Regulation of Viral Intermediate Gene Expression by the Vaccinia Virus B1 Protein Kinase†

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The B1 gene of vaccinia virus encodes a serine/threonine protein kinase that is expressed early after infection. Under nonpermissive conditions, temperature-sensitive mutants (ts2 and ts25) that map to B1 fail to efficiently replicate viral DNA. Our goal was to extend studies on the function of B1 by determining if the kinase is required for intermediate or late gene expression, two events that ordinarily depend on viral DNA replication. First, we established that early viral gene expression occurred at the nonpermissive temperature. By using a transfection procedure that circumvents the viral DNA replication requirement, we found that reporter genes regulated by an intermediate promoter were transcribed only under conditions permissive for expression of active B1. To assay late gene expression, the T7 RNA polymerase gene was inserted into the genome of ts25 to form ts25/T7. A DNA replication-independent late gene transcription system was established by cotransfecting plasmids containing T7 promoter-driven late gene transcription factors and a late promoter reporter gene into ts25/T7-infected cells. Late genes, unlike intermediate genes, were expressed at the nonpermissive temperature. Last, we showed that overexpression of B1 stimulated intermediate but inhibited late gene expression in cells infected with wild-type virus.

DNA viruses have evolved regulatory systems in which early and late stage genes are transcribed from parental and replicated genomes, respectively. Poxviruses have added some unique features to this general strategy. The enzymes needed for transcription of early genes are expressed late in infection, packaged within progeny virions, and activated when the core is released into the cytoplasm following infection (26). In addition, there are two postreplicative stages of poxvirus gene expression, intermediate and late, rather than just one. All of the viral factors required for DNA replication and expression of intermediate genes are products of early genes, whereas the viral factors needed for late gene expression are products of both early and intermediate genes. Thus, the poxvirus life cycle occurs in a temporally ordered sequence: early gene expression → viral DNA replication → intermediate gene expression → late gene expression. The extent to which vaccinia virus proteins interact with one another and/or with cellular proteins to link viral DNA replication with intermediate and late gene expression is not understood.

Studies on the regulation of intracellular processes involved in eukaryotic nucleic acid metabolism have pointed to protein phosphorylation as playing a key role. Poxviruses have acquired analogs of a number of host cell genes to maintain their autonomy from the cell, including two serine/threonine protein kinases (B1 and F10) and a dual-specificity protein phosphatase (H1). The F10 kinase (22) and H1 phosphatase (15) are expressed late in infection, are incorporated into virions, and play major roles in virion morphogenesis (11, 24, 38). Interestingly, the H1 phosphatase has been implicated in early gene transcription (24). Mutant virus particles devoid of H1 phosphatase were unable to transcribe early genes either in vivo or in vitro. This finding led us to investigate the role that phosphorylation might play in the transcription of the other gene classes. The B1 kinase is expressed exclusively early in infection; it localizes to viral replication factories and appears to be a minor component of the virion (3, 23, 34). The B1 kinase associates with (25) and phosphorylates (5, 6) viral late gene transcription factor 4 (VLTF-4; viral protein H5) and two ribosomal proteins in vitro. Temperature-sensitive (ts) vaccinia virus mutants (ts2 and ts25) that map to the B1 gene have a defect in viral DNA replication (9, 28) and consequently would not be expected to express intermediate or late genes. To analyze intermediate and late gene expression, we used a transfection-based assay that mimics the requirement for viral DNA replication (18, 37). Our results show that B1 is essential for the transcription of viral intermediate but not late genes.

MATERIALS AND METHODS

Cells and viruses. African green monkey kidney (BS-C-1) and murine fibroblast (L929) cells were maintained in minimal essential medium (MEM; Life Technologies) supplemented with 10% fetal bovine serum (Life Technologies). The ts mutants ts2 and ts25 were obtained from Richard Condit (8, 9) and grown and titered on BS-C-1 cells. For virological assays, incubations were at permissive (32°C) and nonpermissive (39.5°C) temperatures. Cytosine arabinoside (AraC; Sigma) was used at 44 μg/ml to block viral DNA replication in certain experiments.

