

Multifunctional Non-Coding RNAs Mediate Latent Infection and Recurrence of Herpes Simplex Viruses

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Abstract: Herpes simplex viruses (HSVs) often cause latent infection for a lifetime, leading to repeated recurrence. HSVs have been engineered as oncolytic HSVs. The mechanism of the latent infection and recurrence remains largely unknown, which brings great challenges and limitations to eliminate HSVs in clinic and engineer safe oHSVs. Here, we systematically reviewed the latest development of the multi-step complex process of HSV latency and reactivation. Significantly, we first summarized the three HSV latent infection pathways, analyzed the structure and expression of the LAT1 and LAT2 of HSV-1 and HSV-2, proposed the regulation of LAT expression by four pathways, and dissected the function of LAT mediated by five LAT products of miRNAs, sRNAs, lncRNAs, snRNAs and ORFs. We further analyzed that application of HSV LAT deletion mutants in HSV vaccines and oHSVs. Our review showed that deleting LAT significantly reduced the latency and reactivation of HSV, providing new ideas for the future development of safe and effective HSV therapeutics, vaccines and oHSVs. In addition, we proposed that RNA silencing or RNA interference may play an important role in HSV latency and reactivation, which is worth validating in future.

Keywords: non-coding RNA, small RNA, oncolytic virus, herpes simplex virus, HSV, latent infection, latency-associated transcripts, LATs

Introduction

Herpes viruses are classified into three subfamilies: α , β and γ .¹ The α herpes viruses include herpes simplex virus type 1 and 2 (HSV-1/2). HSV typically goes through lytic infection, latent infection, and reactivation. No effective vaccine has been reported.² HSV infection is very common in human population with an infection rate of approximately 70%. Most HSV infections cause asymptomatic or weak symptomatic illnesses; however, some cause severe illnesses, including viral encephalitis. HSV-1 commonly infects facial areas, such as cold sores, preferentially becomes latent in and reactivates from the trigeminal ganglion (TG) latency; HSV-2 infects genital areas, such as genital herpes, and becomes latent in and reactivated from the sensory dorsal sacral root ganglia.^{3,4} HSV likes to establish long-term latent infections in the nuclei of sensory neurons, which possess a natural protective barrier from immune responses.⁵ The reactivation and recurrence of HSV can be caused by a variety of non-specific stimulation after latent infection, leading to related clinical symptoms, such as herpes labialis, herpangina, herpetic encephalitis, gastrointestinal diseases, esophageal diseases, and genital herpes, with increasing morbidity in young populations.⁶ There are still low levels of lytic gene transcription during the latent stage in mouse models.⁷ When the factors regulating viral latency change, the viral gene expression level increases, and large-scale

reactivation is initiated. Interestingly, HSV is reactivated in response to various factors such as ultraviolet radiation, emotional stress, fever, tissue damage, and immunosuppression. Today, anti-herpes virus drugs can target only lytic infections; therefore, treating latent viral infection remains a challenge.

Since HSV infection can be lytic, inducing rapid and efficient cell killing, HSVs have been used as oncolytic herpes simplex viruses (oHSVs). HSVs can be ideal oHSVs because they have a wide host range, and the lytic virus infection can be controlled by effective anti-herpetic agents.^{8,9} Engineered oHSVs have been reported in many clinical trials.¹⁰ T-VEC is an engineered oHSV with deletion of the genes *ICP34.5* and *ICP47* and insertion of the gene encoding human GM-CSF.¹¹ In 2015, T-VEC was approved to treat the advanced melanoma in US.¹² Although oHSV is an effective treatment for tumors, the challenge is that HSV infection can be latent in the nervous system, prone to reactivate and relapse, which cannot be cured. Therefore, the idea oHSVs should not be able to establish latency. Latency-associated transcript (LAT) is considered to play a very important role in HSV latency and is deleted in some oHSVs to prevent its latency.¹³ However, the mechanisms of HSV latency are not completely understood, which is a barrier to make safe oHSVs.

Therefore, to better treat HSV latent infection and make better oHSVs or other HSV vectors for gene therapy, we need to further understand the mechanisms of HSV latent infection. In this review, we will introduce latest development of research on the lytic and latent infections of HSV, mechanisms of HSV latency, and constructing safe oHSVs, with emphasis on the role of LAT in latent infection. We highlight a broad repertoire of multifaceted structure, expression regulation, expression products (including non-coding RNAs), function of LAT. Highlighting these quantifiable observations will provide a new prospect for better treatment of HSV latency and recurrence, and cancers by oHSV.

Lytic Infection

When HSV enters the human body, it initially infects epithelial tissues or cells for efficient lytic infection, and then spreads to nerve axons to form latent infection in sensory neurons.¹⁴ When HSVs cause lytic infections, HSVs express immediate early genes (IE or α), early genes (E or β), and late genes (L or γ), which play important roles in viral replication.¹⁵ IE genes are the

first transcribed and expressed, and encode five proteins, ICP0, ICP4, ICP22, ICP27, and ICP47.¹⁶ Among them, ICP4 and ICP27 play an indispensable role in virus replication. They are directly related to the synthesis of viral DNA. Once IE proteins have been produced in adequate amounts, infected cells typically transcribe the viral E genes, followed by DNA replication and synthesis of progeny virions. If the IE protein is lacking, the virus does not develop a productive infection.¹⁷

The transcription of the IE genes is activated by VP16, which is a major virion protein of HSV. VP16 acts through the target sequence TAATGARAT, which is present in the HSV IE promoter. TAATGARAT is the binding site of cytokine OCT-1, which is able to bind to the octamer sequence ATGCAAAT.¹⁸ The Oct family of proteins share a DNA-binding structure called the POU domain, which allows for great flexibility in the sequence recognition of various Oct proteins.¹⁹ HCF exists only in the cytoplasm and mainly transports VP16 protein to the nucleus. Therefore, HCF may play an important role in IE gene expression.¹⁷ The complex of VP16/OCT-1/HCF is associated with histone acetyltransferase and demethylase, and interacts with IE gene promoters, preventing the formation of heterochromatin and activating IE gene expression,²⁰ such as ICP0 and ICP4, which allows early and late gene transcription and lytic viral infection to occur. In short, HSV will enter the lytic infection successfully if there is enough VP16 entering the nucleus.²¹

Latent Infection

Latent infection is a typical characteristic of HSV, primarily because the virus limits its own gene expression through its interaction with host cells. During the latent infection, the gene transcription of the virus enters a silent state.²¹ When HSV enters sensory neurons, the VP16/OCT-1/HCF complex cannot form to promote IE gene expression, since the VP16 protein is degraded in the axons of neurons.¹⁷ Instead, the viral genome combines with the nucleosome to form a compact chromatin structure. Histones associated with the promoters of the IE genes are in an inhibitory configuration, preventing IE gene expression. However, the LAT promoter is in the permitted configuration, allowing the transcription of LAT, so that LATs are the only abundant transcription products of HSVs during latent infection of sensory ganglia. LAT plays a crucial role in establishing latency,²² maintaining latency, reactivating the virus from latency, and protecting neurons from apoptosis.^{23,24} HSV lytic and

latent infections are regulated partly by the interaction of different viral gene products.²⁵ Next, we will discuss the pathways and mechanisms of the latent infection of HSV in depth.

