

## NOTES

# Herpes Simplex Virus Type 1 Thymidine Kinase Sequence Fused to the *lacZ* Gene Increases Levels of $\beta$ -Galactosidase Activity per Genome of High-Capacity but Not First-Generation Adenoviral Vectors In Vitro and In Vivo<sup>∇†</sup>

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**Increased transgene expression per vector genome is an important goal in the optimization of viral vectors for gene therapy. Herein we demonstrate that herpes simplex virus type 1 (HSV1) thymidine kinase (TK) gene sequences (1,131 bp) fused to the 3' end of *lacZ* increase transgene expression from high-capacity adenoviral vectors (HCAd), but not from first-generation (Ad) vectors. The woodchuck hepatitis virus posttranscriptional regulatory element (WPRE), in contrast, increased transgene expression levels from Ad but not HCAd vectors. The differential activity of the HSV1 TK gene and WPRE sequences was detected both in vitro and in vivo and suggests potentially different mechanisms of action or the interaction of these elements with vector genomic sequences.**

Adenoviral vectors are effective vectors for gene transfer and gene therapy. However, adenoviral vectors induce dose-dependent innate and adaptive immune responses (24, 33, 34, 41, 46, 50, 51). Decreasing the total dose of therapeutically effective viral vector should result in safer and longer-term gene transfer. To do so, it is desirable to increase the levels of transgene expression per vector genome (vg). This can be achieved through the use of genetic sequences that either increase the number of transcript copies (e.g., stronger promoters) or stabilize the mRNA (e.g., the woodchuck hepatitis virus posttranscriptional regulatory element [WPRE]) (3, 16, 21, 22, 26, 29, 30, 32, 40, 48, 52).

Previously, we reported that intracranial delivery of a first-generation adenoviral vector encoding herpes simplex virus type 1 (HSV1) thymidine kinase (Ad-TK) resulted in unexpectedly high levels of gene expression, widespread distribution, and strong biological activity, suggesting that HSV1 TK has the capacity to upregulate transgene expression (11). Moreover, in the same report we demonstrate that an adenoviral vector with a 60-bp deletion in the coding sequence for the 5' region of TK elicits decreased levels of transgene ex-

pression and biological activity compared to the vector encoding wild-type HSV1 TK (8, 11).

To determine if HSV1 TK gene sequences could upregulate transgene expression when provided in *trans*, we coinjected Ad-TK mixed with an Ad vector expressing the  $\beta$ -galactosidase ( $\beta$ -Gal) reporter gene (49). Our results suggested that the HSV1 TK gene sequence provided in *trans* failed to increase transgene expression from the vectors expressing  $\beta$ -Gal. So far, the use of *cis*-acting HSV1 TK gene elements in viral vectors to increase transgene expression has not been investigated, although elements within the HSV1 TK gene have been shown to act as pre-mRNA processing enhancers to increase expression of intronless genes when tested in plasmids.

In the present work we investigated whether HSV1 TK gene sequences enhance transgene expression from either first-generation adenoviral vectors (Ad) or high-capacity, helper-dependent adenoviral vectors (HCAd) and compared the activity of these gene sequences to the activity of an established posttranscriptional regulatory sequence (i.e., WPRE). To do so, we engineered expression cassettes carrying the reporter transgene (*lacZ*) under the control of the powerful murine cytomegalovirus (mCMV) promoter with potential regulatory sequences linked to the 3' end of *lacZ*. The regulatory sequences tested were as follows: HSV1 TK gene (1,131 bp); the  $\Delta$ TK gene, a truncated form of the HSV1 TK gene (1,072 bp) with a 60-bp deletion downstream of the first initiation codon; and WPRE (594 bp), used as a control posttranscriptional regulatory element (3, 16, 30, 39,

