Bovine herpesvirus 1 regulatory proteins, bICP0 and VP16, are readily detected in trigeminal ganglionic neurons expressing the glucocorticoid receptor during the early stages of reactivation from latency.

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Abstract

Bovine herpesvirus 1 (BHV-1) establishes a life-long latent infection in sensory neurons following acute infection. Increased corticosteroid levels, due to stress, increases the incidence of reactivation from latency. Within minutes, corticosteroids activate the glucocorticoid receptor and transcription of promoters containing a glucocorticoid receptor element. A single intravenous injection of the synthetic corticosteroid dexamethasone consistently induces reactivation from latency in calves. Lytic cycle viral gene expression is detected within 6 hours after dexamethasone treatment of calves latently infected with BHV-1. Cellular transcription factors are induced by dexamethasone in trigeminal ganglionic neurons within 1.5 hours after dexamethasone treatment suggesting they promote viral gene expression during the early phases of reactivation from latency, which we operationally defined as the escape from latency. In this study, immunohistochemistry was utilized to examine viral protein expression during the escape from latency. Within 1.5 hours after dexamethasone treatment, bICP0 and a late protein (VP16) were consistently detected in a subset of trigeminal ganglionic neurons. Most neurons expressing bICP0 also expressed VP16. Additional studies revealed that neurons expressing the glucocorticoid receptor also expressed bICP0 or VP16 at 1.5 hours after dexamethasone treatment. Two other late proteins, glycoprotein C and D, were not detected until 6 hours after dexamethasone treatment and were detected in only a few neurons. These studies provide evidence that VP16 and the promiscuous viral trans-activator (bICP0) are expressed during the escape from latency suggesting they promote the production of infectious virus in a small subset of latently infected neurons.
INTRODUCTION

Bovine herpesvirus 1 (BHV-1) induces clinical disease in the upper respiratory tract, nasal cavity, or ocular cavity of cattle. BHV-1 establishes latency in sensory neurons, but periodically reactivates from latency, and consequently is widespread in cattle (31-33, 36). Infection inhibits cell-mediated immunity (4, 17-19), CD8+ T cell recognition of infected cells (21, 25, 41, 51), and induces apoptosis in CD4+ T cells (12, 74). Two viral regulatory proteins, bICP0 and bICP27, inhibit interferon dependent transcription (6, 22, 33, 61, 62). Infection also erodes mucosal surfaces within the upper respiratory tract, which promotes establishment of bacterial pathogens in the lower respiratory tract (23, 24, 83).

The incidence of BHV-1 reactivation from latency is increased following stressful stimuli that increase corticosteroid levels, reviewed in (34, 37, 56). Regardless of the reactivation stressor, lytic cycle viral gene expression, which is nearly undetectable during latency, must be activated. Administration of the synthetic corticosteroid dexamethasone (DEX) to latently infected calves or rabbits initiate BHV-1 reactivation from latency 100% of the time (28, 31, 32, 35, 36, 60). Six hours after DEX treatment lytic cycle viral RNA expression is readily detected in a subset of trigeminal ganglionic neurons of latently infected calves (73, 76). DEX treatment of latently infected calves induces apoptosis of T cells that persist in trigeminal ganglia (TG) after infection (73). T cells also persist in TG of humans or mice latently infected with HSV-1 (3, 20, 46, 64-66, 69) and are proposed to promote the maintenance of latency (38, 40, 43-45, 59).

Within 3 hours after calves latently infected with BHV-1 are treated with DEX, 11
cellular genes are induced more than ten fold in TG (80). Pentraxin 3, a regulator of innate immunity and neuro-degeneration, is stimulated 35-63 fold at 3 or 6 hours after DEX treatment. Two transcription factors, promyelocytic leukemia zinc finger (PLZF) and Slug are induced more than 15-fold 3 hours after DEX treatment. PLZF or Slug stimulates BHV-1 productive infection 20-fold or 5-fold respectively, and Slug stimulates the late glycoprotein C promoter more than 10-fold. Additional DEX induced transcription factors, SPDEF (Sam-pointed domain containing Ets transcription factor), Kruppel-like transcription factor 15 (KLF15), KLF4, KLF6, and GATA6, stimulate productive infection and certain key viral promoters. The DEX inducible cellular transcription factors are predicted to mediate certain aspects of the early phases of reactivation from latency, including activation of lytic cycle viral gene expression. Latently infected neurons that express detectable levels of lytic cycle viral proteins within six hours after DEX treatment are operationally defined as escaping latency (80).