The Three Pathways to HSV Latent Infection

After the initial infection of a cell with HSV, the viral genome remains in the nonreplicating state or latent state as in sensory neurons, where the HSV genome is not transcribed owing to insufficient IE proteins.¹⁷ It is known that inefficient IE gene expression or function is a prerequisite for the virus to establish latency. The IE gene expression is theoretically regulated by the appropriate viral and cellular signaling pathways and should be hypothetically different in sensory neurons from other cell types. For example, Katz JP et al²⁶ found that the transcription efficiency of the IE gene expression was initially extremely low and E gene expression was not enough to initiate viral DNA synthesis; however, IE and E gene expression rise rapidly once viral DNA synthesis begins. This trigger is attributed to the initiation and extension of DNA synthesis. It was noted that IE genes, especially the IE gene encoding the ICP4 protein is located near the initiation region of DNA replication. Hence, early DNA synthesis may cause structural changes that activate ICP4 transcription. In mice, as with LATs, deletion mutants of HSV-1 ICP4 ORFs, O and P, can establish a significantly more latent infection.²⁷ In addition, the ICP0 expression also regulates the establishment of HSV latent infection. HSV-1 ICP0 defective mutants are approximately 1000 times less effective in establishing an incubation period than wild-type viruses, and this deficiency can be overcome by co-infection with an adenovirus expressing ICP0. This result highlights the pivotal role of ICP0 in determining the outcome of tissue culture cell infections: in the absence of ICP0, the infected cell was transformed into a static state, while in the presence of ICP0, the transformation was prevented or even reversed, leading to lytic infection. Data by Sawtell²⁸ showed that a high level of viral DNA copy number was related to HSV reactivation in the TG. It is hypothesized that transcriptional restoration of a single gene that had not been fully suppressed initiates a chain reaction. Therefore, once ICP0 is synthesized, other genes that have been completely suppressed can be transcribed. Here, we propose three pathways to HSV latent infection below (Figure 1).

(a) When there is adequate IE gene expression, more E genes are expressed, so that a large percentage of neurons may be lysed and subsequently cleared. However, few percentage of neurons may become latently infected likely due to the low copy number of HSV DNA in the neurons. It is difficult to rule out the notion that some of the dormant genomes are retained after viral DNA replication.²⁷

(b) A sufficient amount of IE gene expression can produce a large amount of ICP0 protein. ICP0 activates the JNK signaling pathway, and then AP-1 is phosphorylated by JNK and traffics into the neural nucleus, where AP-1 binds to the transcription regulatory sequence of LAT and activates LAT transcription as we reported recently,¹⁰ leading to latent infection of HSV. At the same time, LAT transcription sequence is complementary to the IE gene encoding ICP0, which continuously consumes ICP0, resulting in the corresponding decrease in IE gene expression. However, ICP0 and LAT increase histone occupancy and heterochromatin levels of lytic genes and block their transcription,¹⁴ causing HSV latency.

(c) Low levels of ICP0 were found to promote the expression of LAT and lytic genes in latently infected ganglion, where the loading of total histone and heterochromatin on HSV genome was increased, maintaining latency. LAT transcription was shown to downregulate lytic gene transcription during latency through the deposition of repressive histone marks on the lytic IE gene promoters.²⁹ Moreover, low levels of ICP0 promoted AP-1 nuclear translocation, which promotes the expression of LAT. Subsequently, many products of LAT (LAT, MiR and ORF) are produced, which will make HSV latent through many mechanisms, including the anti-apoptotic pathway.

Taken together, ICP0 acts as a “switch” in the balance between the lytic and latent states.^{17,30} LATs have a variety of potential roles in promoting latent infection by inhibiting apoptosis, inhibiting the expression of lysed genes and promoting viral latency. This is closely related to the biological structure of LAT as we will describe below.

The Structure and Expression of the LAT

It is believed that the LAT presence indicates HSV latency. LAT plays a key role in latent infection, which is attributed to the genetic structure of LAT. During HSV latent infection, LAT is the only transcription products

