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Human borna disease virus infection impacts host proteome and histone lysine acetylation in human oligodendroglia cells

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Abstract

Background: Borna disease virus (BDV) replicates in the nucleus and establishes persistent infections in mammalian hosts. A human BDV strain was used to address the first time, how BDV infection impacts the proteome and histone lysine acetylation (Kac) of human oligodendroglial (OL) cells, thus allowing a better understanding of infection-driven pathophysiology in vitro.

Methods: Proteome and histone lysine acetylation were profiled through stable isotope labeling for cell culture (SILAC)-based quantitative proteomics. The quantifiable proteome was annotated using bioinformatics. Histone acetylation changes were validated by biochemistry assays.

Results: Post BDV infection, 4383 quantifiable differential proteins were identified and functionally annotated to metabolism pathways, immune response, DNA replication, DNA repair, and transcriptional regulation. Sixteen of the thirty identified Kac sites in core histones presented altered acetylation levels post infection.

Conclusions: BDV infection using a human strain impacted the whole proteome and histone lysine acetylation in OL cells.

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Introduction

Borna disease virus (BDV), a member of the family Bornaviridae in the order Mononegavirales, is a neurotropic, enveloped virus with a non-segmented, negative-strand (NNS) ribonucleic acid (RNA) genome (de la Torre, 1994; Schneemann et al., 1995). BDV replicates in the cell nucleus and persistently infects a wide variety of mammal species including humans (Bode and Ludwig, 2003; de la Torre et al., 1996a; Iwata et al., 1998; Kinnunen et al., 2013; Ibrahim et al., 2002; Koster-Patzlaff et al., 2007), canine disease (Muller et al., 1995), and measles virus (Baczko et al., 1995). BDV is associated with several neurotropic NNS RNA viruses including BDV (Bode et al., 1996; de la Torre et al., 1996b). Laboratory-adapted and wild-type (e.g. human) BDV strains have been shown to differentially affect various host cell types (Li et al., 2013; Poenisch et al., 2009; Williams et al., 2008; Wu et al., 2013). Using a natural human BDV strain appeared particularly advantageous to provide better insight into the pathological consequences of human OL cell infection than laboratory strains from animal origin.
of a bipolar patient should eventually support our understanding of BDV’s impact on neuropathogenesis.

BDV affects the expression of several host mRNA transcripts (Carbone et al., 2001), but the mechanisms remain unclear. The recent discovery of histone lysine acetylation (Kac) as a modulator of gene expression in response to virus infection has brought fresh insight into viral epigenetic regulation (Ferrari et al., 2012). BDV infection has been shown to affect site-specific histone acetylation in cortical neurons in vitro (Suberbielle et al., 2008); however, the histone Kac profile of BDV-infected OL cells remains unknown.

In this study, we hypothesized that BDV infection epigenetically impacts the OL cell proteome through histone Kac. Therefore, using an integrated quantitative proteomic approach assisted by bioinformatic analysis, we comprehensively investigated the proteome profile and constructed a histone Kac atlas of BDV-infected OL cells.

Results

BDV infection of OL/BDV cells

To confirm successful BDV infection of OL/BDV cells, RT-PCR, Western blotting, and immunofluorescence assays were performed to examine the major markers of successful BDV infection – p24 and p40 RNA and protein levels (de la Torre, 1994). Rapid spread of the virus infection was observed in tissue culture; 100% of the cells were infected 14 days post-infection, while 100% of control cells remained non-infected (Supplemental Fig. 1).

Proteomic profiling

We applied SILAC labeling-based proteomics to comparatively quantify the host proteome of OL/BDV cells and control cells. A scheme of the experimental workflow is shown in Fig. 1. In total, 4436 non-redundant proteins were identified, of which 4383 proteins displayed quantifiable differential expression levels in response to BDV infection (Supplemental Table 1). Among these, 1572 proteins displayed a greater than or equal to 1.5-fold increased expression and 165 proteins displayed a lesser than or equal to 1.5-fold decreased expression in response to BDV infection. The proteomic dataset was divided into four quantiles (Q1-Q4) based on the cumulative distribution of SILAC L/H ratios: Q1, less than 15%; Q2, 15–50%; Q3, 50–85%; and Q4, greater than 85%. Enrichment analyses were separately performed in each quantile, and the overrepresented annotations were clustered through one-way hierarchical clustering for comparative analysis.