In this study, we examined bICP0, VP16, gC, and gD protein expression during the escape from latency. bICP0 and VP16 positive TG neurons were detected at 1.5 hours after DEX treatment, but were not readily detected prior to DEX treatment. Consecutive sections were prepared and subsequent studies demonstrated that bICP0 and VP16 were frequently detected in the same neuron. Conversely, gC or gD positive neurons were not detected until 6 hours after DEX treatment, and only a few positive neurons were detected. At 1.5 hours after DEX treatment, bICP0 or VP16 and the glucocorticoid receptor (GR) were detected in the same neuron. These studies suggest that activation of the GR by DEX directly stimulates lytic cycle viral gene expression during the escape from latency.
MATERIALS AND METHODS

Cells and Virus

Bovine kidney (CRIB) cells were grown in Earle’s modified Eagle’s medium (EMEM) supplemented with 10% fetal calf serum, penicillin (10 U/ml), and streptomycin (100 μg/ml). The Cooper strain of BHV-1 was grown in CRIB cells and was used for all studies.

Calf studies

All TG samples from calves used for this study were previously described (80). In brief, BHV-1-free crossbred calves (~200 kg) were inoculated with \(10^7\) PFU of BHV-1 into each nostril and eye as described previously (28, 29, 47, 53-55). Calves were housed under strict isolation and given antibiotics before and after BHV-1 infection to prevent secondary bacterial infections. At 60 dpi, calves were injected intravenously (jugular vein) with 100 mg of DEX. Calves were then transported to the Veterinary Diagnostic lab. Prior to euthanasia by electrocution, calves were heavily sedated with Xylazine. After decapitation, TG were collected, samples from each TG were formalin fixed and then paraffin embedded. The remainder of both TG was minced into small pieces, placed into a single 50 ml conical tube, and the tube placed in a dry ice ethanol bath. TG samples were then stored at -80°C. It took approximately 5 minutes to collect TG, mince the TG, place TG pieces in a 50 ml conical tube, and submerge the tube in a dry ice ethanol bath after decapitation. One calf was decapitated at a time to ensure samples were processed in a timely manner. Calves were decapitated in the same order in which they were injected with DEX to ensure that the time-points after DEX treatment were as close as possible to the designated time point. Three calves / time point were used for these studies.
Experiments were performed in accordance with the American Association of Laboratory Animal Care guidelines and the University of Nebraska IACUC committee.

**Immunohistochemistry**

Immunohistochemistry was performed essentially as previously described (49, 50, 73, 75) using the ABC Kit (Vector Laboratories). In brief, TG from calves were fixed in neutral buffered formalin, and then embedded in paraffin. Thin sections (4-5 µm) were cut and mounted onto slides. Tissue sections were incubated 20 minutes at 65°C followed by two incubations of 10 minutes in xylene and rehydrated in graded alcohols. Tissue sections were then incubated with 3% hydrogen peroxide in PBS (pH 7.4) for 20 minutes at room temperature to block endogenous peroxidase. After 3 washes in TBS (5 minutes each) at room temperature, tissue sections were digested with 40 µl of a “ready to use” Proteinase K solution (53020, Dako) for 20 minutes at 37°C to enhance antigen retrieval. Tissue sections were then blocked with 5% normal serum diluted in TBS containing 0.25% BSA for 45 minutes at room temperature in a humidified chamber.