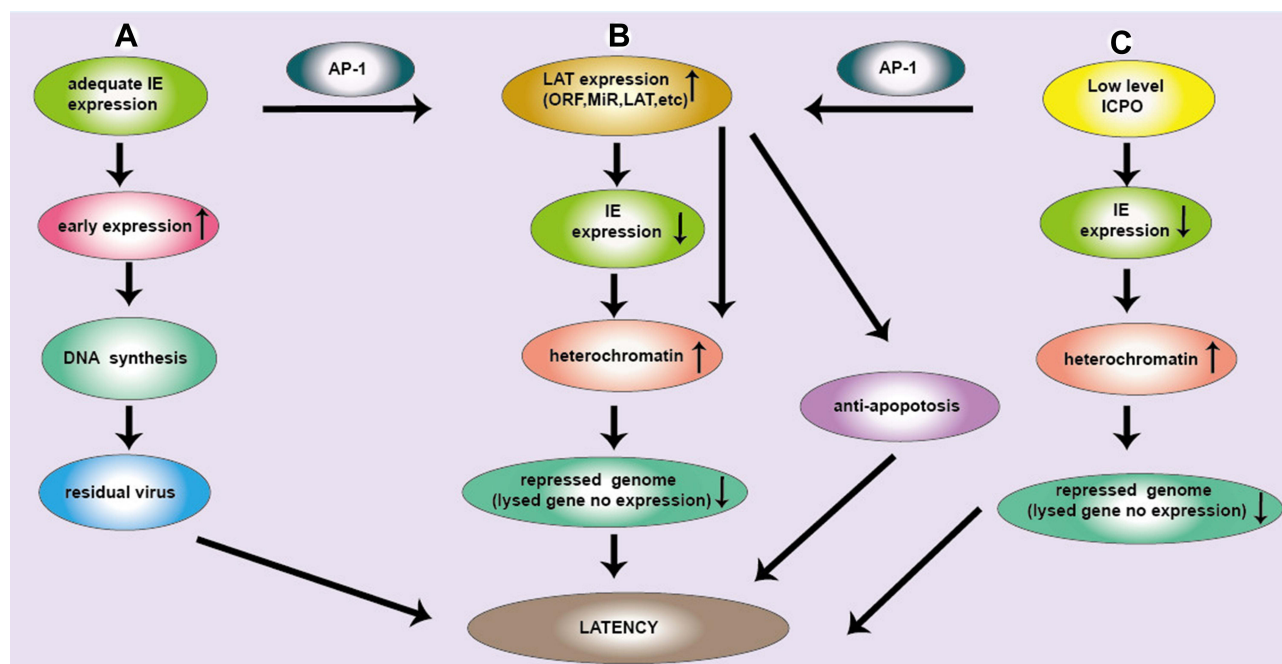


Figure 1 Three pathways of latent infection of HSV. **(A)** IE gene is adequate, the early expression gene is increased and largely neurons are cleaved and subsequently cleared. Due to the low copy number of HSV DNA in neurons, DNA synthesis is reduced and residual viruses are latent infection. **(B)** IE gene expression can produce ICPO protein. ICPO activates the JNK signaling pathway, and AP-1 is phosphorylated, then activate LAT. LAT sequence is complementary to the IE gene encoding ICPO, which continuously consumes ICPO, so decrease IE gene expression. ICPO and LAT increase histone occupancy and heterochromatin levels of lytic genes and block their transcription, causing HSV latency. **(C)** Low levels of ICPO can promote the expression of LAT and lytic genes in latently infected ganglion, where the loading of total histone and heterochromatin on HSV genome was increased, maintaining latency. LAT transcription can downregulate lytic gene transcription during latency. Moreover, ICPO promoted AP-1 nuclear translocation, which promotes the expression of LAT. Subsequently many products of LAT (LAT, MiR and ORF) are produced, which will make HSV latent through anti-apoptotic pathway.

that are highly expressed in the nucleus of host cells. Interestingly, Ying Zhang et al¹⁰ showed that HSV LAT has two intervals, LAT fragment 1 (LAT1) and LAT fragment 2 (LAT2), and their respective promoters, LAT promoter 1 (LAP1) and LAT promoter 2 (LAP2) (Figure 2). LAP1 has been shown to be the critical promoter for activating expression of LAT during latency. LAP2 also has been shown to possess promoter activity and augment HSV-1 LAP1 activity during latency. LAT is also divided into non-major LAT (minor LAT or primary LAT) and major LAT, which are produced by different ways of splicing. HSVs have serotypes HSV-1 and HSV-2, which share many similarities in their morphology, molecular structure and drug sensitivity, but are different in their LAT. HSV-1 LAT1/2 includes a family of transcripts comprising an 8.3 kb primary transcript in low levels and three introns (Figure 2A). Among the three introns, the 2.0 kb LAT was highly expressed in both lytic and latent infection; the 2.0 kb LAT was found only at low levels during latency; the predicted 0.5 kb LAT was not detected.³¹ LAT2 splicing produced a 6.3 kb exon product, which cannot easily be detected, although the 2.0

and 1.5 kb introns can easily be seen in Northern blot analysis.¹⁰ Translation of a protein from the spliced LAT exon mRNA remains to be convincingly shown,³² nevertheless it is clear that the exon 1 region of the LAT is required for protection from apoptosis.^{23,33} The function of the 1.5 kb LAT introns is unclear. However, the 2 kb LAT is upregulated in some cells expressing heat shock proteins.³⁴ Interestingly, the 1.5 kb LAT intron is completely encoded in the 2.0 kb intron and is, thus, called a twintron.³⁵ Both introns are present in the nucleus during latency. The LAT introns can be described as a complex twintron as three overlapping introns (2.0/1.5/0.5kb) can be produced from the LAT transcript. Likewise, HSV-2 LAT2 is transcribed to generate approximately a 9.0 kb of non-primary LAT and a stable 2.2 kb of major LAT. However, transcripts of HSV-2 LAT1 are rarely reported (Figure 2B).

Regulation of LAT Expression by Four Pathways

HSV can establish latency in neurons, thus maintaining a latent state for years, or even a lifetime.³⁶ This