Plasmid constructions and reporter gene assays. Three reporter genes were used in this study: p30/300, pGSR-CAT, and p1KCAT. The construction of p30/300, a plasmid that contains 300 bp of DNA upstream of the vaccinia virus intermediate GSR gene open reading frame (ORF) fused to the lacZ ORF, has been described elsewhere (1). pGSR-CAT was constructed by digesting p30/300 with BamHI and HindIII, completely removing the lacZ ORF, and subsequently

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† This work was dedicated to the memory of Roskey Jennings, whose character was an inspiration to the many people who passed through the Laboratory of Viral Diseases.
inserting a compatible fragment containing the chloramphenicol acetyltrans-
ferase (CAT) ORF in its place. The resultant plasmid (pGSR-CAT) retains the
same 300 bp of GSR upstream sequences as p30/300. A late gene reporter
plasmid, p11K-CAT, was constructed in two steps. First, pGSR-CAT was di-
gested with BamHI, and a synthetic DNA duplex containing 65 bp of sequence
upstream of the 1K ORF was inserted so that only the upstream BamHI site was
regenerated. Second, the resultant plasmid was digested with BamHI and Sall to
completely remove the GSR regulatory sequences. The final plasmid (p11K-
CAT) contains the CAT ORF under the regulation of the 1K late promoter. A
T7 promoter-B1 expression plasmid (pT7-B1) was constructed by PCR amplifi-
cation of the wild-type B1 ORF from vaccinia virus strain WR DNA and inser-
tion of this fragment into the EcoRI and BamHI sites of pSG5 (14). The same
vector expressing a nonsense mutant (pT72A1B1) of the B1 gene was engineered
by inserting a synthetic adapter (5'-AATTCTAGCTAGCTAG-3') into the MscI site
in the B1 ORF. Expression of pT7-B1 would result in a truncated B1 gene
product terminating at the 11th amino acid.

Transfections were carried out with Qiagen column-purified DNAs using the
DOTAP transfection reagent (Boehringer Mannheim) and OptiMEM (Life
Technologies). Typically, 5 μg of covalently closed, circular, supercoiled DNA
was mixed with 15 μl of DOTAP per transfection. In transfection-infection
experiments, cells (approximately 2 × 10⁹ to 3 × 10⁹ per dish) were transducted
3 h prior to infection. The temperature of transfections was kept constant at
37°C in a humidified atmosphere. Inocula were prepared by lysing the cells
at 30°C for 30 min. Extracts were prepared by washing cells in ice-cold phosphate-buffered saline
twice and freeze-thawing three times in 500 μl of 1× reporter lysis buffer
(Promega). Typically, 1 to 10 μl of cell extract was analyzed to remain within the
linear range of enzyme assays. β-Galactosidase assays were conducted as spec-
ified by the manufacturer (Promega) and quantitated using an EL 312E Bio-
Kinetics Reader (Bio-Tek Instruments). CAT assays were done using a fluores-
cently labeled chloramphenicol substrate (FAST CAT yellow; Molecular Probes)
(17, 39). Reaction products were separated by thin-layer chromatography, ana-
yzed on a FluorImager (Molecular Dynamics), and quantitated with Image-
Quant software (Molecular Dynamics).

**Protein analyses.** Radiolmmuno precipitations were conducted by labeling
cells for 30 min with 50 Ci of [35S]methionine (Amersham) per ml in MEM
lacking methionine. Extracts were prepared by washing cells with ice-cold phos-
phate-buffered saline, suspending the cells in 1× radiolmmuno precipita-
tion assay (RIPA) buffer (50 mM Tris-HCl [pH 7.2], 150 mM NaCl, 1% NP-40), and
clarifying the lysate by microcentrifugation for 5 min. Subsequently, anti-vaccinia
virus antibody (BioGenes) was added at 1:500 to the clarified supernatant,
and mixtures were rocked for 4 to 6 h at 4°C. Immune complexes were captured with
50 μl of 20% protein A-Sepharose CL-4B (Pharmacia), and the beads were
washed three times with 1× RIPA buffer containing 0.1% sodium dodecyl sulfate
(SDS). Immune complexes were boiled in SDS-polyacrylamide gel electrophore-
sis (PAGE) loading buffer (0.05 M Tris, 4% SDS, 4% β-mercaptoethanol, 10% glycerol, 0.1% bromophenol blue) for 5 min and microcentrifuged for 30 s. The
solubilized proteins were then resolved on 5 to 20% polyacrylamide gels (21).
Gels were dried and analyzed by autoradiography.

Western blotting was conducted by transferring proteins onto polyvinylidene
difluoride membranes (Millipore), incubating in blocking buffer (0.89 M Tris,
0.02 M EDTA, 1% NP-40, 1% nonfat dry milk), and probing with either anti-vaccinia virus (1:1,000; BioGenes), anti-β-galactosidase (1:500; 5'-
3', Inc.), anti-T7-β-galactosidase (1:500), or anti-T7 polymerase (1:1,500) in blocking
buffer. Immune complexes were detected with an alkaline phosphatase-conju-
gated goat anti-rabbit immunoglobulin G antibody (1:1,500; Life Technologies)
and the Western Blue reagent (Promega).