A peptide specific rabbit antibody was made that is directed against bICP0 (Affinity Bioreagents, Golden, CO) and the antibody was affinity purified. This antibody specifically recognizes bICP0 in infected or transfected cells. A VP16 specific rabbit polyclonal antibody was obtained from Dr. Vikram Misra (University of Saskatchewan, Saskatoon, CA). The gC and gD specific antibodies were obtained from Dr. Shafiqul Chowdhury (Louisiana State University Veterinary School, Baton Rouge, LA). The GR specific rabbit polyclonal antibody was purchased from Santa Cruz Biotechnology (H-300) and was reported to detect the mouse, rat,
and human protein. The designated rabbit polyclonal antibodies or the mouse monoclonal antibodies were used at a 1:500 dilution, incubated overnight in a humidified chamber at 4°C, and the next day washed in TBS (pH 7.6). Biotinylated goat anti-rabbit IgG (Vector Labs, PK-6101) or biotinylated donkey anti-mouse IgG (Vector Labs, PK-6102) was then incubated with the section for 30 minutes at room temperature in a humidified chamber. Next, the avidin-biotinylated enzyme complex was added to slides for 30 minutes at room temperature in a humidified chamber. After 3 washes in TBS, slides were incubated with freshly prepared substrate (Vector Labs, SK-4800), rinsed with distilled water, and counterstained with hematoxylin. Thin sections from mock-infected or latently infected calves were used as a negative control.

**Immunofluorescence to examine GR localization**

CRIB cells were incubated in EMEM supplemented with 2% charcoal-stripped FBS at 37°C for 24 hours and then placed on a glass slide for an additional 24 hours. After 2 hours of DEX treatment (100 nM), cells were fixed in 4% paraformaldehyde for 10 minutes and confocal microscopy performed. The GR primary antibody (catalogue # 3660; Cell Signaling) was diluted 1:250 in PBS with 0.05% Tween 20 and 1% BSA and incubated on coverslips for 2 hours at room temperature. After three washes, coverslips were incubated with Alexa Fluor® 488 Goat Anti-Rabbit IgG (H+L) (A11008; Invitrogen) at a dilution of 1:100 for 1 hour in the dark. After the slides were washed, DAPI (4’,6-diamidino-2-phenylindole) staining was performed to visualize the nucleus. Coverslips were then mounted on slides by use of Gelmount aqueous mounting medium (Electron Microscopy Sciences). Images were obtained with a Bio-Rad confocal laser-scanning microscope (MRC-1024ES).
Western Blot analysis

At the designated times after infection, whole cell lysate was prepared. Cells were washed with phosphate-buffered saline (PBS) and suspended in NP-40 lysis buffer (100 mM Tris \(\text{pH } 8.0\), 1 mM EDTA, 100 mM NaCl, 1% NP-40, 1 mM phenylmethylsulfonyl fluoride, and one tablet of complete protease inhibitor, Roche Molecular Biochemicals, in 10 ml buffer. Cell lysate was incubated on ice for 30 min, sonicated, and then clarified by centrifugation at 10,000 \(x\) \(g\) at 4°C for 15 min. Protein concentrations were quantified by the Bradford assay. For SDS-PAGE, proteins were mixed with an equal amount of 1x sample loading buffer (62.5 mM Tris-HCl \(\text{pH } 6.8\), 2% sodium dodecyl sulfate, 50 mM dithiothreitol, 0.1% bromophenol blue, 10% glycerol) and boiled for 5 min. Proteins were separated in an 8 or 12% SDS-PAGE gel. After electrophoresis, proteins were transferred onto a polyvinylidene difluoride membrane (Immobilon-P; Millipore) and blocked for 4 hours in 5% nonfat dry milk with Tris-buffered saline-0.1% Tween 20 (TBS-T). Membranes were then incubated with primary antibody overnight at 4°C. An antibody directed against β-actin (Santa Cruz Biotechnology, Santa Cruz, CA) was used as a loading control. After 45 minutes of washing with TBS-T, blots were incubated with donkey anti-rabbit horseradish peroxidase-conjugated immunoglobulin G (Amersham Biosciences), which was diluted 1:2000 in 5% nonfat milk in TBS-T. Blots were washed 45 minutes with TBS-T and exposed to Amersham ECL reagents, and then autoradiography performed. Primary antibodies were described above. The GR (Cell signaling, catalogue # 3660), the bICP0, and the VP16 antibody were diluted 1:1,000. The gC and gD antibody were diluted 1:500 in blocking buffer. The secondary donkey anti-rabbit and anti-mouse antibodies were purchased from GE Healthcare.
RESULTS

