Fig. 5D and Table S3). For example, the VtrC protein from Vibrio parahaemolyticus was crystallized in complex with the large hydrophobic molecule taurodeoxycholate (79). Given the association between pUL21 and the lipid transport protein CERT, it is tempting to speculate that pUL16 may also have a role in modifying host or viral lipid membranes, but further biochemical and experimental structural analyses are required to assess to whether the pUL16 N-terminal domain has any capacity to bind and/or transport lipids.

In summary, we demonstrated that pUL21 dephosphorylates and activates the cellular lipid transport protein CERT, stimulating conversion of Cer to SM. Characterizing the solution structure in complex with the PH and START domains of CERT allowed us to identify a single amino acid mutation of pUL21 that disrupts CERT dephosphorylation in infected cells. HSV-1 encoding this pUL21 mutant had similar replication to that experimentally observed for the pUL21N-terminal domain (81) in the 3' untranslated region. The S132A substitution was introduced into this vector using QuikChange mutagenesis (Agilent) according to the manufacturer’s instructions. CERTL1 destined for expression using in vitro transcription/translation system from wheat germ extract was cloned into the plasmid pF3A WG BYDV (Promega) with an N-terminal myc epitope tag. The truncations of CERTL1 were generated by inverse PCR (MR, residues 123–364; STARTL1, residues 365–624) or by introduction of a stop codon (PH, residues 1–122) using QuikChange mutagenesis. For purification following bacterial expression, miniCERTL1 (residues 20–131 plus 351–624) was cloned into pOPTH (77), encoding an N-terminal MAHα tag. The generation of pUL21-Hα and pUL21-GFP was described previously (18), and single amino acid substitutions were introduced by QuikChange mutagenesis. pUL21-Hα was subcloned into pOPTH, and inverse PCR was used to generate Hα-pUL21C, encoding pUL21 amino acids 275 to 535. A plasmid (UK622) encoding mouse PPIγ (UniProt ID: P63087) with an N-terminal GST tag (82) was a kind gift from David Ron (Cambridge Institute for Medical Research). To generate pUL21V382E HSV-1, pEPkan-S containing an I-SceI/KanR selection cassette was used (54).

Mammalian cell culture

Mycoplasma-free spontaneously immortalized human keratinocyte (HaCaT) cells (83), HaCaT cells stably expressing pUL21 (HaCaT21) (18), African green monkey kidney (Vero) cells (American Type Culture Collection; catalog no.: CRL-1586), and HEK293T cells (American Type Culture Collection; catalog no.: CRL-3216) were maintained in Dulbecco’s modified Eagle’s medium (DMEM) with high glucose (Merck), supplemented with 10% (v/v) heat-inactivated fetal calf serum and 2 mM l-glutamine (complete DMEM) with high glucose (Merck), supplemented with 10% (v/v) heat-inactivated fetal calf serum and 2 mM l-glutamine (complete DMEM) in a humidified 5% CO2 atmosphere at 37 °C. For protein purification, Freestyle 293F suspension cells (Thermo Fisher Scientific) were grown in Freestyle 293F medium (Gibco) on a shaking platform (125 rpm) in a humidified 8% CO2 atmosphere at 37 °C.

Doxycycline-inducible stably transfected Freestyle 293F cells expressing StreptI-CERT1 and StreptI-CERT1S132A were generated using a piggyBac transposon-based system (80). A 30 ml suspension culture of Freestyle 293F cells at 1 × 106 cells/ml was transfected with a 5:1 mass ratio of PB–T–CERT1S132A[PBBN:PBase (35 μg total DNA) using Freestyle MAX transfection reagent (Invitrogen) as per the manufacturer’s instructions. After 2 days, the cells were transferred to fresh media supplemented with 500 μg/ml genetin (Gibco), and the drug selection was continued for 2 weeks with media replenishment every 3 days.

**GFP affinity capture**

Monolayers of HEK293T cells grown in 9 cm dishes were transfected with TransIT-LT1 (Mirus) using 7.7 μg of pEGFP-N1 (for GFP alone), pUL21-GFP, or point mutants thereof, following the manufacturer’s instructions. At 24 h post-transfection, the
cells were harvested by scraping into the medium, pelleted (220g, 5 min, 4 °C), washed three times with cold PBS, and lysed at 4 °C in 1 ml lysis buffer (10 mM Tris [pH 7.5], 150 mM NaCl, 0.5 mM EDTA, 0.5% NP-40, and 1:100 diluted EDTA-free protease inhibitor cocktail [Merck]) for 45 min before clarification (21,000g, 10 min, 4 °C). After immunoprecipitation with GFP-Trap beads (ChromoTek) performed in accordance with the manufacturer’s protocol, the samples were eluted by incubation at 95 °C for 5 min in 45 μl 2× SDS-PAGE loading buffer. Input and bound samples were separated by SDS-PAGE and analyzed by immunoblot.

