The early UL31 gene of equine herpesvirus 1 encodes a single-stranded DNA-binding protein that has a nuclear localization signal sequence at the C-terminus

Seongman Kim1, Byung Chul Ahn1, Dennis J. O’Callaghan, Seong Kee Kim*

Department of Microbiology and Immunology, Center for Molecular and Tumor Virology, Louisiana State University Health Sciences Center, Shreveport, LA 71130-3932, USA

A R T I C L E   I N F O

Article history:
Received 30 April 2012
Returned to author for revisions
15 May 2012
Accepted 24 May 2012
Available online 20 June 2012

Keywords:
Equine herpesvirus 1
UL31 protein of EHV-1
Single-stranded DNA-binding protein of EHV-1
EHV-1 DNA replication
Nuclear localization signal

A B S T R A C T

The amino acid sequence of the UL31 protein (UL31P) of equine herpesvirus 1 (EHV-1) has homology to that of the ICP8 of herpes simplex virus type 1 (HSV-1). Here we show that the UL31 gene is synergistically trans-activated by the IEP and the UL5P (EICP27). Detection of the UL31 RNA transcript and the UL31P in EHV-1-infected cells at 6 h post-infection (hpi) as well as metabolic inhibition assays indicated that UL31 is an early gene. The UL31P preferentially bound to single-stranded DNA over double-stranded DNA in gel shift assays. Subcellular localization of the green fluorescent protein (GFP)–UL31 fusion proteins revealed that the C-terminal 32 amino acid residues of the UL31P are responsible for the nuclear localization. These findings may contribute to defining the role of the UL31P single-stranded DNA-binding protein in EHV-1 DNA replication.

© 2012 Elsevier Inc. All rights reserved.

INTRODUCTION

Equine herpesvirus 1 (EHV-1) is a member of the Alphaherpesvirinae subfamily along with varicella-zoster virus (VZV), pseudorabies virus (PRV), bovine herpesvirus 1 (BHV-1), and herpes simplex virus type 1 (HSV-1) (Roizman and Pellet, 2001). As a major pathogen of horses, EHV-1 causes respiratory disease, neurological disorders, and abortion in pregnant mares (Allen and Bryans, 1986; Crabb and Studdert, 1995; O’Callaghan and Osterrieder, 2008). During a lytic infection, EHV-1 gene expression is controlled by viral negative and positive regulatory molecules and is divided into immediate-early (IE), early, and late stages (Ahn et al., 2007, 2010, 2011; Caughman et al., 1985; Charvat et al., 2011; Gray et al., 1987a, 1987b; Holden et al., 1995; Kim et al., 1997, 1999, 2006, 2011; Smith et al., 1992; Zhao et al., 1995). The sole IE gene is expressed first followed by expression of early genes, and then late genes are turned on after viral DNA replication (O’Callaghan and Osterrieder, 2008).

Extensive studies of HSV-1 DNA replication showed that seven viral proteins are essential for HSV-1 DNA replication (Challberg, 1986; Conley et al., 1981; Gao et al., 1993): a HSV-1 single-stranded DNA-binding protein (ICP8), a DNA polymerase (UL30P), a DNA polymerase accessory protein (UL42P), an origin-binding protein (UL9P), and a helicase-primase complex (UL5P, UL8P, and UL52P). Among the seven DNA replication-associated proteins, the roles of single-stranded DNA-binding protein are quite interesting. HSV-1 ICP8 has multi-functions in viral DNA replication. It preferentially binds to single-stranded DNA (Gourves et al., 2000; Lee and Knipe, 1985; Ruyechan, 1983, 1988), mediates strand exchange, inhibits single-stranded DNA digestion, and stimulates double-stranded DNA digestion (Boehmer and Lehman, 1993; Bortner et al., 1993; Dutch and Lehman, 1993; Reuven and Weller, 2005). Furthermore, ICP8 has been reported to modulate the activity of the viral polymerase (UL30P) (Chiou et al., 1985; Hernandez and Lehman, 1990; O’Donnell et al., 1987) and to promote the helicase activity of the viral helicase-primase complex (UL5P/UL8P/UL52P) (Falkenberg et al., 1998; Hamatake et al., 1997). It has been proposed that the C-terminal α-helix is required for the localization of ICP8 to prereplicative sites by binding to viral or cellular factors that target or retain ICP8 at specific intranuclear sites (Taylor and Knipe, 2003). The VZV counterpart of single-stranded DNA-binding protein is encoded by VZV open reading frame (ORF) 29 and is expressed in human ganglia during latency (Kinchington et al., 1988; Lungu et al., 1998; Roberts et al., 1985). It appears in the nucleus during lytic infection but in the cytoplasm of infected neurons during latency (Lungu et al., 1998). In addition, ORF29P influences expression of some viral genes that are regulated by an IE ORF62 protein (ORF62P) (Boucaud et al., 1998; Cohen et al., 2007).
It has been previously demonstrated that the putative catalytic subunit (pORF30) and accessory protein (pORF18) of EHV-1 DNA polymerase are the counterparts of HSV-1 UL30 and UL42, respectively (Allen et al., 1977; Loregian et al., 2006). Activity of ORF30p as a DNA polymerase is stimulated by the ORF18p as an accessory subunit of the viral DNA polymerase. The amino acid sequence of EHV-1 UL31p (previously designated as ICP130) revealed its homology with HSV-1 ICP8 and VZV ORF29p, suggesting that the UL31p may have similar properties of HSV-1 ICP8 or VZV ORF29p in EHV-1 DNA replication. A previous study reported that EHV-1 UL31p nuclear protein bound to both single- and double-stranded DNA when using nuclear cell extracts of EHV-1 infected RK13 cells in gel shift assays (Lewis et al., 1995). However, it is still unknown how UL31 gene is regulated, at which stage the UL31 gene is expressed, which region of the UL31p is responsible for its nuclear localization, and whether UL31p harbors the independent capability to bind to both single- and double-stranded DNAs. In this study, we showed that the EHV-1 UL31 gene is synergistically trans-activated by the IEP and the UL5p (EICP27). Metabolic inhibition assays revealed that UL31 is an early gene. Moreover, the UL31p harbors a nuclear localization signal sequence at the extreme 32 residues of the C-terminus and preferentially binds to single-stranded DNA over double-stranded DNA in gel shift assays.