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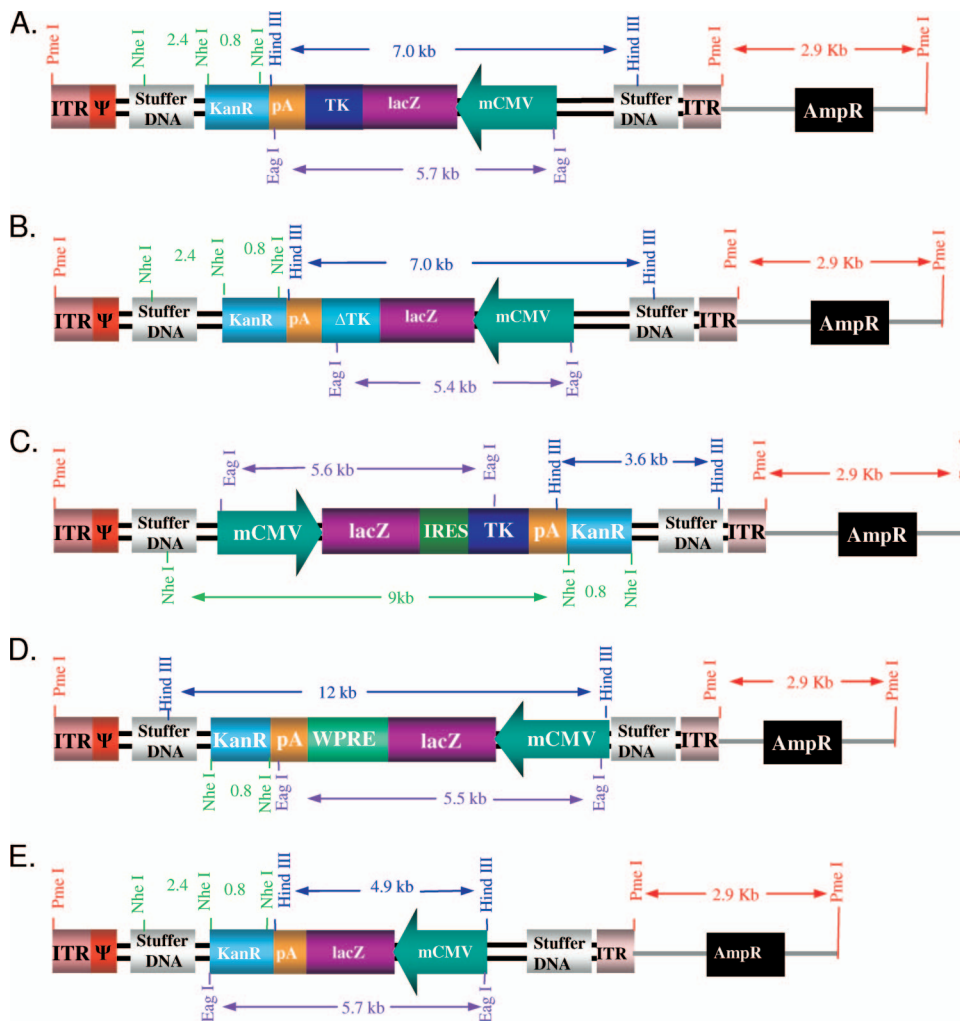


FIG. 1. Linear depiction of the HCAd vectors carrying the various mCMV-driven *lacZ* cassettes containing potential regulatory sequences. The constructs indicate the individual components and the orientations of the cassettes and their promoters. Relevant restriction sites are shown. (A) HCAd-mCMV.βgal.TK; (B) HCAd-mCMV.βgal.ΔTK; (C) HCAd-mCMV.βgal.IRES.TK; (D) HCAd-mCMV.βgal.WPRE; (E) HCAd-mCMV.βgal. Although the cassettes within HCAd are shown, the identical cassettes were utilized in first-generation Ad. For space considerations, these are not illustrated. See the supplemental material for details of vector construction. ITR, inverted terminal repeat; IRES, internal ribosome entry site.

48). A control construct of *lacZ* without a posttranscriptional sequence was also engineered.

The expression cassettes were utilized to generate the first-generation Ad vectors Ad-mCMV.βgal.TK, Ad-mCMV.βgal.ΔTK, Ad-mCMV.βgal.WPRE, and Ad-mCMV.βgal, respectively, using methodologies described by us previously (43). The same expression cassettes (Fig. 1) were also used to generate the high-capacity, helper-dependent adenoviral vectors HCAd-mCMV.βgal.TK, HCAd-mCMV.βgal.ΔTK, HCAd-mCMV.βgal.WPRE, and HCAd-mCMV.βgal using methodologies described by us previously (35, 37). We also generated a fifth HCAd vector, HCAd-mCMV.βgal.IRES.TK, designed to express both β-Gal and TK. Briefly, four of the vectors are completely new and have never been published before: HCAd-mCMV.βgal.ΔTK, HCAd-mCMV.βgal.IRES.TK, Ad-mCMV.βgal.TK, and Ad-mCMV.βgal.ΔTK. Three other vectors, HCAd-mCMV.βgal, HCAd-mCMV.βgal.TK, and HCAd-mCMV.βgal.WPRE, have been used before, but only to assess