**GST pulldown**

GST pull-down experiments were carried out in 96-well flat-bottomed plates (Greiner) using magnetic glutathione beads (Thermo Fisher Scientific) to capture the GST-tagged pUL21 (bait protein). First, 0.5 nmol of purified bait protein was incubated with the beads at 4 °C for 30 min. The beads were washed three times with wash buffer (20 mM Tris [pH 8.5], 200 mM NaCl, 0.1% NP-40, 1 mM EDTA, and 1 mM EDTA) and then incubated at 4 °C for 60 min with various truncations of myc-CERT1L (prey proteins), expressed using the TNT SP6 High Yield Wheat Germ in vitro transcription/translation system (Promega) in accordance with the manufacturer’s protocol, followed by four washes in wash buffer. Protein was eluted using wash buffer supplemented with 50 mM reduced glutathione before being analyzed by SDS-PAGE and immunoblotting.

**Antibodies**

The following antibodies with listed dilutions were used for immunoblotting: rabbit anti-CERT (1:10,000 dilution; Abcam; catalog no.: ab72536), mouse anti-pUL21 (1:50 dilution; clone no.: 1F10-D12) (18) for Figure 1F (note: this monoclonal antibody does not recognize pUL21V382E), rabbit anti-pUL21 (1:5000 dilution (10)) for Figs. 5A and S6A, mouse anti-GAPDH (1:10,000 dilution; GeneTex; catalog no.: GTX28245), mouse anti-Myc (1:4000 dilution; Millipore; catalog no.: 05-724), mouse anti-PP1α (1:1000 dilution; Santa Cruz; catalog no.: sc-271762), rabbit anti-pUL16 (1:2000 dilution) (84), mouse anti-VPS (1:50 dilution; catalog no.: DM165) (85), and rabbit anti-phospho-Akt substrates (1:1000 dilution; Cell Signalling; catalog no.: 9611). Fluorescently labeled secondary antibodies were used at 1:10,000 dilution: LI-COR IRDye 680T donkey anti-rabbit (catalog no.: 926-68023) and goat anti-mouse (catalog no.: 926-68020), LI-COR IRDye 800CW conjugated donkey anti-rabbit (catalog no.: 926-32213), and goat anti-mouse (catalog no.: 926-32210). For immunocytochemistry, mouse anti-pUL21 1:1 (clone no.: L1E4-C10), which was generated in the same immunization experiments as anti-pUL21 (clone no.: 1F10-D12) (18), and Alexa Fluor 488–conjugated goat anti-mouse (1:1000 dilution; Invitrogen; catalog no.: A21236) were used. For visualizing HSV-1 plaques, mouse anti-gD (LP2; 1:50 dilution) (86) and horseradish peroxidase–conjugated rabbit anti-mouse (1:5000 dilution; Diako; catalog no.: P0161) were used.

### HSV-1 pUL21 modulates cellular ceramide transport

Recombinant protein purification following bacterial expression

All recombinant proteins were expressed in Escherichia coli T7 Express lysY/F′ cells (New England Biolabs). Except for GST-PP1γ, cells were grown in 2× TY medium at 37 °C to an absorbance at 600 nm of 0.8 to 1.2 before cooling to 22 °C and inducing protein expression by addition of 0.4 mM IPTG. At 16 to 20 h postinduction, cells were harvested by centrifugation, and pellets were stored at −70 °C until required. For GST-PP1γ, the 2× TY medium was supplemented with 1 mM MnCl2, and the cultures were cooled to 18 °C upon reaching an absorbance of 0.8 at 600 nm, followed by induction using 1 mM IPTG. For all recombinant proteins, cells were resuspended in lysis buffer (see later) at 4 °C before lysis using a TS series cell disruptor (Constant Systems) at 24 kpsi. Lysates were cleared by centrifugation (40,000g, 30 min, 4 °C) and incubated with the relevant affinity resins for 1 h at 4 °C before extensive washing (≥20 column volumes) and elution using the relevant elution buffer (see later). Samples were concentrated and subjected to SEC (see later). Fractions containing the desired protein as assessed by SDS-PAGE were pooled, concentrated, snap-frozen in liquid nitrogen, and stored at −70 °C.

