Human Cytomegalovirus UL135 and UL136 Genes Are Required for Postentry Tropism in Endothelial Cells

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ABSTRACT
Endothelial cells (ECs) are a critical target of viruses, and infection of the endothelium represents a defining point in viral pathogenesis. Human cytomegalovirus (HCMV), the prototypical betaherpesvirus, encodes proteins specialized for entry into ECs and delivery of the genome to the nuclei of ECs. Virus strains competent to enter ECs replicate with differing efficiencies, suggesting that the virus encodes genes for postentry tropism in ECs. We previously reported a specific requirement for the UL133/8 locus of HCMV for replication in ECs. The UL133/8 locus harbors four genes: UL133, UL135, UL136, and UL138. In this study, we find that while UL133 and UL138 are dispensable for replication in ECs, both UL135 and UL136 are important. These genes are not required for virus entry or the expression of viral genes. The phenotypes associated with disruption of either gene reflect phenotypes observed for the UL133/8 NULL virus, which lacks the entire UL133/8 locus, but are largely distinct from one another. Viruses lacking UL135 fail to properly envelop capsids in the cytoplasm, produce fewer dense bodies (DB) than the wild-type (WT) virus, and are unable to incorporate viral products into multivesicular bodies (MVB). Viruses lacking UL136 also fail to properly envelop virions and produce larger dense bodies than the WT virus. Our results indicate roles for the UL135 and UL136 proteins in commandeering host membrane-trafficking pathways for virus maturation. UL135 and UL136 represent the first HCMV genes crucial for early- to late-stage tropism in ECs.

IMPORTANCE
Human cytomegalovirus (HCMV) persists in the majority of the world’s population. While typically asymptomatic in healthy hosts, HCMV can cause significant morbidity and mortality in immunocompromised or naïve individuals, particularly transplant patients and patients with congenital infections, respectively. Lifelong persistence of the virus may also contribute to age-related pathologies, such as vascular disease. One aspect of HCMV infection contributing to complex and varied pathogenesis is the diverse array of cell types that this virus infects in the host. The vascular endothelium is a particularly important target of infection: contributing to viral dissemination and likely leading to CMV complications following transplantation. In this work, we identify two viral gene products required for postentry tropism in endothelial cells. Identifying tropism factors required for replication in critical cell targets of infection is important for the development of strategies to restrict virus replication.

H
uman cytomegalovirus (HCMV) is a ubiquitous herpesvirus with 50 to 99% seroprevalence in the global population. Like all herpesviruses, HCMV persists for the lifetime of the host by way of latent infection (1–3). The persistence of HCMV is asymptomatic in immunocompetent individuals and is characterized by states of subclinical reactivation from latency and low-level virus shedding (1). However, in immunocompromised hosts, HCMV causes significant morbidity and mortality. Of particular concern are stem cell and solid-organ transplant patients, HIV/AIDS patients, and cancer patients undergoing chemotherapy or radiation regimens (4, 5). Additionally, HCMV is the leading cause of infectious disease-related birth defects in the United States, affecting 1 in 150 children born in the United States and most commonly resulting in mild to severe hearing loss (6). While CMV infection of seronegative women during pregnancy poses the most significant risk for severe sequelae in infants (microcephaly, cerebral palsy, and severe hearing loss or cognitive deficits), as many as 75% of congenital infections occur in infants whose mothers were seropositive at the time of conception, indicating that these infections result from reinfecation or reactivation (7). Finally, the persistence of HCMV is increasingly associated with age-related pathologies, even when overt clinical symptoms are absent. Age-related pathologies include vascular disease, immune dysfunction, and frailty (8–13). In spite of the pressing health significance of HCMV infection, the mechanisms underlying infection in multiple tissues and the persistence of the virus in the host are poorly understood.

While highly restricted in host tropism, HCMV infects a wide variety of cell types within the human host. Each cell type offers a unique context that may profoundly affect the outcome of infection: latency or replicative states of infection ranging from smoldering, chronic virus shedding to high-titer virus production (5, 14, 15). The human vascular endothelium is a key target of infection for a myriad of viruses, including HCMV, as a primary site for both hematogenous dissemination and dissemination to organ tissues (1, 16). Infected vascular endothelial cells (ECs) can detach...
and circulate to infect distal tissues (17, 18). Further, ECs are thought to factor importantly into viral persistence (19). Infection and persistence at the vascular endothelium also have the potential to impact tissue integrity. It has been shown that HCMV infection of ECs increases inflammation at the endothelium, promoting monocyte recruitment and extravasation, and increasing vascular permeability (1, 20–25). Further, proinflammatory signaling elicited by the infection of vascular ECs contributes to angiogenesis and vascular disease (8, 26, 27).

HCMV infection of the vascular endothelium and its impact on vascular biology would undoubtedly factor importantly into HCMV-related vascular pathologies. HCMV seropositivity is an important risk factor for graft and solid-organ transplant failure (28). In the case of heart transplantation, HCMV nearly doubles the 5-year rate of cardiac graft rejections as a consequence of transplant vascular arteriosclerosis (29). Additionally, HCMV persistence in healthy individuals is associated with an increased risk of age-related vascular pathologies, including atherosclerosis and restenosis, and correlates positively with greater risks for frailty and cardiac mortality (27, 30–36). In spite of the marked importance of ECs to HCMV infection, little is known about tropism barriers in these cells or the viral factors important for replication specifically in ECs.