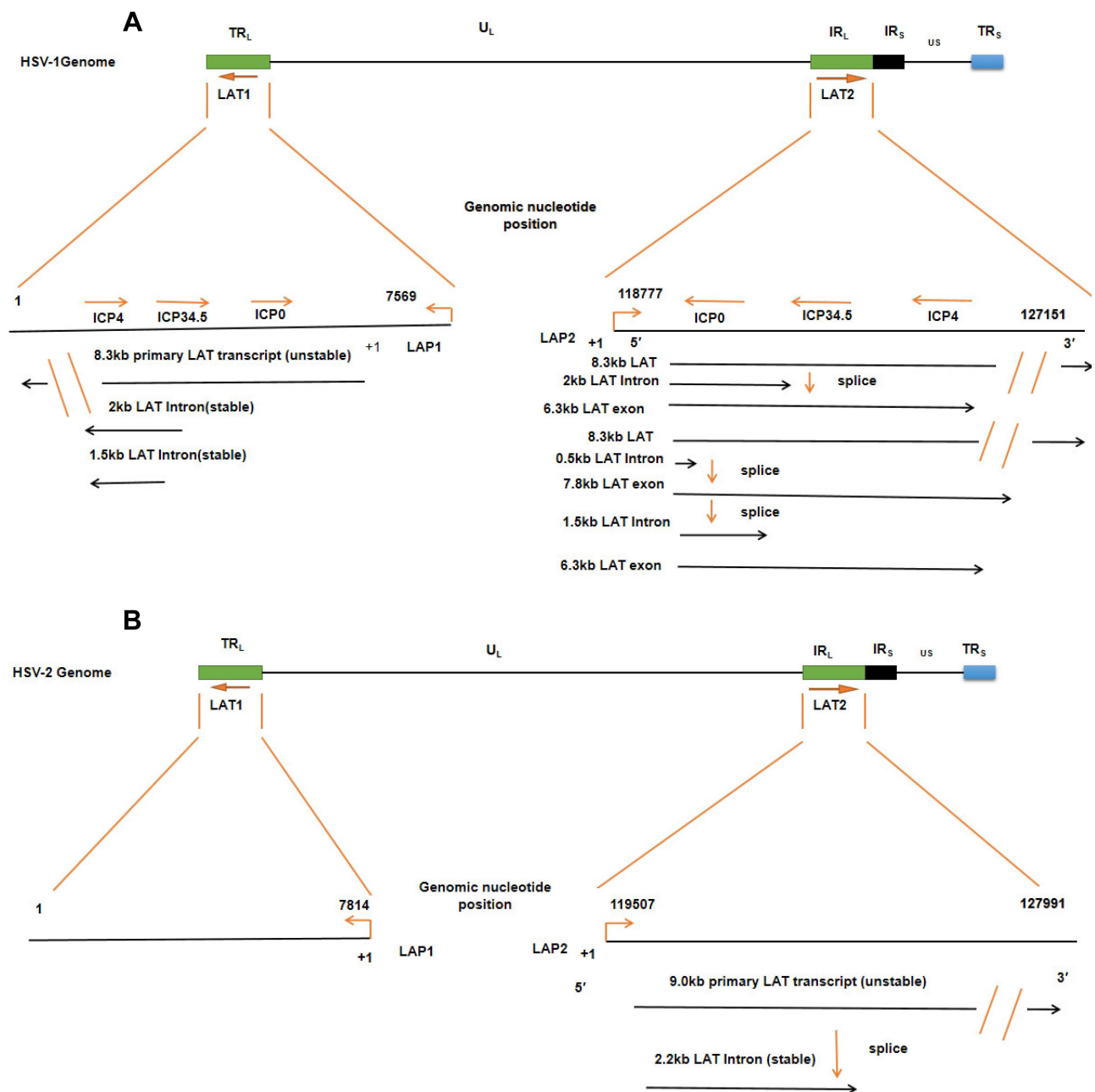


Figure 2 The LAT locus and (twintron) splicing of the HSV genome. **(A)** HSV-1 includes UL and US regions with terminal and internal repeats (TR_L, IR_L, IR_S and TR_S). LAT1 is in TR_L and UL, LAT2 is in IR_L and UL connection region. An TR_L fragment, overlapping ICP0, ICP34.5, and ICP4, is expanded to show LAT1, including 8.3 kb original LAT, and 2 kb and 1.5 kb of the LAT intron. The 8.3 kb LAT2 was spliced into a 2.0 kb intron and a 6.3 kb mRNA or spliced as twintron introns into a 0.5 kb unstable intron and a 7.8 kb RNA. **(B)** HSV-2, like HSV-1, contains the LAT1 and LAT2. Their transcripts include an unstable 9.0 kb primary LAT and a stable 2.2 kb primary LAT.

characteristic is related to the structure, expression and regulation of LATs. Here, we summarize the regulation of LAT expression of HSV (Figure 3) by four pathways.

(A) The ICP0-JNK-AP1-LAT pathway. The envelope glycoprotein gD of HSV binds to the herpes virus entry mediator (HVEM) on cell membrane, and the transmembrane protein HVEM binds to TRAFs in the cytoplasm, then the HSV enters the cell.³⁷ HSV-1 infection results in

the synthesis of IE proteins, such as the ICP0. Subsequently, ICP0 was able to catalyze the K63-linked polyubiquitin chains with the help of the Ubc13ev1A enzyme as it recognizes and interacts with TAK1.³⁸ Then, activated TAK1 phosphorylates MKK6, which subsequently activates the JNK kinase pathway.³⁸ The JNK pathway phosphorylates the AP-1 transcription factor, then the phosphorylated AP-1 translocated into the nucleus to

bind to the LAT promoter sequence and promoted LAT expression, leading to HSV latency.¹⁴

(B) The Sp1-Stat3-ATF3-CRE-LAT pathway. HSV dissociates the capsid and enters the nerve SK-N-SH cells. Sp1 protein is abundant in SK-N-SH cells.³⁹ However, Sp1 protein interacts with a variety of GC rich promoter sequences, and Sp1 recruits STAT3, ATF3, and c-jun complexes.³⁹ In neurons, STAT3 exists in the cytoplasm and travels to the nucleus under neuronal stress.⁴⁰ STAT3 binds to ATF3:c-jun heterodimer in the nucleus without DNA binding activity or phosphorylation. ATF3 is a member of the ATF/CREB transcription factor family.⁴¹ The current consensus is that ATF3 is an immediate early-response gene, a regulator of stress response, and a factor in determining cell fate.⁴² ATF3 binds to a cAMP response element (CRE) with the canonical sequence 5'-TGACGTCA-3' in the form of homodimers or heterodimers with other members of the ATF/CREB family.⁴³ HSV-1 DNA contains eight CRE sites; however, only two of these are in a promoter, specifically in promoter 1 of LAT (LAP1). Results showed that ATF3 bound to both CRE sites in LAP1, formed a homodimer, and fostered increased accumulation of LAT.⁴³ Furthermore, another study found that ATF3 enhanced the expression of LAT,⁴⁴ and played a role in the maintenance of HSV-1 in a latent state.⁴³

(C) The CTRL-CTCF-LAT pathway. When HSV enters the cytoplasm and the capsid breaks away, the IE gene begins to express ICP0. Interestingly, the ICP0 region is flanked by two CTCF binding motifs, namely CTRL1 and CTRL2.⁴⁵ CTRL binds to CTCF and activates LAT promoter (LAP), making the virus latent. Recent studies have shown that the CTCF-insulator's role in lytic gene inhibition of β and γ herpes viruses further supports the possibility that CTCF-insulators also regulate α herpes virus latency via lytic gene repression.⁴⁵

(D) The VP16-LowICP0-LAT pathway. Multi-protein complex of VP16, Oct-1 and HCF initiate IE gene transcription. However, as a virion protein, VP16 cannot be successfully transported to ganglion nuclei with viral genomes because of certain physical losses. In the absence of VP16, multi-protein complex (VP16, Oct-1 and HCF) cannot be formed, thus reducing IE gene transcription.⁴⁶ Therefore, the IE gene encoding the ICP0 protein would be low. Low levels of ICP0 can promote both LAT and lytic gene expression as well as total histone and heterochromatin loading on the HSV genome in latently infected ganglia.¹⁴ In contrast, in latently infected neuron nuclei,

the low levels of ICP0 expression can promote the expression of LAT and the accumulation of H3K27me3/H3K9me3 heterochromatin to silence lytic gene expression.^{47,48} Together, LAT expression may play a pivotal role in the establishment and maintenance of latency through regulation of lytic gene expression.⁴⁹

LAT Function Mediated by Its miRNAs, sRNAs, lncRNAs, sncRNAs and ORFs

The expression of LAT is crucial. LAT has several other expression products, such as microRNA (miRNA), small RNA (sRNA), long non-coding RNA (lncRNA), short non-coding RNA (sncRNA), and LAT open reading frames (ORF). Numerous studies have supported that LATs encode multiple RNAs or ORFs. It is noteworthy that these RNAs or ORFs may play an crucial role in the latency-reactivation cycle.⁵⁰ Here, we summarize the five main types of LAT expression products that are closely related to HSV latency (Figure 3).

