



Functional interactions between herpes simplex virus pUL51, pUL7 and gE reveal cell-specific mechanisms for epithelial cell-to-cell spread

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ABSTRACT

Herpes simplex virus spread between epithelial cells is mediated by virus tegument and envelope protein complexes including gE/gI and pUL51/pUL7. pUL51 interacts with both pUL7 and gE/gI in infected cells. We show that amino acids 30–90 of pUL51 mediate interaction with pUL7. We also show that deletion of amino acids 167–244 of pUL51, or ablation of pUL7 expression both result in failure of gE to concentrate at junctional surfaces of Vero cells. We also tested the hypothesis that gE and pUL51 function on the same pathway for cell-to-cell spread by analyzing the phenotype of a double gE/UL51 mutant. In HaCaT cells, pUL51 and gE function on the same spread pathway, whereas in Vero cells they function on different pathways. Deletion of the gE gene strongly enhanced virus release to the medium in Vero cells, suggesting that the gE-dependent spread pathway may compete with virion release to the medium.

1. Introduction

Assembly of mature, multi-layer herpesvirions occurs by budding of capsids into a cytoplasmic membrane compartment followed by trafficking of the enveloped virion to the cell surface for release, or to junctional surfaces for cell-to-cell spread (CCS) (reviewed in (Johnson and Baines, 2011)). The identity of the cytoplasmic membrane compartment used for final envelopment apparently differs between herpesvirus species, but is derived by modification of host cell structures. Human cytomegalovirus (HCMV) for example, undergoes cytoplasmic envelopment in a discrete assembly compartment constructed by massive reorganization of host Golgi and endosomal membranes (Cepeda et al., 2010; Das et al., 2014; Das and Pellett, 2011; Homman-Loudiyi et al., 2003; Sanchez et al., 2000; Schauflinger et al., 2013; Severi et al., 1988). HSV-1, on the other hand, undergoes cytoplasmic envelopment in multiple locations in the cytoplasm. The nature of the enveloping membrane for HSV-1 is not entirely clear. Secondary envelopment at the *trans*-Golgi network (TGN) has been proposed based on membrane composition of the mature virion, association of capsids with membranes containing TGN markers (Sugimoto et al., 2008; Turcotte et al., 2005). Secondary envelopment at an endosomal compartment is supported by the presence endocytosed horseradish peroxidase in the lumen of enveloping membrane and co-localization of capsids with

transferrin receptor (Hollinshead et al., 2012).

The herpesvirus tegument is a loosely ordered protein layer that lies between the capsid and the envelope (Laine et al., 2015). It consists of at least 20 virus-encoded proteins (reviewed in (Owen et al., 2015)). Tegument proteins are critical for multiple functions late in the virus replication cycle, including assembly of the mature virus particle and trafficking of virus particles for CCS. Interestingly, these functions are not delegated among different sets of proteins, but rather are dual functions of many and, perhaps, most tegument proteins.

The HSV-1 UL51 gene encodes a 244 a.a. palmitoylated tegument protein (Daikoku et al., 1998; Nozawa et al., 2003). A complete deletion of any alphaherpesvirus UL51 gene has not yet been constructed because the UL51 protein coding sequence contains promoter/regulatory sequences for the UL52 gene that encodes one of the helicase/primase subunits of the viral DNA replication apparatus. Alphaherpesvirus UL51 gene function has, therefore been explored by the use of partial deletions that remove most of the protein coding sequence (Klupp et al., 2005; Nozawa et al., 2005; Roller et al., 2014) or by insertion of stop codons a short distance downstream of the initiation codon (Albecka et al., 2017). There are apparent minor differences in the phenotypes obtained with these different approaches, but all of them suggest pUL51 has cell-specific functions in both virion assembly and CCS. Single-step growth in these various mutant viruses is depressed up to 100-fold in

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some cell lines, including Vero (Klupp et al., 2005; Nozawa et al., 2005; Roller et al., 2014; Albecka et al., 2017), and this growth defect has been correlated with accumulation of unenveloped, and sometimes membrane-associated capsids in the cytoplasm (Klupp et al., 2005; Albecka et al., 2017). This suggests that one function of pUL51 is to facilitate curvature or closure of membrane around the capsid/tegument complex in cytoplasmic assembly. Interestingly, however, single-step growth defects were not observed for an HSV-1 partial deletion mutant on HEp-2 cells (Roller et al., 2014), suggesting that the pUL51 assembly function can be complemented by host cell factors in some cell types.

pUL51 of HSV-1 forms a complex with another viral tegument protein, pUL7 (Albecka et al., 2017; Roller and Fetters, 2015). This complex is necessary for incorporation of pUL7 into the mature virion (Roller and Fetters, 2015). A double mutant made by stop codon insertion into UL51 and deletion of UL7 showed a defect in single step growth in Vero and HaCaT cells that was no greater than the defects of the individual deletions, suggesting that pUL51 and pUL7 function on the same pathway, and probably as a complex in assembly (Albecka et al., 2017). The UL51 and UL7 genes are conserved among herpesviruses, and deletion of their homologs in HCMV (UL71 and UL103, respectively) causes defects in formation of the assembly compartment and cytoplasmic envelopment, and results in formation of smaller infection foci, suggesting some conservation of function as well (Ahlgvist and Mocarski, 2011; Schauflinger et al., 2011; Womack and Shenk, 2010).

All alphaherpesvirus UL51 mutants reported so far show profound defects in plaque formation, even in cells in which no single-step growth defect is observed, suggesting that pUL51 plays a critical role in spread between epithelial cells (Klupp et al., 2005; Nozawa et al., 2005; Roller et al., 2014; Albecka et al., 2017). Since the pUL51/pUL7 complex is not displayed on the cell surface, its most likely function in CCS is facilitating delivery of virions to junctional surfaces of cells.

The UL51 gene product is one of several alphaherpesvirus gene products that play an important role in epithelial cell-cell spread (CCS). The best described of these is the gE/gI complex. gI and gE are encoded on the adjacent US7 and US8 genes respectively and, upon translation at the ER, form a stable complex (Johnson and Feenstra, 1987; Johnson et al., 1988). The gE/gI complex is an abundant component of the mature virion envelope, where it functions in immune evasion by interfering with effector functions of IgG by binding to the Fc region (Lubinski et al., 1998; Ndjamen et al., 2014). In addition to its immune evasion function, the gE/gI complex facilitates CCS in epithelial cells by a mechanism that has been reported to require its localization to adherens junctions where it co-localizes with β -catenin (Balan et al., 1994; Dingwell et al., 1994; Dingwell and Johnson, 1998; Farnsworth and Johnson, 2006; McMillan and Johnson, 2001; Wisner et al., 2000). Failure to express gE, or failure of gE/gI to localize to junctions results in formation of small plaques in the presence of neutralizing antibody, and in diminished numbers of virus particles found between cells (Johnson et al., 2001). The mechanism by which gE/gI facilitates spread is not clear, but its spread function can be inhibited by over-expression of the soluble ectodomain of gE, suggesting that an interaction with adjacent cells may be required (Collins and Johnson, 2003).

The localization and CCS function of gE are regulated by viral tegument proteins. A complex of pUL11, pUL16, and pUL21 can form on the cytoplasmic tail of gE, and expression of these proteins is required for gE cell-surface localization (Chadha et al., 2012; Han et al., 2011, 2012; Yeh et al., 2011). Failure to form this complex is associated with defects in CCS and syncytium formation, suggesting that gE/gI CCS function depends on these tegument proteins (Han et al., 2012). A similar set of observations implicates pUL51 in gE CCS function. A UL51 mutant produced by fusion with EGFP at the C-terminus causes failure of gE to accumulate at cellular junctions in Vero cells, and this is correlated with inhibition of syncytium formation and a severe defect in CCS in Vero cells (Roller et al., 2014). pUL51 also interacts in the