Results

UL31 gene trans-activated by the IEP encodes ~120 kDa UL31 protein

The UL31 ORF located within the EHV-1 Ul genomic region encodes a UL31 protein (UL31p) of 1209 amino acids (Fig. 1A) (Telford et al., 1992). The UL31 gene of EHV-1 RacL11 strain was PCR-amplified, cloned, and then sequenced (Kim et al., unpublished data). Comparisons of the DNA sequences of the Ab4p and RacL11 UL31 genes showed seven nucleotide differences (99.8% identity) that resulted in three amino acid differences (99.8% identity). In order to perform functional studies, we cloned and expressed UL31p in RK13 cells as an intact protein as well as a recombinant green fluorescent protein (GFP) fusion protein. An ~120 kDa UL31p was detected in RK13 cells infected with EHV-1 (Fig. 1B, lane 3) or transfected with the plasmid expressing the UL31 ORF (Fig. 1B, lane 4), but not in mock-infected cells (Fig. 1B, lane 2). Moreover, anti-UL31p monoclonal antibody YC3 was also able to detect an ~147 kDa GFP–UL31p fusion protein in cells transfected with the pEGFP-UL31 plasmid (Fig. 1B, lane 5), but not with the pEGFP-N1 control plasmid (Fig. 1B, lane 6). These findings indicated that the UL31 gene is expressed as an ~120 kDa protein. Next, luciferase reporter assays were performed to examine how the UL31 gene is regulated. As shown in Fig. 2, the different regions of the UL31 ORF upstream sequence were cloned into the pGL3-basic plasmid [pUL31(−1423/−3)-Luc, pUL31(−1423/−143)-Luc, pUL31(−1423/−265)-Luc, pUL31(−1033/−143)-Luc, and pUL31(−1033/−3)-Luc]. Reporter plasmids were co-transfected into RK13 cells without effector plasmids, and promoter activities were determined. The maximal luciferase activity was shown from pUL31(−1033/−3)-Luc (Fig. 2B, bar 5) so that this plasmid was used for following luciferase assays. Interestingly, pUL31(−1423/−265)-Luc lacking a tentative TATA box sequence [UL31(−232/−227)] showed very low promoter activity (Fig. 2B, bar 3), suggesting that a TATA box sequence (TATAAT) is required for full activation of the UL31 gene. To determine which EHV-1 regulatory protein trans-activates the

Fig. 1. Location of UL31 gene in EHV-1 genome and detection of UL31 protein. (A) Location of UL31 gene in EHV-1 genome. Gene location was based on previously published Ab4p EHV-1 sequence (Telford et al., 1992). (B) Detection of UL31p from RK13 cells infected with EHV-1 and transfected with plasmid expressing UL31p or GFP–UL31p.