MicroRNAs

MicroRNAs (miRNAs, miRs) are a class of 21 to 24 nucleotides (nt) noncoding RNAs that regulate the stability or the translational efficiency of target messenger RNAs (mRNAs).⁵¹ In transfection assays, miRNAs originating from the LAT region can repress expression of viral lytic proteins.³¹ LAT can promote the establishment of latent infection via LAT-encoded miRNAs.⁵² Firstly, HSV-1-miR-H2, which has been associated with the regulation of latency, binds to viral ICP0 mRNA and inhibits its expression.⁵³ In addition, LAT miR-H2 has been experimentally shown to inhibit the production of ICP0 protein by downregulating ICP0 translation, but not transcription.^{54,55} Some report that LAT miR-H2 functions to decrease HSV-1 pathogenesis and to help maintain HSV latency in Swiss Webster mice.⁵⁶ HSV-1-miR-H6 can target ICP4 mRNA, inhibiting the expression of the transcription factor ICP4 crucial to E and L gene transcription, leading to blocking virus lytic infection and decreasing IL-6 expression; therefore, HSV-1-miR-H6 promotes HSV-1 latency.⁵⁷

Interestingly, miRNA derived from the LAT region helps to inhibit the expression of cleavage function by blocking the synthesis of two viral proteins, ICP0 and ICP34.5, consequently facilitating the maintenance of latency.⁴ Recently, acutely and latently expressed miRNAs encoded by HSV-2 LAT was reported to inhibit ICP34.5

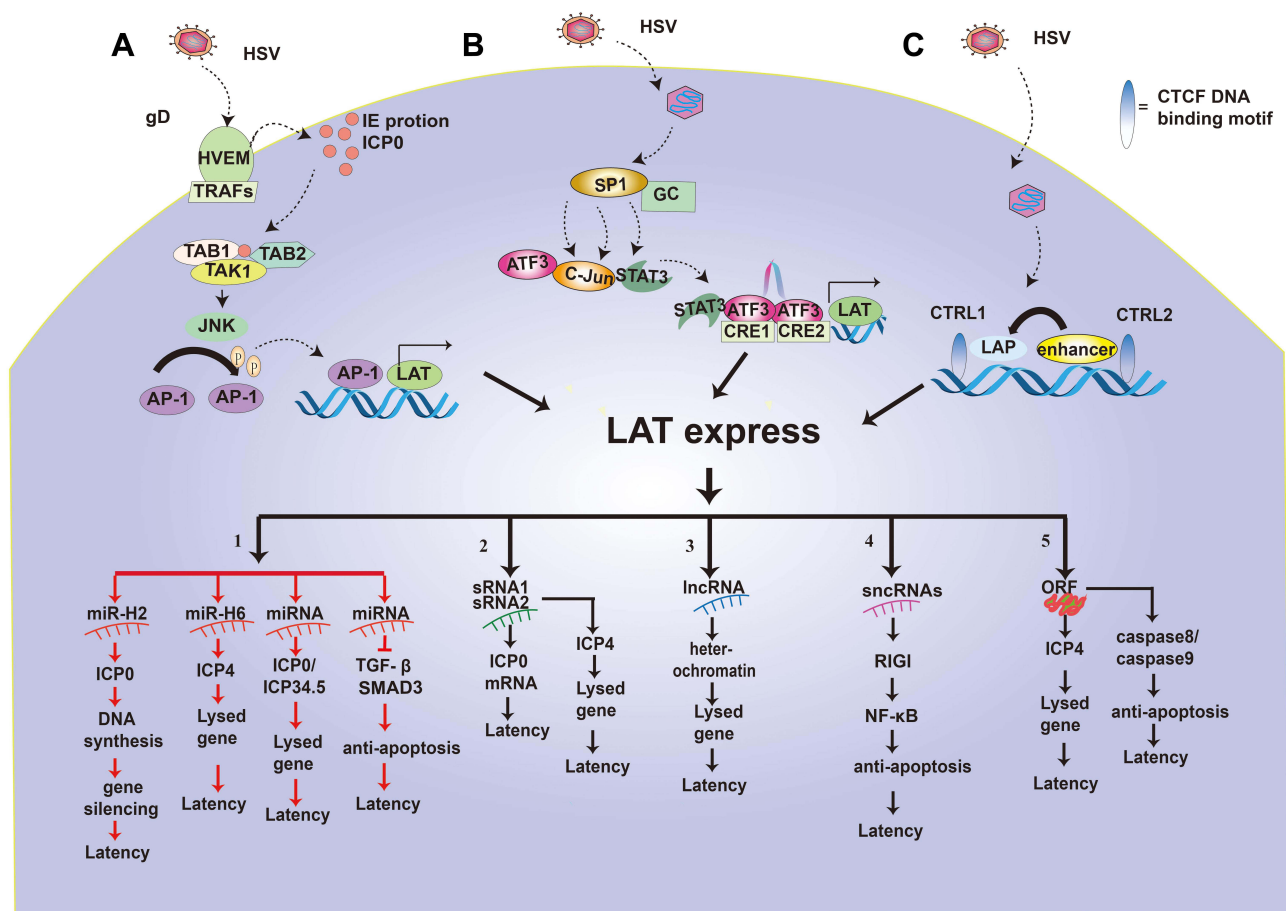


Figure 3 Regulation of LAT expression and function of LAT. (A–C) highlight the pathways that promote the transcriptional expression of LATs. 1–5 shows the five functional mechanisms of LAT latency. 1. LAT miR-H2 can inhibit ICP0 protein so that viral DNA cannot be synthesized, causing the silencing of the lysed gene and HSV latency. LAT miRNA-H6 inhibit the expression of ICP4 and lysed genes. LAT miRNA blocks ICP0/ICP34.5 production and reduces viral DNA synthesis, which in turn inhibits the expression of the lysed gene and renders the virus latent. LAT miRNA can also achieve viral latency by targeting transformed growth factor (TGF)- β and SMAD3 in the TGF- β pathway to resist apoptosis. 2. LAT sRNA1 and sRNA2 can inhibit ICP0 mRNA expression by partial base complementation to maintain latency. LAT sRNA2 can also inhibit the expression of ICP4 and lysed genes to help establish and maintain virus latency. 3. LAT IncRNA can induce facultative heterochromatin to promote lytic gene silencing during latency. 4. LAT sncRNA collaborates with RIG-I to stimulate NF- κ B dependent transcription for anti-apoptosis and latency. 5. LAT ORFs inhibit ICP4 and then down-regulates lysed genes to make the virus latent. At the same time, LAT ORFs inhibit caspase 8 or caspase 9 for anti-apoptosis and latency.