Viral protein expression during productive infection

Initial studies were performed to examine the specificity of BHV-1 antibodies that detect three late viral proteins (gC, gD, and VP16) and the promiscuous viral trans-activator (bICP0). The BHV-1 bICP0 mRNA is expressed as an immediate early and early transcript because it contains an IE and E promoter (77, 78). For these studies, bovine kidney cells (CRIB) were infected with BHV-1 using a moi of 2 pfu/cell. Total cell lysate was prepared at 3, 6, 12, and 24 hours after infection in the absence or presence of the viral DNA polymerase inhibitor PAA. As expected, bICP0 was expressed throughout productive infection and PAA had no effect on expression (Figure 1A, bICP0 panel). VP16 was not readily detected until 6 hours after infection and PAA reduced VP16 protein levels. Expression of the viral glycoprotein gC was not readily detected until 12 hours after infection and its expression were reduced by PAA treatment. Conversely, gD expression was detected earlier than VP16 or gC and its expression was not affected by PAA treatment. These results are consistent with HSV-1 studies examining the kinetics of gC, gD, and VP16. For example, HSV-1 gD is a prototype γ1 gene that is expressed early during infection and its expression is not dramatically altered by PAA (26, 27). Conversely, gC and VP16 are γ2 genes that are expressed relatively late during productive infection, and their expression is dramatically inhibited by PAA (26, 27).

CRIB cells infected for 20 hours were fixed in formalin and then confocal microscopy performed to determine if these antibodies recognized a viral protein in formalin fixed cells. We chose a time late after infection because we knew the late proteins would be expressed at readily detectable levels. As expected, bICP0 was detected in the nucleus of infected cells; but the other...
three viral proteins were primarily localized to membranes in infected cells (Figure 1B). None of
the antisera directed against the respective viral proteins recognized a protein in mock-infected
CRIB cells (Mock panel and data not shown). In summary, these studies demonstrated that the
respective antibodies recognized the expected viral protein by Western Blot analysis and by
confocal microscopy using formalin fixed cells.

Detection of bICP0 and VP16 during the escape from latency
To test whether viral protein expression can be detected during the escape from latency,
bICP0, VP16, gC, and gD protein expression were examined in TG sections by performing
immunohistochemistry (IHC). The rationale for examining bICP0 protein expression is that
bICP0 mRNA is consistently detected during DEX induced reactivation from latency (81), and
bICP0 stimulates productive infection and reactivation from latency (14, 15, 31-33, 37). BHV-1
encoded VP16 may promote the escape from latency because it activates expression of viral IE
genes. HSV-1 encoded VP16, a late transcript, has been reported to be differentially expressed
in neurons relative to cultured cells and consequently may stimulate reactivation from latency
(39, 71). TG neurons were recognized by bICP0 or VP16 antiserum (Figure 2) when calves
latently infected with wt BHV-1 were treated with DEX for 1.5 hours. Certain areas of TG
sections contained bICP0 or VP16 positive neurons whereas other areas did not. As expected,
bICP0 or VP16 positive neurons were not readily detected during latency, and in general fewer
bICP0 or VP16 positive neurons were detected at 3 or 6 hours after DEX treatment relative to 1.5
hours. The number of neurons stained by the gC or gD antiserum was low, and the intensity of
staining was reduced relative to the results obtained with the VP16 or bICP0 antiserum (Figure
3). We were unable to detect gC or gD prior to 6 hours after DEX treatment (data not shown).
In contrast to VP16 or bICP0, ORF2+ neurons were readily detected during latency but not six hours after DEX treatment (67).

bICP0 and VP16 are frequently expressed in the same neuron during DEX induced reactivation

To determine whether bICP0 and VP16 were expressed in the same neuron following DEX treatment, consecutive sections were prepared from TG samples at 1.5, 3, and 6 hours after DEX treatment and each section was stained with the bICP0 or VP16 specific antiserum. At 3 hours after DEX treatment, most VP16+ neurons were also bICP0+ (Figure 4A; double positive neurons denoted by arrows). We also examined 1.5 and 6 hours after DEX treatment, and like 3 hours after DEX treatment, nearly all VP16+ neurons were bICP0+ (Figure 4B).