**RNAse protection assay.** Total RNA was purified from approximately 4 × 10⁶
to 5 × 10⁶ cells using the TRizol total RNA isolation reagent (Life Technolo-
gies), treated with DNase I (5 U for 15 min at 37°C; Ambion), precipitated with
LiCl, and resuspended in H₂O. A T7 promoter-containing template for ribo-
probe synthesis was made by PCR using two oligonucleotides (5'-GTCAAAAAA-
TTGTTAGAACGAC-3' and 5'-TAATGACTAGCTAGCTAGGCTATTAG
AGCAAC-3'; T7 promoter is underlined) and pGSR-CAT as the DNA
template. The 161-nucleotide (nt) antisense riboprobe derived from pGSR-CAT
polymerase transcription of this template should protect the first 96 nt of the
GSR-CAT mRNA. Approximately, 600 pg of uniformly radiolabeled riboprobe
was mixed with 10 μg of total RNA and assayed using an RPA II kit essentially
as described by the manufacturer (Ambion). Protected segments were separated on
5% polyacrylamide—1× Tris-borate EDTA TBE—8 M urea gels and visualized
by autoradiography.
with BS-C-1 cells, which are used more commonly than L929 cells for studies with vaccinia virus. Plasmid p30/300, containing an intermediate gene promoter fused to the lacZ gene, was transfected into cells that were then infected with WR or ts25. As in L929 cells, β-galactosidase synthesis was higher in BS-C-1 cells infected with ts25 than with WR at 32°C but was nearly undetectable at 39.5°C (Fig. 3B and C). We carried out parallel experiments with L929 and BS-C-1 cells in which AraC was added to the medium at the start of infection. In this way, there would be a DNA replication block under all conditions and with both wild-type and mutant viruses. We found that reporter gene expression occurred in the presence of AraC at 32°C (Fig. 3A and C), although the levels were reduced in cells infected with either WR or ts25. At 39.5°C, however, reporter gene expression was specifically blocked in cells infected with ts25 (Fig. 3A and C). We concluded, therefore, that there was a ts defect in intermediate gene expression independent of effects on viral DNA replication.

Inhibition of intermediate gene expression occurred at the transcriptional level. Because the synthesis of early proteins appeared to be unaffected by the ts mutation of B1, i.e., there was no global effect on translation, we suspected that the inhibition of intermediate gene expression occurred at the transcriptional level. To investigate this, we selected CAT as the reporter gene because its mRNA is only 600 nt, compared to 3,000 nt for lacZ mRNA. We considered that if the defect were in RNA elongation or degradation, then synthesis of CAT might be less severely affected than that of β-galactosidase. Such experiments, however, indicated that CAT synthesis was severely inhibited at 39.5°C in ts25-infected L929 cells (Fig. 4A).

An RNase protection assay was performed to determine more directly whether the block in intermediate gene expression occurred at the level of transcription. RNA was purified from BS-C-1 cells that had been transfected with pG8R-CAT.
and infected with either ts25 or WR. We used BS-C-1 instead of L929 cells in this experiment because of the better integrity of RNAs isolated from the former cell type. The riboprobe was designed to quantify steady-state levels of CAT mRNA and also to determine if the mRNAs initiated at the correct position relative to the G8R promoter. A discrete RNA product of predicted size was detected in cells infected with WR or ts25 at 32°C or with WR at 39°C (Fig. 4B). Little or no G8RCAT mRNA was detected in cells infected with ts25 at the nonpermissive temperature, suggesting that in the absence of B1, intermediate mRNAs are either not transcribed or have very short half-lives (Fig. 4B).

Construction of a ts25 mutant that expresses bacteriophage T7 RNA polymerase. We decided to make recombinant virus ts25T7, which retains the ts25 mutation in the B1 gene and expresses the T7 RNA polymerase gene regulated by an early promoter. This virus would allow us to further evaluate the block in intermediate transcription by determining (i) whether a reporter gene regulated by a T7 promoter would be expressed at 39.5°C, (ii) if coexpression of the wild-type B1 gene would reverse the ts phenotype, and (iii) whether expression of late genes was affected at the nonpermissive temperature. Since the assays would be done at 39.5°C and in the presence of AraC, we first needed to determine whether sufficient T7 RNA polymerase would be expressed by ts25T7 under these conditions. A Western blot analysis (Fig. 5A) indicated that at least as much T7 RNA polymerase was synthesized by ts25T7 as by vTF7.3, a well-characterized vaccinia virus WR recombinant containing the T7 RNA polymerase gene (13). At least two cross-reactive species of lower molecular weight were detected; because they are not present in the wild-type virus-infected lysate, we posit that they are cleavage and/or breakdown products of T7 polymerase. As expected, since the T7 gene is under an early/late promoter, blocking late transcription with AraC resulted in less synthesis of full-length as well as truncated T7 proteins. We also needed to establish that ts25T7 could function as a T7 RNA polymerase expression vector in transient assays and retained the ts defect. Cells were transfected with either pG8RCAT or pT7CAT and infected with ts25T7 or vTF7.3. The transfection experiment shown in Fig. 5B indicated that ts25T7 allowed the expression of CAT regulated by an intermediate promoter at 32°C but not at 39.5°C. In contrast, vTF7.3 allowed the expression of CAT at either temperature (Fig. 5B). Significantly, the CAT gene regulated by a T7 promoter was expressed to similar levels at either