expression, whereas two other miRNAs encoded by HSV-1 LAT also mapped antisense to the ICP34.5 region.⁵⁸ In HSV-2 LAT miRNAs, miR-H3 and miR-H4, as well as the corresponding HSV-1 LAT-encoded miRNAs, were processed from the primary LAT and a L/ST that traverses the viral long/short repeat connection targeting ICP34.5.⁵⁹ However, LAT miRNAs are negatively regulated by ICP4.⁵⁴ ICP4 inhibits miRNAs targeted at ICP34.5 and activates lytic virus genes, which suggests that ICP4 plays a critical role in the transition between HSV latency and activation.⁵⁹ Meanwhile, the LAT plays an anti-apoptotic role by encoding miRNA, which prevents neuronal apoptosis and maintains the integrity of the neuron to preserve HSV in a latent state.⁶⁰

In addition, studies have shown that miRNAs can protect nerve cells from apoptosis mediated by CD8+

T cells.⁶¹ During HSV latency, some neurons are surrounded by CD8+ T cells, which are thought to infiltrate neurons to help maintain HSV latency by inhibiting virus reactivation.⁶¹ Surprisingly, when large numbers of CD8+ T cells were injected into cytotoxic lytic granules of latently infected neurons, apoptosis and neuronal death did not occur. Therefore, it has been hypothesized that the protective effect of latently infected neurons from cytotoxic CD8+ T cell killing is attributed to the anti-apoptotic activity of LAT.⁶¹ The ability of LAT to block granzyme B induced apoptosis was certified,⁶² given that granzyme B is the major factor released by CD8+ T cells and responsible for cytotoxic and soluble granulocyte-mediated apoptosis. Granzyme B can directly cleave and activate caspase-3 as well as other proteins known to regulate apoptosis.⁶³ In addition, LAT encoded miRNA

plays an anti-apoptotic role by down-regulating the expression of functional transforming growth factor (TGF)- β and SMAD3 in the TGF- β pathway.⁶⁴ Some results indicate that miRNAs encoded by HSV-1 LAT gene regulates apoptosis of infected cells by modulating TGF- β signaling pathway, leading to the persistence of HSV in a latent form in sensory neurons.⁶⁵

In conclusion, LAT encoded miRNAs make HSV latent through the downregulation of ICP0, ICP34.5, ICP4, TGF- β and SMAD3.

sRNA

HSVs encode small RNAs (sRNA1 and sRNA2), which have extensive secondary structure and a stem-loop structure and are larger than micro-RNAs. LAT sRNA may interact with mitochondria, inhibit the release of pro-apoptotic factors, and enable cells to avoid the fate of apoptosis and may survive, thus giving the HSV an opportunity to be latent in host cells.⁶⁶ Further, LAT sRNA1 and sRNA2 play essential roles in the latency-reactivation cycle because the sRNAs inhibit productive infection and apoptosis, which is consistent with previous studies showing that the same mutation affects its anti-apoptotic function in a large LAT fragment.^{67,68} In addition, sRNA1-LAT and/or sRNA2-LAT inhibit the ICP0 gene expression by partial base complementation to maintain latency.⁵⁰ Surprisingly, when the wild-type LAT sRNA1 was co-transfected with the plasmid expressing LAT sRNA2, the intrinsic pathway of neuro-2A cells apoptosis was blocked, thereby protecting neuro-2A cells from cold-shock induced apoptosis.⁶⁷ Finally, LAT sRNA1 was also predicted to hybridize with VP16 and UL8 mRNA (proteins essential for viral DNA replication), suggesting that LAT sRNA1 may reduce the expression of related viral genes. Further, LAT sRNA2 and miRNA-H6 inhibit the expression of ICP4, which may contribute to establishing and maintaining viral latency via inhibiting the expression of lysed genes and protecting infected neurons from cell death.^{50,67} In short, LAT sRNAs can successfully make HSV latent due to its anti-apoptotic and anti-lytic effects.

lncRNA

HSV LAT is a set of long non-coding RNA (lncRNA), which is the only transcript known to accumulate at a large amount during latency.⁶⁹ lncRNA play a key role in maintaining latent infected neurons.⁴³ During latency, neurons accumulate 1.5/2.0kb of non-coding RNA of LAT. Relevant studies have reported that the transcription of viral noncoding RNAs

can also promote the lytic gene silencing during the incubation period by inducing facultative heterochromatin.⁴⁷ lncRNAs help form heterochromatin in HSV chromosome, inhibit the expression of lytic genes, and make the virus latent.⁴⁷ Furthermore, lncRNAs are also considered to play a critical role in promoting the activation of HSV.⁷⁰ Together, lncRNAs of LAT makes HSV latent or reactivated by modification of heterochromatin.

SncRNAs

The herpes virus entry mediator (HVEM) is one of several cell surface proteins that HSV uses for attachment/entry. HVEM regulates cellular immune responses and can also increase cell survival. SncRNAs of LAT were distributed on HVEM promoter. The increase of HVEM would down-regulate the immune spongy body in the latent microenvironment and increase the survival rate of latent infected cells.⁷¹ Experiments showed that two sncRNAs encoded by LAT can induce the expression of HVEM in latently infected mice in the trigeminal ganglia.⁵⁰ In addition, the effect of LAT on latency was significantly reduced in HVEM-deficient mice. The results showed that LAT regulated latent infection and reactivation of HSV to some extent by upregulating the expression of HVEM mRNA levels,⁷² thereby increasing the survival of latently infected cells and preventing the activation of effector T cells.⁷¹ Therefore, the results indicate that the LAT-HVEM relationship is a new mechanism directly involved in the HSV latency steady state pathway.⁷¹ Interestingly, one study found that LAT sncRNA1 cooperates with the RNA sensor, retinoic acid-inducible gene I (RIG-I), to stimulate IFN- β promoter activity and nuclear factor- κ B (NF- κ B) dependent transcription in human or mouse cells.⁷³ LAT sncRNA2 stimulated RIG-I induction of NF- κ B dependent transcription in mouse neuroblastoma cells (Neuro-2A) but not human 293 cells.⁷³ In Neuro-2A cell model, both sncRNAs work together with RIG-I to suppress cold-shock induced apoptosis.⁷³ Together, two sncRNAs derived from LAT contribute to the establishment of virus latency by combining RIG-I to IFN- β .

