Molecular Features Contributing to Virus-Independent Intracellular Localization and Dynamic Behavior of the Herpesvirus Transport Protein U₅₉

Manuela Pedrazzi¹, Bradley Nash², Olimpia Meucci²,³*, Renato Brandimarti¹,²*

¹Department of Pharmacy and Biotechnology, University of Bologna, Bologna, Italy, ²Department of Pharmacology and Physiology, Drexel University College of Medicine, Philadelphia, Pennsylvania, United States of America, ³Department of Microbiology and Immunology, Drexel University College of Medicine, Philadelphia, Pennsylvania, United States of America

Abstract

Reaching the right destination is of vital importance for molecules, proteins, organelles, and cargoes. Thus, intracellular traffic is continuously controlled and regulated by several proteins taking part in the process. Viruses exploit this machinery, and viral proteins regulating intracellular transport have been identified as they represent valuable tools to understand and possibly direct molecules targeting and delivery. Deciphering the molecular features of viral proteins contributing to (or determining) this dynamic phenotype can eventually lead to a virus-independent approach to control cellular transport and delivery. From this virus-independent perspective we looked at U₅₉, a virion component of Herpes Simplex Virus involved in anterograde transport of the virus inside neurons of the infected host. As the natural cargo of U₅₉-related vesicles is the virus (or its parts), defining its autonomous, virus-independent role in vesicles transport represents a prerequisite to make U₅₉ a valuable molecular tool to study and possibly direct cellular transport. To assess the extent of this autonomous role in vesicles transport, we analyzed U₅₉ behavior in the absence of viral infection. Based on our studies, U₅₉ behavior appears similar in different cell types; however, as expected, the data we obtained in neurons best represent the virus-independent properties of U₅₉. In these primary cells, transfected U₅₉ mostly recapitulates the behavior of U₅₉ expressed from the viral genome. Additionally, ablation of two major phosphorylation sites (i.e. Y₃₂Y₃₃ and S₃₄ES₃₆) have no effect on protein incorporation on vesicles and on its localization on both proximal and distal regions of the cells. These results support the idea that, while U₅₉ post-translational modification may be important to regulate cargo loading and, consequently, virion export and delivery, no additional viral functions are required for U₅₉ role in intracellular transport.

Introduction

Cellular functionality heavily relies on efficient transport of individual components, from single molecules to entire organelles. Even under resting conditions intracellular traffic of these various cellular entities is significant and finely regulated; this is achieved through the activities of specialized molecular motors that move cargoes along cytoskeletal structures [1,2]. The microtubular cytoskeleton offers a great backbone to travel in every direction, from the cell center to the periphery and viceversa. Transport occurring in the same direction as microtubules polymerization, which has been designated “+”, is referred to as anterograde transport, and is usually directed toward the periphery of the cell. Conversely, motion toward microtubules “−” end is defined retrograde, and mostly points to the nucleus [3].

Amongst many other cellular components, special and unwanted cargos are represented by viruses, especially those, like retroviruses and herpesviruses, which must reach the nucleus to complete their replication cycles. As widely acknowledged, viruses are able to efficiently exploit physiological functions (e.g. S phase induction [4]; blockage of MHC-I maturation [5,6]) through the activity of specialized proteins that specifically target cellular factors. Conversely, viral proteins may be seen as tools to both decipher cellular functions and re-program them for different purposes.

