

REVIEW

Investigating the biology of alpha herpesviruses with MS-based proteomics

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Viruses are intracellular parasites that can only replicate and spread in cells of susceptible hosts. Alpha herpesviruses (α -HVs) contain double-stranded DNA genomes of at least 120 kb, encoding for 70 or more genes. The viral genome is contained in an icosahedral capsid that is surrounded by a proteinaceous tegument layer and a lipid envelope. Infection starts in epithelial cells and spreads to the peripheral nervous system. In the natural host, α -HVs establish a chronic latent infection that can be reactivated and rarely spread to the CNS. In the nonnatural host, viral infection will in most cases spread to the CNS with often fatal outcome. The host response plays a crucial role in the outcome of viral infection. α -HVs do not encode all the genes required for viral replication and spread. They need a variety of host gene products including RNA polymerase, ribosomes, dynein, and kinesin. As a result, the infected cell is dramatically different from the uninfected cell revealing a complex and dynamic interplay of viral and host components required to complete the virus life cycle. In this review, we describe the pivotal contribution of MS-based proteomics studies over the past 15 years to understand the complicated life cycle and pathogenesis of four α -HV species from the *alphaherpesvirinae* subfamily: Herpes simplex virus-1, varicella zoster virus, pseudorabies virus and bovine herpes virus-1. We describe the viral proteome dynamics during host infection and the host proteomic response to counteract such pathogens.

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1 Introduction

The *Alphaherpesvirinae* subfamily of the family *Herpesviridae* includes the genera *simplexvirus* and *varicellovirus* [1].

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Abbreviations: α -HV, alpha herpesvirus; **BHV-1**, bovine herpes virus; **BONCAT**, bioorthogonal noncanonical amino acid tagging; **CNS**, central nervous system; **DIGE**, differential in-gel electrophoresis; **E**, early; **HSE**, herpes simplex encephalitis; **HSV-1**, Herpes simplex virus-1; **IE**, immediate early; **L**, late; **MALDI**, matrix-assisted laser desorption/ionization; **MSBP**, MS-based proteomics; **PABP**, poly(A)-binding protein; **PNS**, peripheral nervous system; **PRV**, pseudorabies virus; **PTM**, posttranslational modification; **SILAC**, stable isotope labeling by amino acids in cell culture; **TMT**, tandem mass tag isobaric labeling; **VZV**, varicella zoster virus

These two genera contain several pathogenic species infecting a broad variety of mammals, which for the purpose of this review will be noted as alpha herpesvirus (α -HV). The large double-stranded DNA genome, virion size, and structure (icosahedral capsid, tegument, and glycosylated lipid envelope), and the latency-reactivation cycle are common features of all α -HVs [2, 3]. These viruses are pantropic and neuroinvasive pathogens that establish persistent (latent) infections in the nervous systems of the natural hosts [4]. Upon reactivation, α -HVs cause diverse effects that vary from mild epithelial lesions, to life-threatening necrotizing brain infections and death [5, 6]. The complex and dynamic interplay between virus and host components is an active area of research. The interactions between viral-viral and viral-cell proteins are constantly modified during productive and latent infection. Moreover, the function and localization of viral and cell proteins are regulated at transcriptional, translational, and posttranslational (PTM) levels. Spread of infection in the highly connected nervous system is regulated by directional movement of viral proteins in axons [7]. Spread in the retrograde direction occurs during primary infection

Table 1. Viral species from the *alphaherpesvirinae* subfamily included in the present review. For each virus, the acronym is shown in the left column, and a brief description of the susceptible host, pathogenesis, and life cycle is described in the right column

BHV-1	Bovine herpes virus-1 (BHV-1) infects cattle causing rhinotracheitis symptoms that predispose animals to secondary bacterial infections resulting in 'shipping fever'. BHV-1 is also associated with conjunctivitis, reproductive tract lesions, encephalitis, and fetal infections. Economic losses result mainly from shipping fever, milk drop, loss of body weight, early pregnancy termination, and death. BHV-1 can establish latency within the natural host [103].
PRV	Pseudorabies virus (PRV) is a pathogen of swine that causes important economic losses worldwide. It is also known by its taxonomic name, suid herpesvirus 1, or by its original name, Aujeszky's disease virus. The disease was originally described as "mad itch" because of the intense itching symptoms. The natural host is the swine, although PRV can infect a broad range of vertebrates including cattle, sheep, dogs, cats, goats, chickens, raccoons, possums, skunks, rodents, rabbits, and guinea pigs. PRV is tropic for both respiratory and nervous system tissues of swine. Young swine are severely affected by PRV infection of the CNS. Older swine show symptoms of respiratory disease. Several live attenuated vaccine versions are used in PRV eradication programs [41]. PRV is commonly used by neuroscientists to study neuronal connectivity and function in animal models [21, 104].
VZV	Varicella-zoster virus (VZV) is a ubiquitous human pathogen that spreads from mucosal epithelial cells, where initial acute infection occurs, to the skin via a T-cell-associated viremia, causing varicella (chicken pox) to young age patients. Viremia and cutaneous infection allow transfer of VZV to sensory nerve ganglia, where a latent infection is established in neurons. Zoster (shingles) is caused by VZV reactivation from latently infected neurons, producing a painful vesicular rash (dermatome). Live attenuated VZV vaccines are effective against varicella and zoster [105].
HSV-1	Herpes simplex virus-1 (HSV-1) is a widespread human pathogen that establishes a lifelong reactivateable infection in neurons of the peripheral nervous system of the host. Upon reactivation, HSV-1 particles travel in axons back to epithelial tissue causing oropharyngeal lesions. Occasionally, the infection will spread to the cornea and cause keratitis, or to the central nervous system to cause herpes simplex encephalitis (HSE) with high mortality rates [2]. Currently, no effective vaccine exists, and the only treatment is based on antiviral drugs that inhibit viral replication. HSV-1 is widely used to study neuronal connectivity and function in animal models [40]. Ongoing human clinical trials are testing the efficacy of modified HSV-1 strains as oncolytic vectors [106].

(from infected epithelial cells to the innervating neuron), and in the anterograde direction after reactivation (from infected peripheral nervous system (PNS) neurons to epithelial cells). This pattern of infection moving into and out of the PNS is a hallmark of α -HVs. The viral and host mechanisms that control viral sorting, transport, and egress from axons and dendrites remain partially understood and are active areas of research [8, 9].

In this review, we focus on four α -HV species (see Table 1) where innovative MS-based proteomics (MSBP) methods have contributed toward understanding basic and clinical aspects of these pathogens [10]. We review two main topics: (i) The viral and host proteins required to complete the replication cycle; and (ii) The cellular response to viral infection and the co-evolution of virus and host (see Fig. 1).