Low-passage or clinical strains of HCMV readily infect ECs, whereas laboratory-adapted strains do not (37, 38). The restriction to replication of laboratory strains in ECs may be due to genome rearrangements that occur during passage in cell culture. For example, the ULb region of the genome is lost upon serial passage of the virus in cultured fibroblasts (39–41). The UL128 to UL131A genes are required for HCMV entry into ECs, epithelial cells, leukocytes, and dendritic cells (21, 42–48), but not fibroblasts. However, low-passage-number HCMV strains or laboratory strains repaired for their ability to enter ECs replicate with various efficiencies suggesting the existence of postentry viral factors important to replication specifically in these cell types (21, 46). Accordingly, US16 has been reported to be specifically required for the delivery of the viral genome to the nucleus in ECs and epithelial cells following viral entry (49). Viral genes required for later stages of replication specifically in ECs likely exist but have been more elusive.

We recently demonstrated a requirement for the ULb (UL133/8) polycistronic locus for replication in ECs (50, 51). The UL133/8 locus encodes four genes, UL133, UL135, UL136, and UL138. We originally demonstrated that the entire locus and the individual genes encoded play important and antagonistic roles in the establishment or maintenance of latency and in reactivation (50, 52, 53). While UL133/8 locus genes, particularly UL133, UL138, and one of the isoforms of UL136, are suppressive to viral replication in fibroblasts and CD34+ hematopoietic progenitor cells (HPCs) (50, 52–55), disruption of the UL133/8 locus resulted in a defect in virus replication in ECs, suggesting that UL133/8 genes are required for efficient replication in this cell type (51). Electron microscopic analysis of ECs infected with a virus lacking the UL133/8 locus indicated a postentry, late-stage defect in virus maturation that might be due to increased vesiculation of cytoplasmic membranes and failure to form the viral assembly compartment (VAC), a distinct restructuring of endocytic compartments into a concentric perinuclear organelle in which viral progeny is assembled during HCMV infection (51). UL133/8 mutant virus-infected ECs produced virions that lacked envelopes and had decreased numbers of dense bodies (DBs), host-derived vesicles containing viral tegument proteins. Another intriguing observation is that multivesicular bodies (MVB) induced by HCMV infection become packed with virions and DBs and that the recruitment of these viral products into MVB requires the UL133/8 locus. In contrast, fibroblasts infected with a UL133/8_NULL virus produce essentially wild-type (WT) yields of virus with no defects in virion maturation and no alteration in DBs or intracellular membranes, suggesting that these functions of the UL133/8 locus are EC specific (51). Cell type-specific requirements for the UL133/8 locus illustrate the importance of unique cellular environments to the outcome of infection.

In the present study, we have identified pUL135 and pUL136 as the critical viral gene products encoded within the UL133/8 locus for postentry viral replication in ECs. Viruses lacking UL135 or UL136 exhibited intermediate defects in viral replication relative to that of the UL133/8_NULL virus. The phenotypes associated with the loss of the UL133/8 locus in EC infection are complex and largely segregate between UL135 and UL136 mutant viruses. Both UL135 and UL136 are required for the maintenance of intracellular membrane organization and for efficient formation of the VAC, but distinct defects in virion maturation were observed with UL135 or UL136 mutant virus infection. Our findings demonstrate a cooperative requirement for UL135 and UL136 for efficient viral replication in ECs and indicate roles for pUL135 and pUL136 in commandeering membrane-trafficking pathways at early/late stages during infection in ECs. To our knowledge, these are the first HCMV proteins to be identified as required for postimmediate early (post-IE) stages of infection in ECs. These findings suggest novel mechanisms involving membrane organization or trafficking in mediating tropism specifically in ECs.

**MATERIALS AND METHODS**

**Cells.** Primary human microvascular lung endothelial cells (HMVEC) (Lonza, Walkersville, MD) were cultured using EGM-2MV BulletKit medium (microvascular endothelial cell growth medium 2; Lonza) with 5% fetal bovine serum (FBS) and 100 U/ml penicillin. HMVEC were cultured at 37°C under 5% CO2 and were passaged as described previously (51). **Viruses.** Bacterial artificial chromosome (BAC) clones of the HCMV TB40/E wild-type (WT) and all variant strains express green fluorescent protein (GFP) as a marker of infection. The construction of the UL135STOP variant has been described previously (53). The TB40/E-UL136_GalK construct was generated by replacing the entire UL136 open reading frame (ORF) with a GalK cassette. Briefly, the GalK cassette was amplified by PCR using primers flanked by homologous viral sequences (UL136 GalK Fwd [5’ gcagccacgcctcctcttccaatttctaccgcactagagaaagagcGCTGTTGACAAATTTACGCGCA 3’] and UL136 GalK Rev [5’ gtctcctcttcttaacgcaacttcaccgccgcgagccccttcaccaccGTCAACGTGTGCTT GCCCTT 3’]) (lowercase letters stand for HCMV homology arms) and was recombined into the WT BAC by homologous recombination. GalK-positive recombinants were selected on M63 minimal medium plates followed by MacConkey medium plates containing galactose as the sole carbon source. Multiple independent recombinant viruses were screened by BAC digestion and PCR to confirm the replacement of UL136 with the GalK cassette. Construction of the UL135STOP variant and the UL136STOP (54) and UL135STOP/UL136STOP (53) viruses have been described previously.