U₅₉ is a gene well conserved in the family of alphaherpesviruses, the group of herpesviruses characterized by the ability to establish a lifelong latent infection in the peripheral nervous system of their host. It encodes a small tail-anchored, type II membrane protein. For a long time neglected as a non essential viral product, U₅₉ has recently gained more attention because viruses deleted in the U₅₉ gene show defects in the ability to move in the anterograde direction in the axons and to establish secondary infections in the brains of infected animals [7–12]. Virus anterograde transport occurs inside vesicles; besides being a constitutive component of transported virions, U₅₉ is present on transport vesicles membrane. The impairment shown by deletion viruses implies that U₅₉ participates in the process of virus egress, and that this effect is dependent on the ability of the protein to directly or indirectly regulate the interaction of the viral particle (or of its parts) with the transport machinery. In Pseudo-Rabies Virus (PRV) the role in transport played by U₅₉ has been extensively investigated [7,12–
18] and recently visually demonstrated in vivo by Taylor et al. [19]. However, beside the U9 activity related to virus replication and diffusion in the infected host, it is of great interest to understand the molecular features that confer the protein its ability to drive vesicles transport. Localization studies showed that U9 mostly (but not exclusively) accumulates in Trans Golgi Network [0,20], but it is also detected in more peripheral regions of the cell, as well as at the plasma membrane, in agreement to the assigned transport task deduced from deletion studies.

Starting from the acknowledged role played by U9 in virus transport, we decided to look at U9 stand alone properties using GFP-tagged constructs in a virus-free cellular environment, both in fixed cells and in real time experiments, aimed to investigate if and to which extent U9 properties that in the viral context serve virus transport and infection spread, can be directly ascribed to U9. The results presented here highlight the dynamic behavior of U9, supporting the idea that the viral protein is able to autonomously interact with the cellular transport machinery, in a cell type independent manner. In fact, U9 is always detectable in both proximal and peripheral cellular regions, as well as on the plasma membrane. This ability, which in the viral context supports transport and delivery of viral particles or of its components, is maintained even in the absence of other viral factors. By using truncated forms of the viral protein, we observed that while the U9 trans-membrane domain partially recapitulates the functional behavior of the full length protein, as it is still able to dictate localization of the fused GFP moiety in both cytosolic puncta and at the plasma membrane, it clearly differs from the full length GFP-U9. Notably, the major difference ascribed to the absence of the U9 cytosolic domain is the reduced punctuate staining and the enhanced accumulation of the protein at the plasma membrane. Finally, mutagenesis of key tyrosines and serines in the acidic domain located in the middle of the cytosolic portion of the protein and important for virus transport do not affect U9 behavior, supporting the hypothesis that post-translational modifications represent a tuning system adopted by the virus to regulate cargo loading and, consequently, virion export and delivery; however, and most importantly, they are not required for U9 transport inside the cell.

Results

Construction and analysis of GFP-U9 fusion proteins

The U9 gene in the HSV genome encodes a protein of 90 amino acids [21], highly conserved amongst alphaherpesviruses, with an essential role in anterograde axonal transport [7–12,22–23]. As predicted by its hydrophaticity profile (figure 1A), U9 has been shown to be a tail-anchored type II membrane protein, with the N-terminal domain hanging in the tegument region comprised between virion envelope and nucleocapsid [24–25]. Here we studied the U9 transport properties by tagging the protein with GFP. The U9 gene was cloned into pEGFP-C1, to create the GFP- U9 chimera (figure 1B) named g9. Two more plasmids were constructed, with the same topological organization but with two complementary truncations of the U9 protein (as indicated by red triangles in the sequence in figure 1A); g9-TM (GFP-U9 Trans-Membrane) and g9-ATM (GFP-U9 ATTrans-Membrane), encoding GFP fused to the U9 hydrophobic and hydrophilic domains, respectively (figure 1B). The three constructs, along with the parental plasmid pEGFP-C1, were transfected into MDA cells, and protein contents were analyzed by Western blot using antibodies directed against U9 and GFP (figure 1C). As expected, a band with a relative size corresponding to the GFP proper molecular weight was recognized by the GFP antibody in the GFP only lane (right panel), while no bands in the same region of the gel could be detected with the U9 Ab. Cells transfected with g9 plasmid expressed a protein specifically reacting with U9 and GFP Abs. Both Abs detected a major band of about 42 KDa (corresponding to the expected molecular mass for the fusion protein), and a higher minor band, likely the result of post-translational modifications – in line with reports from infected cells where U9 typically shows a complex migration pattern partially dependent on enzymatic activity [26–30].