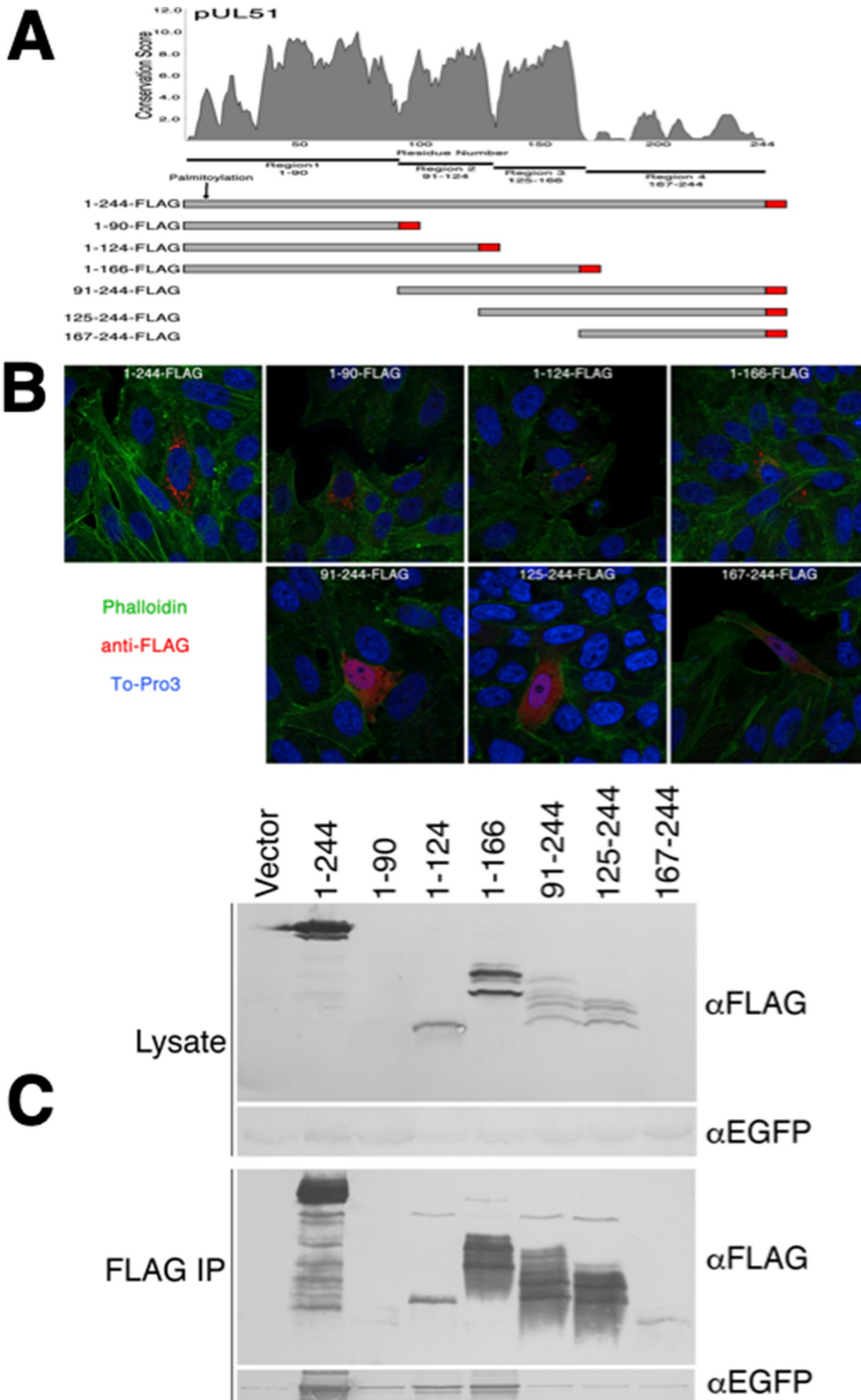
infected cell with gE (Roller et al., 2014). However, pUL51 may have other functions in CCS, since a UL51 deletion has a far more powerful inhibitory effect on CCS than a deletion of gE (Roller et al., 2014).

All of these observations suggest the existence of an epithelial CCS pathway in which delivery of the gE/gI complex to cellular junctions is a central event, regulated by interaction between the cytoplasmic tails of gE, gI, or both with tegument protein complexes. The hypothesis that the pUL51/pUL7 complex, the pUL11/pUL16/pUL21 complex, and the gE/gI complex operate on the same pathway for CCS predicts that multiple mutants that affect formation of multiple complexes will have a phenotype no more severe than that of the single mutants. Here we show that this prediction is borne out for a UL51/gE double mutant in HaCaT cells, but not in Vero cells, and that the behavior of gE with respect to junctional localization is cell type dependent as well.

2. Results

2.1. pUL51 sequences that are necessary and sufficient for complex formation with pUL7 are located between a.a. 30 and 90

pUL51 is composed of two sequence regions – an N-terminal region from amino acids 1–166 that contains sequence that is well conserved among pUL51 homologs and that is predicted to be globular, and a C-terminal region from amino acid 167 to the end of the protein that is poorly conserved and is predicted to be disordered. It has been previously shown that interaction with pUL7 can be mediated by sequences corresponding roughly to the globular domain of pUL51 from residues 29–170 (Albecka et al., 2017; Roller and Fetters, 2015). The N-terminal region can be further subdivided into four conserved regions comprising a.a. 1–20, 21–90, 91–124, and 125–166 (Fig. 1A) (Roller and Fetters, 2015). To see whether a smaller pUL7 interaction region could be defined, we created pcDNA3-based plasmid constructs that express N- and C-terminal truncations of a FLAG-tagged pUL51 protein coding sequence with boundaries corresponding to the pUL51 conserved regions (Fig. 1A). We first tested the subcellular localization of these truncated proteins. Although pUL51 does not have a transmembrane domain it is tightly associated with membranes in infected and transfected cells due to palmitoylation at C9 near the N-terminus (Nozawa et al., 2003). As expected, only the truncation constructs that included the pUL51 N-terminus showed a punctate cytoplasmic distribution with variation in the number and distribution of the puncta due to association with cytoplasmic membranes (Fig. 1B). Those that did not include the N-terminus were invariably distributed diffusely in the cytoplasm and nucleus. We then tested those truncated proteins for ability to interact with EGFP-pUL7 in co-immunoprecipitation assays (Fig. 1C). Expression of the truncated pUL51 constructs was variable, probably due to differing stability of the truncated proteins, and the shortest truncations (1–90 and 167–244) were detectable only in immunoprecipitated samples. Only constructs that contained residues 1–90 of pUL51 reproducibly co-immunoprecipitated EGFP-pUL7 above the vector background, suggesting that these residues were necessary and sufficient for pUL7 interaction. The pull-down by pUL51(1–90), however, was weak, probably due to the very low expression of this construct. Therefore, to confirm and extend this result, we created in-frame fusion constructs in which pUL51-FLAG truncations were fused to the C-terminus of the viral glycoprotein D (gD) and expressed them via transfection with EGFP-pUL7 (Fig. 2). Consistent with our previous report (Roller and Fetters, 2015), expression of EGFP-pUL7 in combination with the control gD-FLAG, resulted in diffuse localization concentrated in the cell nucleus in all transfected cells (Fig. 2A). Co-expression with gD-pUL51(1–90)-FLAG, in contrast, resulted in recruitment of EGFP-pUL7 to cytoplasmic membranes where the two proteins co-localized (Fig. 2B). While co-expression of EGFP-pUL7 and gD-pUL51(1–90)-FLAG always results in strong co-localization of the two proteins on cytoplasmic membranes, EGFP-pUL7 was rarely completely recruited to pUL51-labeled membranes – some was found



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Fig. 1. Localization and pUL7 interactions of pUL51 truncations. (A) Conservation plot of pUL51 protein coding sequence and schematic diagram of FLAG-tagged pUL51 truncations. The plot shows conservation of biochemical properties of amino acids using all available herpesvirus pUL51 homologous sequences aligned using the program MUSCLE (Edgar, 2004). Each residue position receives a conservation score, and scores were averaged over a sliding 5 amino acid residue window. The schematic underneath the plot shows the boundaries of the pUL51 truncations used in this study. The FLAG tag is indicated in red. (B) Localizations of pUL51 truncations in transfected Vero cells. Nuclei are stained with To-Pro3 (blue), actin stress fibers are stained with phalloidin (green) and pUL51 truncations are detected with mouse anti-FLAG (red). Representative images from three independent experiments in which > 50 transfected cells were observed are shown. (C) Co-immunoprecipitation of EGFP-pUL7 with pUL51 truncations. Lysates and immunoprecipitates from Vero cells co-transfected with the indicated pUL51 constructs and with pEGFP-pUL7 plasmid are shown. The top two panels show lysate proteins detected by immunoblot using either anti-FLAG or anti-EGFP. The bottom two panels show FLAG immunoprecipitates detected with either anti-FLAG or anti-EGFP. One of two independent experiments is shown. (For interpretation of the references to colour in this figure legend, the reader is referred to the Web version of this article.)

diffusely in the nucleoplasm in most cells. This may reflect differences in expression levels of the constructs. We made a set of further truncations of the pUL51 N-terminal region and found that truncations that included amino acids 30–90 of pUL51 (Fig. 2B, E, and H) could recruit EGFP-pUL7 to membranes, whereas truncations that lacked any of amino acids 30–90 (Fig. 2C, D, F, and H) did not, suggesting that conserved region 2 of pUL51 is necessary and sufficient for interaction with pUL7.

Interestingly, the gD-pUL51(1–90)-FLAG fusion construct localized differently than the control gD-FLAG. gD-FLAG is found on membranes distributed throughout the cytoplasm and on the cell surface (Fig. 2A). gD-UL51(1–90)-FLAG is found in a reticular distribution in the cytoplasm where it co-localizes completely with the ER marker TRAP α (Fig. 2B). Furthermore, whereas gD-FLAG migrates in SDS-PAGE as a heterogeneous collection of bands typical of a glycoprotein that has matured through the Golgi, gD-UL51(1–90)-FLAG migrates largely as a single species typical of immature glycoproteins (Fig. 2G). All gD fusions that contained amino acids 1–30 of pUL51 co-localized with TRAP α (Fig. 2B, C and D), whereas those that lacked these amino acids did not (Fig. 2E and F). These results suggested that the fusion of gD to the N-terminal 30 amino acids of pUL51 results in retention of gD in the ER.

2.2. Deletion of the C-terminal third of pUL51 completely ablates its CCS function

We have previously described the construction and characterization of a recombinant HSV-1 virus in which amino acids 73–244 of pUL51 were deleted, and found that it showed significant defects in both virus production, virus release to the medium, and cell-to-cell spread in Vero cells (Roller et al., 2014). In a subsequent study, we reported construction of a virus carrying a smaller partial deletion that does not express the poorly conserved C-terminal third of pUL51 (a.a. 167–244) (Fig. 3A line 5) (Roller and Fetters, 2015). In that study, we noted that the virus carrying this smaller deletion, while retaining the interaction between pUL51 and pUL7, appeared to be as deficient in plaque formation as the UL51 Δ 73-244 mutant (Fig. 3A line 3). In order to confirm and quantitate this effect, we performed plaque size assays in the presence of neutralizing antibody (Fig. 4). On Vero cells, both the UL51 Δ 73-244 and UL51 Δ 167-244-FLAG viruses formed plaques significantly smaller than the wild type control ($P < 0.001$ for both comparisons), and these plaques were ~30-fold smaller than those formed by wild-type virus. The spread defect observed for both deletion viruses could be completely complemented by plaque formation on wild-type pUL51-expressing complementing cells, demonstrating that the defect was solely due to the mutation in the UL51 gene.