Sensory neurons that express bICP0 and VP16 frequently express the glucocorticoid receptor

DEX, like the natural corticosteroids, binds and activates the glucocorticoid receptor (GR), reviewed in (13, 63). Since GR is expressed in rat sensory neurons (8), we predicted that bICP0 and VP16 expression may be stimulated directly by DEX in neurons that express the GR. Support for this prediction comes from the finding that the bICP0 E promoter is stimulated by DEX in transient-transfection studies (81). Thus, it was of interest to determine: 1) if the GR is expressed in a subset of bovine TG neurons, and 2) whether VP16+ and bICP0+ neurons express the GR.
Initial studies tested whether commercially available anti-serum recognizes the bovine GR. Antiserum directed against GR (MR-20; Santa Cruz Biotechnology) recognize mouse, rat, and human proteins were used for this study. Western blots revealed that the GR antisera specifically recognized a protein with an approximate molecular weight of 90 kd in bovine kidney cells (CRIB) (Figure 5A, lane B) and mouse cells (lane M), which is the expected size of the GR (13, 63). Confocal microscopy demonstrated that the antisera recognized a nuclear protein in CRIB cells following treatment with DEX (Figure 5B), which occurs when the GR binds DEX. Prior to DEX treatment, the GR was localized throughout CRIB cells. The GR antisera also recognized a subset of TG neurons after treatment with DEX for 3 hours (Figure 5C). In general, the signal localized to the nucleus suggesting the GR was activated after DEX treatment (denoted by arrows). In uninfected bovine TG, the signal was generally disperse and not readily detected when compared to TG sections prepared from calves after DEX treatment. The signal appeared to be more concentrated in few neurons from mock-infected calves (denoted by arrows), which may have been the result of stress during transportation of calves to the necropsy room prior to euthanasia.

Consecutive sections were prepared from calves treated with DEX and studies performed to determine if the GR was expressed in sensory neurons expressing bICP0 or VP16. At 1.5 hours after DEX treatment, GR+ and bICP0+ neurons were detected in certain areas of TG sections (Figure 6A; double positive neurons denoted by arrows). We detected 71 GR+ and bICP0+ neurons: but only 2 that were GR+ but not bICP0+ and no bICP0+ neurons that were GR negative. VP16+ and GR+ neurons were also readily detected at 1.5 hours after DEX treatment (Figure 6B; double positive neurons denoted by arrows). GR+ neurons that were not
VP16+ were occasionally detected at 1.5 hours after DEX treatment (Figure 6B; neurons denoted by closed circles). Seventy-two GR+ and VP16+ neurons were detected; but only two were identified in the same field that were just GR+. At 1.5 hours after DEX treatment, no bICP0+ or VP16+ neurons were detected that were GR negative.

DISCUSSION

In this study, two viral proteins that regulate transcription (VP16 and bICP0) were readily detected in sensory neurons within 1.5 hours after reactivation from latency was initiated with the synthetic corticosteroid DEX. These two viral proteins were frequently detected in the same neuron. In addition, neurons that expressed bICP0 or VP16 also frequently expressed the GR. Conversely, fewer neurons appeared to express the late proteins, gC and gD, before or after DEX treatment. Although one could argue that we were unable to detect gC or gD in TG neurons because the antibodies were not as good as the VP16 or bICP0 antibodies used, the gC and gD antibodies clearly recognized a viral specific protein in formalin fixed cells prepared from productively infected bovine cells. Furthermore, the gC and gD antibodies specifically recognized abundant levels of a viral specific protein during productive infection in a Western Blot.

In general, stress increases corticosteroid levels, and activates the GR, which we predict is a molecular switch that can consistently initiate the escape from latency (see Figure 7 for a schematic of the events that are necessary for reactivation from latency). During the escape from latency, several key events occur that result in lytic cycle viral gene expression. First, specific DEX-induced cellular transcription factors stimulate certain viral promoters and productive infection (80). Secondly, lytic cycle viral gene expression, including bICP0 and VP16, may also
be stimulated directly by an activated GR. Thirdly, DEX represses expression of LR (latency related) gene products (30, 60, 67), which is important because certain LR gene products can inhibit productive infection (2, 14, 30, 79). Finally, apoptosis of T cells in TG that persist during latency occurs following DEX treatment of latently infected calves (73), which may enhance the incidence of reactivation from latency because CD8+ T cells are proposed to maintain latency (7, 38, 40, 43, 44). We predict that the pleiotropic effects of increased corticosteroids levels and activation of the GR lead to the escape from latency.