The g9-TM chimeric protein could be detected only with the GFP Ab, since the U9 Ab is directed against the U9 hydrophilic domain that is missing in the fusion protein [31]. The migration pattern of g9-TM appears very similar to that of GFP, due to the presence of a relatively long extra peptide originated by the translated multiple cloning site sequence in pEGFP-C1. As a consequence, peptides encoded by pEGFP-C1 and g9-TM differ for just 12 amino acids in length. We validated all our constructs by sequencing; more importantly, additional experiments (shown in figure 2 and discussed later) also address this issue and undoubtedly confirm the correctness of the g9-TM construct and the U9 trans-membrane domain contribution to U9 localization and transport properties. Removal of TM domain from U9 in the g9-ATM plasmid results in the lack of 32 aa from the fusion protein, and the bands present in both panels in figure 1C (lanes g9-ATM) migrated as expected (Mr of about 38 KDa).

Localization of U9

Given its role in viral particles transport, the cellular compartments in which U9 accumulates and the contributions of U9 hydrophilic/hydrophobic domains to its localization have been widely studied. Early experiments with PRV U9 and its homologs from other alphaherpesviruses demonstrated that U9 mainly accumulates to the Trans Golgi Network (TGN) but it is also detected in distal intracellular regions as well at the plasma membrane [24–25], which is in agreement with its role in virus transport [7–15,20,26]. These studies utilized specific antibodies against the viral protein and Hemo-Agglutinin or GFP tags. Notably, in the context of viral infection, GFP-U9 can functionally substitute wild type U9, and has been used to trace protein localization in infected cells [19].

Here, g9, g9-ATM, g9-TM, or the control parental plasmid pEGFP-C1, were transfected into MDA cells, and then imaged 16 hours post transfection. In the absence of specific localization signals, GFP expression yielded a diffuse fluorescent staining in both cytoplasm and nucleus (figure 2A–B). Expression of g9 was strongly associated with a bright punctuate pattern (figure 2A–B). Fluorescent signals were completely excluded from nuclei of transfected cells, and widely distributed into the cytoplasm, in discrete puncta. The punctuate staining is typical of vesicles, as expected for a membrane protein involved in intracellular transport and as already reported [8,12,15,24,25]. Puncta were detectable throughout the cytoplasm, and a weak signal was also associated with the plasma membrane. U9 accumulation in the central region of the cell has been associated with markers of the Trans Golgi Network, the putative site of virions assembly and sorting [8,15–16], and this region has been defined as the steady-state site of U9 localization. In all our experiments, this fluorescent pattern was always reproduced, with only the intensity of the fluorescent signal in the perinuclear region possibly diverging. However, not to interfere with physiological variability, we did not attempt to establish stably transfected cell clones, and always performed transient transfections. Nevertheless, labeled
vesicles in distal regions were always detected, even in cells with a heavy perinuclear fluorescent signal.

Cells transfected with g9-TM (figure 2A–B) showed a comparable yet clearly distinguishable fluorescent pattern, comprising both intracellular punctuate staining and plasma membrane fluorescence. Indeed, g9-TM labeling was more pronounced at the plasma membrane, as compared to the full length protein, while intra-cellular punctuate signal was reduced. These data suggest that the 32 aa long trans-membrane domain of US9 is sufficient to drive the insertion of the GFP molecule (which is almost 3 and 8 times larger than US9 and US9-TM, respectively) into the same vesicles already described for the full length protein; however, in the absence of the cytosolic domain, the fusion protein accumulates at the plasma membrane. On the other hand, as expected, removing the US9 trans-membrane domain from the fusion protein dramatically changes the fluorescent pattern. Cells transfected with g9-DTM appeared undistinguishable from cells expressing GFP alone, as the staining was homogenously distributed throughout the cytoplasm and the nucleus. These results clearly confirm that the association of US9 with the vesicular pattern revealed by the fluorescent punctuate labeling depends on the presence of its trans-membrane domain. Moreover, the differences in localization between g9 and g9-TM suggest a role for the US9 cytosolic domain in regulation of protein export to the plasma membrane. In addition to the US9 fluorescent fusion proteins described above, we also generated a US9 chimera with GFP fused to the C-terminal portion of US9, and named it 9g. When transfected in MDA cells, the fluorescent pattern of 9g was undistinguishable from that of g9 (not shown). In this study we opted to use g9 since its GFP moiety has a predictable minor impact on the US9 structure and previous studies showed that 9g does not fully reproduce the behavior of wild type US9 [32].