**Materials.** Viruses were propagated by transfection of 25 μg of BAC DNA plus 3 μg of a plasmid expressing HCMV UL82 (encoding the pp71 protein) into 5 × 10⁵ MRC-5 fibroblasts by electroporation to obtain primary (P0) virus stocks. The secondary virus stocks (P1) were generated by infecting MRC-5 fibroblasts with P0 stocks. The P1 stocks were purified by
Table 1: Real-time PCR primers

<table>
<thead>
<tr>
<th>Gene</th>
<th>Direction</th>
<th>Sequence</th>
<th>UPL probe no.</th>
</tr>
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<tbody>
<tr>
<td>UL123</td>
<td>Fwd</td>
<td>CAGTCGCACCCCTAACTTGT</td>
<td>73</td>
</tr>
<tr>
<td>Rev</td>
<td>ATAAACGGGAGATGTTGATG</td>
<td></td>
<td></td>
</tr>
<tr>
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<td>Fwd</td>
<td>TTGTCCTCGGAAAAGTC</td>
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<tr>
<td>Rev</td>
<td>GGGGGGACCAACAGTAGTACG</td>
<td></td>
<td></td>
</tr>
<tr>
<td>UL69</td>
<td>Fwd</td>
<td>CCTACGACTTCTTGCTTCTC</td>
<td>30</td>
</tr>
<tr>
<td>Rev</td>
<td>CGTCCAGTTGCTGTTCAAA</td>
<td></td>
<td></td>
</tr>
<tr>
<td>UL32</td>
<td>Fwd</td>
<td>TCTTGAGGGGGGAAACCTAC</td>
<td>58</td>
</tr>
<tr>
<td>Rev</td>
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<td></td>
</tr>
<tr>
<td>RNaseP</td>
<td>Fwd</td>
<td>GACGGACTGCGCGAGGTTA</td>
<td>24</td>
</tr>
<tr>
<td>Rev</td>
<td>CCATGCTGAAGTGTCCCA</td>
<td></td>
<td></td>
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*Fwd, forward; Rev, reverse.*

Density gradient centrifugation in a 20% d-sorbitol cushion through a 20,000 rpm in an SW28 rotor (Beckman Coulter, CA) for 80 min at 22°C. Virions were resuspended in IMDM (Isco’s modified MEM [Dulbecco’s modified Eagle medium]) containing 10% BIT (bovine serum albumin, insulin, and transferrin; Stem Cell Technologies) and were stored at −80°C. Virus titers were determined by the 50% tissue culture infective dose (TCID50) method on MRC-5 fibroblasts. Multistep virus growth curves were determined by TCID50 analysis of cell lysates harvested over a time course.

Quantitative PCR. Total viral DNA and viral RNA were isolated from infected cells by using the Zymo Duet kit with on-column DNase digestion for RNA samples according to the manufacturers’ instructions (Zymo Research, Irvine, CA). cDNA was synthesized using a Transcript First Strand cDNA synthesis kit (Roche, Indianapolis, IN). Both viral DNA and viral cDNA were analyzed using advanced relative quantification on a Roche LightCycler 480 instrument according to the manufacturer’s instructions (Roche, Indianapolis, IN). Viral DNA and cDNA levels were normalized to those of the cellular housekeeping genes RNaseP and β-actin, respectively. The primers and corresponding Universal ProbeLibrary probes are listed in Table 1.

Immunoblotting. Immunoblotting was performed as described previously (52). Briefly, protein lysates (15 to 20 μg) were separated on 12% Bis-Tris gels through electrophoresis. Proteins were transferred to 0.45-mesh polyvinylidene difluoride membranes (Immobilon-FL; Millipore, MA). The membranes were then immunoblotted using mouse monoclonal or rabbit polyclonal antibodies directed against each protein. Monoclonal and polyclonal antibodies were detected using fluorescently conjugated secondary antibodies and the Odyssey infrared imaging system (LI-COR, NE). The mouse monoclonal antibodies used in this study include antibodies against IE1/IE2 (clone 3H4; dilution, 1:100; a generous gift from Tom Shenk), UL44 (clone 10D8; dilution, 1:2,500; Virusys), pp28 was used at a 1:20 dilution. The Golgi apparatus was stained with a rabbit monoclonal antibody specific to GM130 (clone EP892Y; Abcam) at a 1:250 dilution. Cells from each infection condition were fixed in 2% formaldehyde in phosphate-buffered saline (PBS) and were then permeabilized. Primary and secondary antibodies were incubated in PBS-Tween 20 (PBS-T) for 1 h. DNA from infected cells was stained with 4′,6-diamidino-2-phenylindole (DAPI) (Molecular Probes) after primary staining. Coverslips were mounted onto slides using Aqua Poly/Mount medium (Polysciences, Inc.). Cells were imaged as lambda stacks by using a Zeiss 510 Meta confocal microscope. Images were unmixed using the Zeiss 510 Meta software, version 4.2, and were processed using Zeiss software.