Fig. 2. UL31 promoter region cloned into luciferase reporter vector and UL31 promoter activity. (A) The UL31 promoter region cloned into luciferase reporter vector. The number indicates nucleotide distance from UL31 ORF first nucleotide (+1). The tentative TATA box sequence is located on UL31(−232/−227). Different regions of UL31 promoter were cloned into luciferase reporter vector as described in Materials and methods. (B) The UL31 promoter activities were examined without effector plasmid as described in Materials and methods. Promoter activity is expressed as relative luciferase units (RLU). Each experiment was carried out in triplicate. Error bars represent the mean ± standard deviation (SD) of three independent experiments.
UL31 gene, further promoter assays were performed using pUL31(–1033/–3)-Luc and effector vectors expressing each of the EHV-1 regulatory proteins. The UL31 gene was trans-activated by the IEP, but not independently by any other EHV-1 regulatory protein (Fig. 3A). Further, additional assays revealed that the IEP synergistically trans-activates the UL31 gene with the UL5P, a homolog of HSV-1 ICP27 (Fig. 3B), whereas IR2P, a negative EHV-1 regulatory molecule (Kim et al., 2006, 2011), interfered with trans-activation of the IEP (Fig. 3B).

**UL31 is an early gene**

Next, to determine in which stage of the EHV-1 replication cycle the UL31 gene is expressed, the synthesis of the UL31 transcript and UL31 protein was examined at various times after infection. The UL31 transcript was not detected at the immediate-early time of either 1 or 3 h post-infection (hpi; Fig. 4A, lanes 2 and 3, respectively) or in mock-infected cells (Fig. 4A, lane 1), but was first detected at 6 hpi and increased significantly in amount at late times of infection at 8 and 12 hpi (Fig. 4A, lanes 5 and 6, respectively). Similarly, the UL31 protein was first detected in small amounts at 6 hpi and was produced in large amounts at 12 hpi (Fig. 4B, lanes 4 and 5, respectively). These findings suggested that UL31 is an early gene. To confirm the assignment of UL31 as a member of the early class, assays employing metabolic inhibitors were performed by using cycloheximide (CHX) to block protein synthesis and phosphonoacetic acid (PAA) to inhibit EHV-1 DNA synthesis. The EHV-1 IE, early thymidine kinase (TK), and late glycoprotein K (gK) genes were used as positive controls. As expected, the IE transcript was expressed in uninhibited and CHX-treated infected RK13 cells (Fig. 4C, IE lanes 2 and 3, respectively). The TK transcript was detected in both uninhibited cells and in PAA-treated cells (Fig. 4C, TK, lanes 2, 4, and 5, respectively) but not in CHX-treated cells even at 6 hpi (Fig. 4C, TK, lane 3) as expected for a gene of the early class. Expression of the gK transcript was not detected at 6 hpi in either uninhibited or CHX-treated infected cells (Fig. 4C, gK, lanes 2 and 3, respectively) and also was not detected at 20 hpi in PAA-treated cells (Fig. 4C, gK, lane 5), confirming that gK is a true late gene as its transcription requires the initiation of viral DNA synthesis. In the case of the UL31 gene, the results were identical to those of the TK gene in that synthesis of the UL31 transcript was readily detected at 6 hpi in uninhibited infected cells and at 20 hpi in PAA-treated cells (Fig. 4C, UL31 lanes 2 and 5, respectively), but was inhibited in CHX-treated cells. Thus, the results of all experiments in Fig. 4 clearly demonstrate that EHV-1 UL31 is an early gene.

**UL31P binds to a single-stranded DNA in a sequence-independent manner**

According to previous studies (Lewis et al., 1995), the UL31P (previously designated ICP130) in nuclear extracts obtained from EHV-1-infected RK13 cells bound to both single- and double-stranded DNAs. However, the amino acid sequence homology of UL31P to the HSV-1 single-stranded DNA binding protein (ICP8) suggests the possibility that the UL31P may have a binding specificity to single-stranded DNA and that the UL31P indirectly binds to double-stranded DNA by nuclear cell extracts obtained from EHV-1-infected cells. To examine this possibility, we purified full-length UL31P as a glutathione S-transferase (GST)-UL31P fusion protein and employed it in DNA-binding assays with a radiolabeled single- and double-stranded DNA probe as described in Materials and methods. Purified GST-UL31 fusion protein

![Fig. 3.](image-url)
(~150 kDa) was detected by western blot using an anti-UL31P monoclonal antibody (Fig. 5A, lane 3). The purified GST-UL31P bound to single-stranded DNA probes and yielded a pattern with multiple bands (Fig. 5B, lanes 3 and 8), but not to double-stranded DNA (Fig. 5B, lane 13), whereas GST protein did not bind to either a single-stranded or double-stranded DNA (Fig. 5B, lanes 2, 7, and 12). Moreover, the addition of anti-UL31P monoclonal antibody YC3 led to a supershift of the single-stranded DNA probe (Fig. 5B, lanes 4 and 9). However, the DNA-protein complex was not supershifted by the addition of EHV-1 IE region 2-specific monoclonal antibody E1.1 (Fig. 5B, lanes 5 and 10). These results indicate that the UL31P binds to a single-stranded DNA.