FIG. 3. The block in intermediate gene expression is independent of DNA replication. BS-C-1 or L929 cells were transfected with p30/300 and subsequently infected with 5 PFU of WR or ts25 per cell in the presence of absence of AraC at the indicated temperature. Lysates were prepared at 24 h after infection and assayed for β-galactosidase activity (optical density at 495 nm [OD₄₀₅]). Experiments were done in triplicate, and standard deviations are indicated by the shaded areas at the tops of the bars. M, mock infected.
temperature by ts25T7 or VTF7.3 (Fig. 5C). Thus, the block in transcription was specific for an intermediate gene promoter.

The next series of experiments were carried out to determine whether the ts25 defect could be overcome by coexpression of a wild-type copy of the B1 gene. Plasmid pT7-B1, containing the B1 ORF adjacent to a T7 promoter, was constructed and shown to program the synthesis of a 34-kDa protein that bound to anti-B1 serum in a coupled reticulocyte transcription-translation system (data not shown). Cells were then transfected with pG8R-CAT and pT7-B1 (containing an intact B1 ORF under T7 promoter control) or pT7ΔB1 (containing a truncated B1 ORF under T7 promoter control) and then infected with ts25T7 or vTF7.3. Transfection of pT7-B1 enhanced intermediate gene expression in cells infected with vTF7.3 at 32 or 39.5°C, whereas transfection with pT7ΔB1 did not (Fig. 6). In contrast, intermediate gene expression was enhanced by transfection of pT7-B1 in cells that were infected with ts25T7 at 32°C but not at 39.5°C (Fig. 6). Apparently, trans expression of B1 enhanced intermediate gene expression in the presence of limiting amounts of active B1 kinase but not in the presence of mutated B1 kinase at the nonpermissive temperature. Either the mutated B1 protein had a dominant negative effect or active kinase was not expressed early enough from the transfected plasmid to prevent an irreversible inhibition of intermediate gene transcription.

Analysis of late gene expression. To extend our analysis to the role of the B1 kinase in late gene expression, we adapted a transfection protocol devised by Keck et al. (18). Late promoters, like intermediate promoters, can regulate transcription of a reporter gene in a transfected plasmid during an AraC block. An additional requirement, however, is that the genes encoding VLTF-1, -2, and -3 (viral genes A1, A2, and G8, respectively) must be cotransfected with the late promoter reporter plasmid. However, because A1, A2, and G8 are of the intermediate gene class, they would be silent during nonpermissive infection of cells with ts25 or ts25T7, and consequently the late transcription factors would not be made. To overcome this problem, we transfected plasmids containing the A1, A2, and G8 ORFs under the control of T7 promoters (18). Late transcription was monitored by cotransfecting a reporter plasmid (p11K-CAT) containing the CAT gene under the control of the viral 11K late gene promoter. The cells were subsequently infected with ts25T7 to provide the T7 RNA polymerase to drive the expression of the three VLTF genes. Under these conditions, CAT synthesis occurred either at 32 or 39.5°C in the presence of AraC, indicating that B1 was not specifically required for transcription of late genes (Fig. 7). Control experiments verified that there was low CAT expression in the absence of transfected VLTF plasmids when cells were infected with ts25T7 at either temperature in the presence of AraC or at 39.5°C in the absence of AraC (Fig. 7).

Knowing that B1 was not essential for late gene expression, we determined the effects of overexpression. Cells were cotransfected with pT7-B1 or pT7ΔB1 and p11K-CAT and subsequently infected with VTF7.3 containing a wild-type B1 gene and T7 RNA polymerase (Fig. 8). The results indicate that trans expression of B1 inhibited late gene expression by nearly 85%.