LAT ORF

All the LATs of HSV-1/2 have multiple ORFs, which encode proteins related to activation and function, and are similar to ICP0 and of great significance in the process of latent infection.⁷⁴ The expression of LAT ORFs can complement the mutation of ICP0, which instead greatly reduces the growth of the virus.⁷⁵ Furthermore, LAT ORFs

can activate wild-type phenotype in the rabbit ocular model.⁷⁶ In addition, LAT ORF overexpression could down-regulate the expression of virus lytic genes by downregulating the ICP4.⁷⁷ Interestingly, LAT ORF has a significant role in the HSV life cycle. LAT ORF can interact with HSV-1 proteins, RNA or DNA, to promote viral growth. Studies have shown that LAT-encoded ORF enhances HSV-1 replication in epithelial and neuronal cells, especially in cells that normally do not allow for virus growth.⁷⁸ Further, LAT-encoded ORF2 can enhance the transactivation of lytic proteins.⁷⁶ Experiment results showed that ORF knocking-down reduced the protective effect of LAT on caspase 8- or caspase 9-induced apoptosis.⁶⁸ Knocking out all eight ORFs blocked the antiapoptotic activity of LAT more significantly and effectively than knocking out only ORFs 1–2 or ORFs 3–8.

In summary, we summarized the function of LAT mediated by LAT encoded miRNA, sRNA, lncRNA, sncRNAs and ORFs, and elaborated on how they regulate

gene expression at both transcriptional and post-transcriptional levels to influence the latency and reactivation of HSVs (Figure 3).

Application of HSV LAT Deletion Mutants in HSV Vaccines and oHSVs

After HSV infection, it may become latent in the ganglia, and then reactivate and relapse under some non-specific stimulation, causing associated clinical symptoms. Because HSVs have a latency and easy activation mechanism, it cannot achieve a radical cure in clinical practice. Therefore, it is necessary to study the latency and reactivation mechanisms of HSVs. Many researchers have paid great attention to the vital role of LATs in latency and recurrence of HSVs for the effective targeted HSV treatment and making safe oHSVs. Here, we have extensively outlined the different effects of LAT deletions from HSVs and oHSVs as shown in Tables 1 and 2.

Table 1 Application of Deleted LAT in HSV

HSV	HSV-1/2	HSV Strain	LAT Mutant	LAT Deletion (nt)	Effect	Ref
HSV	HSV-1	KOS	M3	Deletion of a 937–1074 nt fragment from LAT	Low pathogenicity in 293T cells	[6]
	HSV-1		ΔCTRL2	Removed a 370 bp fragment from within the 2.0kbp LAT intron	Slightly higher levels of LAT expression	[31]
	HSV-1		ΔCTRL2	A 370-bp deletion of the CTRL2 site	Disrupted the establishment of latency	[79]
	HSV-2	333	LAP1	Deletion of LAT1 (3457 to 4081)	Reduced/abolished LAT expression	[85]
	HSV-2	333	LAP2	Deletion of LAT1 (4081 to 4382)	Reduced LAT expression	[85]
	HSV-1	McKrae.	dLAT-cpIAP	Deletion of LAT1 (76 to 1667)	Efficiently reactivation phenotype	[80]
	HSV-1	McKrae	dLAT-ΔH2	Deleted the core LAT promoter and the first 1667 nts of LAT	Increased reactivation phenotype	[56]
	HSV-1	McKrae	dLAT2903	Deletion of LAT1 (–161 to 1667)	Reduced reactivation phenotype and Enhanced apoptosis	[81]
	HSV-1	McKrae	dLAT2903	Deletion of LAT1 (–161 to 1667)	Significantly reduced reactivation	[82]
	HSV-1	McKrae	dLAT2903	Lacks the LAT promoter and the first 1.6 kb of the 5' end of LAT	Reactivation was decreased to approximately 33% of normal.	[3]
	HSV-1	KOS	ΔCTRL2	Deletion of LAT2 (120136 to 120508)	Significantly reduced reactivation	[31]
	HSV-1	McKrae	dLAT2903	Deleted the first 1667 nucleotides (nt) of the LAT transcript (118664–118886).	Reduces latency reactivation by approximately threefold	[29]
	HSV-1	17Syn+	17ΔPst	Deletion of the 118664–118886 core promoter of LAT	Reduces latency reactivation by approximately 3-fold	[49]
	HSV-1	McKrae	dLAT2903	LAT nucleotides (–)161 to +1667	Increased neurovirulence, and slightly more rapid reactivation	[56]
	HSV-1	McKrae	dLAT2903R	Deletion of LAT1 (–161 to 1667) relative to the start of the primary 8.3-kb LAT	Reactivate from latency is significantly reduced.	[83]
	HSV-1	17syn+	LATRz-235	Deletions in the LAT promoter and portions of the 5' exon coding region	Resulting in a two- to threefold decreased reactivation	[84]

Table 2 Application of Deleted LAT in oHSV

oHSV	HSV-I/2	HSV Strain	LAT Mutant	LAT Deletion (nt)	Effect	Ref
oHSV	HSV-1	U87 glioblastoma	DM33	Deleted LAT I (76 to 1667)	Effectively	[86]
	HSV-1	Brain tumors	DM33	Deleted LAT I (remove LAT nucleotides 76 to 1667)	Enhanced safety	[87]
	HSV-1	Superficial cancers and melanoma	HF10	Lacks the expression of LAT I/2	Safe and well tolerated	[88]
	HSV-1	Pancreatic cancer	HF10	Not expression of LAT I/2.	High safety	[89]

These effects have been confirmed in published literature (Table 1). In terms of the HSV-1, one experiment has made a LAT deletion via removing a 937–1074nt fragment from LAT to receive the M3 mutant. The M3 mutant was harvested from the infected human embryonic kidney cell line 293T cells, which showed that it is an attenuated strain with low pathogenicity during both acute and latent infections. The above result supports the use of the M3 mutant as a candidate for the development of an HSV-1 vaccine.⁶ Interestingly, another experiment built a CTRL2 deletion mutant virus by removing a 370-bp fragment from the 2.0-k bp LAT intron to form Δ CTRL2 mutant virus, which showed slightly higher levels of LAT expression during latent infection. However, this was indistinguishable during reactivation.³¹ Conversely, a CTRL2 deletion mutant virus (a 370-bp) resulted in the encroachment of heterochromatin onto the LAT region of the genome to disrupt the establishment of latency, and had lower survival rate.⁷⁹ However, Lee et al³¹ showed that their Δ CTRL2 recombinant can establish an equivalent latent infection in the mouse TG and displayed no replication defect during the lytic infection.