2 Alpha herpesvirus infection and interaction with host proteins

2.1 The composition of alpha herpesvirus virions

Newly assembled α -HV particles egress from infected cells using the host secretory pathway, which is repurposed by viral components. Mature viral particles travel to the plasma membrane inside acidified secretory vesicles to sites of exocytosis where vesicles fuse with the plasma membrane

and release mature virions [11]. The precise global composition of pseudorabies virus (PRV) and herpes simplex virus-1 (HSV-1) mature virions has been well studied by MSBP (Fig. 1A) [12–14]. Both PRV and HSV-1 virions have a conserved capsid structure with many common components [15]. In the HSV-1 study, Loret et al. used highly purified virions and performed in-gel digestion, followed by liquid chromatography and tandem mass spectrometry (nLC-MS/MS). They identified 37 of the 40 previously identified virion components and confirmed the absence of 21 nonstructural proteins [14]. Additionally, four novel virion components were identified (pUL7, pUL23, pUL50, and pUL55). Fifty four percent of all the peptides identified correspond to tegument proteins. In the PRV study [12], Kramer et al. also used highly purified virions and combined two complementary approaches: First they did in-gel digestion and LTQ Orbitrap XL MS, configured with a matrix-assisted laser desorption/ionization (MALDI) source. Second they performed in-solution digestion by filter-aided sample preparation (FASP) and nLC-MS/MS analyses to compensate for sample loss in the first approach. They identified 47 PRV virion proteins, including seven novel viral components (pUL8, pUL20, pUL32, pUL40, pUL42, pUL50, and Rsp40/ICP22). They used proteinase K digestion to exclude nonspecific proteins that could adhere to virions and compared the results with uninfected cell extracts. Additionally, the PRV study identified phosphorylation sites in virion proteins pUL26, pUL36, pUL46, and pUL48. Strikingly, both the HSV-1 and PRV studies identified as many as 49 host