Results

UL135 and UL136 are required for efficient viral replication in endothelial cells. We demonstrated previously that the UL133/8 locus was required for postentry viral tropism across various micro- and macrovascular EC types (50, 51). Infection with a recombinant virus lacking the UL133/8 locus resulted in as much as 1,000-fold-reduced yields of progeny virus compared to that for WT infection. Moreover, the reduction in viral titers was not due to defects in viral entry or in immediate early (IE), early, or late gene expression. Rather, we observed disorganization of intracellular membranes, failure to form a VAC, and defects in the maturation of progeny virus. The requirement for this locus for virus replication was specific to endothelial cells, since the UL133/8 locus was dispensable for replication in primary fibroblasts and primary epithelial cells. The phenotypes associated with UL133/8null infection in ECs are summarized in Table 2. These findings suggested that genes within the UL133/8 locus are required for late-stage tropism in ECs (51).

To define the UL133/8 genes important for replication in ECs, we examined the replication of a series of mutant viruses, each containing a disruption in one of the four viral genes encoded within the locus. UL133, UL135, and UL138 were disrupted by the substitution of stop codons for translation initiation codons, resulting in the UL133STOP, UL135STOP, and UL138STOP recombinant viruses, which fail to express the respective proteins (50, 52, 53). Because UL136 encodes a number of protein isoforms (54), the entire UL136 gene was disrupted by the insertion of a cassette encoding galactosidase K (UL136gal). We examined the replication of each of these viruses in primary human microvascular lung ECs (HMVEC) infected at a low MOI (0.05). Virus yield was measured over a time course (Fig. 1). While viruses containing disruptions in UL133 or UL138 replicated with kinetics and yields similar to those of the WT virus, the UL135STOP and UL136gal Viruses exhibited 100- and 50-fold reductions in progeny virus production. The defect associated with replication in the absence of UL135 or UL136 has been observed with multiple independent clones of each virus. Based on these results, we conclude that UL135 and UL136 are important for efficient viral replication in ECs.
To confirm that the defects in replication are specific to the disruption of UL135 and UL136, we examined the replication of independent mutant viruses containing disruptions in these genes. UL136 is expressed as 5 protein isoforms. We recently mapped the origins of these isoforms, and we have disrupted the expression of all protein isoforms by converting each ATG in the UL136 coding sequence to a stop or alanine codon (54). The resulting UL136\textsubscript{NULL} virus replicates with a defect equal to or greater than that observed for the UL136\textsubscript{GALK} virus (Fig. 1B). We previously demonstrated a requirement for UL135 for replication following transfection of infectious genomic clones in fibroblasts or for reactivation in CD34\textsuperscript{+} HPCs (53). The requirement for UL135 for replication from BAC transfection in fibroblasts could be overcome by subsequent disruption of UL138. Therefore, we analyzed the replication of a virus containing disruptions in both UL135 and UL138 (UL135\textsubscript{STOP}/UL138\textsubscript{STOP}) in order to confirm the importance of UL135 to replication in ECs and to determine if disruption of UL138 could compensate for the loss of UL135 in ECs. The UL135\textsubscript{STOP}/UL138\textsubscript{STOP} virus replicated with a defect equal to or greater than that of the UL135\textsubscript{STOP} virus (Fig. 1B). Further, disruption of UL138 did not compensate for the defect in replication associated with the loss of UL135. Taken together, these results confirm the requirement for UL135 and UL136 for replication in ECs and indicate that the requirement for UL135 in ECs is independent of any effects of UL138.

**Distribution of pUL135 and pUL136 during infection in ECs.** We have previously analyzed the localization of pUL135 and pUL136 in fibroblasts (50, 54). While all proteins encoded in the UL133/8 locus are associated with Golgi membranes and localize to the VAC at late times in infection, the protein encoded by UL135 exhibits cell surface localization early in infection (50) and associates with the cytoskeleton (56). UL136 is expressed as multiple protein isoforms, some of which localize with the Golgi apparatus while others are more diffusely distributed throughout the cytoplasm (54). To determine if the localization of the UL135 or UL136 protein is distinct in ECs, we examined the subcellular distribution of these proteins in infected cells with recombinant viruses expressing a variant of UL135 or UL136 fused with a C-terminal Myc epitope tag (UL135\textsubscript{Myc} or UL136\textsubscript{Myc}), pUL135 localized throughout the cytoplasm and at the cell surface (Fig. 2). Interestingly, we did not observe pUL135 colocalization with Golgi membranes or the VAC, which is the predominant localization in fibroblasts. The distinct localization of pUL135 in ECs may reflect cell type-specific protein functions or differential trafficking of the protein. pUL136 colocalized with the Golgi marker GM130, with some punctate staining throughout the cytoplasm (Fig. 2). The distribution of pUL136 in infected ECs is similar to that during infection in fibroblasts. Further studies are required to define the subcellular distribution of individual pUL136 isoforms during infection in ECs.

**UL135 and UL136 are not required for viral gene expression.** ECs infected with the UL133/8\textsubscript{NULL} virus exhibited no statistically significant deviation for each time point is indicated by error bars.