DEX, as with other natural corticosteroids, specifically bind the GR and mineralocorticoid receptor (MR) (reviewed in (13)) resulting in nuclear localization of the GR or MR (58). Nuclear GR or MR dimers stimulate transcription by binding consensus glucocorticoid response elements {GRE; 5′-GGTACANNTGGTCT-3′} (16, 72). A GR or MR monomer can also stimulate transcription by binding to a GR ½ binding site, TGTTCCT or GGTACA, reviewed in (63). Although it is possible that MR+ TG neurons promote reactivation from latency, we were unable to identify a commercially available antibody that recognizes the MR receptor in bovine cells (data not shown). Within 5 minutes of glucocorticoid treatment, GR enters the nucleus, binds to glucocorticoid response elements, and induces rapid changes in chromatin conformation and transcriptional activation, reviewed in (5, 9). The BHV-1 genome contains 58 GR ½ binding sites in 24 BHV-1 promoters (data not shown) and a previous study demonstrated that DEX stimulates the bICP0 early promoter in transient transfection assays (81). It will be of interest to directly test whether viral promoters are actually bound by a GR during the escape from latency.
In a small subset of latently infected neurons, DEX induced expression of bICP0 and/or VP16 is predicted to stimulate lytic cycle viral gene expression, productive infection, and production of infectious virus in a small subset of latently infected neurons (Figure 7). Most neurons that escape latency do not produce infectious virus because they lack factors necessary for productive infection; consequently these neurons re-establish latency (60). This study suggested that expression of late genes is one bottleneck that must be overcome to produce infectious virus. It may not be necessary to produce high levels of late proteins; however expression of all late proteins necessary for producing infectious viruses would appear to be necessary. Several sub-types of sensory neurons exist in TG (82), and certain subtypes of TG neurons are more permissive for HSV-1 and HSV-2 productive infection (1, 48, 82). Consequently, we predict that only certain neuronal subtypes, which are latently infected, can escape latency and have the necessary factors to activate expression of viral genes necessary to produce infectious virus. An unanswered question that is raised as a result of these studies is: what is the threshold of corticosteroids that must be achieved to induce successful reactivation from latency? Every day mammals encounter increases in corticosteroids, but those that are latently infected with their respective alpha-herpesvirinae subfamily member do not always successfully reactivate from latency, as judged by shedding of infectious virus.

Published studies concluded that the normal cascade of viral gene expression in cultured cells is different than what occurs during the escape from latency. For example, a late viral promoter (gC) is trans-activated by DEX inducible cellular transcription factors (SPDEF and Slug) (80) and Notch1 or Notch3 (79). Furthermore, the bICP0 E promoter, but not the bICP0 and bICP4 IE promoter (IEtu1), is activated during reactivation from latency (81). Finally, this
study demonstrated that VP16 (a late viral gene), but not gC or gD, is detected in TG neurons within 1.5 hours after latently infected calves are treated with DEX. Several studies also suggest that the normal cascade of HSV-1 gene expression is different during reactivation from latency. For example, E gene expression and DNA replication is proposed to occur prior to IE gene expression (42, 52, 57, 68). Another study concluded that expression of a late HSV-1 gene (VP16) promotes the exit from latency (70). During explant induced reactivation from latency, viral gene expression was reported to be “disorganized” (10, 11). It does not appear that BHV-1 reactivation from latency is a random process in cattle because we consistently detected VP16 and bICP0, but not gC or gD protein expression in TG neurons within 1.5 hours after DEX treatment.

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Figure Legends

Figure 1. Analysis of BHV-1 protein expression in productively infected cells.

Panel A: Western blot analysis comparing the expression of BHV-1 genes following infection of bovine CRIB cells. Monolayers of CRIB cells were mock infected or infected with BHV-1 (moi=2) in the presence or absence of 400 μg/ml of phosphonoacetic acid (PAA). At the designated times after infection (hours), whole cell lysate was prepared. A total of 50 μg of protein was separated in an 8% SDS-PAGE gel and viral proteins were detected by Western blot analysis using antibodies specific for the designed BHV-1 genes. β-actin protein levels were analyzed in the respective samples as a loading control. The position of molecular weight markers for the respective panels is denoted. Molecular weight mass of the markers is shown on the right (kDa). The approximate molecular weight (kDa) of bICP0 is 97, VP16 is 65, gC is 95, and gD is 77.