To determine whether the interaction of UL31P with a single-stranded DNA is sequence-specific, three different sequence DNA probes were employed in gel shift assays: a 46-mer oligonucleotide corresponding to nucleotide positions 1-23 of the EHV-1 IE gene promoter (Fig. 5C, lanes 1-4), a 40-mer spanning the EHV-1 TK gene promoter (positions -290 to -349 nt relative to the translational start codon of TK ORF; Fig. 5C, lanes 5-8), and a 60-mer spanning the simian virus 40 (SV40) early promoter (positions -60 to -169 nt relative to the transcription start site; Fig. 5C, lanes 9-12). The GST-UL31P was able to bind to all three single-stranded DNA probes with multiple bands (Fig. 5C, lanes 3, 7, and 11), suggesting that the DNA-binding ability of the UL31P is not dependent on DNA sequence. The presence of multiple bands in gel shift assays may be due to the cooperative binding of the UL31P to single-stranded DNA as shown in binding of HSV-1 ICP8 (EHV-1 UL31P homolog) to single-stranded DNA (Dudas and Ruyechan, 1998; Mapelli et al., 2000; Ruyechan, 1983) and interaction of single-stranded DNA with the ICP130 in nuclear extracts obtained from EHV-1-infected cells (Lewis et al., 1995). To further confirm the binding specificity of the UL31P to the single-stranded DNA, competition experiments were performed using approximately 1-, 10-, and 100-fold molar excesses of unlabeled competitors (Fig. 5D). The formation of the GST-UL31P(IE(+)) DNA complex was inhibited in the presence of increasing amounts of three single-stranded DNA competitors IE(+), IE(−), and SV40E(+), respectively (Fig. 5D, lanes 4–5, 7–8, and 12–14). However, the double-stranded DNA competitor IE(+)/IE(−) did not inhibit the protein-DNA complexes in the presence of increasing amounts of the competitor (Fig. 5D, lanes 9–11). These observations suggest that the UL31P has binding specificity for a single-stranded DNA.

**UL31P harbors a nuclear localization signal in the extreme 32 residues of the C-terminus**

Since the UL31P plays an essential role in viral DNA replication and is a DNA-binding protein, it may harbor nuclear localization signal sequences. To identify a sequence of the UL31P responsible for nuclear localization, the plasmids of pGFP-UL31 encoding different regions of the UL31 protein (Fig. 6A) were transfected into RK13 cells, and the distribution of the green fluorescent protein (GFP) was determined under fluorescence microscopy as described in Materials and methods. The protein expression level of all GFP-UL31 constructs was confirmed by western blot using an anti-rabbit polyclonal GFP antibody (Fig. 6B and D). As shown in Fig. 6C, the full-length GFP-UL31P(1–1209) and GFP–UL31P(601–1209) were localized in the nucleus, but the control GFP was distributed in both cytoplasm and nucleus. GFP–UL31P(1–600) and GFP–UL31P(301–900) were distributed in cytoplasm, suggesting that the nuclear localization signal (NLS) sequence is located within residues 601–1209 of the UL31P. The UL31P(601–1209) was further dissected to identify the NLS sequence (Fig. 6D). The GFP–UL31P(801–1209) and GFP–UL31P(1001–1209) fusion proteins (Fig. 6C) as well as GFP–UL31P(1101–1209) (Fig. 6E) were localized in the nucleus, indicating that the NLS sequence of the UL31P is located within the C-terminal 109 amino acid residues. To define the essential sequences of UL31P NLS, N-terminal sequences of UL31P(1101–1209) and C-terminal sequences of UL31P(801–1209) were deleted as described in Materials and methods. The deletion of C-terminal 9 and 34 residues of the UL31P(801–1209) resulted in the failure of nuclear localization of the protein (Fig. 6E, 801–1200 and 801–1175), but only UL31P(801–1209) containing the intact C-terminal sequence showed nuclear localization of GFP (Fig. 6C). Furthermore, N-terminal sequence of the UL31P(1101–1209) was deleted to determine the minimal NLS sequence required for its nuclear localization. Deletion of N-terminal 71 and 77 amino acid residues of the UL31P(1101–1209) [GFP–UL31P(1172–1209) and GFP–UL31P(1178–1209)] did not affect the nuclear localization of GFP–UL31P (Fig. 6E). However, deletion of the 83 and 89 amino acid residues [GFP–UL31P(1184–1209) and GFP–UL31P(1190–1209)] led to GFP distribution in both cytoplasm and nucleus (Fig. 6E), indicating that the minimal NLS sequence of the UL31P is located within the C-terminal 32 residues (aa 1178–1209).

To further determine the functional relevance of the NLS of UL31P homologs, we compared the C-terminal 50 amino acid residues of EHV-1 UL31 ORF harboring its NLS sequence with those of UL31P homologs encoded by three other alphaherpesviruses. Multiple sequence alignment showed that the extreme C-terminus of EHV-1 UL31P has approximately 42.6% and 44% amino acid identity with that of HSV-1 ICP8 and PRV UL29,
respectively, but exhibited only 10% homology with that of VZV ORF29 that contains the NLS domain within its N-terminus (Fig. 7). The minimal NLS sequence of the UL31P is similar to that of HSV-1 ICP8.