BDV infection affects transcription factors

We found 201 transcription factors in the whole OL cell proteome (Supplemental Table 2), of which 84 transcription factors showed significantly increased expression and 11 showed significantly decreased expression in response to BDV infection. Only 30 of these significantly differentiated transcription factors were mapped onto the KEGG database. The highest-ranking canonical KEGG pathways are listed in Table 1.

Bioinformatic analysis

We analyzed the quantifiable proteome data set for three enrichment gene ontology (GO) categories: biological process, molecular function, and cellular compartment (Fig. 2A–C). Using Interpro domain enrichment analysis, we analyzed the domain features of those enriched proteins dysregulated by BDV infection (Fig. 2D). We performed a pathway clustering analysis through the Kyoto Encyclopedia of Genes and Genomes (KEGG) (Fig. 2E). Using a manually curated CORUM database, we performed enrichment analysis on protein complexes and used k-means clustering to identify the specific subgroups that were most impacted by BDV infection (Fig. 2F). We obtained two protein complexes (the amyloid precursor protein mitochondrial translocase APP-TOMM40 and IFIP35-NMI) enriched in Q4 that are primarily related to mitochondrial protein transport and interferon signaling, respectively (Lee et al., 2012; Zhou et al., 2000), and 17 protein
complexes enriched in Q1 that are primarily related to DNA replication, DNA repair response, and chromosome shape regulation. The results are summarized in Table 2.

**BDV infection affects histone Kac**

As BDV infection impacts transcription factor expression and histone Kac regulates gene transcription, we investigated the effect of BDV infection on site-specific histone Kac. We applied SILAC-based quantitative proteomics to comparatively profile histone Kac in OL/BDV cells and control cells. A total of 30 Kac sites in core histones were identified (Fig. 3A), which covers almost all reported Kac sites in mammalian core histones. The representative spectra of well-known histone Kac peptides, including the peptide spectra of H1.2K16ac, H2BK20ac, H3K18ac, and H4K5ac, are provided (Fig. 3B–E). Furthermore, the sequences of identified Kac peptides in core histones and all corresponding quantitative Kac profiles in response to BDV infection are summarized in Table 3. As indicated, the Kac levels in 15 of 30 sites were significantly decreased (e.g., H2AK5ac, H2BK5ac, H3K14ac, and H4K5ac), whereas the Kac level in one site, H2BK15, was significantly increased in response to BDV infection. The Kac levels in the other 14 sites showed no significant changes.

To further validate the different histone acetylation profiles, Western blot analysis was performed with histone Kac sequence-specific antibodies. Consistent with the quantitative results (Table 2), BDV infection significantly decreased the Kac level in H2AK5ac, H2BK5ac, H2BK20ac, H3K14ac, H3K18ac, and H4K5ac (Fig. 4). Consistent with the MS findings, we observed an increased H2BK15ac level and unchanged H4K8ac level post-BDV infection (Fig. 4).

**Validation of histone acetyltransferase and deacetylase expression by western blotting**

Histone acetylation is dynamic and regulated by histone acetyltransferases (HATs) and histone deacetylases (HDACs). From the proteomic data, the expression of several HATs (ELP3, BRD4) and HDACs (SIRT1, HDAC7) was significantly altered by BDV infection. To validate these changes, two HATs (GCN5 and PCAF) and eight HDACs (SIRT1, SIRT2, HDACs 1, 2, 3, 4, 5, and 7) were selected for Western blotting (Fig. 5). Two HATs – GCN5 (p = 0.039) and PCAF (p = 0.029) – were found to be significantly down-regulated, and four HDACs – SIRT1 (p = 0.015), SIRT2 (p = 0.005), HDAC4 (p = 0.002), and HDAC7 (p = 0.033) – were found to be significantly upregulated, in OL/BDV cells relative to control cells. There was no significant dysregulation observed in HDACs 1, 2, 3, or 5 (p > 0.05).