2.3. Mutation of either pUL51 or pUL7 impairs accumulation of gE at cell junctions

pUL51 and gE have both been shown to play important roles in epithelial cell-to-cell spread, and we have previously demonstrated a physical interaction between them that suggests that their functions in cell-to-cell spread might be coupled (Roller et al., 2014). It has also been previously reported that gE in infected cells concentrates at

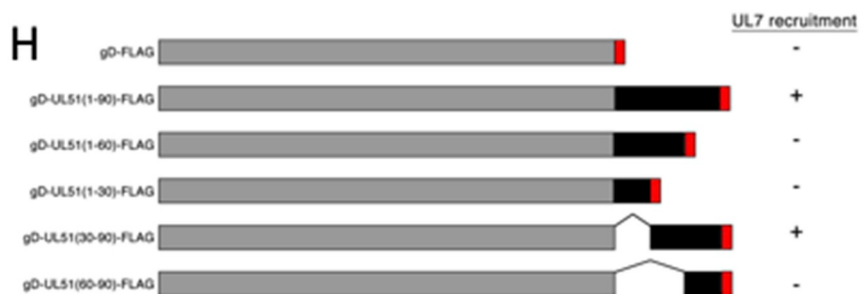
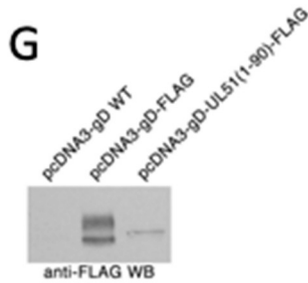
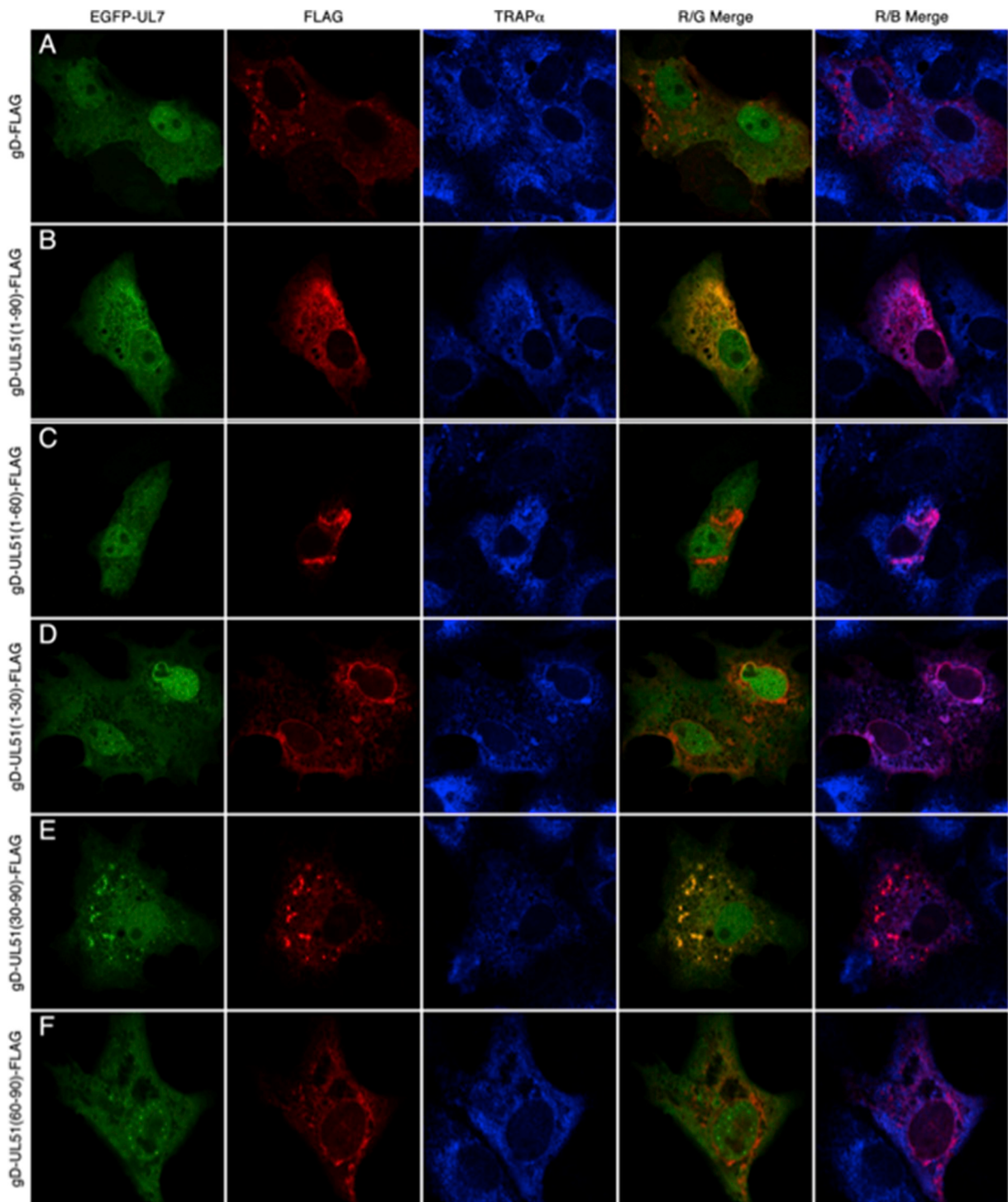
cellular junctions where it co-localizes with β -catenin, suggesting a localization to adherens junctions (Dingwell and Johnson, 1998), and that mutation of pUL51 impairs localization of gE to junctional surfaces of Vero cells (Roller et al., 2014). We have also shown that pUL51 forms a complex with pUL7, and that pUL7 also functions in cell-to-cell spread (Roller and Fetters, 2015).

In order to determine whether the UL51 Δ 167-244 mutant or a UL7 null mutation also disrupt gE localization, we first generated a UL7-null virus (Fig. 3A line 7), and demonstrated that it expresses no detectable pUL7 (Fig. 3C). We then infected cells with wild-type, UL51 mutant, or UL7 mutant viruses for 12 h, and then visualized gE and β -catenin localization (Fig. 5). Although Vero cells form adherens junctions (Krummenacher et al., 2003), they do not form polarized epithelia and do not form a continuous belt of β -catenin staining. Rather, we observed β -catenin staining in numerous small patches at the junctional surfaces of uninfected cells (Fig. 5A and D) that visually define those junctional surfaces. We observed no difference in the staining intensity or distribution of β -catenin 12 h after infection with any of the viruses (Fig. 5A, E, I, and M).

As previously reported, in wild-type virus-infected cells, gE concentrates at the nuclear membrane, on aggregates of membranes in the cytoplasm, and at the junctional surfaces of cells (Roller et al., 2014). Here, we observed that, although gE and β -catenin both concentrated at junctional surfaces, they were not uniformly distributed and there was very little co-localization between patches of β -catenin and gE staining (Fig. 5H, boxed area). Both the UL51 Δ 167-244 and UL7-null mutant viruses showed almost no gE immunofluorescent staining at junctional surfaces (Fig. 5I–P), demonstrating that both deletions impair gE junctional localization in infected cells. Quantitation of gE junctional staining (Fig. 5Q) showed that both UL51 deletion and UL7-null viruses showed a similar degree of inhibition.

2.4. pUL51 and gE double deletions have different spread phenotypes in HaCaT and vero cells

The importance of both gE and pUL51 in epithelial cell-to-cell spread, the physical interaction between gE and pUL51, and the effect of pUL51 deletion on gE localization suggested the hypothesis that pUL51 and gE function on the same pathway for cell-to-cell spread and that function of gE in spread might be partially or fully dependent upon pUL51. To test this hypothesis, we created a pair of independently constructed recombinant mutant viruses deficient in pUL51 spread function and in gE expression by combining the UL51 Δ 167-244 mutation with a deletion in the US8 gene that abrogates expression of gE (Fig. 3A, Line 6). Immunoblot analysis showed that these viruses express, as expected, a truncated, FLAG-tagged pUL51, and fail to express detectable gE (Fig. 3B). Also as noted previously, the truncated pUL51 is expressed at a considerably lower level than the wild-type protein (Roller et al., 2014). Previous results evaluating the effects of pUL51 deletion showed cell-specific effects on virus replication, spread and release to the medium (Roller et al., 2014). Specifically, we observed that both growth and spread defects were considerably greater in Vero than in HEP-2 cells, suggesting that the functions of pUL51 in Vero and human epithelial cells might differ. We therefore evaluated the growth and spread phenotypes of these viruses on Vero cells and on the human



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Fig. 2. Interaction of pUL7 with gD-UL51 truncation fusions. (A–F) Digital images of cells that are immunofluorescently stained and detected by confocal microscopy are shown. The gD fusion construct used is indicated to the left of each panel. EGFP-pUL7 is shown in green, FLAG staining in red, and staining for the ER marker TRAP α is shown in blue. Red/green merge images show co-localization between gD fusions and EGFP-pUL7. Red/blue merged images show co-localization between gD fusions and TRAP α . Representative images from two independent experiments in which > 50 transfected cells were observed are shown. (G) Immunoblot detection of gD fusions using anti-FLAG antibody. (H) Summary schematic showing the structures of the gD fusions used for this experiment, and their ability to recruit EGFP-pUL7 to membranes. (For interpretation of the references to colour in this figure legend, the reader is referred to the Web version of this article.)