Discussion

The 78 genes of EHV-1 are regulated by EHV-1 regulatory molecules (O’Callaghan and Osterrieder, 2008) at immediate-early, early, and late stages (Caughman et al., 1985; Gray et al., 1987a). After the expression of immediate-early and early genes, viral DNA replication occurs followed by the expression of the late genes (O’Callaghan and Osterrieder, 2008). Elucidating the functions of replication-associated proteins is a key to understanding the mechanisms of DNA replication and the pathogenesis of viral diseases. There is little known about viral proteins associated with EHV-1 replication except that ORF30P and ORF18P were previously identified as a DNA polymerase and its accessory subunit, respectively (Loregian et al., 2006). Considering the diverse role of HSV-1 ICP8 and VZV ORF29P in the replication of HSV-1 and VZV, respectively, it is still necessary to further characterize the UL31 gene and the UL31P for a better understanding of EHV-1 replication.
Promoter assays indicated that the UL31 gene is trans-activated only by the IEP in synergistic fashion with the UL5P (EICP27). In addition, detection of the UL31 RNA transcript and the UL31P at the early stage of viral infection as well as metabolic inhibition assays revealed that the UL31 gene, like the EHV-1 representative early gene TK, is an early gene. We also observed that a consensus IEP-binding site (IEBS; 5’–ATCGT-3’) is located at 455-bp upstream of a tentative TATA box within the UL31 promoter region. This consensus IEBS may be a cis-acting element required for IEP-mediated trans-activation of the UL31 gene although we did not provide any evidence of the interaction of the IEP with the IEBS sequence in this study.

Although previous studies (Lewis et al., 1995) reported that the UL31P (previously designated ICP130) binds to both single- and double-stranded DNAs in gel shift assays using EHV-1-infected cell nuclear extract, sequence analysis of EHV-1 UL31P showed a homology to single-stranded DNA-binding proteins, such as HSV-1 ICP8 and VZV ORF29P that have multiple functions.
in DNA replication and gene regulation (Cohen et al., 2007; Cohrs et al., 2002; Reuven et al., 2003; Reuven and Weller, 2005; Stallings et al., 2006). These observations imply that the certain protein(s) in EHV-1-infected nuclear cell extract may be involved in binding of the UL31P to the double-stranded DNA. To examine this possibility, we employed purified GST–UL31P fusion protein in binding assays. Similar to HSV-1 ICP8, which binds to a single-stranded DNA (Gourves et al., 2000), the UL31P preferentially bound to a single-stranded DNA in a sequence-independent manner. This result supports that the UL31P is a single-stranded DNA-binding protein, and also suggests that the binding of UL31P to double-stranded DNA may be due to the additional viral and/or cellular protein(s).

HSV-1 ICP8 and VZV ORF29P that harbor DNA-binding activity are localized in the nucleus (Cohrs et al., 2002; Taylor and Knipe, 2003). It was also shown that the UL31P, a single-stranded DNA-binding protein, is localized in the nucleus, suggesting that the UL31P harbors nuclear localization signal sequence. Dissection of the UL31P fused to the GFP revealed that the essential NLS sequence of UL31P is located at the C-terminus of the protein and the extreme 32 amino acid residues (aa 1178–1209) of UL31P is responsible for its nuclear localization. According to previous results, HSV-1 ICP8, like UL31P, harbors a NLS located in the C-terminal 28 amino acid residues (Gao and Knipe, 1992), whereas VZV ORF29P contains noncanonical NLS sequence located in amino acids 9–154 of the N-terminus for each UL31P homologs. The NLS sequences of HSV-1 ICP8 and VZV-1 ICp8 are underlined.