**Discussion**

The goal of this study was to analyze the effects of BDV infection on the proteome and histone Kac profiles of OL cells through a SILAC-based quantitative proteomic approach in order to gain insight into BDV pathogenesis by using a natural human virus strain. To date, several studies have addressed virus-induced changes on the host proteome (e.g., BDV, EBV, HBV, HCV, HIV-1, and SARS-CoV) (Zhou et al., 2011), but minimal data has been published on virus-induced changes in histone Kac. To our

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**Fig. 2.** Enrichment and clustering analysis of the quantifiable proteomic data set. Quantifiable proteins were classified by gene ontology annotation based on (A) molecular function, (B) cellular compartment, and (C) biological process. In each category, the differential quantifiable proteins were divided into four quantiles based on the cumulative distribution of SILAC L/H ratios: Q1, less than 15%; Q2, 15–50%; Q3, 50–85%; and Q4, greater than 85%. An enrichment analysis was separately performed in each quantile for diverse categories, and the overrepresented annotations were clustered through one-way hierarchical clustering for comparative analysis. Quantifiable proteins were also annotated based on (D) the PFAM domain database, (E) the KEGG pathway database, and (F) the CORUM protein complex database.
knowledge, this is the first SILAC-based quantitative proteomic study to assess the response of human OL cells to human strain BDV Hu-H1 infection and the first comprehensive atlas of host cell histone acetylation change in response to BDV infection. Previous studies have shown that BDV infection impacts gene transcription in OL cells (Carbone et al., 2001; Koster-Patzlaff et al., 2007), while this study found many regulated transcription factors by BDV infection, which linked gene transcription and the differential proteomic expression towards BDV infection. In view that histone acetylation is generally considered a marker involved in activating gene expression, and our data suggested altered histone site-specific acetylation in response to BDV infection, it appears reasonable to consider a link of altered histone acetylation, gene transcription and subsequently proteomics. (Supplemental Fig. 2)

Histone Kac

Previous studies have revealed virus-induced changes in some histone Kac sites. For example, adenovirus small early region 1a (e1a) protein induces H3K18 hypoacetylation by restricting some HATs (Horwitz et al., 2008). With respect to BDV, Suberbielle et al. (2008) revealed histone modifications, including H2BK5 and H2BK20, in BDV laboratory strain He/80 infected-neurons.

Here, 30 histone Kac sites were identified, of which half presented with decreased acetylation and only one site, H2BK15, presented with increased acetylation. Histone modification at the N-terminal plays a central role in chromatin remodeling and transcriptional regulation. Here, most of the quantifiable Kac sites were at N-terminals of core histones, suggesting pronounced epigenetic modulation of BDV-infected OL cells.

Some specific functions of histone site acetylation have already been verified. For example, global hypoacetylation at H3K18 is observed in prostate carcinomas with poor prognosis which suggests that processes resulting in global H3K18 histone acetylation may be linked to oncogenic transformation (Horwitz et al., 2008). H4K16 acetylation regulates cellular lifespan because the level of H4K16 acetylation increased with age examined in yeast cells (Dang et al., 2009). Deregulated H4K12 acetylation may represent an early biomarker of an impaired genome-environment interaction in the aging mouse brain because aged mice display a specific deregulation of H4K12 acetylation and fail to initiate a hippocampal gene expression program associated with memory consolidation during learning. Moreover, restoration of physiological H4K12 acetylation reestablishes the expression of learning-induced genes and leads to the recovery of cognitive abilities (Peleg et al., 2010). Further studies should investigate the biological function of the histone specific-site acetylation altered by BDV infection.

HATs and HDACs

Histone acetylation levels are determined by the combined activities of HATs and HDACs. HATs have been shown to preferentially acetylate specific histones and/or specific lysines (Turner, 2000). HDACs through deacetylation of histones has been associated with decreased transcriptional activation and proteomic expression; for example, non-small cell lung cancer A549 cells treated with suberoylanilide hydroxamic acid (SAHA), a pan HDAC inhibitor, resulted in histone Kac and proteome changes (Shahbazian and Grunstein, 2007; Wu et al., 2013). Here, the dysregulation of several HATs and HDACs was accompanied by significant proteomic change, suggesting a link between histone Kac and proteomic expression in BDV-infected OL cells (Supplemental Fig. 2).