HaCaT keratinocyte cell line.

In HaCaT cells, as previously reported, deletion of gE significantly ($P < 0.01$) inhibits cell-to-cell spread of virus in the presence of neutralizing antibodies so that mutant virus plaques are roughly three-fold smaller than the wild-type control (Fig. 6A) (Dingwell and Johnson, 1998). Deletion of a.a. 167–244 of pUL51 had a more dramatic effect on spread, so that plaques were roughly 10-fold smaller than wild type. The double deletion plaques, however, were no smaller than the plaques formed by the UL51 single deletion, suggesting that, in HaCaT cells, UL51 and gE function on the same cell-to-cell spread pathway. The effects of the single and double deletions on plaque size are completely due to effects on cell-to-cell spread, since neither the UL51 Δ 167–244 alone, the gE deletion alone, nor the double deletions had any significant effect on single-step virus growth or release of virus to the medium (Fig. 6B and C).

The result in Vero cells was different in several interesting ways (Fig. 7). As previously reported, the effects of the UL51 single deletion

on plaque formation in the presence of neutralizing antibody were far more dramatic than on HaCaT cells, such that the plaques formed by UL51 deletion viruses consisted of only a few tens of cells (Fig. 7A). Over three experiments, the UL51 deletion caused a roughly 40-fold reduction, and the gE deletion caused a roughly three-fold reduction in plaque size (Fig. 7A, graph). Second, the double mutants formed significantly smaller plaques ($P < 0.01$) than either of the single mutants, and the defect (about 120-fold) was the product of multiplication of the effects of the single deletions. In order to confirm this, a multi-step growth experiment was used for comparison of the UL51 single mutant and the double deletions (Fig. 7B). As expected, the UL51 Δ 167–244 single deletion was associated with a dramatic defect in multi-step growth such that by 72 h, virus yield is diminished by three log orders of magnitude compared to the wild-type control. The two double deletions were further impaired such that by 72 h, virus yield was diminished 5-fold compared to the UL51 Δ 167–244 single deletion. The enhanced impairment in cell-to-cell spread seen in the double mutants

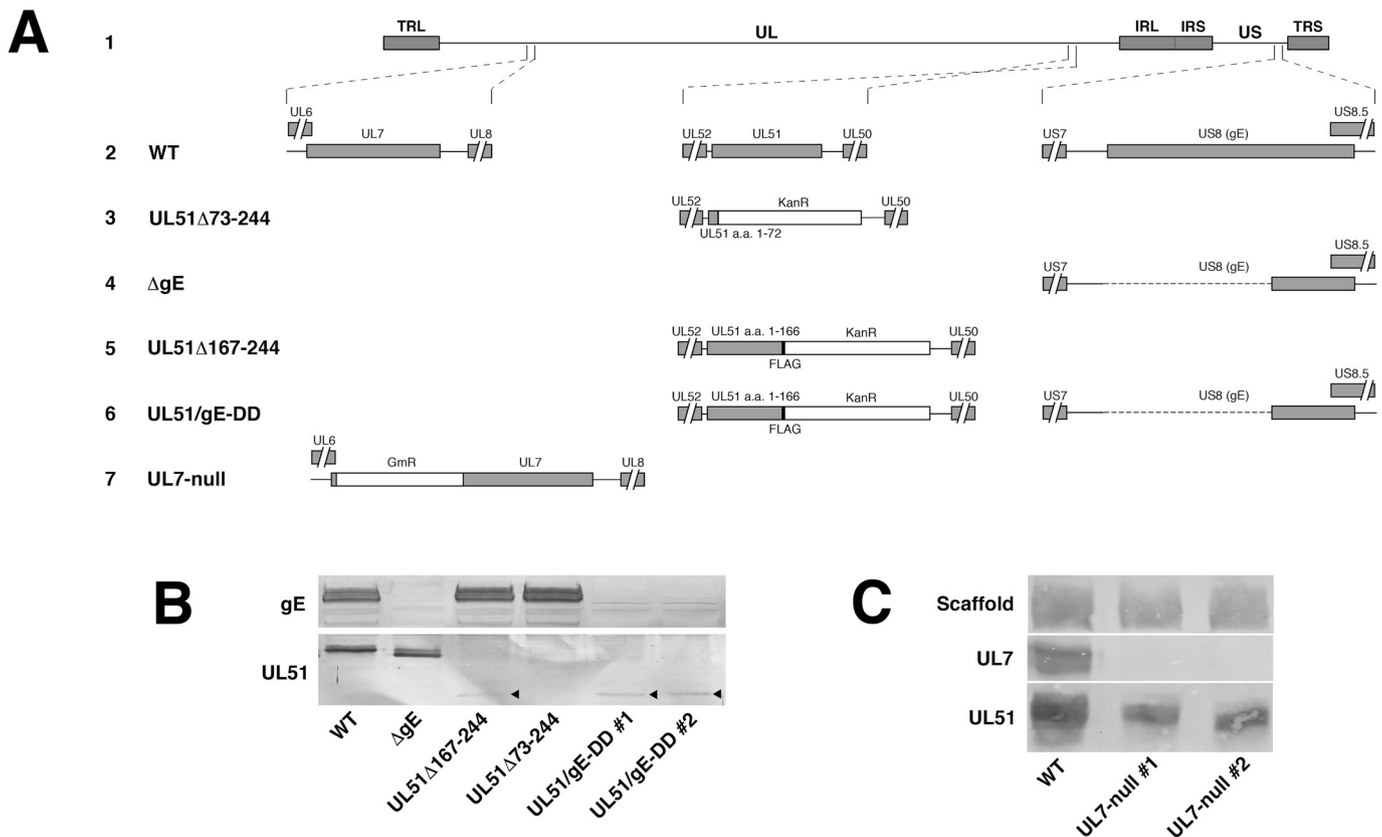


Fig. 3. Schematic diagrams of recombinant viruses used in this study and expression analysis of deletion viruses. (A) Schematic diagram of the HSV-1(F) genome (line 1) and of the recombinant viruses constructed for this study. The positions of the UL7, UL51 and US8 (gE) loci with respect to the unique long (UL) and unique short (US) sequences and the inverted repeats that flank them (TRL and IRL, and IRS and TRS, respectively), are shown in line 1. Expansions of the arrangements of the wild-type sequences in the regions of UL7, UL51 and US8 are shown in line 2. Lines 3 through 7 show altered sequences in recombinant viruses used in this study (Line 3) UL51 Δ 73-244 carries a stop codon and Kanamycin resistance cassette (KanR) in place of the sequences coding for amino acids 73–244 of pUL51 and has been previously described (Roller et al., 2014). (Line 4) The gE-null virus has a deletion of the sequence coding for amino acids 1–335 and has been previously described (Roller et al., 2014). (Line 5) In UL51 Δ 167-244-FLAG, sequences coding for amino acids 167–244 of pUL51 are replaced by an in-frame FLAG epitope tag, a stop codon and Kanamycin resistance cassette. This virus has been previously described (Roller and Fetters, 2015). (Line 6) The UL51/gE-DD virus contains the sequence alterations for both the UL51 Δ 167-244-FLAG and gE-null viruses, and was constructed by altering the UL51 locus in the previously constructed gE-null virus. (Line 7) The UL7-null virus contains a gentamycin resistance cassette (GmR) insertion immediately following the stop codon for the UL6 coding sequence. This change places a stop codon in the GmR cassette in frame with the UL7 coding sequence and is predicted to result in expression of only the first seven amino acids of pUL7. (B).

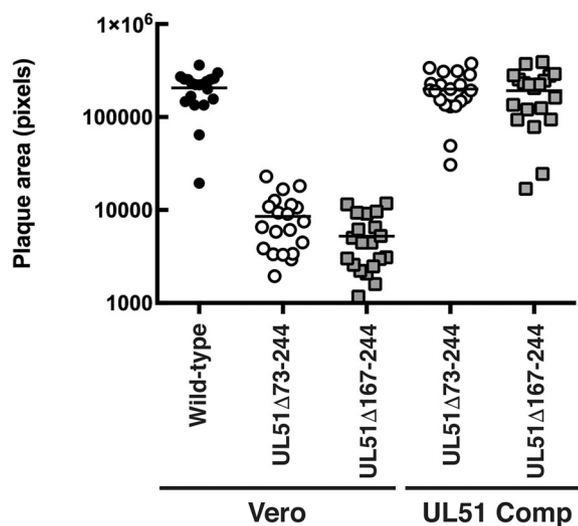


Fig. 4. UL51 deletions show similar UL51-dependent cell-to-cell spread defects. Confluent Vero or UL51 complementing cell monolayers were infected at low multiplicity with wild-type (black circles) UL51 Δ 73-244 (open circles) or UL51 Δ 167-244 (gray squares), and after two days of incubation in the presence of neutralizing antibody, plaques were immunostained, and plaque areas measured. Statistical significance of differences between samples was measured using one-way ANOVA and Fisher's least significant difference test.

was not due to an enhanced defect in virus replication. The double deletions replicated no worse in single-step growth than the UL51 Δ 167-244 single deletion (Fig. 7C).