genome levels in the development of a novel method for the quantitation and titration of Ad vectors (39), even though their biological activity was neither evaluated nor reported previously. The expression levels of only two vectors (Ad-mCMV.βgal and Ad-mCMV.βgal.WPRE) were described by our group in detail previously (3, 15), and these vectors are included here only as controls. Thus, in this paper we report four completely novel vectors never described before and, for the first time, the biological activity for seven vectors. A total of nine vectors were used in this work.

Integrity of the cassettes was confirmed by sequencing (data not shown). All vectors were titered in parallel for blue forming units (BFU) and vgs and were certified free of contaminating lipopolysaccharide and replication-competent adenovirus as described before (12, 39, 43). The vectors obtained were titered, and the values obtained are shown in Table 1.

To determine the optimal, nonsaturating vector dose, we generated a dose-response curve using the cell lines CNS1

TABLE 1. Titer of viral preparations by BFU and vgs

Vector	BFU/ml	vgs/ml
Ad-mCMV.βgal	$1.64 \times 10^{11}$	$2.21 \times 10^{12}$
Ad-mCMV.βgal.TK	$3.28 \times 10^{11}$	$1.60 \times 10^{12}$
Ad-mCMV.βgal.ΔTK	$1.32 \times 10^{12}$	$2.70 \times 10^{13}$
Ad-mCMV.βgal.WPRE	$5.12 \times 10^9$	$4.97 \times 10^{10}$
HCAAd-mCMV.βgal	$4.27 \times 10^9$	$3.68 \times 10^{11}$
HCAAd-mCMV.βgal.TK	$7.66 \times 10^9$	$6.96 \times 10^{10}$
HCAAd-mCMV.βgal.ΔTK	$1.01 \times 10^{11}$	$4.37 \times 10^{12}$
HCAAd-mCMV.βgal.IRES.TK	$2.46 \times 10^{11}$	$1.29 \times 10^{13}$
HCAAd-mCMV.βgal.WPRE	$2.44 \times 10^{11}$	$3.63 \times 10^{12}$

(Lewis rat glioma), GL26 (C57BL/6 mouse glioma), J3T (canine glioma), Cos7 (primate kidney, as a control), IN859 (human glioma), and U251 (human glioma) (data not shown) at multiplicities of infection (MOIs) ranging from 30 to 1,000 genomes per cell. Based on the linear range of expression for all vectors, an MOI of 300 vgs/cell was chosen for further detailed analysis. Expression of β-Gal and β-Gal activity were tested in all cell lines as described elsewhere (1, 3, 7, 47). For all subsequent data analysis, results were standardized and reported as β-Gal activity per vg. All data were analyzed by one-way analysis of variance (ANOVA) followed by the Tukey-Kramer multiple comparison test.

When carried within a first-generation Ad vector, the HSV1 TK gene sequences did not increase β-Gal activity per vg in any of the cell lines used for the study (Fig. 2, left). The enhancement provided by WPRE was similar to results reported by others (25, 36).

However, when contained within HCAAd vectors, the HSV1 TK gene sequence provided increases in β-Gal activity per genome that ranged from 2.9- to 9.3-fold. However, no increased β-Gal activity per vg was detected using the ΔTK gene sequence in HCAAds. The inability of the ΔTK gene sequence to enhance transgene expression from the HCAAd vector backbones could indicate that the 60 bp at the 5' extreme of the HSV1 TK gene is required for the enhanced expression provided. We also compared the β-Gal activity per vg mediated by infection with HCAAd-mCMV.βgal.IRES.TK to determine if the HSV1 TK gene sequence separated from the β-Gal transgene by an internal ribosome entry site or the HSV1 TK protein potentially expressed from an internal ribosome entry site could increase levels of transgene expression. With this vector, no increased β-Gal activity was observed.