Two Gcn5 mammalian HAT subclasses have been described: GCN5 and p300/CREB-binding protein-associated factor (PCAF). They are transcriptional coactivators and possess intrinsic histone acetylase activity, providing a direct link between hyperacetylated chromatin and transcriptional activation. GCN5 and PCAF specifically promote acetylation on H3K9, H3K14, H4K8, and H4K16. Deletions of GCN5/PCAF in cells reduces acetylation on H3K9 and correlates with nuclear receptor target gene activation at some level (Jin et al., 2011). The GCN5/PCAF have a preference for H3K14 (Grant et al., 1999), and also acetylate H4K8 and H4K16 to a relatively slight degree (Kuo et al., 1996; Schiltz et al., 1999). By Western blotting, we found that the expression levels of GCN5 and PCAF were downregulated, consistent with the decreased acetylation of H3K14 and H4K16 in response to BDV infection. Interestingly, acetylation of H3K9 and H4K8 was not decreased here, which linked gene transcription and the differential proteomic expression towards BDV infection.

Table 2

| Biological process | GO-molecular functions | GO-cellular compartment | Protein domain | KEGG pathway | Protein complex |
|--------------------|------------------------|-------------------------|----------------|-------------|----------------|
| Q4 (the quantile with high L/H ratio) | Chromosomal segregation and cell division, primary and macromolecular metabolic processes | Nucleic acid processing | Nucleus, chromosomes, ribosomes, and minichromosome maintenance (MCM) complex | Nucleic acid processing (e.g., helicase, nucleotide-binding, RNA recognition, exonuclease, and transcription regulation [SNF2-related and COMM domains]) | Nucleic acid, cell cycle, ribosomal processes, purine and pyrimidine metabolism |
| Q1 (the quantile with low L/H ratio) | Multiple stimuli (e.g., extracellular, stress, oxygen levels, inorganic substances), lipid metabolism, extracellular structure organization, reactive oxygen species and organic hydroxy species metabolic responses, immune and defense responses | Membranes (e.g., Golgi apparatus, endosomes, MHC class molecules, organelle membranes, plasma membrane, nuclear outer membrane) | Immune competence (e.g., MHC class molecules and immunoglobulins) | Phosphatidylinositol signaling, GABAergic synapse, cell adhesion molecules (CAMs), glycerophospholipid metabolism, and immune response pathways (e.g., antigen processing and presentation, autoimmune thyroid disease, allograft rejection, graft-versus-host disease) | Two protein complexes related to mitochondrial protein transport and interferon signaling | 17 protein complexes related to DNA replication, DNA repair response, chromosome shape regulation |
Fig. 3. Analysis of histone lysine acetylation (Kac). (A) The illustration of identified histone Kac sites in OL cells in response to BDV infection. The identified sites in core histones were numbered and highlighted. (B) MS/MS spectra of a tryptic peptide ion histone H1.2K16 acetylated peptide _ (ac)SETAPAAPAAPAEPK(ac)APVK. (C) MS/MS spectra of a tryptic peptide ion histone H2BK20 acetylated peptide_AVTK(ac)VQK. (D) MS/MS spectra of a tryptic peptide ion histone H3K18 acetylatedpeptide_K(ac)QLATK (ac)AAR. (E) MS/MS spectra of a tryptic peptide ion histone H4K5 acetylated peptide _GK(ac)GGK(ac)GLGK.
BDV infection has a profound impact on membrane-associated molecules (Gonzalez-Dunia et al., 1998; Honda and Tomonaga, 2004). In this study, western blotting analysis showed that the expression level of HDACs 4 and 7 were increased in BDV-infected OL cells, while few changes were observed in HDACs 1, 2, 3, and 5. These results suggest that class II HDACs may pose specific changes in histone lysine acetylation (Kac) sites.