In order to determine whether the double mutant spread defect was due to a generalized virus release defect, the titer of culture supernatant virus was also measured during the single-step growth experiment. Similar to what was previously reported for a larger UL51 deletion, the UL51 Δ 167-244 single deletion showed an impairment in virus release in Vero cells compared to the wild-type control (Roller et al., 2014). Surprisingly, the double mutants both showed enhanced efficiency of virus release to the medium compared to the UL51 single deletion, and the single gE deletion showed enhanced release efficiency compared to the wild-type control (Fig. 7D). Comparing wild-type virus to the gE deletion, the effect was most dramatic (greater than one log order of magnitude) at the earliest time points, and diminished as the efficiency of wild-type virus release rose over time. For the UL51 mutants, the double deletions showed greater than a log order of magnitude enhancement of release compared to the single UL51 mutant at all time points tested.

3. Discussion

The mechanism of herpesvirus cell-to-cell spread is still poorly understood, but several lines of evidence suggest that it reflects specific trafficking of virus particles to junctional surfaces of cells where virus may infect an adjacent cell in a space that is sterically protected from neutralizing antibody and that is confined enough that essential interactions with the host cell (e.g., envelope protein receptor interactions) may occur rapidly and efficiently. This model accounts for the necessity of the HSV entry apparatus for CCS (Cai et al., 1988; Forrester et al., 1992; Ligas and Johnson, 1988; Roop et al., 1993), and suggests that viral and cellular factors will participate in sorting of virions to junctional surfaces of cells.

Identification of viral factors with roles in epithelial CCS has been problematic because all have additional functions in virion assembly. The phenotypes of complete deletion mutations, therefore, are complex, and exploring CCS mechanisms with such mutants is difficult because an earlier step or steps in virion assembly and egress is also inhibited. Exploring the CCS function of such factors would be greatly aided by

characterization of mutants with specific defects in that function. UL51 Δ 167-244 mutant described here has no detectable defect in virus production or release to the medium in HaCaT cells, but is strongly impaired for CCS, making it a valuable tool for further studies of the CCS mechanism in these cells.

pUL51 and pUL7 have both virion assembly and cell-to-cell spread functions (Klupp et al., 2005; Nozawa et al., 2005; Roller et al., 2014; Albecka et al., 2017). The phenotype of single and double deletions of these genes suggest that they function on the same pathways for assembly and spread, and it is likely that they function as a complex (Albecka et al., 2017). We have previously shown that a truncated pUL51 that is missing amino acids 167–244 could interact with pUL7, and Albecka et al. showed that pUL51 sequences necessary for interaction with pUL7 were located from a.a. 29–170 (Albecka et al., 2017; Roller and Fetters, 2015). We show here that residues 30–90 of pUL51 are both necessary and sufficient for interaction with pUL7. We also observed that fusion of gD to UL51 truncations that contain amino acids 1–30 results in retention of gD in the ER suggesting the existence of an ER retention signal near the N-terminus of pUL51. Residues 1–30 of pUL51 do not contain a dibasic motif suggesting that ER retention either occurs indirectly, by way of interaction with some protein that does contain a retrieval motif, or by some other mechanism. Interestingly, native pUL51 is retained on cytoplasmic membranes by palmitoylation at cysteine 9 (Nozawa et al., 2003). While most palmitoyl transferase enzymes are found on Golgi membranes, some are localized to the ER (Gorleku et al., 2011), and it is possible that the gD fusions that contain amino acids 1–30 are palmitoylated, and that this may contribute to their retention in the ER.

Localization of gE to junctional surfaces has been previously shown to be necessary for gE function in epithelial CCS in HaCaT cells (Farnsworth and Johnson, 2006). gE co-localizes with β -catenin in HaCaT cells, suggesting localization to adherens junctions (Dingwell and Johnson, 1998). Here we have observed that, in Vero cells infected with wild-type virus, gE concentrates preferentially at the junctional surfaces of cells, but does not co-localize with β -catenin (Fig. 3). This may suggest either that, in Vero cells, gE targets to junctional complexes that are not adherens junctions (e.g., desmosomes), or that it recognizes some other structural feature specific to junctional surfaces.

We have previously shown that pUL51 interacts with gE in infected cells, and that expression of a pUL51-EGFP fusion causes mislocalization of gE (Roller et al., 2014). Here we show that in cells infected with UL51 partial deletion or UL7-null mutants, gE does not concentrate at junctional surfaces of cells. Our results, therefore, suggest that one function of pUL51 and pUL7 in CCS is to facilitate junctional localization or retention of the gE/gI complex. pUL51 is a cytoplasmically-oriented membrane-associated protein (Nozawa et al., 2003) and pUL7 is also located in the cytosol (Roller and Fetters, 2015; Nozawa et al., 2002). It seems plausible, therefore, that the pUL51/UL7 complex facilitates gE localization by marking gE-containing vesicles for trafficking from the Golgi apparatus or trafficking endosomes to the junctional cell surface. Alternatively, pUL51 might facilitate retention of gE at junctional surfaces by preventing its retrieval and recycling. It is tempting to speculate that this trafficking function requires the interaction between pUL51 and gE/gI.

In one intuitively attractive model, epithelial CCS results from trafficking of nascent virions for release at the basolateral surfaces of cells in a polarized epithelium. The belt of tight and adherens junctions just below the apical surface of the epithelium would then provide protection from immune effectors. This model suggests that CCS trafficking mechanisms should be most important in cells that form polarized monolayers, and less important or completely ineffective in non-polarized cells. Our results show nonetheless that the effects of UL51 mutations that affect CCS are far more potent in non-polarized Vero cell monolayers than in confluent HaCaT monolayers. This suggests the possibility that targeting of virions for CCS is more precise than simple sorting to basolateral cell surfaces. Virions may be targeted to specific

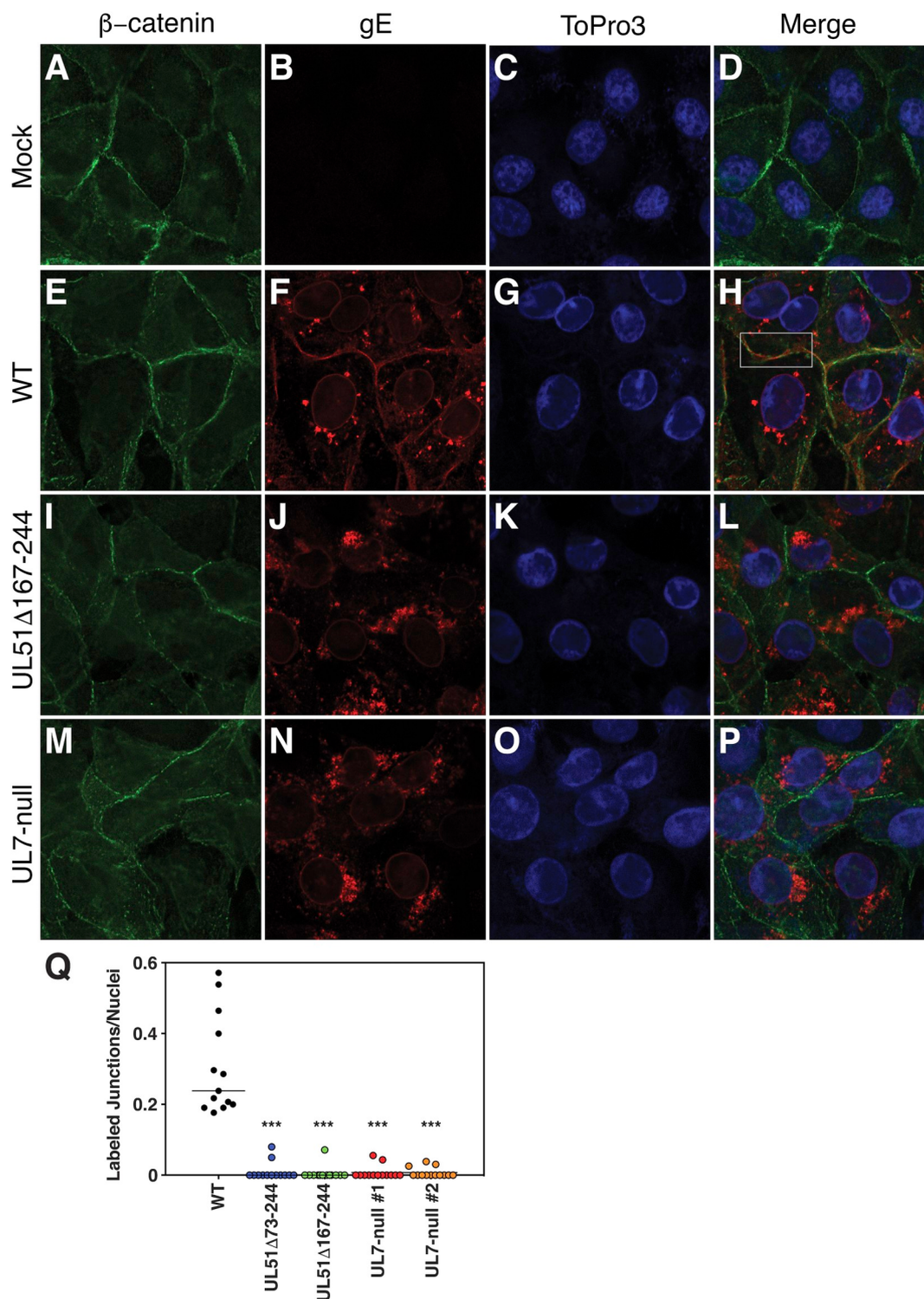


Fig. 5. gE mislocalization in UL51 and UL7 mutant-infected cells. (A–P) Vero cells that had been infected with the indicated viruses for 12 h were fixed and immunostained for β -catenin (green), gE (red) or TO-PRO-3 to stain DNA (blue). The white outlined box in panel H indicates apposition of two cells with minimal colocalization of gE and β -catenin. (Q) Quantitation of junctional gE staining. For 20 images, each containing at least 20 cells, apposing surfaces of adjacent cells that stained for gE were counted and divided by the total number of nuclei in the image. For example, in panel F, four apposing surfaces that stain for gE would be counted and divided by the seven nuclei in the image. Each point represents one such image. Statistical significance of differences between samples was measured using one-way ANOVA. (For interpretation of the references to colour in this figure legend, the reader is referred to the Web version of this article.)

junctional complexes that form regardless of whether the cell monolayer is effectively polarized. We observed that gE is not uniformly distributed at junctions between Vero cells, but rather is distributed in patches that do not correspond to concentrations of β -catenin (Fig. 5). Whether these concentrations of gE correspond to concentrations of

other host cell junctional components is not yet clear.