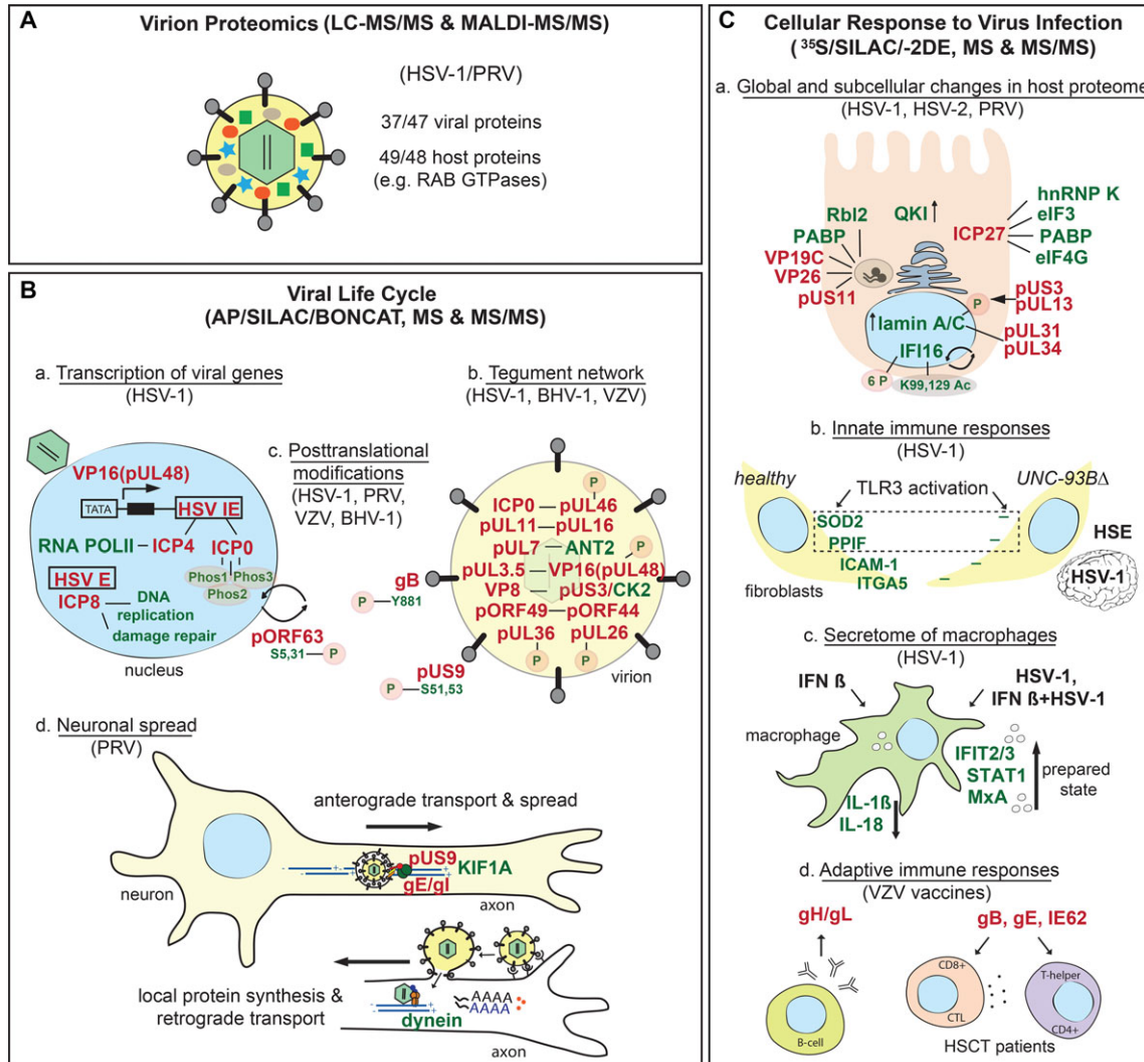


Figure 1. Use of MSBP in understanding alpha herpesvirus biology. (A) Understanding the complex structure of α -HV virions. HSV-1 and PRV virions are comprised of several viral and cellular proteins. (B) Understanding the viral life cycle. (a) Upon α -HV infection and entry, viral IE genes are transcribed in the nucleus with the action of tegument protein VP16 (pUL48) and its cellular interacting partners. IE proteins (e.g. ICP0 and ICP4) interact with several cellular and viral proteins. Cellular interacting partners of ICP4 and E protein ICP8 were identified by MSBP. (b) MSBP has also contributed to understanding the complex tegument network. Several viral and cellular proteins were found to interact with each other and some of them (e.g. pUL26, pUL36, pUL46, pUL48) contained novel PTMs. (c) Two viral envelope proteins (PRV pUS9 and VZV gB) were phosphorylated (P), this PTM is important for the protein function. PTM analyses revealed three phosphorylation regions (Phos1-3) in HSV-1 ICP0, which is a multifunctional nucleocytoplasmic shuttling protein (depicted by arrows). Similarly, VZV protein pORF63 was modified by phosphorylation, which possibly regulates its subcellular regulation. (d) How newly made PRV virions are sorted into axons during neuronal infection was clarified with the identification of KIF1A and pUS9 interaction by MSBP. This interaction is stabilized by two other glycoproteins, gE and gI. The rapid response after PRV entry into axons was also characterized by MSBP, and several host proteins were found to be locally synthesized in axons. Some of these proteins aid retrograde transport of viral particles. (C) Characterizing the changes in the host cell proteome after α -HV infection. (a) Proteins regulating translation (e.g. PABP, hnRNP K, eukaryotic initiation factor 3, eukaryotic initiation factor 4G, QKI), transcription (e.g. Rbl2), and nuclear proteins (e.g. lamin A/C) enabling capsid assembly and egress were characterized by MSBP. Innate immune responses involving DNA sensor IFI16 were identified upon HSV-1 infection by MSBP. (b) Differences in innate immune responses between healthy individuals and HSE patients with UNC-93B mutations were studied detecting differential regulation of SOD2, PPIF, ICAM-1, and ITGA5 in fibroblasts. (c) IFN- β treatment with or without HSV-1 infection in macrophages showed that IFN- β alarms cells by increasing the secretion of proteins such as IFIT2 and 3, STAT1, and MxA (prepared state). HSV-1 proteins counteract these antiviral responses by decreasing the secretion of cytokines such as IL-1 β and IL-18. (c) MSBP was used to characterize the adaptive immune responses to the VZV vaccine Zostavax, and to identify immunogenic proteins to develop subunit vaccines for HSCT patients. Viral and host proteins are shown in red and green, respectively. Interactions between viral and cellular proteins are depicted as a horizontal line. Increase/decrease in protein levels are shown with arrows. All other abbreviations and references are detailed in the text.

proteins that were incorporated into virions, including several Rab GTPases that are relevant for viral trafficking and egress from infected cells (Fig. 1A). The role of these low copy number host proteins in virions is a subject of research and debate. These proteins could function immediately after viral entry into cells, perhaps facilitating establishment of the infection. For example, the presence of host translation initiation and elongation factor in virions could jump start the production of new viral components [13]. Another idea is that virions may incorporate host proteins from one cell type that are not present in a different cell type perhaps allowing a wider infectious range.

Studies on the virion composition of bovine herpes virus (BHV-1) are ongoing (<http://vivo.usda.gov/display/NIFA-1001199-PROJ>).

2.2 Viral and host protein interactions during viral transcription, translation, and replication

After fusion-mediated cell entry, α -HV capsids and some tegument proteins are actively transported by dynein to the cell nucleus where viral DNA is delivered and transcribed. Viral genes are transcribed in three kinetic groups: immediate early (IE), early (E), and late (L) [4, 5]. The timing of gene expression is regulated at transcriptional and post-translational levels by viral and host proteins. The HSV-1 genome encodes five IE genes, called infected-cell polypeptides or ICP's. ICP0, ICP4, ICP22, ICP27, and ICP47 proteins have multiple regulatory and immunomodulatory functions and are essential for a productive infection [16]. IE genes are transcribed efficiently before viral protein synthesis or viral replication primarily because the incoming virion carries the tegument protein VP16 (UL48), which is a transcription activator (Fig. 1B, panel a). After synthesis, the IE proteins modulate the transcription of early viral genes (E), which are required for viral DNA replication, transcription of late viral genes (L), and assembly of progeny virions. IE proteins can either activate or repress transcription of most viral genes. To do so, the IE proteins must interact with the host transcription machinery [17]. Since α -HVs do not encode their own RNA polymerase, the host RNA polymerase II complex (RNA pol II) must be repurposed for transcription of viral genes. Several host general transcription factors (GTF) along with RNA pol II and the viral IE protein ICP4 assemble on viral DNA promoters to form the preinitiation complex (PIC). ICP4 is a 350 kDa homodimeric phosphoprotein containing a DNA binding domain, a nuclear localization signal, and regions that modulate the transcription of viral and host genes including its own repression. ICP4 stabilizes the formation of PICs on viral promoters and interacts with GTFs [18]. Affinity purification (AP) of ICP4 followed by MSBP enabled identification of the proteins involved in this process. Forty-six proteins that co-purify with ICP4 were identified by LC-MS/MS, including 11 GTF components of the TFIID complex and four components of the Mediator complex. TFIID is a protein complex

that includes TATA-binding proteins (TBP) which engage essentially all HSV promoters. Mediator is a large multicomponent complex that can facilitate the entry of RNA pol II into the PIC. Mediator interacts with VP16, further demonstrating the relevance of these interactions to recruit all the necessary components to transcribe IE viral genes. Determining the precise number of cell proteins required for efficient cell and viral replication is an area of active research [19, 20].

Viral IE genes not only play a pivotal role in the precise timing of transcription of E and L viral genes, but they also have multiple functions in regulating both productive and latent infections. Most IE genes are promiscuous transactivators and modulate transcription and translation of host genes, occasionally leading to cytotoxic effects [21]. ICP27 is an HSV-1 IE gene product that co-purifies with translation initiation factors such as poly A binding protein (PABP), eukaryotic initiation factor 3, and eukaryotic initiation factor 4G. In these complexes, ICP27 stimulates or inhibits viral and host mRNA translation (Fig. 1C, panel a) [22]. Viral protein functions, including those of the viral IE proteins, are also regulated by PTMs [23]. PTM sites in viral proteins are readily identified by MSBP (see Section 2.4). The ICP0 IE protein contains multiple phosphorylation sites. MS studies enabled the identification of three regions within or adjacent to transactivation domains of the protein. In total, 11 phosphorylation sites were detected in regions Phos 1, Phos 2, and Phos 3 (Fig. 1B, panel a). These sites were mutated to alanine by site-directed mutagenesis to determine their relevance in ICP0 function after viral infection [24]. Mutations in the phosphorylation sites of ICP0 affected its subcellular localization. The most dramatic case was observed when residues in Phos 3 were mutated and ICP0 localization switched from being almost exclusively nuclear to localize both in the nucleus and cytoplasm. Interestingly, only mutations in region Phos 1 disabled the ability of ICP0 to complement ICP0-null viral mutants.