Panel B: Subcellular localization of viral proteins in productively infected cells. Confocal microscopy was performed to detect BHV-1 proteins in CIRB cells infected with BHV-1
(moi=2) for 20 hours and then fixed with formalin. Mock infected CIRB cells served as a control. Fixed cells were incubated with antibodies specific for the designed viral genes. An Alexa Fluor 488 goat anti-rabbit secondary antibody (green) or Alexa Fluor 633 goat anti-mouse antibody (red) was used to detect viral protein expression. Images were obtained with Bio-Rad confocal laser-scanning microscope (MRC-1024ES) with excitation/emission at 488/520 nm. Nuclear DNA (blue) was stained with DAPI (4′,6-diamidino-2-phenylindole). Merged images are shown and they are representative of three independent experiments.

Figure 2. bICP0 and VP16 are expressed in sensory neurons following DEX treatment to induce reactivation from latency.

IHC was performed using the bICP0 or VP16 antibody as described in the materials and methods. Arrows denote neurons that were recognized by the respective antibody. Magnification is approximately 400x.

Figure 3. The glycoproteins gC and gD are weakly expressed in sensory neurons following DEX treatment to induce reactivation from latency.

IHC was performed using the gC or gD specific antibody as described in the materials and methods using TG sections from latently infected calves (latency), 1.5, 3, or 6 hours after DEX treatment. Arrows denote neurons that were recognized by the respective antibody. Magnification is 400x.

Figure 4. bICP0 and VP16 are frequently expressed in the same neuron following DEX induced reactivation from latency.
Panel A: Consecutive sections were cut from formalin fixed paraffin embedded TG from calves latently infected with BHV-1 that were treated with DEX for 3 hours. IHC was performed using the bICP0 antibody on one section and the VP16 antibody was used to stain the consecutive section. Arrows denote neurons that were recognized by the respective antibody. Magnification is 400x.

Panel B: The number of bICP0+ and VP16+ neurons (double positive) is shown for 1.5, 3, and 6 hours after DEX treatment (black columns). The number of VP16+ neurons (white columns) or bICP0+ (grey columns) are also shown at the designated times after DEX induced reactivation from latency.

Figure 5. Detection of GR in bovine kidney cells (CRIB)

Panel A: Bovine cells (CRIB) or mouse neuroblastoma cells (Neuro-2A) (30 ug protein/lane) were electrophoresed in a SDS PAGE gel, and Western Blot analysis performed using the commercially available GR antibody (Santa Cruz: H-300). Lanes denoted B were cell lysate prepared from bovine kidney cells (CRIB) and lanes denoted M were cell lysate from mouse neuroblastoma cells (Neuro-2A).

Panel B: Cells were treated with DEX and immunofluorescence performed as described in the materials and methods. As controls, certain cultures were not treated with DEX. Localization of the GR was examined by immunofluorescence. Nuclear DNA was identified by DAPI staining.

Panel C: IHC was performed with TG samples from uninfected calves and three hours after DEX treatment using procedures described in the materials and methods. Arrows denote the GR+ neurons. Magnification is approximately 400x.
Figure 6. GR+ neurons frequently express bICP0 or VP16 during the escape from latency.

Consecutive sections were cut from formalin fixed paraffin embedded TG from calves latently infected with BHV-1 that were treated with DEX for 1.5 hours. IHC was performed using the bICP0 antibody on one section and the GR antibody on the adjacent section (Panel A). IHC was performed using the VP16 antibody on one section and the GR antibody on the adjacent section (Panel B). Arrows denote neurons that were recognized by the respective antibody. Closed circles denote GR+ neurons that were VP16 negative. Magnification is 400x.

Figure 7. Putative steps that occur during DEX induced escape from latency.

For details, see the discussion.
Figure 7

Stress (DEX)

GR activated

Viral gene expression stimulated by DEX-induced cellular transcription factors and GR
Repressed LR gene expression
Infiltrating lymphocytes undergo apoptosis

Escape from latency (many neurons)

Re-establish latency
No virus shedding (Most neurons)

All late genes expressed
Infectious virus produced (Rare neurons)