| Kac site | Modified sequence | Normalized ratio L/H (11) |
|----------|-------------------|--------------------------|
| H1.2K16ac | (ac)SETAPAAPAAAPPCCPacAPVK | 0.5683 |
| H1.2K45ac | ASGPVSELT(ac)AVWAASK | 0.9315 |
| H1.2K84ac | LGLK(ac)SILVSK | 0.8309 |
| H1.2K89ac | SILVSK(ac)GTLVQGTK | 1.2392 |
| H1.2K96ac | GTLVQGTK(ac)GTGASC3FK | 0.6828 |
| H2AK5ac | G(ac)EQGGGK(ac)R | 0.4577 |
| H2AK9ac | G(ac)QGGGK(ac)R | 0.4577 |
| H2BK5ac | PEPAK(ac)SAPARK(ac)R(ac)GSK | 0.3748 |
| H2BK11ac | PEPAK(ac)SAPARK(ac)R(ac)GSK | 0.8559 |
| H2BK12ac | K(ac)SSTK(ac)R(ac)AVTK | 1.2412 |
| H2BK15ac | GSK(ac)K(ac)AVTK(ac)QK | 1.5352 |
| H2BK16ac | K(ac)AVTK(ac)AQK | 0.8097 |
| H2BK20ac | AVTK(ac)VKQ | 0.5944 |
| H2BK23ac | KAVTK(ac)QK(ac)R | 1.1076 |
| H2BK43ac | ESYSYTVYK(ac)VLK | 0.4037 |
| H2BK108ac | LLLPGELAK(ac)HAYSEGK | 0.6612 |
| H2BK120ac | AVTK(ac)YTSSK | 0.5863 |
| H3K5ac | K(ac)STGGK(ac)APR | 1.0686 |
| H3K14ac | KSTGGK(ac)APR | 0.9965 |
| H3K18ac | K(ac)SXLTK(ac)KAR | 0.5660 |
| H3K21ac | QLATK(ac)QAR | 0.9600 |
| H3K56ac | YQK(ac)STELLR | 0.5650 |
| H3K79ac | EIAQFK(ac)TDLR | 0.6652 |
| H3K122ac | VTMPKR(ac)DIQLR | 0.5615 |
| H4K5ac | G(ac)GK(ac)GGLK(ac)GGAK | 1.2583 |
| H4K12ac | GLGK(ac)GGAK | 0.9165 |
| H4K16ac | GLGK(ac)GGAK(ac)R | 0.4638 |
| H4K31ac | DNIQGTK(ac)PAIR | 0.5159 |
| H4K91ac | TYTAMDVYVVLK(ac)R | 0.3895 |

Nuclear proteins and DNA replication

As the host cellular chromatin inhibits viral gene expression and replication by suppressing DNA accessibility, viruses that enter and persist in the nucleus (such as BDV) have evolved chromatin-associated mechanisms to efficiently propagate the viral genome (Lieberman, 2006). For example, BDV’s viral ribonucleoprotein (RNP) directly interacts with the mitotic host chromosome using core histones as a docking platform (Matsumoto et al., 2012). BDV infection was found to differentially impact on nuclear and chromosomal proteins enriched in DNA replication and repair, transcription regulation, and chromosomal shape regulation in Q1, most of which were downregulated in response to BDV infection.

Metabolic pathways

Through KEGG analysis, a wide variety of pathways (e.g., phosphatidylinositol signaling, GABAergic synapse, CAMs, glycerophospholipid metabolism, immune response pathways, nucleic acid processes, cell cycle, ribosomal processes, purine metabolism, and pyrimidine metabolism) were identified as the most significantly altered set of host biological pathways. Consistent with the current KEGG findings in phosphatidylinositol signaling, glycerophospholipid metabolism, and purine/pyrimidine metabolism, our previous metabonomic profiling study in BDV-infected OL cells revealed significant perturbations in myo-inositol, α-glucose, acetate, pyruvate, and nicotineamide adenine dinucleotide (NAD) (Huang et al., 2012). Thus, these proteomic data and the previous metabonomic data also based on human BDV Hu-H1 strain, mutually support each other.

Conclusions

Using SILAC-based quantitative proteomics coupled with bioinformatic analysis, this study is the first to reveal the host proteome and histone Kac profiles in BDV-infected OL cells using a natural human virus strain. BDV infection appears to preferentially dysregulate membrane, nuclear, and chromosomal host protein expression while affecting metabolic pathways, immune response, DNA replication, DNA repair, and transcription regulation. BDV infection was found to affect histone acetylation of specific lysine residues. Moreover, BDV infection affected the expression of many transcription factors, several HATs and HDACs. As histone Kac epigenetically regulates gene transcriptional activation, the differential acetylation of specific lysine residues may have impacted the changes in the host proteome profile.