The lysis buffer for pUL21-GST (20 mM Tris [pH 8.5], 300 mM NaCl, 0.5 mM MgCl2, 1.4 mM β-mercaptoethanol, and 0.05% Tween-20) and GST-PP1γ (50 mM Tris [pH 7.5], 500 mM NaCl, 1 mM MnCl2, 0.5 mM MgCl2, 1.4 mM β-mercaptoethanol, and 0.05% Tween-20) was supplemented with 200 to 400 U bovine DNase I (Merck) and 200 μl EDTA-free protease inhibitor cocktail (Merck). Cleared lysates were incubated with glutathione Sepharose 4B (Cytiva), washed with wash buffer (20 mM Tris [pH 8.5, pUL21-GST; pH 7.5, GST-PP1γ], 500 mM NaCl, 1 mM DTT, plus 1 mM MnCl2 [GST-PP1γ only]), and the proteins were eluted using wash buffer supplemented with 25 mM reduced glutathione. SEC was performed using a HiLoad Superdex 200 16/600 column (Cytiva) equilibrated in 20 mM Tris (pH 8.5), 500 mM NaCl, 1 mM DTT (pUL21-GST) or 50 mM Tris (pH 7.5), 100 mM NaCl, and 1 mM DTT (GST-PP1γ). pUL21-H6, pUL21V382E-H6, and Hc-pUL21C were purified in Tris buffer at pH 8.5 in 500 mM NaCl, and Hc-minICERT1L was purified in Tris buffer at pH 7.5 in 150 mM NaCl. Lysis buffer (20 mM Tris, 20 mM imidazole, NaCl, 0.5 mM MgCl2, 1.4 mM β-mercaptoethanol, and 0.05% Tween-20) was supplemented with 200 to 400 U bovine DNase I (Merck) and 200 μl EDTA-free protease inhibitor cocktail (Merck). Cleared lysates were incubated with nickel–nitrilotriacetic acid agarose (Qiagen), washed with wash buffer (20 mM Tris, 20 mM imidazole, and NaCl) and eluted using elution buffer (20 mM Tris, 250 mM imidazole, and NaCl). SEC was performed using a HiLoad Superdex 200 (pUL21-H6 and pUL21V382E-H6) or 75 (Hc-pUL21C and Hc-minICERT1L) 16/600 column equilibrated in 20 mM Tris, NaCl, and 1 mM DTT.

Recombinant protein purification following mammalian cell expression

StrepII-CERT10 used for phosphatase assays was purified following transient transfection of Freestyle 293F cells, as
described before (18). StreptII-CERTL\textsuperscript{p} and StreptII-CERTL\textsuperscript{S132A} were purified from stably transfected Freestyle 293F cells following induction with 2 μg/ml doxycycline (Fisher) for 72 h. Next, cells were harvested by centrifugation (220g, 5 min, 4 °C) and washed once with ice-cold PBS before being resuspended in ice-cold lysis buffer (with phosphatase inhibitors, StreptII-CERTL\textsuperscript{p} only) (100 mM Tris [pH 8.0], 150 mM NaCl, 0.5 mM EDTA, 1 mM DTT [10 mM tetrasi-dium pyrophosphate, 100 mM NaF, and 17.5 mM β-glycerophosphate]). Cells were lysed by passage through a 23G needle six times, and lysates were clarified by centrifugation (40,000g, 30 min, 4 °C). The supernatants were sonicated at 50% amplitude for 60 s using a sonicating probe (MSE), and the supernatants were passed through a 0.45 μm syringe filter (Sartorius). StreptII-tagged proteins were captured using 1 ml StrepTrap HP column (Cytiva) that had been pre-equilibrated in wash buffer (100 mM Tris [pH 8.0], 150 mM NaCl, 0.5 mM EDTA, and 1 mM DTT). After extensive washing (20 column volumes), the protein was eluted using wash buffer supplemented with 2.5 mM desthiobiotin. Pooled eluate was applied in wash buffer (100 mM Tris [pH 8.0], 150 mM NaCl, 0.5 mM EDTA, and 1 mM DTT). After extensive washing (20 column volumes), the protein was eluted using wash buffer supplemented with 2.5 mM desthiobiotin. Pooled eluate was applied to a Superose 6 10/300 GL column (Cytiva) equilibrated in ITC buffer (20 mM Tris [pH 8.5], 500 mM NaCl, 0.5 mM Tris(2-carboxyethyl)phosphine [TCEP]), and fractions containing StreptII-CERTL as assessed by SDS-PAGE were pooled, concentrated, and used for downstream applications.