E genes are expressed after the IE proteins engage the host transcription machinery. E genes have several functions including the reduction of IE gene expression, DNA replication and recombination, and activation of L gene transcription. The viral replication proteins include pUL30 (DNA polymerase), pUL29, and pUL42 (DNA-binding proteins), ORI-binding proteins (pUL9), and the helicase/primase complex (pUL5, pUL8, and pUL52). MSBP studies on ICP8 (encoded by UL29 gene) revealed an extensive list of interacting cellular partners that may be relevant for viral replication. ICP8 is a 128 kDa nonstructural multifunctional zinc metalloprotein that binds single-stranded DNA and localizes to viral replication compartments in the infected cell nucleus. ICP8 is involved not only in viral DNA replication, but also in repression of transcription of parental viral genomes and stimulating transcription of L genes. A study from Taylor and Knipe identified 49 host proteins and 6 viral proteins that interact with ICP8 [25]. Most of the host proteins identified are involved in DNA replication and damage repair, nonhomologous and homologous recombination, and chromatin

remodeling (Fig. 1B, panel a). The authors showed that the cellular nonhomologous end-joining machinery (NHEJ) inhibits viral replication in a mutant murine fibroblast line with a disrupted NHEJ pathway. Some of these cell proteins are recruited by ICP8 to viral replication compartments [25, 26]. After translation of the last wave of viral L proteins, all the players required for assembly of viral particles are available. The many viral and host components required for virion assembly need to be in the right place at the right time and in the precise stoichiometry to complete this elaborate process. Precise targeting of virion components to the diverse cell compartments required for α -HV assembly is only now being understood.

2.3 Alpha herpesvirus tegument proteins and their relevance in infection

As discussed previously, the most abundant proteins in the virion are found in the tegument. It is a proteinaceous complex that fills the space between the nucleocapsid and viral envelope membrane. Such complex tegument is unique to herpesviruses and is comprised of a variety of viral and host proteins, all involved in many viral processes, either immediately after infection or during assembly of the virions. It is not well understood how the tegument components assemble into newly synthesized virions. MSBP has been essential to define and analyze this unusual collection of proteins (Fig. 1B, panel b). Michael et al. showed that deletion of the PRV tegument protein pUL21 resulted in a drastic decrease in the incorporation of three tegument proteins, pUL46, pUL49, and pUS3, into mature virions [27]. The attenuated vaccine strain, PRV Bartha, contains missense mutations in pUL21 and in the unique short (US) region and also failed to package pUL46, pUL49, and pUS3 in virions. In a similar study, the authors used stable isotope labeling with amino acids (aa) in culture (SILAC) and MSBP, to show that the composition of the virion changes when tegument proteins pUS3, pUL47, pUL49, or glycoprotein E (gE) are not expressed. When gE was not expressed, mutant virions had significant reductions in the pUL46, pUL48, and pUL49 tegument proteins. The authors also observed an increase in the amount of a smaller isoform of pUL48 and in host actin in the gE-null virions [28]. In HSV-1, the tegument protein pUL46 modulates the transactivating activity of VP16 (pUL48), however it was not known if other cell or viral proteins interact with pUL46 or if it has additional roles early in infection. Lin et al. showed that HSV-1 pUL46 is phosphorylated in at least 23 sites and specifically interacts with the viral protein ICP0, an E3 ubiquitin ligase [29]. They also demonstrated that pUL46 is partially degraded in a proteasome-mediated manner by ICP0 (Fig. 1B, panel b). This finding is the first report of a viral protein being targeted for degradation by another viral protein during HSV-1 infection. Interestingly, pUL46 was found in a phosphorylated form in mature PRV virions [12].

pUL11 is an HSV-1 tegument protein comprised of 96 amino acids, and it localizes to the cytoplasmic faces of

nuclear- and Golgi-derived membranes in infected cells. This suggests a role for pUL11 in virion secondary envelopment. MSBP studies with UL11 mutants revealed that the protein is myristylated and palmitylated near the N terminus, and these modifications stabilize membrane-binding and mediate Golgi-targeting. pUL11 localizes only to the Golgi membranes but not to nuclear membranes in transfected cells, strongly suggesting that pUL11 interacts with additional viral proteins upon infection. Loomis et al. demonstrated that pUL11 interacts with pUL16, a protein involved in nucleocapsid assembly, and that this interaction is independent of other viral proteins (Fig. 1B, panel b) [30].

Tegument protein networks involving HSV-1 pUL7 and the cell mitochondrial protein adenine nucleotide translocator 2 (ANT2) have been identified by MSBP (Fig. 1B, panel b) [31]. UL7 mutants are able to grow in tissue culture but have cell–cell spread and replication defects. ANT2 is part of the permeability transposition pore (PT) complex and exchanges ATP and ADP on the inner mitochondrial membrane. In apoptotic conditions, ANT2 can form a nonspecific pore. The relevance of this interaction is yet to be determined.

The pUL3.5 tegument protein is not conserved in all α -HVs: PRV, VZV, and BHV-1 genomes contain UL3.5 homologs while HSV-1 and HSV-2 genomes do not. The roles of the pUL3.5 protein in replication may not be conserved: PRV pUL3.5 is required for virion egress, but VZV mutants lacking pUL3.5 (pORF57) grow well in cell culture with normal kinetics. On the other hand, BHV-1 pUL3.5 protein can complement the egress defect of a PRV UL3.5 mutant, suggesting that they have similar functions in egress. Lam and Letchworth identified the BHV-1 UL48 gene product (BHV-1 α -transinducing factor; α BTIF) as a pUL3.5 interacting partner (Fig. 1B, panel b) [32]. α BTIF is the HSV-1 VP16 homolog, a transactivator protein that is incorporated into the virion and promotes transcription of IE genes. Although the authors did not demonstrate the transactivating activity of α -BTIF, they showed pUL3.5 and α -BTIF co-localization in the Golgi of infected cells that depends on 20aa on the N-terminus of pUL3.5.

Another BHV-1 tegument protein studied by MSBP is the major tegument protein VP8. The homolog of VP8 in HSV-1 and PRV is VP13/14 (UL47). Labiuk et al. demonstrated that VP8 is phosphorylated by the viral kinase pUS3 and cellular kinase CK2 and that they co-localize to the nucleus of infected cells (Fig. 1B, panel b) [33].

In VZV, the nonessential tegument protein, pORF49, functions as a cell tropic factor in cell culture. ORF49 null mutants produce small plaques and have low infectious virus yield. The interaction between pORF49 and pORF44 is eliminated by mutation of a single amino acid of pORF44 [34]. The authors identified pORF49 residues 41–44 as the pORF44-binding interface. The phenotype of ORF49 delta 41–44 was comparable to that of ORF49-null mutants, suggesting that the pORF49-44 interaction is important for VZV infection and efficient production of progeny virus. The homolog of pORF49 in HSV-1 is the nonessential protein pUL11, which

is involved in secondary envelopment of virions. In HSV-1, the pORF44 homolog is pUL16, which functions in viral entry and nuclear and cytoplasmic egress. MSBP has been essential to unveil not only the complex composition of α -HV tegument but also the plethora of functions that tegument proteins perform during infection. Tegument proteins include all three kinetic groups of proteins (IE, E, L), suggesting their essential role in every step of infection (Fig. 1B, panel b).