Table 1

<table>
<thead>
<tr>
<th>Plasmid</th>
<th>Primer name (sequence [5’–3’])</th>
</tr>
</thead>
<tbody>
<tr>
<td>pUL31(−1423/−3)–Luc</td>
<td>UL31-F1 (ggctcgagagcctcttggtagttgctc)</td>
</tr>
<tr>
<td>pUL31(−1423/−143)–Luc</td>
<td>UL31-F1/UL31-R3 (gggaagcttggtggggaatacccta)</td>
</tr>
<tr>
<td>pUL31(−1423/−265)–Luc</td>
<td>UL31-F1/UL31-R2 (tttaacctacagctgcgctgccga)</td>
</tr>
<tr>
<td>pUL31(−1033/−143)–Luc</td>
<td>UL31-F2 (ttctcgagagctcttggtagttgctc)</td>
</tr>
<tr>
<td>pUL31(−1033/−3)–Luc</td>
<td>UL31-F2/UL31-R4 (ggttgcggccgcttagccacgctccacgctcca)</td>
</tr>
<tr>
<td>pCMV–UL31</td>
<td>UL31ORF-R (ggaagcttggtggggaatacccta)</td>
</tr>
<tr>
<td>pEFP–UL31orf5</td>
<td>Ultralink-R (tttgtgctcagcccccacagc)</td>
</tr>
<tr>
<td>pEFP–UL31orf1</td>
<td>SpeI–linker-R (gtttgcggccgctcaggttaataaagtttatggc)</td>
</tr>
<tr>
<td>pEFP–UL31orf2</td>
<td>UL31ORF-R (gtttgcggccgcttagagcatgtcaaaggtgagc)</td>
</tr>
<tr>
<td>pEFP–UL31orf3</td>
<td>SpeI–linker-F (tttgtgctcagcccccacagc)</td>
</tr>
<tr>
<td>pEFP–UL31orf4</td>
<td>UL31ORF-R (gtttgcggccgcttagagcatgtcaaaggtgagc)</td>
</tr>
<tr>
<td>pEFP–UL31orf5</td>
<td>SpeI–linker-R (gtttgcggccgctcaggttaataaagtttatggc)</td>
</tr>
<tr>
<td>pEFP–UL31orf6</td>
<td>UL31ORF-R (gtttgcggccgcttagagcatgtcaaaggtgagc)</td>
</tr>
<tr>
<td>pEFP–UL31orf7</td>
<td>SpeI–linker-F (tttgtgctcagcccccacagc)</td>
</tr>
</tbody>
</table>

Fig. 7. Alignment of the C-terminal 50 amino acids of EHV-1 UL31P with those of alphaherpesvirus homologs. The amino acid sequences of EHV-1 UL31 ORF and its homologs were aligned by use of CLUSTALW2 multiple sequence alignment program (http://www.ebi.ac.uk/Tools/es/cgi-bin/clusterw2). EHV-1 UL31 (AB48 strain, GenBank ID: NC001491; Racl1 strain, Kim et al., unpublished data); HSV-1 ICP8 (GenBank ID: BAe78520); PRV UL29 (GenBank ID: NC006151); VZV ORF29 (GenBank ID: ABW00912). The numbers indicate the amino acid position relative to the N-terminus for each UL31P homologs. The NLS sequences of EHV-1 UL31P and HSV-1 ICp8 are underlined.
the UL31P may have additional role(s) in EHV-1 DNA replication. Our findings such as the regulation of the early UL31 gene, NLS localization of the UL31P, and a single-stranded DNA-binding function of the UL31P would give an insight into further studies for elucidating the mechanism of the UL31P in viral DNA replication.

Materials and methods

Cell and virus

Rabbit kidney RK13 cells were maintained with Eagle’s minimal essential medium supplemented with 100 units/ml of penicillin, 100 μg/ml of streptomycin, nonessential amino acids, and 5% fetal bovine serum. Pathogenic RacL11 was used as wild-type EHV-1.

Construction of the plasmids

PCR products were amplified using a pRacL11 EHV-1 BAC template, Accuprime pfx polymerase (Invitrogen, Carlsbad, CA), and appropriate primers (Table 1). PCR products were digested with appropriate enzymes and cloned into the appropriate vector using standard methods (Sambrook et al., 1989), and were confirmed by sequence analysis. Primer design was based on the published sequences (Telford et al., 1992). The UL31 promoter regions were PCR amplified as shown in Fig. 2A. PCR products of UL31 (–1423/–3), UL31 (–1423/–143), UL31 (–1423/–265), UL31 (–1033/–143), and UL31 (–1033/–3) were digested with XhoI and HindIII (named pUL31 (–1033/–3)-Luc, pUL31 (–1423/–143)-Luc, pUL31 (–1423/–265)-Luc, pUL31 (–1033/–143)-Luc, and pUL31 (–1033/–3)-Luc, respectively).

The UL31 ORF PCR product was digested with EcoRI and NotI, and cloned into pEGFP-N1 (BD Bioscience, San Jose, CA) digested with SpeI enzyme site cloned into pEGFP-N1 (BD Bioscience, San Jose, CA) digested with EcoRI and NotI (named pCMV–UL31). To create a SpeI enzyme site digested with SpeI and AflII (named pGFP–UL31P(1196–1209), pGFP–UL31P(1193–1209), pGFP–UL31P(1192–1209), pGFP–UL31P(1184–1209), pGFP–UL31P(1178–1209), and pGFP–UL31P(1172–1209), respectively).