### FIG 1

UL135 and UL136 are required for replication in endothelial cells. (A) HMVEC were infected with the TB40/E-WT, -UL133/8\textsubscript{NULL}, -UL133\textsubscript{STOP}, -UL135\textsubscript{STOP}, -UL135\textsubscript{STOP}/UL136\textsubscript{GALK} or -UL135\textsubscript{STOP}/UL138\textsubscript{STOP} virus at an MOI of 0.05. Virus yields were measured in lysates harvested over a time course (measured in days postinfection) by TCID\textsubscript{50} on fibroblasts. The values plotted are averages for 3 to 4 independent experiments. The standard deviation for each time point is indicated by error bars.
significant difference in gene expression at the immediate early, early, or late stage of viral infection from ECs infected with the WT virus (51). However, because we have observed requirements for individual UL133/8 locus genes that are not realized in the context of deletion of the entire UL133/8 locus (53), we analyzed the accumulation of viral genomes, transcripts, and proteins in HMVEC infected at an MOI of 0.1 with the WT, UL135STOP, or UL136GalK virus. The accumulation of viral genomes and transcripts was measured at 1, 6, and 12 dpi by quantitative PCR (Fig. 3). The 1-dpi genome data are graphed alone in the Fig. 3A inset in order to better represent input genomes. HCMV genomes, measured using a primer pair specific to the UL99 gene, accumulated similarly over time in cells infected with the WT or UL135STOP virus, while cells infected with the UL136GalK virus accumulated slightly higher levels of genomes by 12 dpi. In agreement with the analysis of viral genomes, UL136GalK virus infection also resulted in greater accumulation of UL123/IE1 (Fig. 3B), UL69 (Fig. 3C), and UL32/ pp150 (Fig. 3D) transcripts than infection with the WT or UL135STOP virus during the 12-day time course. These results suggest that the defect in replication observed in ECs infected with UL135 or UL136 mutant viruses is not due to a failure to replicate viral genomes or to express viral genes. Indeed, viral gene expression was enhanced during infection with the UL136 mutant virus.

We then examined representative IE (IE1 [72 kDa] and IE2 [86 kDa]), early (UL44), and late (pp28) viral proteins over a time course by immunoblotting (Fig. 4A). In each infection, proteins at each stage of infection accumulated with similar kinetics and to levels similar to those of WT virus infection. Quantification of IE1 and pp28 bands over 3 independent experiments indicated that there was no statistically significant difference in the accumulation of IE or late proteins between WT virus infection and infection with either mutant virus (Fig. 4B and C). These data are consistent with the defect associated with UL133/8NULL virus infection in ECs. The absence of a substantial defect in viral gene expression in HMVEC infected with these mutant viruses indicates a requirement for UL135 and UL136 in the maturation of progeny virus, as was observed with the UL133/8NULL virus.

UL135 and UL136 are important for efficient formation of the viral assembly compartment. The VAC is a virus-induced reorganization of intercellular cytoplasmic membranes into a concentric perinuclear compartment important for virus maturation (57–62). Our previous work demonstrated a requirement of the UL133/8 locus for maintaining intracellular membrane structures and for the formation of the VAC (51). In the absence of the UL133/8 locus, a number of cellular trafficking markers associated with the VAC are dispersed throughout the cell (51, 58, 59). In the present study, we examined the formation of the VAC in HMVEC infected with a WT, UL135 mutant, or UL136 mutant virus. Infected cells were labeled with antibodies specific to the Golgi marker GM130 or the late protein pp28, both known to mark the VAC (59). Both pUL135 and pUL136 were required for efficient VAC formation (Fig. 5). Mutant viruses lacking either UL135 or UL136 resulted in the dispersal of Golgi membranes and the VAC in approximately 50% of infected cells. We also analyzed VAC formation in cells infected with UL133 or UL138 mutant viruses to ensure that the requirement for VAC formation was limited to UL135 and UL136. UL133 and UL138 mutant viruses formed the VAC similarly to the WT virus. These results indicate that UL135 and UL136, but not UL133 and UL138, are important for membrane organization during HCMV infection of ECs.

UL135 and UL136 are required for the maturation of progeny virions. We previously characterized late-stage defects in virion maturation and egress in ECs infected with the UL133/8NULL virus by using transmission electron microscopy (TEM) (51). To explore the roles of UL135 and UL136 in the maturation of virions, we infected HMVEC with TB40/E-WT, -UL135STOP, or UL136GalK at an MOI of 4 and processed the cells for TEM at 5 dpi,
a time point at which the level of infectious virus production is high.

We observed striking defects in the maturation of virus particles in ECs infected with the \textit{UL135} \textit{STOP} or \textit{UL136} \textit{GalK} virus relative to that with the WT (Fig. 6). Interestingly, the phenotypes associated with \textit{UL133/8} \textit{NULL} infection segregated between those for \textit{UL135} \textit{STOP} and \textit{UL136} \textit{GalK} infections (Table 2). WT infection results in the production of virus particles containing an encapsidated viral genome, tegument, and envelope (Fig. 6A). HMVEC infected with the \textit{UL135} \textit{STOP} virus accumulated two types of cytoplasmic virus particles in addition to a minority of virions that appeared morphologically normal: nonenveloped capsids and particles with abnormal envelopes that appeared to wrap the capsid loosely (Fig. 6B). Only 27% of the virus particles in cells infected with the \textit{UL135} \textit{STOP} virion morphologically resembled WT virions. We previously reported a similar envelopment defect for \textit{UL135} \textit{STOP} infection in fibroblasts and a corresponding increase in the number of genome-containing particles required to form an infectious unit (53). Figure 7B shows representative \textit{UL135} \textit{STOP} virus particles in infected fibroblasts, where the envelopment defect is marked. Further, in fibroblasts infected with the \textit{UL135} \textit{STOP} virus, we had noted an accumulation of noninfectious enveloped particles (NIEPs); however, NIEPs were far less pronounced in ECs infected with the \textit{UL135} \textit{STOP} virus (data not shown). Sixty-five percent of the capsids that accumulated in the cytoplasm of ECs infected with the \textit{UL136} \textit{GalK} virus lacked envelopes entirely, a phenotype similar to the \textit{UL133/8} \textit{NULL} infection phenotype in ECs (Fig. 6C). Nonenveloped capsids were not observed in fibroblasts infected with the \textit{UL136} \textit{GalK} virus (Fig. 7C). In ECs, nonenveloped or aberrantly enveloped virus particles in \textit{UL136} \textit{GalK} and \textit{UL135} \textit{STOP} infection, respectively, did not appear to be fully tegumented (Fig. 6).