**Mutagenesis of viral genomes and generation of recombinant HSV-1**

All HSV-1 strain KOS viruses used in this study were reconstituted from a bacterial artificial chromosome (BAC) (87), and the mutated strain was generated using the two-step Red recombination method (54) with the following primers:

Forward: 5’-CGGCTCGTAGCCGGTGACACACAGGCCACGCTGCTAGCAGTACCCGACGACCCAGTAGATGAGCCGACTACTATAGGGG

Reverse: 5’-CGTTGATGCGATCGGCAAGACTGGTCGTCGGGCGCGCTTTTCGTAGACACAGGCGGTCGTCAGCAGTACCCGACGACCCAGTAGATGAGCCGACTACTATAGGGG

The generation of pUL21 deletion mutant (ΔpUL21) was described previously (18). To generate the p0 stocks, Vero cells were transfected with the recombinant BAC DNA together with pGS403 encoding Cre recombinase (to excise the BAC cassette) using TransIT-LT1 (Mirus) following the manufacturer’s instructions. After 3 days, the cells were scapped into the media, sonicated at 50% power for 30 s in a cup-horn sonicator (Branson), and titrated on Vero cell monolayers. The subsequent stocks were generated by infecting either Vero (HSV-1 WT) or HaCaT pUL21 cells (HSV-1 mutants) at an MOI of 0.01 for 3 days. The cells were then scraped and isolated by centrifugation at 1000g for 5 min. Pellets were resuspended in 1 ml of complete DMEM supplemented with 100 U/ml penicillin, 100 μg/ml streptomycin, and freeze/thawed thrice at ~70 °C before being aliquoted, titered on Vero cell monolayers, and stored at ~70 °C until required. The presence of the desired mutation in the reconstituted virus genomes was confirmed by sequencing the pUL21 gene.

**HSV-1 pUL21 modulates cellular ceramide transport**

Metabolic labeling was performed using subconfluent (60–80% confluence) HaCaT or HaCaT21 cells grown in a 6-well plate. For analysis of stable cell lines, the cells were pretreated for 30 min with complete DMEM containing 1 μM N-(3-hydroxy-1-hydroxymethyl-3-phenylpropyl)dodec-anamide (HPA-12; Tokyo Chemical Industry) dissolved in 0.1% dimethyl sulfoxide (DMSO; Merck), or 0.1% DMSO alone, and HPA-12 or DMSO was retained at the same concentrations throughout the subsequent incubation steps. For infection, cells were infected (see later) 14 h before metabolic labeling.

For metabolic labeling, cells were washed twice with warm PBS before incubation in 500 μl prewarmed DMEM with 1% (v/v) Nutridoma (Merck) supplemented with 5 μM alkyn-Sph (Cayman Chemical) for 5 min (pulse). Next, cells were washed twice with warm PBS, and 1 ml of prewarmed DMEM with 1% (v/v) Nutridoma was added to each well, followed by incubation at 37 °C for the indicated times of chase. Alkyn-Sph was stored as a 3.3 mM ethanolic stock solution at ~20 °C.

At the indicated times of chase, the plate with cells was transferred onto the ice, washed twice with 1 ml ice-cold PBS, scraped into 300 μl of ice-cold PBS, and transferred into appropriate 1.5 ml microcentrifuge tubes containing 600 μl of methanol. To each tube, 150 μl of chloroform was added, followed by vigorous vortexing. The precipitated protein was pelleted (20,000g, 1 min) at room temperature (RT). The organic supernatant was transferred to separate 2 ml tubes containing 300 μl chloroform. About 600 μl of 0.1% acetic acid in water was subsequently added to each tube to induce formation of two phases. Following extensive vortexing, the phases were separated by centrifugation (20,000g, 1 min, RT), and the lower phase was transferred to a new 1.5 ml microcentrifuge tube. This lipid-containing solvent phase was dried in a UniVapo centrifugal vacuum concentrator (UniEquip) at 30 °C for 20 min.

If required for SDS-PAGE analysis, the protein pellet was solubilized with 5 μl 10% (w/v) SDS, diluted in 75 μl lysis buffer (20 μM Tris 7.5, 150 mM NaCl, and 1% [v/v] Triton X-100), boiled for 10 min, and sonicated at 75% power for 60 s in a cup-horn sonicator. The protein samples were mixed with 5× loading buffer before being subjected to SDS-PAGE analysis.