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of transcription myocyte enhancer factor 2 activity, involved in muscle development and neuronal survival (Lomonte et al., 2004). In addition, HDAC4, HDAC5, and HDAC7 are known to shuttle muscle development and neuronal survival (Lomonte et al., 2004). These results suggest that class II HDACs 4 and 7 may pose specific changes in histone lysine acetylation (Kac) sites.

| Kac site | Modified sequence | Normalized ratio L/H (11) |
|----------|-------------------|--------------------------|
| H1.2K16ac | (ac)SETAPAAPAAAPPCCPacAPVK | 0.5683 |
| H1.2K45ac | ASGPVSELT(ac)AVWAASK | 0.9315 |
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| H1.2K89ac | SILVSK(ac)GTLVQGTK | 1.2392 |
| H1.2K96ac | GTLVQGTK(ac)GTGASC3FK | 0.6828 |
| H2AK5ac | G(ac)EQGGGK(ac)R | 0.4577 |
| H2AK9ac | G(ac)QGGGK(ac)R | 0.4577 |
| H2BK5ac | PEPAK(ac)SAPARK(ac)R(ac)GSK | 0.3748 |
| H2BK11ac | PEPAK(ac)SAPARK(ac)R(ac)GSK | 0.8559 |
| H2BK12ac | K(ac)SSTK(ac)R(ac)AVTK | 1.2412 |
| H2BK15ac | GSK(ac)K(ac)AVTK(ac)QK | 1.5352 |
| H2BK16ac | K(ac)AVTK(ac)AQK | 0.8097 |
| H2BK20ac | AVTK(ac)VKQ | 0.5944 |
| H2BK23ac | KAVTK(ac)QK(ac)R | 1.1076 |
| H2BK43ac | ESYSYTVYK(ac)VLK | 0.4037 |
| H2BK108ac | LLLPGELAK(ac)HAYSEGK | 0.6612 |
| H2BK120ac | AVTK(ac)YTSSK | 0.5863 |
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| H4K31ac | DNIQGTK(ac)PAIR | 0.5159 |
| H4K91ac | TYTAMDVYVVLK(ac)R | 0.3895 |
Materials and methods

Virus strain

The human strain BDV Hu-H1 was used throughout the study and kindly provided by Professor Hanns Ludwig (Free University of Berlin, Berlin, Germany). This strain dated back to 1996 and could be recovered from freshly isolated white blood cells of a female bipolar I disorder patient in Germany who was admitted to hospital during a severe depressive episode. The isolation procedure had required laborious co-cultivating of patient’s PBMCs with human oligodendroglia (OL) cells. After at least 10–12 blind passages which did not exhibit viral growth, infectious human virus had finally been harvested and subsequently passaged to further characterization and storage. The isolation protocol of BDV strain Hu-H1 and two other isolates have been described in full detail in the original publication (Bode et al., 1996). Strain Hu-H1 and other two isolates were originating from PBMC samples which were testing positive for BDV RNA and protein. Further molecular characterization had revealed that corresponding RNAs (of PBMC and isolate) displayed identical nucleotide sequences, proving the authenticity of the human isolates. Moreover, the isolates
displayed few but meaningful point mutations which differentiated them from each other and from laboratory strain V and a second lab strain (He/80) (de la Torre et al., 1996b). Further biological characterization had revealed that strain Hu-H1 and other two human strains displayed differing pathogenicity in animal experiments, namely inducing behavioral changes in rabbits but no deadly disease as compared to str. V (Bode et al., 1996).

Own recent experiments of our research group could demonstrate that the above described human strain Hu-H1 inhibited proliferation and supported apoptosis, whereas lab. Strain V did quite the opposite (Li et al., 2013). This remarkable difference was pivotal to again choose strain Hu-H1 for this study, apart from the fact that wild-type viruses are supposed to reflect natural pathogenicity patterns at the cellular level which might be lost in highly adapted laboratory strains.