2.4 PTMs during alpha herpesvirus infection

MSBP is a powerful approach to globally identify PTMs in complex settings like an infected cell. Here we provide some examples illustrating the relevance of PTMs in α -HV infection. The HSV-1 genome encodes more than 74 proteins, and most of them are modified after translation. So far, the most abundant PTMs detected on viral proteins are phosphorylation and glycosylation. Viral infection also affects cellular protein PTM [35]. A recent study of the HSV-1 proteome identified 10 novel ubiquitylation and 95 new phosphorylation sites in a macrophage cell line lysate 8 h after infection [23]. Kramer et al. detected phosphorylated PRV proteins pUL26, pUL36, pUL46, and pUL48 in purified mature virions (Fig. 1B, panel b) [12]. Authors speculate that PRV protein phosphorylation might activate viral proteins for specific functions at very early stages of infection. When infected cells are treated with inhibitors of DNA replication, both the protein abundance and the degree of PTMs change [23]. The IE gene product ICP0 contains 11 phosphorylation sites across three regions Phos 1, Phos 2, and Phos 3, as detailed in section 2.2 [24]. When the phosphorylated residues in Phos 1 are mutated to alanines, ICP0 is unable to complement an ICP0 null HSV-1 mutant (Fig. 1B, panel a). PTMs are also important for efficient axonal transport of PRV in infected neurons. The viral phosphoprotein pUS9 contains two serine phosphorylation sites (S51 and S53) (Fig. 1B, panel c) [36]. When these residues are mutated to alanines, anterograde neuronal transport of PRV particles is impaired both *in vitro* and *in vivo*.

In VZV, Oliver et al. identified a novel immunoreceptor tyrosine-based inhibition motif (ITIM) in the cytoplasmic domain of viral glycoprotein B (gB) [37]. gB is a conserved protein across α -HVs, and it is essential for fusion-mediated viral entry into susceptible cells. The authors showed that the tyrosine in position 881 of gB (Y881) is phosphorylated and that this PTM is essential for gB function (Fig. 1B, panel c). When VZV gB Y881 is mutated to alanine, cell–cell spread is reduced *in vitro*, and skin infection is severely impaired *in vivo*. The Y881 phosphorylation-dependent cell–cell fusion mechanism may involve novel intracellular signaling through the gB cytoplasmic motif that is essential for skin pathogenesis. This study also provides evidence that ITIMs have functions different from their known role in immune modulation. Another example of VZV PTM function involves pORF63. pORF63 is a 278aa IE protein, important for maintenance of viral latency, and shares homology with the HSV-1

IE protein ICP22. Mueller et al. identified six phosphorylation sites in pORF63 including the novel sites Serine5 and Serine31 (Fig. 1B, panel c) [38]. The authors did not analyze the contribution of these PTMs to VZV infection, but suggested based on bioinformatics analysis, that a novel cellular kinase could be phosphorylating these residues. PTMs add another layer of regulation for interactions of viral and host proteins. They can affect protein function, subcellular localization, interaction with other proteins and stability.

2.5 Neuroinvasion by alpha herpesvirus

After acute infection of epithelial tissue, newly made α -HV particles invade PNS axons innervating such tissues and undergo fast transport toward the distant cell body. To accomplish this long distance transport, viral nucleocapsids must engage microtubule-dependent motors to move from axons to neuron cell bodies (retrograde motion). After reactivation from a latent infection, particles must reverse direction and move into axons for transport back to the peripheral tissues (anterograde motion) [7, 39–41]. Understanding how efficient invasion of axons is stimulated and how viral particles engage different motors for directional transport has been a challenge. However, recent work with MSBP has made significant contributions toward identifying the complicated interaction between viral and host components required for viral spread within and between neurons. Koyuncu et al. used bioorthogonal noncanonical amino acid tagging (BONCAT) and LC-MS/MS to show that PRV infection of axon induces rapid, local protein synthesis, including proteins involved in cytoskeletal remodeling, intracellular trafficking, signaling, and metabolism [42]. The rapid translation of axonal mRNAs appears to be required for efficient PRV retrograde transport and infection of cell bodies (Fig. 1B, panel d). Strikingly, the axonal damage response in uninfected neurons also induces local synthesis of similar proteins. There is evidence that damage signals and viral particles compete for retrograde transport. The hypothesis is that a pre-existent host response is repurposed for efficient PRV neuroinvasion.

α -HV genomes do not encode motor proteins and rely on the transport machinery of their hosts to spread efficiently in cells. The motors used for both retrograde and anterograde transport of viruses in neurons and other polarized cells have been identified [43, 44]. Affinity purification and MSBP studies revealed that newly made PRV virions in transport vesicles engage the microtubule motor protein Kinesin-3 (KIF1A) for sorting into axons and subsequent anterograde transport (Fig. 1B, panel d) [45]. Kinesin-3 interacts with the small viral membrane phosphoprotein pUS9, to move PRV particles that are contained in a transport vesicle. pUS9 is essential for efficient *in vivo* axonal sorting and anterograde transport of PRV and BHV-1 [46, 47]. Two additional viral glycoproteins, gE and gI, are needed to facilitate or stabilize the interaction between Kinesin-3 and PRV US9 (Fig. 1B, panel d) [48]. The role that

HSV-1 pUS9, gE, and gI play in anterograde transport of viral particles is under study [49, 50].

3 Cell response to viral infection

3.1 Global proteomic profiling of alpha herpesvirus-infected cells

Viral infection alters the proteomic profile of host cells. Some responses reflect the cell autonomous, intrinsic early warning signals in response to infection (danger signals), while others reflect the directed interactions of viral proteins to produce a productive environment for viral replication. How many of these responses are general, and how many are specific both to virus and to cell type? Systematic studies of viral transcription profiles are useful, but they cannot report on protein composition, stability, or modifications (see for example [35]). Early studies compared the mock-infected and HSV-1-infected cell proteome using ³⁵S pulse-labeling and 2DE [51]. While HSV-1 infection strongly inhibited host protein synthesis, the synthesis of a number of acidic proteins and basic ribosome proteins was maintained or stimulated. The specific identities and functions of most of these dynamic proteins were not known. Now, MS and protein sequence databases enable the analysis of the complex virus–host interactions.