**DISCUSSION**

Several studies demonstrated that the B1 kinase plays an important role in viral DNA replication temperature (8, 9, 28).
However, in some cell lines this effect seemed insufficient to entirely account for the reduction in virus titer and plaque formation. For example, at 39.5°C, ts25 directed the synthesis of up to 60% of control levels of DNA but only 15% of control levels of viable progeny and no distinct plaques in BS-C-40 cells (28). This is in contrast to other studies conducted on ts vaccinia virus mutants, where a decrease in viral DNA synthesis resulted in a concomitant reduction of comparable magnitude in viral titers (33). Therefore, we considered that B1 might have another role downstream of viral DNA replication. In this report, we provide evidence that intermediate gene transcription is blocked at the nonpermissive temperature and, unlike the defect in DNA replication, is similarly restricted in mouse and monkey cells.

We first showed that early gene expression was unaltered in cells infected with ts25 at the nonpermissive temperature, whereas late gene expression was completely blocked. Because this defect would result from inhibition of viral DNA synthesis, we relied on transfection experiments to specifically analyze the expression of intermediate and late genes at the nonpermissive temperature. Although a reporter gene regulated by an intermediate promoter was expressed in cells infected with wild-type vaccinia virus even in the presence of a potent inhibitor of DNA synthesis, expression did not occur at 39.5°C in cells infected with ts25. This result provided evidence that B1 was required for intermediate gene expression independent of any effect on DNA replication. Furthermore, the block occurred at the level of transcription. By constructing a recombinant virus that contained the ts25 defect and a T7 RNA polymerase gene, we demonstrated that a reporter gene regulated by a T7 promoter was expressed under nonpermissive conditions, ruling out a global defect in plasmid-based transcription. Moreover, by cotransfecting the intermediate genes encoding late transcription factors under T7 promoters, we could demonstrate that B1 was not required for transcription of a reporter plasmid with a late promoter. Transfection experiments further showed that overexpression of wild-type B1 stimulated intermediate gene expression from 1.5- to 3.5-fold but severely inhibited late gene expression in cells expressing functional B1. Failed attempts to reverse the block in intermediate transcription by expression of wild-type B1 at 39.5°C suggested that the ts B1 protein may be trans-dominant to the wild-type protein. Alternatively, if the timing of B1 synthesis is crucial, then expression from the T7 promoter, which depends on prior synthesis of T7 RNA polymerase, may occur too late.

We are uncertain as to how the B1 kinase is involved in intermediate gene expression. In vivo studies have shown that intermediate genes are transiently expressed from 1 to 4 h after infection (2). Biochemical analyses have indicated that at least six proteins (viral intermediate transcription factors [VITFs]) are required for transcription; these include the multisubunit...
viral RNA polymerase (35), viral capping enzyme (16, 36), VITF-1 (viral protein E4 [29]), VITF-3 (a complex of the viral A8 and A23 proteins [31]), and an unidentified cellular factor termed VITF-2 (30). An in vitro transcription system consisting solely of recombinant proteins has not been derived; thus, the possibility of additional factors or posttranslational modification of known ones remains. For example, the B1 kinase may be associated with a partially purified component of the transcription system or may have phosphorylated a viral or cellular factor prior to purification. Another possibility is that the viral H5 protein is involved in intermediate transcription. This protein, identified as VLTFT-4 (19), is the only VLTFT synthesized prior to DNA replication (20). The fact that H5 is phosphorylated in vitro by the B1 protein kinase (5) and is underphosphorylated by a B1 ts mutant at the nonpermissive temperature (4) makes H5 an attractive candidate for further investigation.

Additional studies indicate that the H5 protein interacts with viral proteins encoded by the A18 and G2 genes, which are involved in transcription elongation (7), and the A20 gene (25), involved in DNA replication (12). The situation is even more complex because a recent report of charge-to-alanine mutagenesis of the H5 ORF produced a ts mutant that has a block in viral morphogenesis with no apparent defect in either viral DNA or protein synthesis (10). Whether the B1 kinase is directly involved in this additional role of the H5 protein is unknown.

The B1 gene is conserved among many but not all poxviruses. Molluscum contagiosum virus encodes homologs of the F10 protein kinase and the H1 phosphatase but does not contain a B1 homolog (32). One explanation for its absence from the molluscum contagiosum virus genome is that it may be compensated by a host cell protein kinase. In this regard, there are two human putative serine/threonine protein kinases, VRK1 and VRK2, with a high degree of sequence similarity to the B1 kinase (27).

Taking into consideration the complexity of the cellular pathways that are regulated by protein phosphorylation and dephosphorylation, it is not surprising that the B1 protein kinase has multiple effects on the vaccinia virus life cycle. Clearly, more work is required to identify all of the viral and cellular factors that are regulated by the B1 protein kinase.

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