Cell line and viral infection

Human OL cells (a cell line derived from fetal human oligodendrocytes, 112 passages) and BDV Hu-H1 strain (77 passages in OL cells) (Bode et al., 1996) were used. Like the virus strain, human OL cells were kindly supplied by Professor Hanns Ludwig. Persistently-infected OL cells and non-infected OL cells were cultured in Dulbecco’s modified Eagle’s medium (DMEM; Hyclone, Logan, Utah, USA) with 10% fetal bovine serum (FBS, Hyclone), 100 U/ml penicillin, and 100 μg/ml streptomycin (Hyclone) within a humidified incubator (5% CO2, 37 °C) and were passaged when they reached 90% confluence by trypsinization (Hyclone).

Stock viral solution was prepared and titrated as previously described (Huang et al., 2012). Briefly, OL cells washed twice with serum-free DMEM were sub-divided in one 10-cm dish and infected with BDV Hu-H1 at a multiplicity of infection (MOI) of 1.0 (800 μl of virus stock). Cells were then stored in a humidified incubator (5% CO2, 37 °C) for two hours with gentle shaking for 15 min. Excess virus was removed by washing with 5 ml of serum-free DMEM before bathing the cells in 10 ml of culture medium (10% FBS in DMEM). Thereafter, OL cells were passaged 5 to 6 times until all cells were infected with BDV Hu-H1. The now persistently-infected cells (OL/BDV cells) and non-infected OL cells (control cells) were kept under the same conditions for the remainder of the study.

RNA extraction and RT-PCR for BDV detection

Total RNA subjected to reverse transcription polymerase chain reaction (RT-PCR) using avian myeloblastosis virus reverse transcriptase (AMV RT) (Promega, Madison, WI, USA) was extracted from OL/Hu-H1 cells and control cells with Trizol (Trizol LS; Invitrogen, Carlsbad, CA, USA). For BDV phosphoprotein 24 (p24) and nucleoprotein (p40) gene amplification, PCR was performed according to the manufacturer’s protocols except for the modified annealing temperatures of 50 °C and 61.5 °C (GoTaq® Green Master Mix, Promega, Madison, WI, USA). The PCR products were separated by 2% agarose gel electrophoresis and stained with ethidium bromide. The primer sequences were as follows: p40, OR (5′–GCTGGTTTCCTTGACACTTG-3′), OF (5′–GATCAAAGCAGGAGCCGAGCAG-3′) (size: 545 bp); and p24, OR (5′–CTCCATCATGTCCTTCATG-3′), OF (5′–TGATCTCAGACCCAGAGC-3′) (size: 224 bp).

Imunofluorescence assay for BDV detection

Both OL/BDV cells and control cells were grown on six-well dishes for 30 min at room temperature and fixed with 4% paraformaldehyde followed by permeabilization for 5 min in 0.4% Triton X-100. Thereafter, both lines were rinsed with PBS and blocked with 5% (w/v) skimmed milk solution for one hour at 37 °C. Overnight incubation with anti-recombinant BDV-specific p24 or p40 protein primary monoclonal antibody (both diluted

Fig. 5. Validation of histone acetyltransferase (HAT) and histone deacetylase (HDAC) expression by Western blotting. (A) Western blots of two HATs (GCN5 and PCAF) and eight HDACs (SIRT1, SIRT2, HDACs 1, 2, 3, 4, 5, and 7) with β-actin used as a control. (B) GCN5 (p = 0.039) and PCAF (p = 0.029) were found to be significantly downregulated, while SIRT1 (p = 0.015), SIRT2 (p = 0.005), HDAC4 (p = 0.002), and HDAC7 (p = 0.033) were found to be significantly upregulated, in OL/Hu-H1 cells relative to control cells. There was no significant dysregulation observed in HDAC1 (p = 0.196), HDAC2 (p = 0.169), HDAC3 (p = 0.520), or HDAC5 (p = 0.881).
1:500; GenScript, Piscataway, New Jersey, USA) at 4 °C was followed by one hour of incubation with secondary antibodies at room temperature. After three PBS washes, immunofluorescence was detected by phase-contrast microscopy.