**Click reaction and TLC of labeled lipids**

Lipid standards (16:0(alkyne)–18:1 PC [Avanti], pacFA GalCer [Avanti], d18:1(alkyne)–6:0 Cer [Cayman Chemicals], 18:0(alkyne) Sph [Cayman Chemicals]) or extracted lipids (aforementioned) were resuspended in 20 μl of 1:1 chloroform:ethanol. To each tube, 1 μl of Tris(1-benzyl-4-triazolyl) methyl)amine solution (2.5 mM in DMSO; Merck), 10 μl of Tetraakis(acetonitrile)copper(I) tetrafluoroborate (10 mM in acetonitrile; Merck), and 1 μl of 3-azido-7-hydroxycoumarin solution (1 mM in ethanol; Jena Bioscience) was added. After brief mixing, the reaction was dried in a UniVapo centrifugal vacuum concentrator (UniEquip) for 10 min at 45 °C.


**HSV-1 pUL21 modulates cellular ceramide transport**

Lipids were dissolved in 20 μl of 65:25:4:1 chloroform:methanol:water:acetic acid, of which 8 μl was spotted on a 20 × 10 cm HPTLC Silica gel 60 plates (Merck). When dried, the plate was developed for 5 cm with 65:25:4:1 chloroform:methanol:water:acetic acid, dried again, and developed for 9 cm with 1:1 hexane:ethylacetate. Labeled lipids were visualized via UV illumination using a G:BOX Chemi XX9 (Syngene), bands were quantified using the Image Studio Lite software (LI-COR) with local background subtraction, and statistical tests were performed using Prism 7 (GraphPad Software, Inc).

**Virus infections**

Monolayers of indicated cells were infected by overlaying with the appropriate viruses and diluted in complete DMEM to the specified MOI. The time of addition was designated 0 hpi. Cells were incubated with the inoculum at 37 °C in a humidified 5% CO₂ atmosphere for 1 h, rocking the tissue culture plate housing the monolayers every 15 min, before complete DMEM supplemented with 100 U/ml penicillin, 100 μg/ml streptomycin was added to dilute the original infection medium fivefold. Infected cells were incubated at 37 °C in a humidified 5% CO₂ atmosphere and harvested at the specified times.

**Immunoblotting**

For analysis of protein expression and phosphorylation, cells were washed two times with ice-cold 50 mM Tris (pH 8.5), 150 mM NaCl, and scraped into 100 μl of ice-cold lysis buffer (50 mM Tris [pH 8.5], 150 mM NaCl, 1% [v/v] Triton X-100, 1% [v/v] EDTA-free protease inhibitor cocktail [Merck], 10 mM tetrasodium pyrophosphate, 100 mM NaF, and 17.5 mM β-glycerophosphate). After 30 min of incubation on ice, the lysates were sonicated (two 15 s pulses at 50% power) in a cup-horn sonicator followed by centrifugation at 20,000g for 10 min. For all samples, protein concentrations were determined using a bicinchoninic assay (Pierce), and normalized protein lysates were analyzed by SDS-PAGE. For enhanced separation of phosphorylated proteins, 7% (w/v) acrylamide gels contained 50 μM MnCl₂ and 25 μM PhosTag reagent (Wako) where indicated. Separated proteins were then transferred to nitrocellulose membranes using the MiniPROTEAN system (Bio-Rad) and analyzed by immunoblotting and signals being detected using Odyssey CLx Imaging System (LI-COR). For quantification of CERT phosphorylation relative abundance, the signal for CERT-O band alone and for all CERT bands (CERT-O + CERT-P) were quantitated using Image Studio Lite software with local background subtraction. Statistical tests were performed using Prism 7.

**Analytical SEC of miniCERT̄₁−pUL21C complex**

Analytical SEC experiments were performed using Superdex 200 10/300 GL column equilibrated in 20 mM Tris (pH 8.5), 500 mM NaCl, and 1 mM DTT. To allow complex formation, 200 μl of 140 μM H₆-p miniCERT₁ and 680 μl of 55.4 μM H₆-pUL21C were mixed and incubated for 15 min at RT prior to injection.

**ITC**

ITC experiments were performed at 25 °C using a MicroCal PEAQ-ITC automated calorimeter (Malvern Panalytical). Proteins were transferred into ITC buffer (20 mM Tris [pH 8.5], 500 mM NaCl, and 0.5 mM TCEP) either by SEC or extensive dialysis prior to experiments. Titrants (StrepII-CERT₁, StrepII-CERT₁ S132A, or H₆ miniCERT₁) were titrated into titrates (pUL21-H₆, pUL21 V832E-H₆, or H₆-pUL21C) using either 19 × 2 μl or 12 × 3 μl injections (Table S1). Data were analyzed using the MicroCal PEAQ-ITC analysis software (Malvern Panalytical) and fitted using a one-site binding model.

