



Restarting Lytic Gene Transcription at the Onset of Herpes Simplex Virus Reactivation

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ABSTRACT Herpes simplex virus (HSV) establishes a latent reservoir in neurons of human peripheral nerves. In this quiescent state, the viral genome persists as a circular, histone-associated episome, and transcription of viral lytic cycle genes is largely suppressed through epigenetic processes. Periodically, latent virus undergoes reactivation whereby lytic genes are activated and viral replication occurs. In this Gem, we review recent evidence that mechanisms governing the initial transcription of lytic genes are distinct from those of *de novo* infection and directly link reactivation to neuronal stress response pathways. We also discuss evidence that lytic cycle gene expression can be uncoupled from the full reactivation program, arguing for a less sharply bimodal definition of latency.

KEYWORDS JNK signaling, episome, herpes simplex virus, heterochromatin, histone methylation, latency, neurotropic viruses, reactivation, transcriptional regulation

Often we think of viruses as self-serving agents that aggressively replicate to the detriment of the infected cell, but in truth many establish a long-term relationship with their hosts, thereby ensuring a continuous presence and prolonged transmission. Herpesviruses offer a clear example, having evolved a dedicated strategy, termed latency, that limits the pathogenic consequences of infection and avoids immune clearance. For herpes simplex virus (HSV), latency is restricted to postmitotic neurons in the peripheral nervous system (1). Latently infected neurons serve as lifelong reservoirs from which infectious particles can emerge (reactivate) from time to time. In some individuals, repeated reactivation gives rise to painful oral or genital lesions, corneal scarring, nerve inflammation, and even life-threatening encephalitis.

WHAT IS LATENCY?

Herpesvirus latency is defined as the persistence of viral DNA in the absence of infectious viral particles with the potential for these latent genomes to reactivate. For HSV, this refers to latency at the level of individual ganglia because in reality only a fraction of individual infected neurons will reactivate, even in response to a strong stimulus. The clearest molecular hallmark of HSV latency is the expression of several noncoding RNAs known as the latency-associated transcripts (LATs). However, levels of LAT expression also vary between neurons, and HSV latency can be established in the absence of LAT expression, indicating that the transcripts themselves are not required for latency but do contribute to neuronal survival and modulate the efficiency of reactivation (2–4).

In the neuronal nucleus, the 150-kb double-stranded DNA HSV genome exists as a closed circle loaded with regularly spaced nucleosomes. Histones assembled on the viral lytic gene promoters are decorated with posttranslational modifications associated with transcriptional repression (Fig. 1), specifically histone H3 trimethylation at lysine 27 (H3K27me3) and histone H3 di- and trimethylation at lysine 9 (H3K9me2/3) (5–7).

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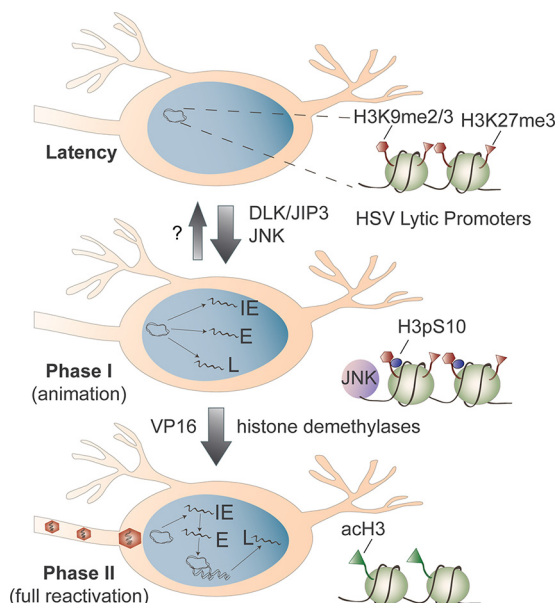


FIG 1 Stepwise reversal of host-mediated epigenetic silencing of the HSV latent genome. During latency, the histones associated with HSV lytic promoters, such as histone H3 trimethylation at lysine 27 (H3K27me3) and histone H3 di- and trimethylation at lysine 9 (H3K9me2/3), are enriched for epigenetic silencing modifications, resulting in a stable chromatin state refractory to transcription by RNA polymerase II. Activation of neuronal stress response pathways triggers DLK/JIP-3-mediated activation of JNK and phosphorylation of histone H3 serine 10 (H3pS10) adjacent to the H3K9me2/3 mark. This dual modification (the methyl-phospho switch) displaces repressive factors and renders the chromatin permissive for simultaneous transcription of the viral IE, E, and L gene mRNAs characteristic of phase I (animation). Viral regulatory factors, such as VP16, synthesized during phase I can then associate with the viral IE promoters and recruit cellular demethylases to remove heterochromatic marks along with histone acetyltransferases that promote hyperacetylation of the viral chromatin. This increases the transcriptional capabilities of viral genes, achieving sufficient levels to initiate HSV DNA replication and assembly of new infectious particles.