Unlike increased transgene expression provided by WPRE when carried by first-generation Ad vectors, there was no effect of WPRE on transgene expression when it was carried by the HCAAd vector (Fig. 2, right). The same shuttle vector was used to generate both Ad and HCAAd vectors, and sequence analysis was used to confirm that the sequences of the WPREs in both cassettes were identical (data not shown). Although WPRE has recently been used by several groups in HCAAd vectors in conjunction with tissue-specific promoters (6, 18, 20), a direct comparison of transgene expression levels from HCAAds with and without the WPRE has not been performed. Our data indicate that, under our experimental conditions, and compared side by side, WPRE failed to increase transgene expression from HCAAd.

We further tested the effects of the HSV1 TK sequence on

transgene expression from HCAAd and Ad vectors in vivo. Adult female C57BL/6 mice with body weights between 18 and 25 g were used. Five mice per group ( $n = 5$ ) were injected with  $1.0 \times 10^6$  vgs of vectors that had shown increased transgene expression due to either the HSV1 TK gene or WPRE and their respective control vectors (i.e., HCAAd-mCMV.βgal, HCAAd-mCMV.βgal.TK, and HCAAd-mCMV.βgal.WPRE and Ad-mCMV.βgal, Ad-mCMV.βgal.TK, and Ad-mCMV.βgal.WPRE) into the striatum as described previously (43, 47). For immunohistochemistry, 5 days postinjection, animals were perfused with 100 ml oxygenated Tyrode's solution followed by a 4% paraformaldehyde solution. Brains were serially sectioned using an electronic Vibratome (Leica) to obtain 50-μm free-floating sections. Serial sections were immunoreacted using a rabbit polyclonal anti-β-Gal primary antibody (diluted 1:1,000; generated in our laboratory [2, 42–44]). A series of sections spanning the entire injection site (and containing β-Gal-expressing cells), separated from each other by 250 μm, were quantified by unbiased quantitative stereology using Microbrightfield software on a Zeiss upright microscope.

We obtained a 2.64-fold increase of β-Gal-expressing cells per vg in the brains of mice injected with Ad-mCMV.βgal.WPRE into the striatum, compared to the level obtained with the control vector Ad-mCMV.βgal (Fig. 3a and b, left). As shown in the in vitro experiments, the HSV1 TK gene sequence within Ad failed to increase transgene expression per vg in vivo. We found a 1.8-fold increase in the number of β-Gal-expressing cells per vg in the brains of mice stereotactically injected with HCAAd-mCMV.βgal.TK into the striatum, compared to the number in brains injected with the control vector HCAAd-mCMV.βgal (Fig. 3a and b, right); as in the in vitro experiments, the WPRE within HCAAd failed to increase transgene expression per vg in vivo.

This experiment was repeated, but injected brains were homogenized and β-Gal enzymatic activity was measured (Fig. 3c). Results from these experiments were comparable to those in which the transgene-expressing cells were counted, i.e., the WPRE increased β-Gal activity per genome from Ad and the HSV1 TK gene increased β-Gal activity per genome from HCAAd (Fig. 3c). Data were analyzed by one-way ANOVA followed by the Tukey-Kramer multiple comparison test.

In previous publications authors have characterized elements found within the HSV1 TK gene that could act as post-transcriptional enhancers to increase expression from intronless genes. In detail, Liu and Mertz described a 119-bp element from the HSV1 TK gene that can act in *cis* to provide expression to intronless genes when carried upstream of a desired gene; this was done using in vitro transfections of plasmids (25). Guang et al. further demonstrated that a sequence, named the pre-mRNA processing enhancer, derived from the HSV1 TK gene enhances both polyadenylation and nucleocytoplasmic export of intronless mRNAs; this work also utilized only in vitro assays of plasmid DNA transfections (17). Finally, Otero and Hope have shown that sequences from the HSV1 TK gene behave similarly to WPRE by enhancing levels of expression of another construct; this work was also restricted to in vitro plasmid DNA transfections (36). Intriguingly, although these papers strived to isolate minimal elements conferring increased gene expression, they also showed that a longer HSV1 TK gene provided the same benefits; this longer