In addition to the viral envelope proteins that are essential for entry, epithelial CCS in HSV is affected to a greater or lesser degree by mutations in tegument proteins (pUL51, pUL7, pUL21, pUL16, pUL11, and pUL49 (VP22)) (Roller et al., 2014; Albecka et al., 2017; Han et al.,

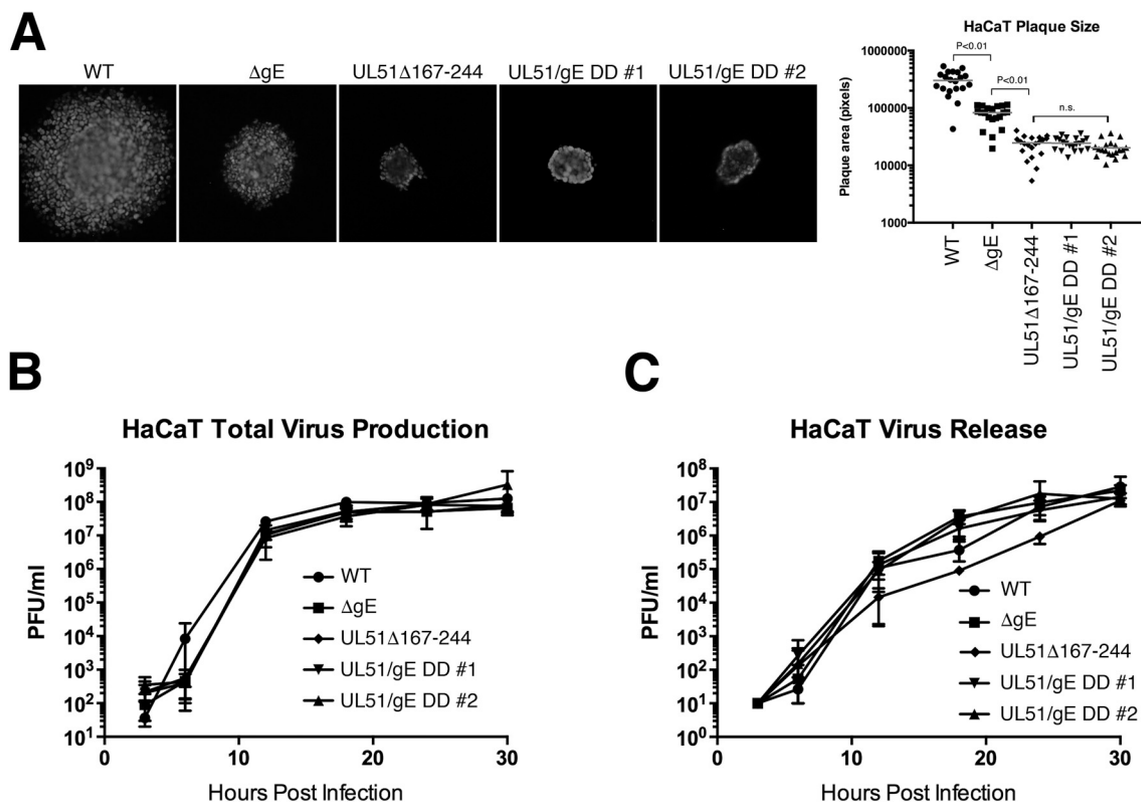


Fig. 6. Growth, spread and release of UL51 and gE mutants on HaCaT cells. (A) Double mutants show no enhanced spread defect compared to a UL51 single mutant. Digital images are shown of representative plaques formed by 48 h on HaCaT cells in the presence of neutralizing antibody. The graph at the right plots areas of 20 randomly chosen plaques for each virus. Statistical significance of differences between samples was measured using one-way ANOVA. (B) Single-step growth of single and double mutant viruses on HaCaT cells. Each point is the mean of three independent experiments, and error bars indicate the range of values. (C) Virus released to the medium in the single-step growth experiments shown in (B).

2012; Duffy et al., 2006). All of these proteins that function in CCS have been shown to interact directly or indirectly with gE (Roller et al., 2014; Han et al., 2011, 2012; Maringer et al., 2012; Stylianou et al., 2009), suggesting that the cytoplasmic domain of the gE/gI complex would be implausibly crowded if all of these interactions occurred simultaneously. Furthermore, several of these proteins (pUL51, pUL7, and the complex of pUL11, pUL16 and pUL21) have been shown to be required for proper localization of gE to junctional surfaces of cells (Roller et al., 2014; Han et al., 2011). It is possible that some of these interactions are sequential rather than simultaneous and are important for different stages of gE trafficking to and retention at junctional surfaces of cells.

The previously published evidence that pUL51 and gE interact in infected cells, and the evidence presented here that pUL51 and pUL7 expression is necessary for localization of gE to junctional surfaces of cells, suggested a simple model in which gE/gI and the pUL51/pUL7 complex function on the same CCS pathway. In this model, pUL51/pUL7 is required for delivery or retention of gE at cell junctions, and gE would be non-functional in the absence of pUL51/pUL7 due to improper localization in the cell. This model predicts that a pUL51/gE double deletion would have a CCS phenotype no more severe than the worst of the two single mutations. This prediction was borne out in HaCaT cells (Fig. 6), suggesting that pUL51 is necessary for gE/gI CCS function and that both proteins function on a single pathway for delivery of virions for CCS as depicted in Fig. 8A. Nonetheless, it appears that pUL51 must contribute more to CCS in HaCaT cells than facilitating gE function, since the UL51 mutation had a more powerful inhibitory effect on CCS than a gE deletion. The nature of this additional function is unclear, but does not reflect defects in virus replication or release to the medium.

In contrast to their behavior in HaCaT cells, it appears that gE and

pUL51 may function independently in CCS in Vero cells (Fig. 8B). The CCS phenotype of the double mutant in Vero cells was significantly more severe than the phenotype of a UL51 deletion alone, and was roughly the product of multiplication of the effects of the two single mutations. This suggests that there are genetically distinct gE- and pUL51-dependent pathways for spread in Vero cells. Although Fig. 8B depicts these as entirely separate pathways, it seems likely that they have substantial overlap in the other viral and cellular factors and cellular organelles that are involved. The independence of gE and pUL51 CCS function in Vero cells was surprising, since gE does not concentrate at junctional surfaces in Vero cells infected with the UL51 deletion mutant (Fig. 5). This suggests the possibility that the gE/gI CCS function in Vero cells is not dependent on concentration at junctional surfaces. The previously published evidence suggesting that junctional gE localization is necessary for gE function in CCS was based on correlation between junctional localization and CCS for gE cytoplasmic tail partial deletions observed in HaCaT and ARPE-19 cells (Farnsworth and Johnson, 2006). Whether the same result might be obtained in Vero cells is an interesting subject for further investigation. Further evidence that pUL51 and other tegument factors that affect CCS may not always function on the same pathway comes from the behavior of a double UL51/UL11 deletion in PRV (Klupp et al., 2005). This double deletion showed a more severe plaque formation defect than either of the single mutants despite having a single step growth defect no more severe than the UL11 deletion.

In both HaCaT and Vero cells, a UL51 mutation has a more powerful inhibitory effect on CCS than a gE deletion. This suggests that, in both cell types, pUL51 very likely has an additional, gE-independent CCS function. The nature of this function in HaCaT cells is unclear, but the observation that pUL51 inhibits both CCS and virus release to the medium in Vero cells suggests a role for the pUL51/pUL7 complex in