Expression of the LATs is thought to modify the chromatin on the viral genome to promote stable but ultimately reversible silencing (8). Repressive chromatin can act as a barrier to positively acting transcription factors, and RNA polymerase II and silencing are augmented by viral microRNAs expressed from the LAT locus that act cooperatively with neuronal miR-138 to suppress translation of key viral mRNAs (9, 10). These blocks result in very reduced levels of viral antigen and prevent viral replication, protecting infected neurons from both intrinsic and acquired immune defenses.

MECHANISMS OF HSV LYTIC GENE EXPRESSION DURING THE FIRST PHASE OF HSV REACTIVATION ARE DISTINCT FROM THOSE OF *DE NOVO* INFECTION

Although the endpoint of reactivation is similar to that of a *de novo* infection, the start points are very different. When HSV infects a permissive cell, it delivers the viral genome together with tegument proteins that ensure high levels of viral gene expression. The tegument factors include VP16, a transcriptional activator that is corecruited with cellular transcription factors HCF-1 and Oct-1/POU2F1 to a VP16-response sequence found in the promoters of the viral immediate early (IE) gene (11). Synthesis of the IE proteins is necessary for early (E) gene expression, which in turn allows HSV DNA replication and late (L) gene expression. Expression of viral genes at each stage requires both viral and cellular proteins, including histone demethylase enzymes that remove repressive chromatin modifications deposited into the genome immediately following infection (12–15).

Reactivation incurs two major differences from *de novo* infection. First, the viral genome is associated with a more compact chromatin structure (16), and second, tegument factors such as VP16 are likely absent. Even if transcription occurs, host and viral microRNAs suppress translation of stimulatory viral factors, such as ICP0.

So how is latency reversed, and how is the switch from one epigenetic state to the other linked to environmental inputs? Our recent work has uncovered a conceptually simple answer in the form of a two-stage reactivation program initiated by a preexisting epigenetic switch (14, 17). The viral response begins with a generalized burst of HSV gene transcription, termed phase I or animation, resulting in the *de novo* synthesis of many viral proteins, including the missing tegument factors (17). This discovery came from *in vitro* studies using primary neurons isolated from peripheral ganglia of prenatal rats or postnatal mice (18, 19). In these models, reactivation can be deliberately induced by interruption of signaling by neurotrophic factors, such as nerve growth factor (NGF), through inhibition of phosphoinositide 3-kinase (PI3K), AKT, mammalian target of rapamycin complex 1 (mTORC1), and/or cap-dependent protein synthesis (20, 21).

Phase I corresponds to a transient wave of viral lytic gene transcription not observed during acute infection. In phase I, IE gene expression is not dependent on the viral transactivator VP16 and the expression of both viral E and L genes occurs even when viral protein synthesis is inhibited, indicating that synthesis of the IE proteins is not required. Likewise, L gene expression is not affected by inhibition of viral DNA replication. Similar disordered patterns of gene expression occur during *ex vivo* reactivation triggered by axotomy in combination with neurotrophin deprivation (22), cementing the notion of a unique mechanism for transcriptional activation from latent HSV genomes. Activation of the late VP16 promoter has also been found to occur independently of IE gene expression in an *in vivo* model of HSV reactivation (23). Finally, phase I gene expression occurs in the presence of histone demethylase inhibitors (14), indicating that viral transcription takes place without removal of repressive histone modifications.

CO-OPTION OF A NEURONAL STRESS PATHWAY FOR PHASE I OF REACTIVATION

At the onset of reactivation, viral proteins are absent and activation of HSV genes must rely on cellular activities. In the studies described above, reactivation was triggered by inhibiting PI3K signaling. This and other triggers, such as withdrawal of trophic support, axotomy, and heat shock, are known inducers of c-Jun N-terminal kinase (JNK) signaling. In most cells, JNKs play an important role in stress response pathways, but in neurons they also control dendritic arborization and synaptic plasticity (24). In response to nerve injury, JNKs are redirected through mobilization of JNK-interacting protein 3 (JIP3) and dual-leucine-zipper kinase (DLK). This can be induced by inactivation of AKT, a negative regulator of DLK, which occurs when neurotrophin signaling is interrupted. Recent studies showed that activation of JNK by DLK/JIP3 is required for phase I, indicating that activation of the JNK stress response can initiate HSV reactivation (14). Although JNK signaling is important in HSV lytic replication (25), the mechanism by which JNK functions to initiate phase I is distinct because JNK activation by DLK/JIP3 is not required during *de novo* infection.