Antrobus et al. combined 2DE with LC-MS to perform the first global proteomic analysis of HSV-1 infected cells [52]. Uninfected and infected Hep-2 whole cell extracts were analyzed with 2DE, identifying 103 protein spot changes. Subsequent LC-MS/MS of these spots identified a total of 144 individual proteins involved in a variety of biological processes, including DNA replication, chromatin remodeling, pre-mRNA splicing, mRNA stability, and the ER stress response. Similar whole-cell proteomic analysis of PRV-infected bovine kidney cells has also been reported [53]. Proteins from infected and uninfected cell extracts were fractionated by affinity solid-phase extraction prior to 2DE and MALDI-MS/MS. The use of SILAC allowed differentiation and quantification of isotopically modified peptides that are chemically identical, while prefractionation of whole cell extracts enhanced yield as well as resolution of proteins. Changes were observed for 103 protein spots, including 55 individual proteins that are representatives of stress response proteins, constituents of the nuclear lamina, heterogeneous nuclear ribonuclear proteins (hnRNPs), ribosome proteins, proteins involved in trafficking and transport, as well as other biological processes. Only a small portion of host cell proteins was altered during early stages of infection by either HSV-1 or PRV. This finding is consistent with the earlier study using only 2DE analysis [51]. Both studies showed that the expression levels of lamin A/C and hnRNP K were affected by infection. The HSV-1 pUS3 [54–56] and HSV-2 pUL13 [57] protein kinases can phosphorylate lamin A/C, while HSV-1 pUL31 and pUL34 can cause conformational changes to lamin A/C (Fig. 1C, panel a). These changes affect the subcellular localization of lamin

A/C and may be involved in regulating viral genome replication and transcription [59], targeting pUL31 and pUL34 to lamina, and regulating viral capsid nuclear egress [58]. The shift of lamin A/C to more acidic isoforms during PRV infection indicates that HSV-1 and PRV infection affect common biological processes [53]. Other changes after HSV-1 infection involve hnRNP K, which forms a complex with ICP27 and host p32 and CK2 proteins. CK2 then phosphorylates the other three complex components in an ICP27-dependent manner. The formation of this complex may disrupt or modulate pre-mRNA splicing, facilitating splicing-independent nuclear export of viral transcripts [60–62]. hnRNP K also modulates HSV-1 egress from infected cells [63]. It is striking that no other proteins are shared among the lists of proteome changes for either PRV or HSV-1. While technical differences may explain this lack of congruence, it may be that cells sense the two α -HVs differently or that these distantly related viruses have evolved different interactions for efficient replication. Global proteomic studies have provided us with the fundamental tools to examine and address such specific differences between viruses and infected cells.

3.2 Subcellular proteomic profiling of alpha herpesvirus infected cells

Monitoring the global proteomic profile in host cells often obscures subtle alterations in subcellular fractions such as the cytoplasm, ribosomes, and nucleus [51–53]. Studies utilizing fractionation methods have expanded our knowledge of the host response to α -HV infection.

Cytosolic, microsomal, and nuclear proteomic changes were analyzed separately in HSV-1-infected human hepatoma cells. In one study, biochemically fractionated cytosolic and microsomal samples were analyzed by 2D-DIGE and nLC-MS/MS [64]. In this method, proteins from mock-infected and infected cells were differentially labeled and fractionated in the same gel. The expression levels of 12 cytosolic and 6 microsomal proteins involved in apoptosis, signal transduction, and ER-associated degradation pathway, were altered. In another study, Sanchez-Quiles et al. analyzed nuclear isolates from HSV-1-infected hepatoma cells by two separate approaches, 2D-DIGE and tandem mass tag isobaric labeling (TMT) [65]. The two approaches identified 24 and 38 proteins, respectively, with significant changes. These two techniques complemented each other in that 2D-DIGE provided distinct information about the different isoforms and PTMs of the same proteins, while the TMT approach enabled better resolution and detection. Many of the identified proteins play roles in cell cycle regulation and RNA homeostasis. One protein, quaking (QKI), was further validated to promote viral protein synthesis. Only one protein, RuvB-like 2 (RBL2), was common in both studies, showing increased expression in the microsomal fraction at 4hpi and reduced expression in the nucleus at 8hpi. RBL2 is an antiapoptotic ATPase and putative DNA helicase (Fig. 1C, panel a) [66] whose synthesis is increased after HIV infection [67]. RBL2 expression also

interferes with influenza replication [68]. The function of RBL2 is not clear during α -HV infection.

Despite the fact that different HSV-1 strains and cells were used, the sampling times were not uniform, and the proteomic methods were not identical, global and subcellular proteomic profiling gave similar results [52, 64, 65]. The expression levels of four nuclear proteins (lamin A/C, cofilin-1, hnRNP C1/C2, and NonO/P54) and three cytosolic proteins (hnRNP K, UPF0160 protein MYG1, and FK506 binding protein) were consistently altered after HSV-1 infection.

The proteome of fractionated ribosomes changes after HSV-1 infection. Greco et al. identified ribosomal and viral proteins using N-terminal sequencing and MS [69]. The viral proteins VP19C, VP26, and multiple phosphorylated forms of pUS11, as well as a host protein poly(A)-binding protein 1, were the only nonribosomal proteins consistently associated with the ribosome after infection (Fig. 1C, panel a). Subsequent studies also showed that the HSV-1 IE protein, ICP27 interacted with PABP, eukaryotic initiation factor 3 (eIF3), and eukaryotic initiation factor 4G (eIF4G) [22]. Together, these studies demonstrate the value of MSBP in determining the proteins that regulate protein synthesis and the selective translation of proteins after HSV-1 infection.

Fractionating cellular compartments has been extended to the axons of neurons infected with α -HVs. The efficient invasion by α -HVs of peripheral axons after infection of epithelial surfaces is poorly understood. Most viruses that infect epithelial surfaces do not infect axons that innervate that tissue. It is well known that early in development, axons can autonomously respond to various guidance stimuli as the early nervous system is established. In addition, when damaged, axons can respond quickly by synthesizing new proteins. Indeed, a subset of mRNAs and the complete protein synthesis machinery are localized to axons in uninfected neurons [70]. Local translation in axons, not in the distant cell body, regulates growth cone navigation and integrity during development and regeneration. Such retrograde communication of axons with the cell body in adult neurons plays a key role in axonal damage signaling [71–76]. As mentioned in section 2.5, Koyuncu et al. used primary neurons cultured in chambers to physically separate axons from cell bodies [42]. Authors reported that PRV infection of isolated axons also induces rapid and local protein synthesis. They identified the newly synthesized axonal proteins using BONCAT followed by MS/MS (Fig. 1B, panel d). These nascent proteins include participants in cytoskeleton remodeling, intracellular trafficking, signaling, and energy metabolism pathways. Interestingly, this *de novo* protein synthesis is a requisite for transport of virus particles in axons. The two levels of fractionation—physically collecting only axons and BONCAT to purify and enrich for newly made proteins—combined with highly sensitive detection of MS, enabled analysis in molecular detail. While proteomic profiling of subcellular compartments provides insight into local changes within cells after α -HV infection, additional studies comparing different subcellular compartments, cell types, and α -HV species are needed.

3.3 Host immune response after alpha herpesvirus infection

An effective host immune response will contain and clear most viral infections [77]. The first line of defense is the cell autonomous, intrinsic defensive response within the first infected cell where pathogen sensing converge on the production of inflammatory cytokines such as type I interferon (IFN). The endocrine and paracrine signaling of inflammatory cytokines activates the expression of a plethora of cellular genes that function in controlling the virus infection in the infected cell, and in protecting surrounding uninfected cells from infection. The inflammatory cytokines also serve as important regulators of the adaptive immune response where T-lymphocytes and B-lymphocytes elicit specific antiviral mechanisms through cellular and humoral immune responses, respectively. Many investigators have studied the role of immune responses in controlling productive α -HV infection or promoting a quiescent latent infection. Here we review the studies where MSBP has provided insight into the immune mechanisms that regulate α -HV infection.