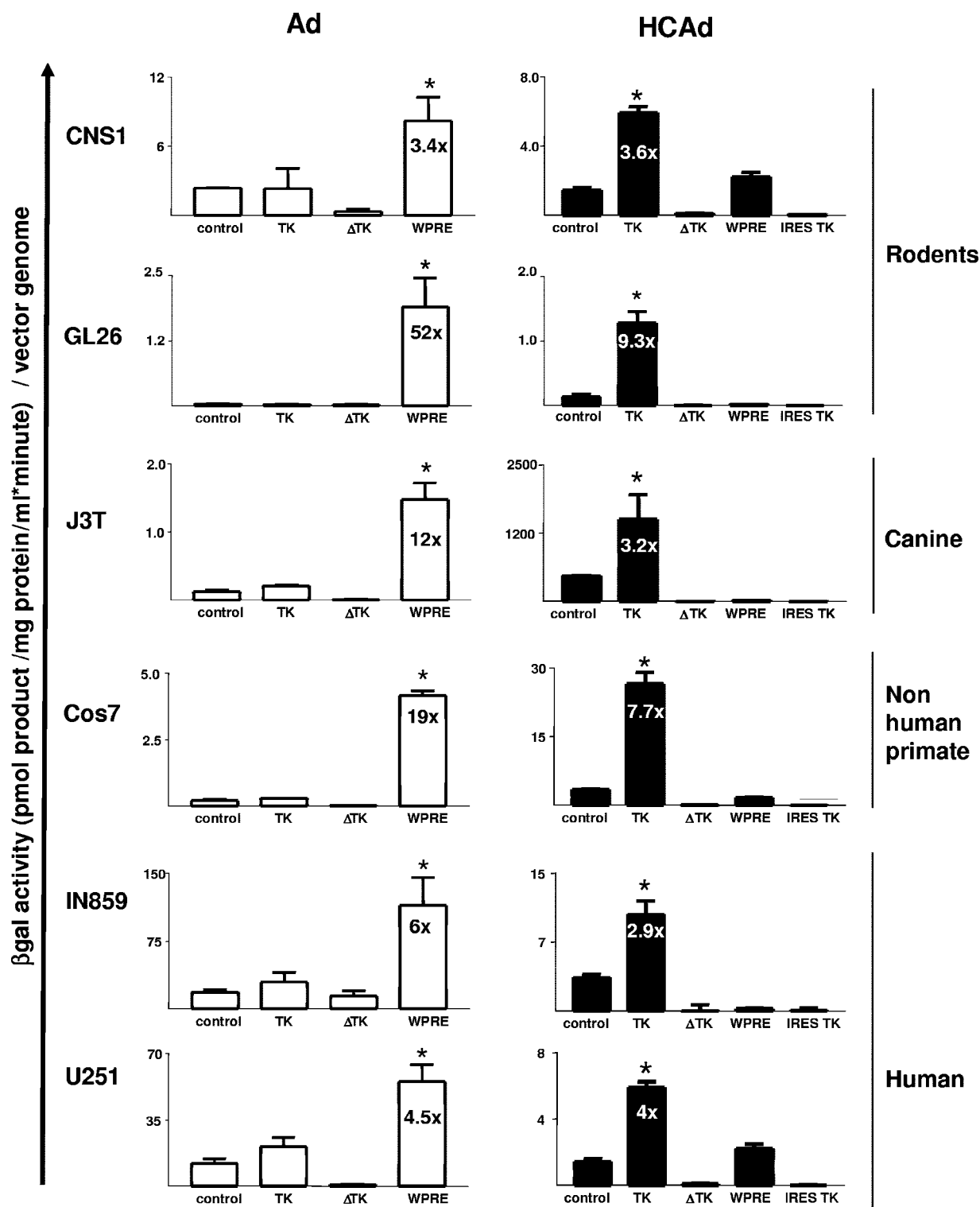


FIG. 2. In vitro transgene expression is upregulated by the HSV1 TK gene sequence carried in HCAd (right) but not in Ad (left), while WPRE increased expression in Ad (left), but not in HCAd (right). Rat (CNS-1), mouse (GL26), and dog (J3T) glioma cells, monkey kidney cells (Cos7), and human glioma cells (IN859 and U251) were infected at an MOI of 300 vgs/cell. Cells were incubated for 72 h, and transgene expression was determined by  $\beta$ -Gal activity assay of cell lysates. Experiments were performed in triplicate. Numbers of vgs were determined for all the viral preparations using quantitative PCR. Bars represent the means  $\pm$  standard errors of the means of  $\beta$ -Gal activity, calculated as  $o$ -nitrophenol produced (pmol)/(sample protein content [mg/ml]  $\times$  incubation time (min), per inoculated vg. Data were analyzed by one-way ANOVA followed by the Tukey-Kramer multiple comparison test. \*,  $P < 0.05$  versus control group (cells infected with the corresponding vector bearing the mCMV- $\beta$ -Gal cassette).