To further investigate the biochemical properties that confer US9 the described intracellular punctuate distribution, we analyzed the presence of the chimeras in the cytosolic and the membrane fractions [33,34]. The electrophoretic profile of proteins transiently expressed from GFP, g9, g9-TM, and g9-DTM plasmids and separated in 12% polyacrylamide gels were incubated with US9 and GFP antibodies. Closed symbols indicate bands corresponding to the different fusion proteins (as schematically indicated in B). M, in kDa $\times 10^3$ is reported on the left. doi:10.1371/journal.pone.0104634.g001

**Figure 1. Construction and analysis of GFP-US9 chimeras.** A) Kyte-Doolittle hydropathicity profile of US9 aminoacid sequence (aa reported under the plot). B) schematic topological orientations of GFP-US9 fusion proteins. The darker area inside the red ribbon, corresponding to the US9 full length sequence, indicates the position and the approximate extension of the TM domain. The two truncated chimeric US9 forms g9-ATM and g9-TM were generated by splitting the US9 aa sequence in the middle of the 6 arginine motif, as pointed by the arrowheads in the sequence in A. C) Western blot analysis of GFP-US9 fusion proteins expressed in transfected cells. Two identical membranes containing lanes with proteins from MDA cells transfected with the indicated constructs and separated on 12% polyacrylamide gels were incubated with US9 and GFP antibodies. Closed symbols indicate bands corresponding to the different fusion proteins (as schematically indicated in B). M, in kDa $\times 10^3$ is reported on the left.
Figure 2. Intracellular localization and membrane association of US9. A) MDA cells transfected with GFP, g9, g9-TM (GFP fused the US9 transmembrane domain), and g9-ΔTM (GFP fused the US9 hydrophilic domain) were imaged under a fluorescence microscope. Nuclei were stained with Hoechst 33342 (blue) and images were merged in B). C) Proteins extracted from MDA cells expressing the same constructs described in A) were divided in post-nuclear soluble (S), and membrane (M) fractions, separated on SDS Page, and revealed with GFP antibody. Bar = 10 μm.

doi:10.1371/journal.pone.0104634.g002
U9 localization is cell type independent

In this set of experiments, meant to investigate U9 ability to maintain its typical localization pattern independently from other viral factors, we introduced the g9 plasmid, carrying the full length gene fused to GFP, into several cell types derived from different species, including rat primary neurons (RN) and cells of human origin. All cells were transiently transfected and fixed 16 hours post transfection. Images representative of the localization patterns for 7 cell types, HOS, MDA, HEP-2, Saos, Vero, HA, and RN, have been chosen and assembled in figure 3. As already shown in figure 2 for MDA cells, U9-associated fluorescent signals were completely excluded from nuclei of transfected cells, and widely distributed into the cytoplasm, in bright puncta in all cell types tested. Taking in account the morphological differences between these cells, the punctate staining was similar in all cell types, with a more dense distribution in a perinuclear region, which has been previously defined as the U9 steady-state residence and well characterized with the aid of specific TGN markers [8]. Labeled vesicles in distal regions were always detected, even in cells with a heavy perinuclear fluorescent signal, as clearly seen in figure 3. While the steady-state accumulation of U9 at the TGN is not informative about the behavior of distal vesicles and putative U9 transport properties, the presence of a comparable punctate pattern in all g9-transfected cells suggests that this feature is solely dictated by U9 sequence, supporting our hypothesis of an autonomous, stand alone transport property of U9.