We observed striking differences between DBs accumulating in cells infected with the \textit{UL135} or \textit{UL136} mutant virus and those in WT-infected cells (Fig. 8A to C). DBs are vesicles containing viral tegument proteins. Infection of HMVEC with the \textit{UL135} \textit{STOP} virus resulted in an average of 2-fold fewer DBs than WT infection (Fig. 8B), a finding similar to the reduced presence of DBs in \textit{UL133/8} \textit{NULL} infection. In \textit{UL136} \textit{GalK} virus-infected ECs, DBs accumulated to the same numbers as in WT infection but were strikingly larger (Fig. 6C and 8C). Measurements of CT indicated that DBs in \textit{UL136} \textit{GalK} infection were on average 2.5 times larger than DBs in WT infection. We have not observed any alterations in DB formation in fibroblasts infected with the \textit{UL135} \textit{STOP} or \textit{UL136} \textit{GalK} virus (Fig. 7) (53). To determine if the accumulation of dense bodies is the result of altered accumulation of proteins that form dense bodies, we analyzed the levels of two DB/tegument proteins, pp65 and pp150, in the context of infection (Fig. 9A). The levels of pp65 and pp150 in multiple experiments were quantified and are shown in Fig. 9B and C, respectively. Infection with the \textit{UL135} \textit{STOP} virus resulted in pp65 and pp150 levels 2- to 3-fold lower than those in WT infection. In contrast, \textit{UL136} \textit{GalK}
Mock signaling (63–67). MVB are induced during HCMV infection of other secretory vesicles. MVB have important functions in cells: the fusion of MVB with lysosomes results in the degradation of virus-infected cells accumulated viral proteins to levels similar to those for WT infection despite the higher level of accumulation of viral transcripts in UL136GalK infection (Fig. 3B to D). Therefore, while reductions in pp65 and pp150 levels may contribute to the reduction in the number of DBs in UL135STOP infection, the enlarged DBs in UL136GalK infection do not result simply from increased protein levels. These results suggest that there is a defect in the expression of at least some late genes during infection with UL135STOP virus that become apparent after 96 hpi (comparing time courses in Fig. 4 and 9). The late stage defects associated with UL135STOP infection may be related to a defect in IE2 expression associated with UL135STOP infection in fibroblasts (53).

MVB are large vesicles formed by the fusion of endosomes and other secretory vesicles. MVB have important functions in cells: (i) the fusion of MVB with lysosomes results in the degradation of MVB contents; (ii) MVB serve as a platform for intracellular signaling; and (iii) MVB trafficking to and fusion at the cell surface results in the secretion of exosomes that may affect intracellular signaling (63–67). MVB are induced during HCMV infection of ECs. The MVB in infected ECs typically contain virus progeny and DBs in addition to intraluminal vesicles (51). The UL135STOP virus matures at late stages of infection in ECs.

**DISCUSSION**

The vascular endothelium represents a key target for myriad viruses as a gateway for viral dissemination throughout the human host (68). HCMV is no exception; vascular ECs represent a site of infection critical to viral dissemination, persistence, and pathogenesis (1, 18, 20, 21, 24, 25, 69, 70). Understanding the mechanisms underlying HCMV replication in key targets of infection such as ECs is crucial for the development of new therapeutics to control HCMV spread and the associated disease in the host. Given the importance of ECs as a target of infection, cytomegaloviruses encode a number of genes to ensure access to and tropism for these cells (42, 44, 47, 71, 72). In addition to the genes involved in entry, two genes that function in postentry steps in the replication of cytomegaloviruses in ECs have been identified previously. The HCMV US16 gene is specifically required for the delivery of viral genomes and tegument proteins to the nucleus following entry into ECs (49), and the murine CMV M45 gene contributes to viral replication specifically in ECs and macrophages by inhibiting apoptosis (73). These studies suggest that HCMV has evolved functions to overcome specific tropism barri-