Induction of the JNK stress response pathway results in activation of multiple transcription factors that might be involved in HSV reactivation. Less easily understood is how activation permits transcription of lytic genes associated with histones that carry repressive lysine modifications. Although many studies focusing on histone modifications use antibodies generated against individual modifications, the reality is much more complex, with multiple combinations of histone modifications acting together to control gene expression (26). One way in which gene expression can still be initiated even when histones carry repressive lysine modifications is through phosphorylation on the neighboring serine residue, a process known as a histone methyl/phospho switch (27, 28). This switch is used during phase I and is dependent upon JNK activity and correlated with JNK recruitment to viral promoters (14). Therefore, activation of the JNK pathway results in a chromatin state permissive for transcription, even though repressive histone lysine modifications are maintained.

THE SECOND PHASE OF REACTIVATION CLOSELY RESEMBLES *DE NOVO* INFECTION

Even if all latently infected neurons are exposed to the stress, only a subset will undergo productive reactivation. One idea is that in the reactivating population, sufficient quantities of key viral proteins are made in phase I to cross a threshold that allows a second wave of viral gene expression termed phase II or the synthesis phase, which culminates in the amplification of viral DNA and production of infectious virus (17). Viral gene expression in phase II resembles the cascade of viral gene expression observed during *de novo* infection. Expression of the L genes is dependent on viral DNA replication, and the stimulatory effect of VP16 is evident. As already mentioned, VP16 is recruited together with host cofactors, such as HCF-1, to the IE promoters and facilitates the recruitment of additional chromatin-remodeling and histone-modifying proteins (29). The requirement for VP16 in phase II argues that chromatin remodeling is critical for full reactivation. Likewise, the activities of cellular H3K27 demethylases (UTX/KDM6A and JMJD3/KDM6B) and the H3K9 demethylase (LSD1/KDM1) are also required for the transition to full reactivation (14, 29–31). Interestingly, explant reactivation, which involves significant physical trauma (axotomy) to the neurons, produces new virus much faster than reactivation through PI3K inhibition (32). Explant/axotomy activates multiple stress responses simultaneously (33), and this might compress the biphasic program, accelerating the onset of phase II.

LATENT INFECTION *IN VIVO* IS DYNAMIC

Early studies implied that the outcome of neuronal infection is essentially binary: viruses either express lytic mRNAs and attempt to replicate or express the LATs and become latent (34, 35). However, new findings argue for a less rigid distinction. Use of a tracing method to permanently mark neurons that have expressed lytic genes revealed latently infected neurons with evidence of prior lytic promoter activation (36). Likewise, low levels of lytic mRNA can be detected in the ganglia of latently infected mice, and a recent study found lytic mRNAs in more than two-thirds of latently infected neurons (37). These and other observations challenge the either/or view of HSV latency and raise exciting questions that will direct future research. For example, are these “lytic” mRNAs functional, and do they influence latency, neuronal survival, or some aspect of reactivation? IE protein ICPO, for instance, stimulates LAT expression during latency and alters the chromatin composition of the latent genome (38). Lytic gene activity may also be sensed by the neuron, reinforcing the expression of host antiviral factors that help maintain latency.

CONCLUSIONS AND FUTURE DIRECTIONS

These exciting advances highlight the value of modeling HSV latency and reactivation in both cultured neurons and infected animals. The cultured-neuron models have shown that continuous neuronal signaling is necessary to sustain the repressive chromatin state of the viral episome and expression of latency-associated RNAs. Importantly, this provides new ideas for how transcription might be initiated from epigenetically silenced chromatin in the absence of viral factors. Release from epigenetic silencing through a bivalent histone mark (the methyl/phospho switch) is appealing because it directly links viral transcription to neuronal stress response pathways known to induce reactivation. Host transcription factors activated by the same stress pathways will also benefit from the altered chromatin state through increased access to binding sites and greater RNA polymerase processivity. Although these studies have focused on histone H3 serine 10 (H3S10) phosphorylation, stress-associated kinases can also phosphorylate histone H3 serine 28 (H3pS28), which is immediately adjacent to H3K27me3 and represents a second methyl/phospho switch, triggering gene activation from cellular promoters through polycomb displacement (28). Whether this contributes to the initiation of phase I remains to be determined.

Clearly, there has been much progress, but important questions remain. For instance, what limits the progression from phase I/animation to phase II and full reacti-

vation? What are the specific chromatin-remodeling events that need to occur in each step, and what factors are required? How do changes in the higher-order structure of the episome (chromatin looping) observed during reactivation (39) correlate with these events? How do levels of reactivation and chromatin silencing vary between different types of neurons and between HSV-1 and HSV-2? Lastly, we should not forget that neurons are highly specialized cells and often use nuclear functions differently from other cell types. We have just begun to explore epigenetic processes in a neuronal context, so there will doubtless be more surprises along the way.

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