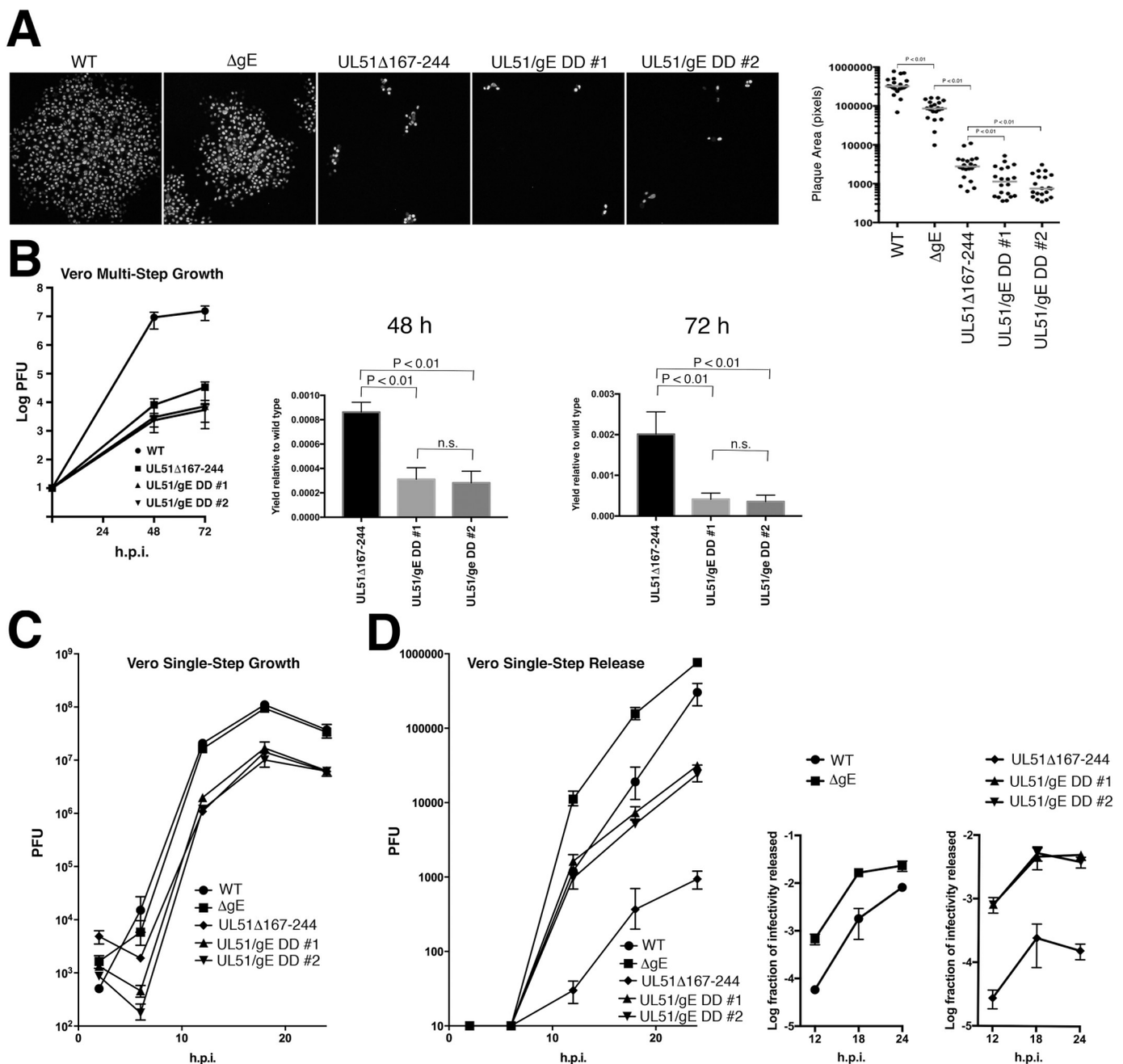


Fig. 7. Growth, spread and release of UL51 and gE mutants on Vero cells. (A) Double mutants show enhanced spread defect compared to single mutants. Digital images are shown of representative plaques formed by 48 h on Vero cells in the presence of neutralizing antibody. The graph at the right plots areas of 20 randomly chosen plaques for each virus. Statistical significance of differences between samples was measured using one-way ANOVA and Fisher's least significant difference test. There was no significant difference between the two isolates of the double deletions. (B) Double mutants show an enhanced spread defect compared to the single UL51 mutant in multi-step growth. The left-hand graph shows the multi-step growth curve. Each point is the mean of three independent experiments, and error bars indicate the range of values. The two right hand graphs compare the values obtained for the UL51Δ167-244 mutant and for the two double mutants at 48 and 72 h post infection. Error bars indicate the standard deviation. Statistical significance of differences between samples was measured using one-way ANOVA. (C) Single-step growth of single and double mutant viruses on Vero cells. Each point is the mean of three independent experiments, and error bars indicate the range of values. There were no significant differences between WT and ΔgE, and between UL51Δ167-244 and the two double mutant isolates. (D) Virus released to the medium in the single-step growth experiments shown in (C). The graph on the left shows the PFU released to the medium over time. The two smaller graphs to the right show the efficiency of release to the medium calculated as PFU in medium divided by total culture PFU for each virus at each time point. Statistical significance of differences between samples at each time point was measured using one-way ANOVA. Differences between WT and ΔgE and between UL51Δ167-244 and the double mutant viruses were significant ($P < 0.001$) at all time points. There were no significant differences between the two double mutant isolates.

virion export from the cell on either pathway. Our results are consistent with a model for virion trafficking in Vero cells in which pUL51 (and perhaps pUL7) are required for transition of virion transport intermediates onto a general pathway for export from the cell. Once on that pathway, these intermediates arrive at a sorting point for release to the

medium or transport to cell junctional surfaces for CCS. At that sorting point, the virions may take one of three pathways: (i) a gE-dependent pathway for CCS; (ii) a pUL51/pUL7-dependent pathway for CCS; or (iii) a pathway for release to the medium. Our data further suggest that the gE-dependent pathway for spread, and the pathway for release to

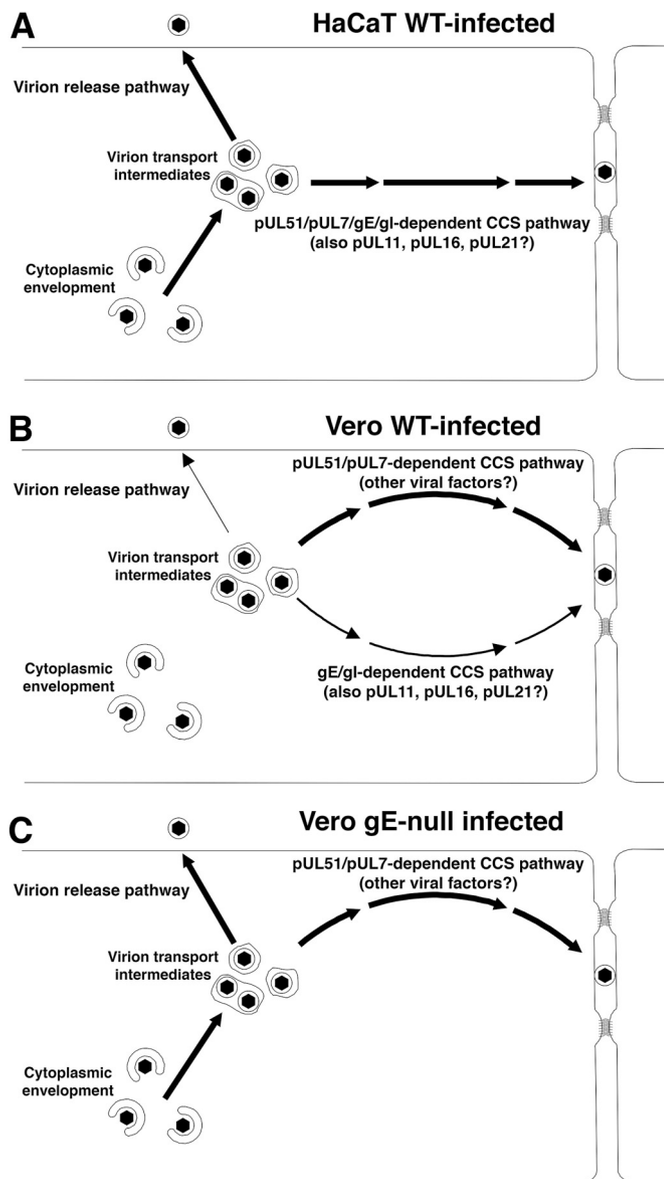


Fig. 8. Schematic diagrams of spread and release pathways in HaCaT (A) and Vero (B and C) cells infected with wild-type (A and B) or gE-null viruses (C). Thickness of arrows indicates the likely relative usage of pathways based on the effect of mutations that affect each one. In WT virus-infected HaCaT cells (A), virus is trafficked for CCS on a pathway that depends upon both gE and pUL51. Neither gE nor pUL51 participates in virus release to the medium. In WT virus-infected Vero cells (B), gE and pUL51 operate independently in CCS. The pUL51-dependent pathway is evidently more important for CCS (as indicated by the thicker pathway arrow), and pUL51 participates in the virus release pathway as well. We depict the gE and pUL51 pathways with two completely different pathway arrows, but the gE-dependent and pUL51-dependent CCS pathways may share other viral and cellular factors. In Vero cells infected with gE-null virus (C), The gE-dependent CCS pathway is abrogated, diminishing CCS, but the virion release pathway is enhanced.

the medium may be competitive, since deletion of gE and consequent elimination of the gE-dependent CCS pathway, results in more efficient release of virus to the medium (Fig. 8C).

The molecular basis for cell-specific differences in CCS is unknown, but HSV likely uses a wide range of cellular trafficking factors to move virions from the site of assembly to the plasma membrane. While many of these may be common to virion release and spread pathways, some cellular factors that have a specific function in HSV epithelial CCS have been identified. Inhibition or knock-out of protein tyrosine phosphatase

1b diminishes HSV CCS of HaCaT and mouse embryonic fibroblasts without affecting virus replication or release (Carmichael et al., 2018), suggesting that it facilitates spread. The cellular proteins extended synaptotagmins 1 and 3, in contrast, negatively regulate CCS and virus release in Vero cells (El Kasmi et al., 2018). Furthermore, pUL51 function in CCS in HaCaT and Vero cells is specifically impaired by a mutation that prevents phosphorylation at S184, suggesting that cellular kinases might also differentially regulate CCS (Kato et al., 2018). Differing levels of expression or activity of these proteins in different cell types might contribute to the differences observed here.