**FIG 4** Accumulation of IE, early, and late viral proteins in infected HMVEC. (A) HMVEC were infected at an MOI of 2 with TB40E-WT, -UL135STOP, or -UL136GalK. Protein lysates were harvested over the time course indicated (in hours postinfection [hpi]) and were analyzed by immunoblotting using antibodies specific to the 72-kDa IE1 and 86-kDa IE2 proteins, the early protein UL44, the late protein pp28, and β-actin. β-Actin served as a loading control. (B and C) Quantification of the average IE1 (B) and pp28 (C) protein levels (normalized to those for actin) from three independent experiments. The standard deviations are indicated. Student's t test indicates that there is no significant difference between WT and mutant viruses at each time point.
UL135 and UL136 are required for efficient formation of the viral assembly compartment. HMVEC were infected at an MOI of 2 with TB40/E-WT, -UL133STOP, -UL135STOP, UL136GalK, or UL138STOP. At 144 h postinfection, cells were processed for indirect immunofluorescence microscopy using monoclonal antibodies specific to the Golgi marker GM130 and the late viral protein pp28. Nuclei were stained by DAPI. Both the UL135STOP and UL136GalK viruses failed to reorganize the intracellular membranes into the VAC. A merge of all three images is shown at the right. Bars, 10 μM.
ers and to navigate unique cell type environments. Our present study identifies two additional gene products, UL135 and UL136, and their roles in mediating intracellular membrane organization as important for the later stages of replication in ECs. To our knowledge, these are the first post-immediate early stage viral determinants of tropism for ECs to be identified.

We reported previously that EC infection with a virus lacking the entire UL133/8 locus, the UL133/8NULL virus, resulted in dramatic disruption of intracellular membranes and abnormal virion morphology (51). The defect in virion morphology might be secondary to the disruption of intracellular membrane organization. The phenotypes observed during infection with the UL133/8NULL virus are consistent with these hypotheses.
virus in ECs segregated between the phenotypes for infection with UL135 or UL136 mutant viruses (Table 2)—implying that the functions of UL135 and UL136 are distinct, albeit cooperative. Both UL135 and UL136 were required for efficient formation of the VAC during infection in ECs (Fig. 5). Dispersal of Golgi membranes and the VAC was observed in ~50% of ECs infected with either the UL135STOP or the UL136GalK virus. The dispersal of the VAC is more pronounced in ECs infected with the UL133/8NULL.

FIG 8 Dense body formation impacted by the disruption of UL135 or UL136. HMVEC infected with the TB40/E-WT (A), -UL135STOP (B), or -UL136GalK (C) virus were fixed, embedded, and sectioned for TEM at 5 dpi. Representative micrographs are shown to illustrate the accumulation of DBs (arrows). Bars, 2 μm.

FIG 9 Enlarged DBs are not due to increased accumulation of DB proteins. (A) HMVEC were infected at an MOI of 2 with the TB40/E-WT, -UL135STOP, or -UL136GalK virus. Protein lysates were harvested over the time course indicated (in hours postinfection [hpi]) and were analyzed by immunoblotting using antibodies specific to the pp150 and pp65 proteins. β-Actin served as a loading control. (B and C) Quantification of the average pp65 (B) and pp150 (C) protein levels (normalized to actin) in three independent experiments. The standard deviations are indicated. Student’s t test indicated that there was no significant difference between WT and mutant viruses at each time point.
FIG 10  pUL135 is required for the incorporation of viral products into MVB. HMVEC were infected at an MOI of 4 with the TB40/E-WT (A), -UL135STOP (B), or UL136GαK (C) virus. At 5 days postinfection, cells were fixed, embedded, and sectioned for TEM. Representative micrographs are shown to illustrate the accumulation of MVB (arrows) and the incorporation of virions (arrowheads) and DBs (open arrowheads) into MVB. Bars, 500 nm.

virus, lacking both UL135 and UL136, and further work is required to understand how these proteins function cooperatively to maintain membrane organization for VAC formation. It is possible that the dispersal of intracellular membranes observed during UL133/8STOP, UL135STOP, or UL136GαK infection in ECs is an intermediate step in normal VAC formation and that UL135 and UL136 are required for the reorganization of membranes for VAC formation. Studies at higher resolution will be required to determine if the VACs formed in a portion of the mutant virus-infected ECs are truly equivalent to those formed in WT infection. This requirement for pUL135 and pUL136 in restructuring host membranes to form the VAC is consistent with a role for these proteins in membrane trafficking and reorganization. Neither the UL133/8 locus nor UL135 and UL136 are required for VAC formation in fibroblasts, indicating unique membrane-trafficking requirements in ECs. The HCMV tegument proteins pUL48, pUL94, and pUL103 have been shown to be required for VAC formation in fibroblasts (74); however, it is not known if these proteins are required in ECs. Defining the mechanisms by which viral proteins function to mediate VAC formation will contribute to our understanding of cellular trafficking pathways in ECs.

Distinct defects in virion maturation, possibly secondary to the defects in membrane organization, were observed in UL135 and UL136 mutant virus-infected cells. For both mutant infections, as many as 75% of the virions in infected ECs deviated from the WT in their morphology. The majority of virions lack a secondary envelope altogether in both UL135STOP virus- and UL136GαK virus-infected ECs (Fig. 6), a phenotype commonly observed in UL133/8NULL infection of ECs. Nearly half of the virions that acquired an envelope in UL135STOP Virus-infected ECs were abnormal; the envelope was loosely wrapped, and the virions appeared to lack tegments (Fig. 6B). Not surprisingly, UL135STOP virions could not be banded on glycerol tartrate gradients (data not shown). One intriguing possibility is that these mutant viruses have allowed us to capture an intermediate in virion maturation due to the stalling of the process at a point following the egress of the nucleocapsid into the cytoplasm and prior to full tegumentation and envelopment of the capsid. The formation of DBs was also altered in cells infected with UL135 or UL136 mutant viruses: UL135STOP infection produced fewer DBs, and UL136GαK produced enlarged DBs (Fig. 8). While little is known about the formation of DBs and their contribution to infection, they are thought to be an intermediate in the maturation process and may contribute to secondary rounds of infection by delivering a payload of tegument proteins (75). No alteration in DB formation was seen for either UL135 or UL136 mutant viruses in fibroblasts (Fig. 7). A similar phenotype of enlarged DBs was observed in fibroblasts infected with a mutant virus containing a disruption in UL97, a viral kinase important for early and late stages of infection (76), but it is not known if that phenotype exists in EC infection.