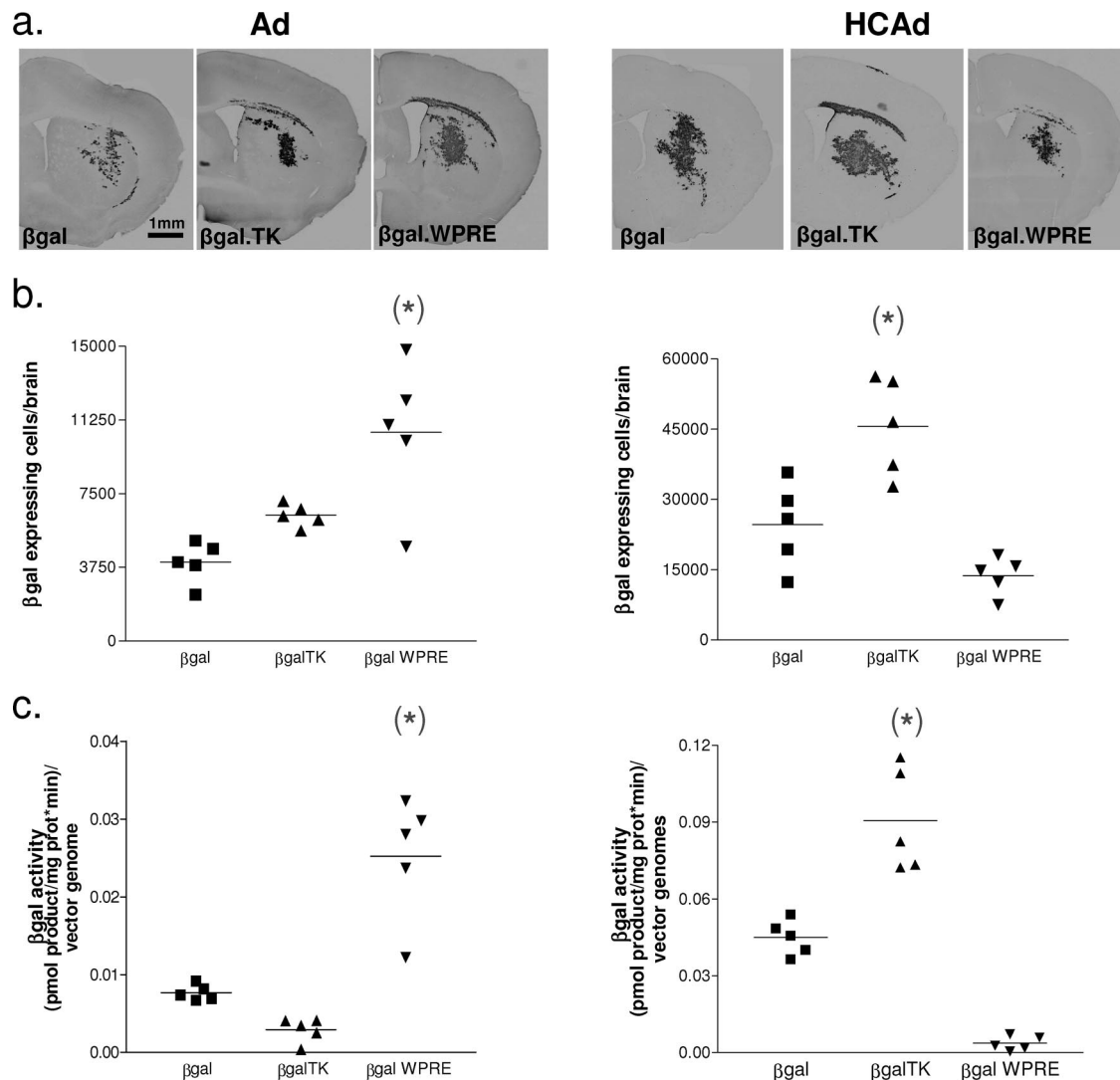


FIG. 3. In vivo transgene expression is upregulated by the HSV1 TK gene sequence carried in HCAd vectors, but not in Ad vectors. Five mice per group were stereotactically injected with  $1.0 \times 10^6$  vgs of each of the high-capacity adenoviral vectors HCAd-mCMV. $\beta$ gal, HCAd-mCMV. $\beta$ gal.TK, and HCAd-mCMV. $\beta$ gal.WPRE and with Ad-mCMV. $\beta$ gal, Ad-mCMV. $\beta$ gal.TK, and Ad-mCMV. $\beta$ gal.WPRE. (a) Low-power microphotographs showing  $\beta$ -Gal immunoreactivity in representative striatal sections from mice injected intracranially with the different vectors; (b) Numbers of  $\beta$ -Gal-expressing cells per brain. The  $\beta$ -Gal staining was performed 5 days after injection, when animals were perfusion fixed, and transgene expression was determined by immunocytochemistry. The data are expressed as  $\beta$ -Gal-expressing cells per mouse striatum.  $\beta$ -Gal-expressing cells were quantified utilizing Stereo Investigator software, version 5.00 (Microbrightfield, Inc., Colchester, VT). (c) Five mice per group were stereotactically injected with  $1.0 \times 10^6$  vgs of each of the high-capacity adenoviral vectors HCAd-mCMV. $\beta$ gal, HCAd-mCMV. $\beta$ gal.TK, and HCAd-mCMV. $\beta$ gal.WPRE and with Ad-mCMV. $\beta$ gal, Ad-mCMV. $\beta$ gal.TK, and Ad-mCMV. $\beta$ gal.WPRE. The enzymatic activity ( $\beta$ -Gal activity in brain lysates) was measured 3 days after intracranial injection of vectors, following animal perfusion with oxygenated Tyrode's solution. After testing for normality, all data were analyzed by one-way ANOVA followed by the Tukey-Kramer multiple comparison test. \*,  $P < 0.05$  versus control group, injected with a  $\beta$ -Gal-encoding vector.

HSV1 TK gene sequence, moreover, seems to be devoid of the initial 60 bp, which we determined to be required to provide increased expression from HCAd.

Taken together, our in vitro and in vivo results demonstrate that the HSV1 TK gene sequence increases levels of transgene expression per vg from high-capacity but not first-generation adenoviral vectors. Our results also suggest that the 60 bp at the 5' extreme of the HSV1 TK gene may have an important role in the enhancement of transgene expression by our full-length HSV1 TK gene, since their deletion abolishes the en-