3.3.1 Early sensing of alpha herpesvirus DNA in infected cells

The first and foremost step of mounting an appropriate immune response is sensing the invading pathogen through multiple pattern recognition receptors. Knowledge of how and where the nucleic acid of DNA viruses, such as herpesvirus and poxvirus, is detected within the cell was limited until the discovery of DNA sensors over the past decade. The DNA sensors, γ -IFN inducible protein 16 (IFI16), was first revealed during a proteomic search of vaccinia virus (VACV) DNA-binding partners [78]. IFI16 also detects HSV-1 DNA both in the nucleus and the cytoplasm. This interaction leads to expression of pro-inflammatory cytokines such as IFN- β , chemokine ligands 10 (CXCL10), interleukin 6 (IL-6) and tumor necrosis factor (TNF). While herpesvirus genomes replicate in the nucleus, poxvirus DNA is replicated in the cytoplasm. Therefore, IFI16 must be localized in both the nucleus and the cytoplasm. Indeed, Li et al. discovered that while the induction of IFN- β production by HSV-1 relies on a nuclear localization sequence (NLS) in the IFI16 protein, this NLS is not required for detection of VACV DNA [79]. Combining immunoaffinity purification with MS, Li and colleagues further identified six phosphorylation and nine acetylation (Ac) sites in the IFI16 protein sequence (Fig. 1C, panel a). Strikingly, acetylation of two lysines (K99 and K129) within the NLS inhibited nuclear localization of IFI16. These analyses were the first to suggest that localization of a DNA sensor (IFI16) could be regulated by PTM. It will be important to know what host and viral gene products aid in regulating precise intracellular localization. Thus, further MSBP studies are required to reveal these specific mechanisms.

3.3.2 The innate immune system controls herpes simplex encephalitis

Herpes simplex virus encephalitis (HSE), a disease caused by the rare infection of the CNS by HSV, may result from deficiency of type I and III IFN production in response to Toll-like receptor 3 (TLR3) activation during HSV-1 infection [80]. This conclusion follows from five genetic etiologies of HSE (UNC-93B, TLR3, TRAF3, TRIF, TBK1) that are all involved in TLR3 signaling. The story is more complicated because not all HSE patients harbor deficiencies in these five genes. Moreover, 30% of HSE patients do not have deficiencies in TLR3 induced type I and III IFN production. These observations prompted Pérez de Diego et al. to use MSBP to search for other factors regulating the TLR3 signaling pathway or other independent pathways that may be associated with HSE susceptibility [80]. The proteomic changes in fibroblasts from healthy and HSE patients were compared after TLR3 activation. The HSE fibroblasts were obtained from patients with UNC-93B deficiency or an unknown deficiency and analyzed with SILAC/MS (Fig. 1C, panel b). The presence of superoxide dismutase 2 (SOD2) and peptidyl-prolyl cis-trans isomerase F (PPIF) was detected in fibroblasts from both healthy individuals and those from the patient with unknown deficiency but not in fibroblasts from the patient with UNC-93B deficiency. SOD2 is an antioxidant enzyme strongly increased after TLR3 activation in macrophages and HSV-1 infection in neurons. It may be involved in protecting these cells from oxidative stress and apoptosis [81, 82]. PPIF is a part of the mitochondrial permeability transition pore in the inner mitochondrial membrane and is a key protein regulating neuronal cell death [83–85]. These genes might be the downstream effectors in the TLR3 pathway which are defective in patients with genetic predisposition to HSE. Furthermore, the expression of ICAM-1 (an intercellular adhesion molecule) and ITGA5 (integrin alpha 5) were significantly increased only in healthy individuals and not in any HSE patients, suggesting their role in TLR3-independent pathways that affect HSE susceptibility [86] (Fig. 1C, panel b). Variation on a population scale has posed a formidable barrier for the analysis of this data because large variations are observed even among healthy individuals. In the simplest case, we do not have a precise definition of what it means to be “healthy”. Proteomic analysis of large cohort studies may help to address these challenges [80].

3.3.3 The alpha herpesvirus secretome after infection of macrophages

The complex set of macromolecules secreted by cells after infection (the secretome) affect local and global responses. For instance, pro-inflammatory cytokines produced and secreted from infected cells, will not only inform and prepare uninfected neighboring cells, but also attract innate and adaptive immune cells to the site of infection. Traditional methods

of examining the secreted components of an infected cell have been limited in scope and sensitivity. Miettinen et al. did a global profiling of the secretome from IFN- β treated and HSV-1-infected human primary macrophages [87]. They used 4-plex labeling of primary amino groups in intact proteins using isobaric tags for relative and absolute quantitation (iTRAQ) combined with LC-MS/MS analysis. They found that IFN- β treatment, HSV-1 infection, and IFN- β priming prior to HSV-1 infection, increased the secretion of 46, 161, and 399 human proteins, respectively. These observations indicate that cells exposed to IFN- β , were probably in a “prepared” state rather than a fully responsive state (Fig. 1C, panel c). Once these “prepared” cells were infected, there was a massive increase in the amount and type of secreted proteins that are likely to control the subsequent spread of infection to other cells. Another interesting finding was that 80% of the secreted proteins were known exosomal proteins. This non-classical vesicle-mediated protein secretion system appears to be activated by IFN- β priming or HSV-1 infection. Proteins related to immune and inflammatory responses, IFN-induced proteins, and endogenous danger signal proteins (such as IFIT2, IFIT3, STAT1, and MxA) were secreted. They also noted a depletion of IL-1 β and IL-18 in HSV-1-infected macrophages, suggesting that inflammasome function may be specifically antagonized (Fig. 1C, panel c) [87]. Secretome profiling has been done for cells infected by other viral families [88, 89]. It will be interesting to compare the secretome of different types of cells under various infectious conditions to define the cell, tissue, and system level responses in both natural and nonnatural hosts.

3.3.4 Adaptive immune responses to alpha herpesvirus infection

The activation of adaptive immune responses is essential for effective control of α -HV during both productive and latent infection [90]. Although most α -HV infections induce a robust humoral immune response, the role of this response in controlling infections is not clear. Many studies point out specific antibody interactions with essential viral processes that may contribute to effective control of infection. For example, a specific antibody reconstructed from a Zostavax-immunized male patient, targeted the VZV gH/gL complex, that is important in cell–cell fusion [91]. Since cell–cell spread of VZV relies in part on activities of glycoproteins gB, gH, and gL on the surface of infected cells, this antibody from an immunized patient suggests that it might function in limiting the spread of infection (Fig. 1C, panel d). The efficacy of antibodies is limited by their avidity. Pasman et al. showed enhanced potency of an antibody against BHV-1 by creating a single-chain variable antibody, whose variable regions (antigen recognizing regions) of the light and heavy chain were covalently linked [92]. Q-TOF MS analysis of the single-chain variable antibody suggested that it formed a multimer, which was later shown to increase the potency of this antibody. This hold promise for

direct transfer of antibodies to infected individuals (passive immunity).