**MALS**

MALS data were collected immediately following SEC (SEC–MALS) by inline measurement of static light scattering (DAWN 8+; Wyatt Technology), differential refractive index (Optilab T-rEX; Wyatt Technology), and UV absorbance (1260 UV; Agilent Technologies). Samples (100 μl) were injected onto an Superose 6 increase (all except H₆-pUL21C) or Superdex 75 increase (H₆-pUL21C) 10/300 GL columns equilibrated in 20 mM Tris (pH 8.5), 500 mM NaCl, and 0.5 mM TCEP at 0.4 ml/min. The protein samples were injected at the concentration of 2 mg/ml. Molecular masses were calculated using ASTRA 6 (Wyatt Technology) using a protein dn/dc of 0.186 ml/g.

**SAXS**

SAXS experiments were performed in batch mode (H₆-pUL21C and H₆-miniCERT₁) or inline following SEC (H₆-pUL21C–H₆ miniCERT₁) at EMBL-P12 bioSAXS beamline (PETRAIII) (88, 89).

For SAXS in batch mode, scattering data (I(θ)) versus s, where $s = 4\pi\sin(θ)/λ$ nm⁻¹, 2θ is the scattering angle, and λ is the X-ray wavelength, 0.124 nm) were collected from the indicated batch samples and a corresponding solvent blank (20 mM Tris [pH 8.5], 500 mM NaCl, 3% [v/v] glycerol, and 1 mM DTT). The scattering profiles of H₆-miniCERT₁ (1.73, 2.59, 3.46, and 3.71 Å) were measured in continuous-flow mode using an automated sample changer (30 μl sample at 20 °C; 1 mm path length). The sample and buffer measurements were measured as 100 ms data frames on a Pilatus 6M area detector (Dectris) for total exposure times of 2.1 and 2.6 s, respectively. For H₆ miniCERT₁, the resulting SAXS curves were extrapolated to infinite dilution.

For SEC–SAXS experiments, the scattering data were recorded using a Pilatus 6M detector with 1 s sample exposure times for a total of 3600 data frames spanning the entire course of the SEC separation. Proteins were mixed at 1:1 molar ratio, incubated on ice for 15 min, and then concentrated using a centrifugal concentrator to the desired concentration as estimated from absorbance using extinction coefficients (90) calculated assuming a 1:1 molar ratio. About 40 μl of purified
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The structure of HSV-1 pUL16 (UniProt ID: P10200) in complex with pUL21C (residues 275–535, UniProt ID: F8RG07) was predicted using a locally installed version of AlphaFold-Multimer, version 2.2.2 (52). Structural similarity searches and superpositions were performed using the DALI web server (53). Molecular images were generated using PyMOL.

**Differential scanning fluorimetry**

Differential scanning fluorimetry experiments were carried out using Viia7 real-time PCR system (Applied Biosystems) and 1x Protein Thermal Shift dye (Applied Biosystems). The assay buffer (20 mM Tris [pH 8.5], 500 mM NaCl, and 1 mM DTT) was mixed with dye stock solution and protein solution (H6-pUL21 or H6-pUL21V382E) in an 8:1:1 ratio, giving 1 ng protein in a final volume of 20 μl, and measurements were performed in triplicate. Samples were heated from 25 to 95 °C at 1 °C per 20 s, and fluorescence was monitored at each increment. The melting temperature ($T_m$) is the inflection point of the sigmoidal melting curve, determined by nonlinear curve fitting to the Boltzmann equation using Prism 7.

**In vitro dephosphorylation assays**

Dephosphorylation assays were performed upon 0.5 μM of purified CERTΔ9 using a fixed concentration of GST-PP1y (25 nM) in the presence of H6-pUL21 or H6-pUL21V382E at the indicated concentrations. Reactions (50 μl) proceeded in assay buffer (150 mM NaCl, 20 mM Tris [pH 8.5], 0.1% Tween-20, and 1 mM MnCl₂) for 30 min at 30 °C before being stopped by the addition of 50 μl of 2× SDS-PAGE loading buffer and boiling at 95 °C for 5 min. Samples were analyzed by SDS-PAGE using 7% (w/v) acrylamide gels supplemented with 25 μM PhosTag reagent and 50 μM MnCl₂, and the protein was visualized using InstantBlue Coomassie stain (Expedeon). To measure the ratio of CERTΔ9 to total CERT, Coomassie-stained gels were scanned using an Odyssey CLx Imaging System (LI-COR). The signals detected in the 700 nm channel for the CERTΔ9 band alone and for all CERT bands (CERTO + CERTΔ9) were quantitated using Image Studio Lite software with local background subtraction.