Western and northern blot analyses

90% confluent RK13 cells were infected with EHV-1 at a multiplicity of infection (moi) of 5 and transfected with pCMV–UL31, pEGFP–UL31 or pEGFP–N1 plasmid using a Lipofectin (Invitrogen) according to the manufacturer's directions. After 48 h incubation, total cell extracts were prepared from virus-infected or plasmid-transfected RK13 cells and applied to western blot analyses. To determine the expression level of the UL31P at indicated times post infection, RK13 cells were infected with RacL11 at an moi of 5. Whole cell lysates were separated by a sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE), and then transferred to a nitrocellulose membrane (Ambion, Austin, TX). The UL31P was detected by using an anti-UL31P monoclonal antibody YC3 (a gift from Dr. Caughman, Medical College of Georgia) as primary antibodies and an anti-mouse IgG[Fc]-alkaline phosphatase conjugate (Promega) as a secondary antibody. In addition, the GFP-UL31 fusion proteins were detected with an anti-rabbit polyclonal GFP antibody (Santa Cruz). Protein was visualized by incubating the membrane containing blotted proteins in the AP conjugate substrate (AP conjugate substrate kit, Bio-Rad Laboratories) according to manufacturer's directions.

To detect the UL31, gK, TK, and IE transcripts, total RNA was isolated from mock- and virus-infected cells by using a RNeasy Mini kit (Qiagen, Valencia, CA). Total RNA was separated on 3 or 5% denaturing acrylamide gel (8M Urea) and transferred to a nylon membrane (Ambion) by using a semi-dry electroblotter (Bio-Rad Laboratories). The PCR amplicons of UL31 gene (primers, 5’-ctc gcg caa gcc agc ccc tca tca-3’/5’-tgc gct gct cat gag ggg tat tag aga-3’), gK gene (primers, 5’-ctc gcg caa gcc agc ccc tca tca-3’/5’-gag ggt gct gct cat gag ggg tat tag aga-3’), gK gene (primers, 5’-tta gga cga gga gga gga cca taa gag ccc-3’/5’-ttt gattag cta cct gct gat taa agg-3’), and IE gene (primers, 5’-ctc tac gag ttc gag aac gac gag cca taa gag ccc-3’/5’-ctt gtt gct gtt gag cca taa gag ccc-3’), as well as pSVSPORT1 DNA (Gibco, BRL, MD) was added to the reporter plasmid.Reporter plasmid with or without effector plasmid(s) were transfected at 48 h post transfection (hpt), normal growth medium was added, and luciferase activities were measured by a luciferase assay kit (Promega) and a POLARStar OPTIMA plate reader (BMG LABTECH Inc., Bury, NC) according to the manufacturer's directions.

Luciferase reporter assay

Promoter activity was examined as described in previous studies (Kim et al., 2006). Briefly, 70% confluent RK13 cells were prepared in 24-well plates and 0.1 μM of reporter vector and 0.3 μM of effector vector were used for cotransfection. Reporter vectors were pUL31 (–1423/–3)-Luc, pUL31 (–1423/–143)-Luc, pUL31 (–1423/–265)-Luc, pUL31 (–1033/–143)-Luc, and pUL31 (–1033/–3)-Luc. The EHV-1 regulatory protein-expressing vectors such as pSVIE, pSVIR, pCMV–IR4 (EICP22), pSVULS (EICP27), and pCMV–ETiF (Ahn et al., 2007) as well as pCMV–UL31 were used as effector vectors. 6 μl of lipofectin (Invitrogen) was mixed with 300 μl of Opti-MEM medium (Gibco, BRL, Gaithersburg, MD), and incubated for 45 min at room temperature. Reporter plasmid with or without effector plasmid(s) were mixed with 300 μl of Opti-MEM medium (Gibco, BRL), and total DNAs were adjusted to the same amount with the pSVSPORT1 DNA (Gibco, BRL). The mixture was combined and incubated at room temperature for 15 min. One-third of the total mixture was transferred into each of three wells of the RK13 cells. At 5 h post transfection (hpt), normal growth medium was added, and luciferase activities were measured by 48 hpt by using a luciferase assay kit (Promega) and a POLARStar OPTIMA plate reader (BMG LABTECH Inc., Bury, NC) according to the manufacturer's directions.

Expression and localization of the UL31P

To detect UL31P expression and localization, RacL11-infected RK13 cells were fixed with 3 M NaOH for 10 min at room temperature and then neutralized by adding an equal volume of 1 M Tris–HCl (pH 7). Prehybridization,
hybridization and washing were performed using the NorthernMax Kit (Ambion) followed by autoradiography using a phosphor image screen and the molecular imager FX system (Bio-Rad Laboratories).