**Cell culture and labeling**

To profile protein expression and histone Kac in response to BDV infection, OL/BDV cells and control cells were labeled by the SILAC Protein Quantitation Kit (Invitrogen, Carlsbad, CA) according to manufacturer's instructions. OL/BDV and control cells were separately grown in DMEM supplemented with 10% FBS and either the light isotopic forms of [U-12C6]-L-lysine and [U-12C6,15N4]-L-arginine or the heavy isotopic forms of [U-13C6]-L-lysine and [U-13C6,15N4]-L-arginine, respectively. After six generations of passages, the heavy labeling efficiency of [U-13C6]-L-lysine and [U-13C6,15N4]-L-arginine in control cells was evaluated by mass spectrometer analysis to confirm a greater than 99% labeling efficiency. Then, cells were continuously expanded in SILAC medium until reaching the desired population size. Finally, cells in each heavy- or light-labeled pool were harvested separately and combined in equal amounts.

**Affinity enrichment of histone lysine acetylated peptides**

Prior to affinity enrichment of histone Kac peptides, anti-acetyllysine antibody agarose beads (PTM Biolabs, Chicago, IL) were washed twice with ice-cold PBS. To enrich lysine acetylated peptides, histone tryptic peptides dissolved in NETN buffer (100 mM NaCl, 1 mM EDTA, 20 mM Tris–HCl, 0.5% NP-40, pH 8.0) were incubated with prewashed antibody agarose beads at 4 °C overnight with gentle shaking. The beads were washed four times with NETN buffer and twice with ddH2O. The bound peptides were eluted from the beads with 0.1% trifluoroacetic acid. The eluted fractions were combined and vacuum-dried and then analyzed by high-performance liquid chromatography with tandem mass spectrometry (HPLC – MS/MS).

**LC-ESI-MS/MS analysis**

Peptides were dissolved in solvent A (0.1% formic acid in 2% acetonitrile), directly loaded onto a reversed-phase column (360 μm OD × 75 μm ID), packed in-house with 3 μm C18 beads (Reprosil-Pur C18-AQ, Dr. Maisch), and eluted with a linear gradient of 5–30% solvent B (0.1% formic acid in 98% acetonitrile) for 35 min at a constant flow rate of 300 nL/min on an EASY-nLC 1000 Ultra performance liquid chromatography (UPLC) system. Eluted peptides were electrosprayed into LTQ-Orbitrap Elite spectrometers (Thermo Scientific) operating in a data-dependent mode that acquired MS/MS spectra for 20 most intense ions. Full MS (m/z range: 350–1300) was acquired using Fourier transform MS (FTMS) in the Orbitrap at 240,000 resolution at 400 m/z, and MS/MS was acquired using collision-induced dissociation (CID) in the LTQ-Orbitrap Elite at 35% normalized collision energy. A lock mass ion from ambient air (m/z 536.165369) was used for internal calibration of all full-scan measurements with an Orbitrap detector. Dynamic exclusion was enabled with a repeat count of 1, repeat duration of 5 s, exclusion duration of 60 s, and exclusion mass width of ± 10 ppm relative to the reference mass.

**Western blotting**

Cell extracts and Western blots were performed as previously described (Liu et al., 2014). Briefly, monolayers of OL/BDV and control cells were lysed in standard lysis buffer and sonicated on ice. Then, 10 μg lysates were separated by 10% sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) and transferred onto a polyvinylidene fluoride (PVDF) membrane. The membranes were incubated overnight at 4 °C with primary antibodies (i.e., anti-recombinant BDV p24 antibody, anti-recombinant BDV p40 antibody, both diluted 1:500; GenScript; anti-H2AK5ac, H2BK5ac, H2BK15ac, H2B2K20ac, H3K14ac, H3K18ac, H4K5ac and H4K8ac antibodies, all diluted 1:2000; PTM Biolabs, Chicago, IL; anti-GCN5, PCAF, SIRT1, SIRT2, HDACs 1, 2, 3, 4, 5, and 7 antibodies, all diluted 1:1000; CST, Beverly, MA, USA). All membranes were washed and incubated with their respective horseradish peroxidase-coupled immunoglobulin G (IgG). After extensive washing, antibody-detected protein bands were visualized by enhanced chemiluminescence (ECL) and exposed to autoradiography film.

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**Appendix A. Supporting information**

Supplementary data associated with this article can be found in the online version at http://dx.doi.org/10.1016/j.virol.2014.06.040.

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