This peculiar trait (i.e. distal localization) of U9 phenotype is better visualized in cells characterized by long protrusions. This is obviously the case of neurons (figure 3-RN), the cell type in which the U9 virus transport function becomes critical. From an experimentally relevant viewpoint, a clear distal staining can be seen in MDA cells. As shown in figure 3 (MDA hm), in g9-transfected MDA cells, the typical punctuate fluorescent staining is detectable in long cellular protrusions and accumulates in large plates contacting neighbor cells.

Dynamic properties of U9

Data accumulated in PRV and HSV U9 studies showed that U9 is involved in virus transport inside the cells [7,12,35], and deletion mutants are defective in anterograde-dependent infection spread [10,11,13,15,19,36]. As expected, U9 localization strictly depends on the integrity of microtubular cytoskeleton (Figure S1). We were interested in understanding the dynamic properties of U9 in the absence of other viral factors. Therefore, we imaged the transfected live cells to look at the localization of the protein in vivo. MDA cells were transfected with g9, and imaged 16 h post-transfection. The results of these experiments are shown in figure 4 as well as in movies S1 and S2 (Supplemental Online Material). These real time imaging studies confirmed the typical punctuate staining seen in g9-transfected cells (Figure 4A). The green box in the figure frames the region of the cell (figure 4B) that has been followed over time. Selected frames from the complete movie S1 have been arranged in figure 4D and clearly show the dynamic properties of g9, with labeled puncta moving in the cytosol of expressing cells. Some movements appeared to be short in distance, with no preferential direction. However, some fluorescent spots did travel long distances, and these movements align with the longitudinal axis of the analyzed cell protrusion. The series of panels in figure 4D are sequential frames of g9 transfected cells in which two arbitrary chosen vesicles have been tracked with the aid of green and red triangles (Movie S2). The images clearly show long traits of linear movements of g9 containing vesicles. Different labeled vesicles move in anterograde (green triangles) or retrograde (red triangles) motions, and some even appear to change the direction of movement over time. Figure 4C reconstructs the trajectories of the two vesicles described above. The time frame for this dataset was less than 30 seconds, but we do not intend to draw any conclusion about the possible speed of U9 vesicles, as in the many movies we shot there was always a great variability in both speed and direction. Nevertheless, these results undoubtedly indicate that U9 can associate with vesicles transported inside the cell, even in the absence of other viral factors or other viral-dependent modifications, and that its sequence is sufficient to target GFP to the same transport vesicles.

Transfected U9 recapitulates viral U9 behavior

U9 role in vesicles transport has been extensively investigated with the intent to define its contribution to virus transport. Thus it is now established that U9 is necessary for virus anterograde transport in axons of infected neurons [7,9–12,15,22,35]; likewise, related virus-dependent post translational modifications of U9 have been identified [18,20,26,27]. However, from our perspective (i.e. determining the autonomous ability of U9 to act as a transport protein) it was important to compare the behavior of GFP-U9 expressed from a transfected plasmid to the one present in the context of virus infection, in distal regions of the cell. While TGN resident proteins have been used to positively ascertain the identity of the central region in which U9 accumulates [8], no unambiguous markers of distal vesicles have been identified. Early (Rab4, Rab5) and late (Rab7) endosomal markers showed little colocalization with viral transport vesicles [37]; in other studies, clathrin-like coats were only occasionally associated with L-particle and virion assembly sites [38]. In order to avoid any uneven estimation of the correspondence between stand alone and viral U9-labeled pattern, we employed a different approach and decided to use viral U9 as specific marker of these peripheral vesicles. However, this kind of analysis is constrained by the way viruses replicate and assemble, as explained below, and for this reason our comparison strategy included both a direct and an indirect analysis, whose results are shown in figures 5 and 6. For these studies, we replaced the GFP sequence in g9 with the gene coding a red fluorescent protein (RFP), and obtained a new construct, named r9, whose fluorescent distribution does not differ from that of g9 (as shown in figure 7, panel B). We also introduced the GFP-U9 chimeric gene from g9 in the viral genome to replace
the wild type U₉ gene, and the resulting recombinant virus expressing GFP-U₉ was named Fg₉.