**Metabolic inhibition assays**

Metabolic inhibition assays were performed using virus-infected RK13 cells as previously described (Gray et al., 1987b; Zhou et al., 1997). Briefly, 85% confluent RK13 cells were infected with EHV-1 at an moi of 5 and incubated at 37°C in the presence of or absence of the protein synthesis inhibitor cycloheximide (CHX, 100 μg/ml; Sigma-Aldrich Corp. St. Louis, MO) for 6 h, or in the presence of the viral DNA replication inhibitor phosphonoacetic acid (PAA, 200 μg/ml; Sigma-Aldrich Corp.) for 20 h followed by total RNA isolation using a RNeasy Mini Kit (Qiagen, Valencia, CA). Presence of RNA transcripts were determined by northern blot assays using total RNA and probes specific to each of IE, UL31, gK and TK RNA transcripts.

**Gel shift assay**

The DNA-binding assay and purification of glutathione S-transferase-UL31 fusion protein (GST-UL31P) were conducted as previously described (Kim et al., 1995). A 46-mer DNA oligonucleotide and its complement spanning the EHV-1 IE promoter region (positions –18 to –63 nt relative to the transcription start site, 5'-cca cta ggg gaa cag cca aac ttc ctc gta gta taa agc acc t-3') and 5'-agg ctc tgg ata cta cca ggg agt ttt gcc ttc ccc cta ggt g-3'), a 60-mer spanning EHV-1 thymidine kinase promoter region (positions –290 to –349 nt relative to the transcriptional start site of TK ORF, 5'-tac tcg cgg tgt tca tat ttt tgg aac gag acg acc ttt gta cct gta tta ggc acc aca-3'), and a 60-mer spanning SV40 early promoter (positions –10 to –69 nt relative to the transcription start site, 5'-agg tcc gcc ctg tct ccc ccc cat cgc tga tca att ttt att tat gca ggc gag ggg-3') were used in gel shift assays. Each oligonucleotide and its complement were annealed and either end-labeled with [γ-32P]dATP (New England Nuclear Corporation) and T4 polynucleotide kinase (Promega) according to the manufacturer's directions. The DNA-binding reactions were carried out in a total volume of 20 μl containing approximately 1 ng of radiolabeled single- or double-stranded DNA probes (2 × 10^6 cpm/ng, 0.1 μg of poly(dI-dC) as a nonspecific competitor, binding buffer (20 mM HEPES-KOH [pH 7.9], 0.5 mM dithiothreitol, 10% glycerol, 0.1 mM EDTA, 0.025% NP-40, 25 mM KCl, 2 mM MgCl2), and 100 ng of indicated protein. The DNA binding reaction mixtures were incubated at room temperature for 20 min. In the supershift assay, an anti-UL31P monoclonal antibody YC3 or EHV-1 IE region 2-specific monoclonal antibody E1.1 was added to the indicated reactions, and the reactions were incubated for another 20 min at room temperature. For the competition assay, 1-, 10-, 100-fold molar excess of unlabeled DNA competitor was added to the binding reaction. After the reaction, 5 μl of loading buffer (200 mM HEPES-KOH [pH 7.9], 50% [vol/vol] glycerol, 0.02% bromphenol blue) was added, and the reaction samples were separated by electrophoresis using a 3.5% polyacrylamide gel with 0.5 × Tris–borate–EDTA running buffer and 2.5% glycerol [vol/vol] for 2.5 h at 200 V followed by autoradiography using a phosphor image screen and molecular imager FX system (Bio-Rad Laboratories).

**Direct fluorescence microscopy**

70% confluent RK13 cells on cover slips were transfected with GFP–UL31 fusion protein expressing plasmids or pEGFP-N1 control plasmid using a Matra A reagent (IBA Biotechnology, Goettingen) according to manufacturer's instructions. Briefly, three micrograms of plasmid DNA was diluted to 200 μl of complete media (Mediatech, Inc., Manassas, VA) and mixed with 3 μl of MATra A reagent, and the mixture was incubated at room temperature for 20 min. Cells were washed with complete media once, and one milliliter of complete media was added to DNA-MATra A mixture. After the addition of 1.2 ml of the mixture to each well, the plate was incubated on the magnet plate for 15 min. The mixtures were removed and replaced with 3 ml of normal growth media followed by a 24 h incubation at 37°C in a humidified CO2 incubator. The cells were washed twice with PBS at 24 hpt, fixed with 4% paraformaldehyde for 15 min at room temperature, washed, and permeabilized with 0.2% Triton X-100 in PBS for 2 min followed by three PBS washes. Mounting solution containing DAPI, Gold Antifade (Invitrogen), was applied to the RK13 cells, and the location of GFP within the cell was examined by a fluorescence microscopy (Leica DMBI 6000 microscope).

**Acknowledgments**

We thank Mrs. Suzanne Zavecz and LaShunta Barrow for excellent technical assistance. This research was supported by Agriculture and Food Research Initiative Competitive Grant 2008-35204-04438 from the USDA National Institute of Food and Agriculture, and by COBRE grant GM103433 from the National Institute of General Medical Sciences of the NIH.

**References**