**Plaque size analysis**

Confluent monolayers of Vero or HaCaT cells in 6-well tissue culture plates were infected with 100 plaque-forming units of the indicated virus diluted in complete DMEM to a final volume of 500 μl. Following the adsorption for 1 h at 37 °C in a humidified 5% CO₂ atmosphere, rocking the plate every 15 min, the cells were overlaid with plaque assay media (DMEM supplemented with 0.3% high viscosity carboxymethyl cellulose, 0.3% low viscosity carboxymethyl cellulose, 2% [v/v] fetal calf serum, 2 mM L-glutamine, 100 U/ml penicillin, and 100 μg/ml streptomycin) and incubated for further 48 h. Next, cells were fixed with 3.7% (v/v) formal saline for 20 min, washed three times with PBS, and incubated for 1 h with mouse anti-gD (LP2) (86), diluted 1:50 in blocking buffer (PBS supplemented with 1% [w/v] bovine serum albumin and 0.1% H₆-pUL21C–H₆-miniCERTΔ₉ (4.3 mg/ml) was injected at 0.35 ml/min onto a Superdex 200 Increase 5/150 GL column (Cytiva) equilibrated in 20 mM Tris (pH 8.5), 500 mM NaCl, 3% (v/v) glycerol, and 1 mM DTT. SAXS data were recorded from a single peak (frames 1245–1296 s), and solvent blank was collected from pre-elution frames (frames 247–411). Primary data reduction was carried out using CHROMIXS (91), and 2D-to-1D radial averaging was performed using the SASFLOW pipeline (92). Processing and analysis of the SAXS output was performed using the ATSAS 3.0.2 software package (93). The extrapolated forward scattering intensity at zero angle, I(0), and the radius of gyration, R₉, were calculated from the Guinier approximation (lnI(s) versus s², for sR₉ < 1.3). The maximum particle dimension, D_max, was estimated based on the probable distribution of real-space distances, p(r), which was calculated using GONI (94). A concentration-independent estimate of molecular weight was determined using a Bayesian consensus method (95). All structural parameters are reported in Table S2.

Ab initio modeling was performed using GASBOR (49), and figures show the models that best fit their corresponding SAXS profiles (lowest χ²). Pseudoatomic modeling was performed using CORAL (96). For H₆-miniCERTΔ₉, crystal structures of the CERT PH (Protein Data Bank [PDB] ID: 4HHV) (24) and START (PDB ID: 2E3M) (25) domains were used, the latter including predicted secondary structure as modeled by I-TASSER (97) for intron 11 (residues 275–535) using BLASTP (100) for simplexviruses (taxid: 10294). Truncated sequences (accession numbers 371–396), present only in CERTL. The miniCERTL 23 amino acid linker and the purification tag (MAH₆) were modeled using CORAL. The model with the best fit was superimposed onto the PH:START crystal structure (PDB ID: 5JJD) (46) by aligning the START domains using PyMOL, version 2.5.2 (Schrödinger). For H₆-pUL21C, the N-terminal (MAH₆QD) and C-terminal (HGQSV) elements missing from the crystal structure (PDB ID: 5ED7) (51) were modeled using CORAL. For H₆-pUL21C–H₆-miniCERTΔ₉, the heterodimer was modeled using the SAXS curve of the complex using a fixed conformation of H₆-pUL21C and mobile CERT domains, with unstructured elements modeled as aforementioned. All the models were assessed for fit to the corresponding SAXS profiles (χ² and CorMap) using CRYSTOL (98). Molecular images were generated using PyMOL.

**Sequence analysis, deep learning structure prediction, and structural analysis**

Sequences of the C-terminal domain from pUL21 homologs were identified by querying the National Center for Biotechnology Information nonredundant protein sequence database (99) with HSV-1 pUL21 (UniProt ID: F8RG07) residues 275 to 535 using BLASTP (100) for simplexviruses (taxid: 10294) and PSI-BLAST with two iterations for alphaherpesviruses (taxid: 10294). Truncated sequences (accession length <80% of HSV-1 pUL21) or those containing ambiguous amino acids (“X”) were manually removed. Sequences were aligned using COBALT (101) and are provided as supporting information (Data S1).
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Tween-20. Cells were washed thrice in PBS and incubated for 30 min with horseradish peroxidase–conjugated rabbit anti‐mouse antibody (DaKo; catalog no.: P0161) diluted 1:5000 in blocking buffer, followed by two subsequent PBS washes and one wash with ultrapure water. Plaques were visualized using the TrueBlue peroxidase substrate in accordance with the manufacturer’s instructions (Saracare). Plaques were scanned at 1200 dpi, and plaque areas were measured using Fiji (102, 103).