The existence of cell-specific pathways for CCS is not surprising, given that HSV may infect multiple cell-types, in various manifestations of disease, including epidermal keratinocytes, mucoepithelial cells, corneal epithelial cells, and neurons. HaCaT cells are considered a relevant model of skin keratinocytes, since they are derived from these cells, and retain the ability to differentiate into a stratified squamous epithelium in appropriate conditions (Boukamp et al., 1988; Schoop et al., 1999). Whether the pathways of spread observed in Vero cells are typical of any of the other cell types important for HSV pathogenesis is a subject for further exploration. The observation that multiple deletions of spread genes can produce profound, additive spread defects in some cell types suggests that such multiple deletions might form the basis for development of safe and effective anti-HSV vaccines.

4. Materials and methods

4.1. Cells and viruses

Vero, HaCaT, and UL51 complementing cells were maintained as previously described (Roller et al., 2014). All viruses were derived from the HSV-1 strain (F) BAC described in (Tanaka et al., 2003). The properties of HSV-1(F) have been previously described (Ejercito et al., 1968; Roller et al., 2000). The construction of the UL51Δ73-244, UL51Δ167-244, and gE-null viruses has been described previously (Roller et al., 2014).

4.2. Plasmid constructs

The pcDNA3 pUL51-FLAG expression construct was previously described (Roller and Fetters, 2015). Inserts for construction of plasmids that express pUL51 truncations were constructed by amplification from pcDNA3 pUL51-FLAG using the primers shown in Table 1. PCR products were digested with *EcoRI* and *XhoI* restriction enzymes and then ligated into *EcoRI*-*XhoI*-digested pcDNA3.

pRR1407, for expression of C-terminally FLAG-tagged gD was constructed by amplification of gD coding sequences from HSV-1(F) using the forward primer (5'-GATCAAGCTTGGTGCCTCCGGTATGGGGG-3') and the reverse primer (5'-CATGCTCGAGTACTTATCGTCATCGTCTTGTAGTCGGATCCCCCGGGGAATTCGTAACAAGGGCTGGTGCGAGG-3'), digestion of the resulting PCR product with *HindIII* and *XhoI*, and ligation into *HindIII*/*XhoI*-cut pcDNA3. pRR1407 was then used as the backbone for Gibson assembly cloning of all gD-UL51-FLAG fusions using the New England Biolabs Gibson Assembly Master Mix according to the manufacturer's instructions. Each plasmid was assembled from two PCR products; one comprised of vector and gD-FLAG, and the other amplified from pcDNA3 pUL51-FLAG. The sequences of primers used for these amplifications are shown in Table 1.

The plasmid that expresses EGFP-UL7 was constructed by amplification of UL7 coding sequences from HSV-1 (F) DNA using the forward primer (5'-GCTCAAGCTTCGATGGCCCGCGACG-3') and the reverse primer (5'-TCGAGAATCAACAAAAGTATAAACAGCGACGACG-3'), digestion of the resulting PCR product with *HindIII* and *EcoRI*, and ligation into *HindIII*-*EcoRI*-cut pEGFP-C1 (Clontech).

Table 1
Primer sequences for cloning of pUL51 truncations.

Insert	Forward Primer	Reverse Primer
pUL51 truncations in pcDNA3		
pUL51 (1–90)-FLAG	GTACGAATCCATGGCTTCTCTTCGCGGGCTATATG	GGCCCTCGAGTCACTTATCGTCATCGTCTTTGTAGTCGGGGTGGTGGCGCGC
pUL51 (1–124)-FLAG	GTACGAATCCATGGCTTCTCTTCGCGGGCTATATG	GGCCCTCGAGTCACTTATCGTCATCGTCTTTGTAGTCATGTACATCTGCAGGATGGTGG
pUL51 (1–166)-FLAG	GTACGAATCCATGGCTTCTCTTCGCGGGCTATATG	GGCCCTCGAGTCACTTATCGTCATCGTCTTTGTAGTCGAAGCGGAGAGGCCAAGGG
pUL51 (91–244)-FLAG	GTACGAATCCATGGGGCTCGAGGCCCGCCAC	GGCCCTCGAGTCACTTATCGTCATCGTCTTTGTAGTCTTGACCCAAAACACACGGAGCTG
pUL51 (125–244)-FLAG	GTACGAATCCATGGGGTTCGCGGGGAACC	GGCCCTCGAGTCACTTATCGTCATCGTCTTTGTAGTCTTGACCCAAAACACACGGAGCTG
pUL51 (167–244)-FLAG	GTACGAATCCATGGGGTTCGCGGGGAACC	GGCCCTCGAGTCACTTATCGTCATCGTCTTTGTAGTCTTGACCCAAAACACACGGAGCTG
pUL51 truncations fused to gD		
pUL51(1–30)-FLAG	CCAGCCCTGTTTTACATGGCTTCTCTTCGCGGGC	TCATCGTCTTTGTAGTCTCCGAGGGCGGAACGGC
pUL51(1–60)-FLAG	CCAGCCCTGTTTTACATGGCTTCTCTTCGCGGGC	TCATCGTCTTTGTAGTCTCCAGGACCCAGGGC
pUL51(1–90)-FLAG	CCAGCCCTGTTTTACATGGCTTCTCTTCGCGGGC	CCTATCGTCATCGTCTTTGTAGTCTCCGAGGGTGGTGGCGCGC
pUL51(30–90)-FLAG	CCAGCCCTGTTTTACATGGCGGAGCCGCGGCTG	CCTATCGTCATCGTCTTTGTAGTCTCCGAGGGTGGTGGCGCGC
pUL51(60–90)-FLAG	CCAGCCCTGTTTTACATGGACACCCGACGCTCGTGAAG	CCTATCGTCATCGTCTTTGTAGTCTCCGAGGGTGGTGGCGCGC

4.3. Construction of recombinant viruses

A double mutant recombinant carrying the UL51 Δ 167–244 and gE-null deletions was constructed using the gE-null BAC as the parent and modifying it to carry the UL51 partial deletion as previously described (Roller et al., 2014). Two independent isolates were constructed, and were recovered by transfection of the BAC DNAs into UL51 complementing cells. A UL7-null recombinant was constructed by insertion of a gentamicin resistance (GmR) cassette into the UL7 protein coding sequence immediately following the stop codon for the UL6 coding sequence. Insertion of the GmR cassette places a stop codon in-frame with the UL7 reading frame 10 nucleotides following the insertion junction. The primers used for insertion of the GmR cassette were 5'-GGGGGCATCGGGCACCGGGATGGCCGCCGACGCGCCGACGA TGAgctcgatcctagggataaacagg-3' and 5'-CGATGGCCTGCTTGAGGATG GTGGCGGCCGACCCctctagaggcgcggttg-3'. In each case, lowercase nucleotides correspond to sequence from the inserted GmR cassette. Virus construction was performed as described by Tischer et al., using the HSV-1 strain F BAC in bacterial strain GS1783 (kind gift of Greg Smith) (Tischer et al., 2010). The UL7-null recombinant virus was rescued and amplified on Vero cells. Proper structure of the recombinant BAC and virus at the UL7 locus were confirmed by PCR amplification and sequencing of the gene region.

4.4. Plaque size assay

Plaque size assays were performed in the presence of neutralizing antibody as previously described (Roller et al., 2014). Since plaque areas are not always normally distributed, statistical analysis was performed by one-way ANOVA using the non-parametric Fisher's least significant difference test as implemented in Prism 7 (GraphPad Software, Inc.).

4.5. Single-step growth measurement

Measurement of replication of HSV-1(F), UL51 Δ 167–244 and double mutant viruses on Vero and HaCaT cells after infection at high multiplicity (5 PFU/cell) was performed as previously described (Roller et al., 2000). Virus release efficiency was calculated as PFU in the culture medium at 24 (Vero) or 48 (HEp-2) h.p.i./PFU produced in the total culture at that time point.

4.6. Immunoblotting

Nitrocellulose sheets bearing proteins of interest were blocked in 5% non-fat milk plus 0.2% Tween 20 for at least 2 h. The membranes were probed either with a 1:1000 dilution of mouse M2 anti-FLAG

(SIGMA), a 1:5000 dilution of rabbit anti-EFP (gift of Craig Ellermeier), a 1:1000 dilution of a rabbit polyclonal antiserum raised against a UL51-GST fusion protein (Roller et al., 2014), a 1:1000 dilution of a rabbit polyclonal antiserum raised against gE (kind gift of H. Friedman), a 1:1000 dilution of rabbit polyclonal antiserum raised against pUL7 (kind gift of Yasushi Kawaguchi), or a 1:2000 dilution of mouse monoclonal antibody directed against the HSV-1 scaffold protein (AbD Serotec) followed by reaction with alkaline phosphatase-conjugated secondary antibody (Sigma).

4.7. Indirect immunofluorescence

Immunofluorescent detection of pUL51-FLAG or gD-pUL51-FLAG constructs was performed as previously described using 1:1000 mouse M2 anti-FLAG monoclonal antibody (SIGMA) (Roller and Fetters, 2015). DNA was fluorescently stained using TO-PRO-3 (Invitrogen) at 1 μ M and filamentous actin was stained using 1 μ M phalloidin-488 (Invitrogen) during the secondary antibody incubation. HSV-1 gE and cellular β -catenin were detected using 1:1000 diluted rabbit polyclonal antiserum raised against gE and 1:1000 diluted mouse monoclonal anti- β -catenin (Invitrogen), respectively (Bjerke et al., 2003; Reynolds et al., 2001).

Acknowledgments

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