hancement described. Additionally, our data indicate that HSV1-derived sequences that regulate transgene expression from HCAd may differ from those previously shown to increase expression from intronless genes in plasmids (25, 36).

Immune responses to viral vectors constitute one of the major limitations of gene therapy. Recent improvements in vectors to overcome immunological challenges, such as the development of strong promoters, PEGylation of viral capsids, and the HCAd vector system, have improved the safety profile of viral vectors (4, 9, 10, 14, 31). Additional increases in trans-

gene expression per vg would allow further reduction of the viral vector doses needed to achieve a therapeutic efficacy. WPRE has been systematically evaluated as a posttranscriptional regulatory element to increase transgene expression in adenoviral (3, 5, 30, 48), retroviral (19, 23), lentiviral (13, 29, 32, 40), and adeno-associated virus vectors (26, 27, 38, 45). Although WPRE has been used in HCAd vectors, transgene expression was not compared with that by the non-WPRE-containing vector version (6, 18, 20). In these various vector systems, the WPRE has been shown to increase transgene expression when tested with constitutive or cell-type-specific promoters. In these models and vectors, WPRE increased expression by 3- to 10-fold, similar to the increased expression from first-generation Ad vectors observed in our experiments. However, when carried in the HCAd vector backbone, WPRE failed to enhance transgene expression. One potential explanation could be that adenoviral sequences and/or gene products, which are absent from HCAd vectors, are necessary for WPRE to upregulate transgene expression (28). Whether differential interactions between adenoviral vector sequences and/or gene products and elements such as WPRE or the HSV1 TK gene are necessary for increased transgene expression remains to be determined.

In summary, while our data do not directly prove that the HSV1 TK gene sequence acts as a posttranscriptional regulatory element, the enhancement of HCAd-mediated transgene expression in the presence of the HSV1 TK gene sequence both in vitro and in vivo suggests that the HSV1 TK gene sequence could act as a posttranscriptional regulatory sequence when used in conjunction with HCAd vector systems. These data demonstrate the utility of a novel HSV-1-derived sequence that can act to increase transgene expression in the background of high-capacity adenoviral vectors.

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