Single-step (high MOI) virus growth assays

Virus growth assays were performed in technical duplicate for each independent experiment. Monolayers of HaCaT or Vero cells were infected with the indicated viruses diluted in complete DMEM to an MOI of 10. The time of virus addition was designated 0 hpi. After adsorption for 1 h at 37 °C in a humidified 5% CO₂ atmosphere, rocking the plates every 15 min, extracellular virus particles were neutralized with an acid wash (40 mM citric acid, 135 mM NaCl, 10 mM KCl, pH 3.0) for 1 min, and cells were then washed three times with PBS before being subsequently overlaid with complete DMEM. At designated times postinfection, cells were harvested by freezing the plate at −70 °C. When all plates were frozen, samples were freeze/thawed one more time before scraping and transferring to 1.5 ml tubes and storing at −70 °C until titration. Titrations were performed on monolayers of Vero cells. Serial dilutions of the samples were used to inoculate the cells for 1 h, followed by overlaying with DMEM containing 0.3% high-viscosity carboxymethyl cellulose, 0.3% low-viscosity carboxymethyl cellulose, 2% (v/v) fetal bovine serum, 2 mM L-glutamine, 100 U/ml penicillin, and 100 μg/ml streptomycin. Three days later, cells were fixed in 3.7% (v/v) formal saline for 20 min, washed with water, and stained with 0.1% toluidine blue. Statistical tests were performed using Prism 7.

Virus release assays

Monolayers of HaCaT or Vero cells were infected as described previously for single-step virus growth assays, infections being performed in technical duplicate or triplicate for each independent experiment. At 12 hpi, the media (500 μl) were harvested to 1.5 ml Eppendorf tubes, centrifuged for 5 min at 1000 g to remove any detached cells, and 300 μl of the supernatant was carefully transferred to fresh tubes and stored at −70 °C until titration. The cells were overlaid with 500 μl of fresh DMEM, and the plates were immediately frozen at −70 °C. Titrations were performed on monolayers of Vero cells as described previously. Statistical tests were performed using Prism 7. Figure 6C was generated using SuperPlotsOfData (104).

Immunocytochemistry

Cells grown on #1.5 coverslips were infected at an MOI of 1 as described previously. At 14 hpi, cells were washed with PBS and incubated with freezing cold (−20 °C) methanol for 5 min at −20 °C. Coverslips were washed with PBS, followed by incubation with blocking buffer (1% [w/v] bovine serum albumin in PBS) for 30 min at RT. Primary antibodies (aforementioned) were diluted in blocking buffer and incubated with coverslips for 1 h. Coverslips were washed 10 times with PBS before incubation for 45 min with the secondary antibodies (aforementioned) diluted in blocking buffer. Coverslips were washed 10 times in PBS and 10 times in ultrapure water before mounting on slides using Mowiol 4-88 (Merck) containing 200 nM 4′,6-diamidino-2-phenylindole. Images were acquired using an inverted Olympus IX81 widefield microscope with a 60× Plan Apochromat N oil objective (numerical aperture of 1.42) (Olympus) and Retiga EXi Fast1394 interline CCD camera (QImaging).

Data availability

The SAXS data measured for each concentration, with an accompanying report, are available in the Small Angle Scattering Biological Data Bank (SASBDB) (105), entries SASDNB7, SASDNC7, and SASDND7. The AlphaFold-Multimer model of the HSV-1 pUL16–pUL21C complex, with associated raw output and quality statistics, has been deposited in the University of Cambridge Data Repository (https://doi.org/10.17863/CAM.88729).

Supporting information—This article contains supporting information (18, 50, 51, 73, 92, 106–111).

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**Abbreviations**—The abbreviations used are: BAC, bacterial artificial chromosome; Cer, ceramide; DMEM, dulbecco’s modified eagle’s medium; DMSO, dimethyl sulfoxide; ER, endoplasmic reticulum; GST, glutathione-S-transferase; HEK293T, human embryonic kidney 293T cell line; hpi, hours postinfection; HPTLC, high-performance TLC; HSV, herpes simplex virus; ITC, isothermal titration calorimetry; MOI, multiplicity of infection; MR, middle region; NEC, nuclear egress complex; PC, phosphatidylcholine; PDB, protein data bank; PH, pleckstrin homology; PP1, protein phosphatase 1; RT, room temperature; SAXS, small-angle X-ray scattering; SEC–MALC, size-exclusion chromatography with inline multiangle light scattering; SM, sphingomyelin; Sph, sphingosine; START, steroiodogenic acute regulatory–related lipid transfer; TCEP, Tris(2-carboxyethyl)phosphine; TGN, trans-golgi network.

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