Deletion of LAT in HSV affects not only HSV latency but also virus reactivation. One study has shown that deletion of LAT1 (76 to 1667) can effectively reactivate phenotype.⁸⁰ Similarly, deletion of the core promoter of LAT and the first 1667 nts of LAT can increase the reactivation phenotype.⁵⁶ Inversely, deletion of LAT (–161 to 1667) reduced the reactivation phenotype and enhanced apoptosis, resulting in the decrease in HSV reactivation rate.⁸¹ Other studies have shown that deletion of LAT can significantly reduce HSV reactivation.^{31,82,83} In addition, studies have also shown that deletion of LAT can significantly reduce reactivation to about 33% of the normal level.³ Similarly, partial LAT deletion mutation has been shown to lead to an approximately threefold increase of latency in the TG of wild-type mice infected with LAT(+)

virus compared with mice infected with LAT(–) virus.^{29,49,84} On the contrary, another experiment has constructed a LAT deletion mutant virus by removing a –161 to +1667 fragment to form the dLAT2903 mutant virus, which showed increased neurovirulence and slightly more rapid reactivation.⁵⁶ In HSV-2, removing LAT1/LAT2 significantly reduced LAT expression during productive infections and abolished LAT expression during latency.⁸⁵

As the causal infectious virus, deletions of the LAT gene of HSVs will significantly reduce the latency and reactivation of the viral infection, which will alleviate the pain of clinical herpes patients. As oHSVs, there is growing interest about the genetic modification of LAT to make an effective and safe oHSVs for cancer treatment (Table 2). Recent studies have shown that modifying the LAT gene may significantly improve the targeting and safety of oHSVs for cancer therapy.^{86–89} For example, the oHSV DM33 was made by deleting γ 34.5 and LAT genes, which removed LAT nucleotides 76 to 1667. DM33 was safe and effective in treating human U87 glioblastomas and brain tumors.^{86,87} Moreover, oHSV HF10 lacks the expression of LATs, showing a strong safe and well-tolerated effect in the treatment for superficial cancers, melanoma and pancreatic cancer.^{88,89}

We look forward to developing a better therapeutic regimen based on the experimental application of HSV and oHSV by removing LAT through in-depth study, to give full play to its therapeutic potential. However, more clinical trials are needed to drive the development of HSV LAT-based diagnostic tests and therapeutic interventions to benefit patients with herpes and tumors.

Conclusion and Perspectives

In summary, latent infection of HSV is a multi-step complex process. Here, we have systematically reviewed the latest development of the multiple pathways to HSV latency, and

the structure and expression, regulation, and function of the LAT, and the genetic analysis and engineering of LAT. LAT performs a wide range of function by different molecular mechanisms, including transcriptional and post-transcriptional regulation. Significantly, we first summarized the three HSV latent infection pathways, systematically analyzed the structure and expression of the LAT1 and LAT2 of HSV-1 and HSV-2, proposed the regulation of LAT expression by four pathways, and dissected the function of LAT mediated by five LAT products of miRNAs, sRNAs, lncRNAs, snRNAs and ORFs. We further analyzed that application of HSV LAT deletion mutants in HSV vaccines and oHSVs. Deleting LAT significantly reduced the latency and reactivation of HSV, providing a new idea for the future development of HSV vaccines and safe oHSVs.

In different heterogeneous tissues, the translation level of LAT ORFs are strictly limited in time and space. Whether LAT can be translated into protein is uncertain. For example, studies have shown that LAT gene may encode a protein with a latency-related function, as has been demonstrated for bovine herpes virus.⁹⁰ Currently, there has been one report of a LAT-encoded protein in latently infected primary neuron cultures,⁹¹ but it has not been confirmed. Fortunately, HSV-1 LAT is consistently detected in abundance during latency and in the neurons of infected humans, mice, and rabbits.⁹² In latency models, the abundance of the LAT RNAs increases during the establishment of latency and can be detected in neurons of ganglia from the sacral, thoracic, and lumbar regions of recently deceased humans.⁹³ These reports provide us with the knowledge about the future application in establishing latency and reactivation models to further uncover the veil of HSV latency and reactivation.

It is well established that LAT controls apoptosis,⁹⁴ which likely occurs via the type I interferon (IFN) pathway.⁹⁵ The type I IFN pathway consisting mainly of IFN- α and - β is activated by several HSV-1 glycoproteins.⁹⁵ However, the HSV-1 LAT transcript was shown to delay several IFN- α subtypes, including IFN- α 1, IFN- α 4, IFN- α 5, IFN- α 6, and IFN- α 14, as well as IFN- β , during an acute infection in vitro, and IFN- β and IFN- γ in vivo.⁹⁶ In addition, LAT is responsible for CD8⁺ T cells immune exhaustion, which is also essential for latency establishment.⁹⁷ Previous reports have shown a LAT-dependent elevation of the immune exhaustion markers CD4, CD8- α , CD8- β , PD-1, and Tim-3 as well as their associated cytokines IL-2, IL-21, IFN- γ , and TNF- α .⁹⁸ Therefore, LAT interacting with immune system is considered to be related to HSV latency, which deserves more investigation in depth.

For restraining LAT latent infection, many promising vaccine approaches are being explored in animals and humans, such as modifying the LAT gene to make its latent period longer or reduce its reactivation for the appropriate vaccine type against HSVs.^{99,100} The M3 virus, which has a partial LAT deletion, was harvested from the infected human embryonic kidney cell line 293T cells and exhibited the lowest pathogenicity during acute and latent infection. Thus, LAT mutant is an ideal candidate for HSV vaccine development. In addition, the new CRISPR/Cas9 technology can now be used to make engineered HSVs for better vaccine and oHSV development.

In addition, RNA silencing and RNA interference have been discovered in plants, animals and human beings.^{101,102} Although it is shown that sRNA1-LAT and/or sRNA2-LAT inhibit the ICP0 gene expression by partial base complementation to maintain latency,⁶¹ its underlining mechanism is not known. We propose that RNA silencing or RNA interference may play an important role in HSV latency and reactivation, which is worth validating in future.

Acknowledgments

This work was partly supported by grants from the National Natural Science Foundation of China (81872412 to XHW). We thank Guangzhou Key Medical Discipline Construction Project (CSZ) and Yangtze University Fellowship to graduate student ZY.

Disclosure

The authors report no conflicts of interest in this work.

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