Cellular immune responses mediated by T cells limit viral spread and are thought to be required to maintain a latent infection in immune-competent individuals. Immune-deficient or compromised individuals are more prone to α -HV reactivation. For example, VZV reactivates in 25–40% of patients after allogeneic hematopoietic stem cell transplantation (HSCT) [93]. Boosting cellular immunity by the current live VZV vaccine is not appropriate for these immunocompromised patients. A possible solution arises from MSBP study by Kleemann et al. Using HPLC and MS, they identified VZV proteins gB, gE, and IE62, as potent activators of both CD4+ and CD8+ T cells from HSCT patients upon onset of VZV reactivation (Fig. 1C, panel d) [93]. These viral components may be used to activate specific T cells for passive transfer or for use as subunit vaccines suitable for immune-compromised patients.

Single cell mass cytometry is a powerful method. This technology is similar to conventional flow cytometry but allows simultaneous labeling of over 40 proteins per cell by metal-isotope labeled antibodies. It is possible to determine individual cell states and phenotypes based on single-cell mass differences [94]. This method was used to study human tonsil T cells infected with VZV [95]. These cells spread VZV to the skin of infected hosts to cause varicella. In contrast to the idea that VZV has a tropism for skin tropic T cells, these authors found that VZV infection induces or enhances these properties in a variety of T cells by increasing the expression of relevant surface proteins regardless of the original profile of the cell. The increased expression of such surface proteins was triggered by Zap70 and Akt signaling pathways in a T-cell receptor (TCR)-independent manner. The use of single-cell mass cytometry changes our view of proteomic data from a population level to single-cell level, providing important information about the heterogeneity within the infected tissue.

4 Nonproteomic use of MS in alpha herpesvirus research

MS is also used to identify molecules other than proteins involved in α -HV infections. For example, direct comparison of MALDI-TOF spectra generated from uninfected, HSV-2, or VZV infected cells, showed distinct differences in many mass species that can be used for detection and specific identification of infected cells [96]. In addition, PCR coupled with MS to identify viral nucleic acids from cerebrospinal fluid samples has proven to be a valuable tool for early diagnosis of neurological disorders correlated with infection by neurotropic viruses such as HSV-1 and VZV [97]. MS facilitated the analysis of the chemical composition of *Thymus capitata* extracts, which have a potential noncytotoxic antiviral effect against BHV-1 [98].

Maybe one of the more challenging and useful nonproteomic MS technologies is to measure changes in small molecule metabolites before and after virus infection. Vastag et al. discovered marked differences in the metabolome after human cytomegalovirus (HCMV) or HSV-1 infection of primary fibroblasts [99]. HCMV infection increases components driving the tricarboxylic acid cycle (TCA) and fatty acid biosynthesis, while HSV-1 infection promotes pyrimidine biosynthesis. Such studies provide novel targets for antiviral drug discovery because when these metabolic pathways were altered by inhibitors, infectious virus production was affected.

MS techniques can also be used to identify complex oligosaccharides. Liu et al. used MS to characterize the structure of a heparan sulfate octasaccharide that binds to gD [100]. Viral glycoprotein gD uses cell surface heparan sulfate as the cell receptor to infect susceptible cells. The authors used MS to analyze a cell heptasulfated octasaccharide that binds to gD with an affinity of 18 μ M.

5 Conclusion and perspective

We reviewed the state of the art of proteomic methods used in the last 15 years to study α -HVs. The literature is limited to studies done with HSV-1, PRV, VZV, and BHV-1 (Table 1). We reviewed work done to identify the viral and host proteins important for the transcription of viral genes and the recruitment of host RNA polymerase II. We also reported studies defining the proteome of mature extracellular HSV-1 and PRV virions. A curious finding is that the number of host proteins in virus particles exceeds that of the viral components. The function of these host proteins is not well understood. We reviewed reports that analyzed the proteome of neurons and axons infected with different α -HVs to understand the mechanisms of retrograde and anterograde transport. Finally, we discussed how PTMs of host and viral proteins change after infection (Fig. 1A and B).

We also reviewed the literature that used global MSBP analysis of infected cells as well as subcellular organelles after α -HV infection. In particular, we examined work describing the immediate intrinsic defense responses of cells. MSBP has contributed extensively to understanding how cells sense and react to incoming α -HV DNA, how innate immune signaling may affect HSE pathogenesis, how the secretome components of an infected cell change, and how antibodies and T-cell activation may limit viral infection (Fig. 1C).

Many of the reports described new interacting partners of a specific viral protein. The most common technique was to isolate the protein of interest (and interacting partners) by affinity purification, followed by LC-MS/MS. Affinity purification of fusion proteins labelled with green fluorescent protein (GFP) with a GFP antibody, is a powerful procedure to preserve protein complexes and minimize nonspecific interactions prior to analyses. GFP tagged proteins can be readily visualized in living cells, enabling determination of the relationship of

phenotype to localization. Using GFP antibodies broadens the range of protein targets that can be purified when specific antibodies against the protein of interest are not available [101]. This technology is also useful when purifying proteins from virus infected cells where the GC content is low, as found for VZV [102]. We note that many of the papers reviewed here study the composition of a protein interacting network without detailed validation of the function of such interactions. A list of viral interactors is a strong start, but without functional investigation by other techniques (e.g. viral or cell mutants, RNAi, shRNA, qPCR, microarrays, sequencing, among others), the significance is uncertain.

The vast majority of published MS studies with α -HVs have used epithelial cells and fibroblasts. Considering that neurons play significant roles in the life cycle of most α -HVs, it is important that more proteomic studies of infected neurons in vitro, in vivo, or ex vivo are completed. Only a few studies use MSBP analysis of α -HV infection of neurons [42,45,48,107]. These studies all focused on productive infection. Analyzing samples obtained from latently infected and reactivating neurons will be revealing.

In summary, MSBP is a relatively new field of study that has already contributed to the understanding of α -HVs. As the technology related to MS and subcellular fractionation evolves and improves, we expect to see more MSBP studies that will capture the changing composition of organelles and subcellular structures in α -HV-infected cells and tissues. MSBP studies of single infected cells will enable the analysis of spatiotemporal changes after viral infection with higher resolution. Finally, these studies produce an enormous amount of information, so it is important that proteomics data are made available in public databases. The data will then enable comparison of results by different laboratories, accelerating data analysis, validation, and fostering collaborations.

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