Confocal microscopy studies in RNs infected with Fg₉ and transfected with r₉ were performed to analyze the distribution of plasmid-driven (red) and viral (green) U₉ (see top left panels r₉ and Fg₉ in figure 5; the two images are merged in the top right panel). Most vesicles appear yellow in the merge panel, demonstrating substantial, though incomplete, colocalization of the two chimeric proteins. Higher magnifications of different regions of the cell better define the level of colocalization. At the cell body, both in the area of higher expression/accumulation and where individual vesicles can be detected, a widespread yellow signal indicates the presence of colocalized U₉ and double-labeled vesicles. Nevertheless, zones/vesicles individually labeled with either the red or the green fluorophore can be spotted, indicating the presence of non-colocalizing fluorescent U₉ (figure 5, panel 1 merge). Idetical conclusions can be drawn from the magnified distal region in panels 2 (figure 5, bottom row).

The presence of these vesicles is not surprising, as viral infection normally generates a multiplicity of vesicles, including empty transport vesicles (i.e. containing no virion), as well as light particles (subviral assemblies, with tegument and envelope proteins, but no capsid). Moreover, U₉ is a constitutive component of the virion and consequently, in infected cells, it is also present inside the U₉-labeled transport vesicle. Finally, irrespective of the source of the tagged U₉ gene, the plasmid-encoded viral protein present in the infected cell is functionally undistinguishable from the one expressed from the viral genome.

This experimental approach based on the co-expression in the same cell of green and red U₉ allows canonical co-localization analysis. However, positive results shown in figure 5 are biased by the fact that both proteins (independently of their source) are exposed to the same infected molecular environment. For this reason we also compared the localization of stand alone and viral U₉ in different cells. Here we used tubulin as a “reference” molecule to better monitor U₉ distribution. Even in the absence

Figure 3. GFP-U₉ localization. Cells shown here are representative of several experiments and all share the typical GFP-U₉ pattern, i.e. a bright punctuate signal present throughout the cytoplasm and partially concentrated in a peri-nuclear region. Additionally, GFP-U₉ can be seen at the plasma membrane, while no fluorescent signal is present in the nuclear region. In the panel labeled MDA hm (higher magnification), the microphotograph focuses on the extension tip protruding from one transfected cell and reaching the body of one untransfected cell. GFP-U₉ can be detected in vesicles present in this distal region as well as on the plasma membrane. In green: GFP-U₉ fluorescence; nuclei are stained in blue with Hoechst 33342. Bar = 10 μm.
doi:10.1371/journal.pone.0104634.g003
of a direct interaction with U₉, several lines of evidence support the choice of tubulin: intact microtubular cytoskeleton is required for virus axonal spread [39]; KIF1A, a microtubule-dependent kinesin-3 motor, interacts with PRV U₉, in the presence of other viral factors [17]; isolated HSV-containing organelle structures are capable of efficiently bind purified microtubules in vitro (approximately 62% of the organelles bound) [37]. Finally, we show here that the depolymerizing agent Nocodazole disrupts U₉ localization (Figure S1).

The top left panel of figure 6 shows the already described punctuate signal clearly detectable both at the cell body and throughout neurites in a g9-transfected neuron (with cell morphology rendered in black & white in the second top panel). The two boxes in the left panels frame the regions of the cell magnified in the two top right panels, arbitrarily chosen to show U₉ localization at cell body and neurites extremities. A different magnification of the same cell is also present in figure S2. Besides the Golgi region, where individual vesicles cannot be detected due to U₉ high concentration and expression level, virus-independent U₉ is frequently found in close proximity to cytoskeleton microtubules, especially in distal regions.