While there are important distinctions, some of the defects associated with the disruption of UL135 in ECs are reminiscent of phenotypes we have observed during infection of fibroblasts with the UL135STOP virus (53). The UL135STOP bacterial artificial chromosome clone is defective in initiating infection following its electroporation into fibroblasts (53). Once produced, virus stocks have an increased particle-to-PFU ratio (a greater number of genome-containing particles is required to form an infectious unit) but replicate with little defect relative to the WT virus when fibroblasts are infected with equivalent multiplicities. In contrast, infection with the UL135STOP virus stock results in a 100-fold defect in replication in ECs relative to WT infection (Fig. 1A), indicating a greater requirement for UL135 for replication in ECs than in fibroblasts. In agreement with the defect in ECs, the UL135STOP virus is defective for reactivation from latency in CD34+ HPCs even at MOIs that result in WT virus yields in fibroblasts (53). The UL135STOP defect in reconstituting viral replication from BAC transfection in fibroblasts and much of the defect in virion maturation can be overcome by the additional disruption of UL138 (53). However, the defect in UL135STOP replication in ECs was not reduced by the additional disruption of UL138 (Fig. 1B). Aberrant envelopment of UL135STOP virions is observed in both fibroblasts and ECs. Taken together, these findings suggest that while UL135 is important for viral replication across different cell types, unique, EC-specific functions exist that are independent of the antagonistic relationship between UL135 and UL138.

The fate of virus particles found within MVB is not yet entirely clear. While MVB could represent a dead-end pathway for the virus if they were to fuse with the lysosome, MVB may be a vehicle for transport out of the cell by exocytosis (77). Viral hijacking of MVB is an important route of entry and egress for enveloped viruses and has recently been shown to be a means by which viruses may alter intercellular communication and inflammation by redesigning exosomes (65, 77–87). HIV proteins interact with a
number of components of ESCRT-I and ESCRT-III complexes to facilitate budding at the plasma membrane (88), yet MVB appear to be the site of assembly for HIV in monocytoid-derived macrophages (89, 90). Within the herpesvirus family, ESCRT components and MVB are a means of maturation and egress for herpes simplex virus 1 (HSV-1) (64, 91, 92). Specifically, secondary envelopment of HSV-1 requires VPS-4, a component of the ESCRT-III machinery required for the generation of intraluminal vesicles in MVB (92). Similarly, HCMV and human herpesvirus 6 (HHV-6) virions have been observed within MVB (51, 93, 94), and HCMV replication depends on the ESCRT proteins VPS-4 and CHIMP-1 (95). MVB and recycling endosomal markers, including Rab11 and transferrin, localize within the VAC, further suggesting a role for these structures in assembly and egress (58). Intriguingly, ESCRT and MVB have also been implicated in the envelopment and egress of viruses traditionally classified as non-enveloped. The picornavirus hepatitis A virus hijacks an envelope from cellular membranes in a VPS-4B- and Alix-dependent fashion (78).

In agreement with a role for MVB in virus maturation and egress, our studies demonstrate that MVB are induced during HCMV infection in ECs and become densely packed with virions and DBs (Fig. 10). In contrast to those in WT infection, MVB in ECs infected with the UL135STOP virus were devoid of virions and DBs (Fig. 10B). This finding suggests that virions and DBs are actively recruited into MVB by the virus, a process that may require UL135. Further evidence that HCMV encodes mechanisms to modulate MVB biogenesis comes from the disruption of the gene encoding the UL71 tegument protein, which results in enlarged vesicles resembling MVB with virions clustered on the limiting membrane (96). In work from two groups, UL71 mutant viruses exhibit dramatic alteration of intracellular membranes, altered VAC formation, and defects in virus egress (96, 97). In addition to their role in egress, MVB are increasingly being recognized as important organelles for intra- and intercellular signaling (63, 98) through their ability to attenuate signaling or secrete signaling molecules in exosomes, respectively. MVB have been shown to be important for intercellular signaling during Epstein-Barr virus infection of B cells (67, 79, 80). Along these lines, it is intriguing to consider how UL135 may alter exosome composition in HCMV-infected ECs and to speculate on the potential impact of this manipulation on virus dissemination and inflammation of the endothelium. It was reported recently that pUL135 and UL136 and the mechanisms by which they ensure tropism in ECs is critical to understanding how viruses target and modulate infection in key cell types in the host. These questions have important implications for understanding how infection of the vascular endothelium impacts inflammation of the endothelium and vascular pathologies such as atherosclerosis, as well as how we might control HCMV persistence and chronic virus shedding by limiting infection of ECs.

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