This fluorescence distribution is basically reproduced in RNs infected with Fg9, the recombinant HSV-1(F) virus in which the U₉ coding sequence was replaced by the GFP-U₉ chimeric gene present in g9. The Fg9-infected cell in figure 6 (left bottom panel) and Figure S2 shows a very similar fluorescent pattern, with fluorescent vesicles accumulated both in perinuclear region and distributed throughout neurites. As for transfected g9 in the top panels, two representative regions were chosen (framed boxes in left two panels) and visualized at higher magnification in the two right bottom panels of figure 6. Analysis of these images clearly show the presence of U₉-associated vesicles and their relationship to immuno-labeled microtubules architecture, with the overall general distribution comparable to that of GFP-U₉ in the absence of virus shown in the top panels. Comparison in figure 6 is indirect, as plasmid- and virus-encoded U₉ are present in different cells – i.e. in a uninfected or infected molecular environment, respectively. Consequently, no actual co-localization can be assessed. Nevertheless, figure 6 shows the real stand alone phenotype of U₉.

Taken together, results from the experiments depicted in figures 5 and 6 indicate that the general distribution of stand alone and viral U₉ is highly comparable, with a large, major population of vesicles containing both plasmid- and virus-derived U₉, and the minor presence of vesicles incorporating only one of the two chimeric fluorescent proteins. All double and single labeled U₉ vesicles show no preferential localization and are randomly distributed along the described punctuate pattern of either infected or transfected cells, supporting the idea that U₉ properties do not depend upon other viral contributions.

Localization of mutated U₉

The cytosolic portion of the U₉ protein contains an acidic domain important for viral trans-neuronal spread [14,20]. Of the
18 residues of serine, threonine, and tyrosine scattered throughout the protein (accounting for 20% of total amino acids), two tyrosines (Y32Y33) and two serines (S34 and S36) are part of this acidic cluster. Substitutions of these residues with alanine have been shown to have an impact on virus transport and spread [18,20]. As already mentioned, US9 is heavily post-translationally modified, and enzymes responsible for phosphorylation of tyrosines and serines in the acidic cluster have been identified or postulated, in vitro or in silico [18,27,30,40].

Given this background, it was of absolute relevance to investigate whether these modifications are required to the stand alone molecular properties of US9. We constructed two new US9 fluorescent chimaeras, in which tyrosines 32 and 33 or serines 34 and 36 where substituted by alanines, named g9yyaa and g9aea, respectively. The localization patterns of the two US9 mutants were analyzed and compared in figure 7 with the RFP-US9 wt fluorescent staining. RNs co-transfected as outlined showed the typical vesicles distribution already described. Both g9yyaa (figure 7C) and g9aea (figure 7D) localize in puncta scattered throughout neurites and are also present in an intensively fluorescent area surrounding the nuclei. Merging the images obtained by wild type (red) and mutants (green) US9 localization generates a perfect yellow pattern (figure 7C and 7D, right panels). From these results we can conclude that these US9 modifications, shown to affect virus transport and infection spread in PRV models, do not modify US9 behavior, supporting the hypothesis that US9 does not require additional viral factors in order to exert its transport function.

**Discussion**

Efficient and specific transport of single molecules and multi-molecular complexes to different cellular compartments and structures or to the outside environment is a key feature of eukaryotic cells. Viruses - unwanted travelers that must efficiently move inside the infected cell in order to reach specific compartments and complete their replication cycle [41] - use the highly efficient and specialized cell transport system. Identification and analysis of viral proteins contributing to (or determining) this dynamic phenotype can eventually lead to a virus-independent
approach to control cellular transport and delivery. From this perspective we looked at US9, a viral protein highly conserved amongst alphaherpesviruses, with a critical role in anterograde spread of the virus [7–12,19]. Herpes Simplex Virus is a lifelong guest of infected hosts, in which it becomes and remains latent in dorsal root ganglion sensory neurons [42]. Given the peculiar replication cycle of the virus and the morphology of the infected neurons, HSV must traverse very long distances inside the cell, in both retro- and antero-grade directions, making it a good potential tool to understand (and conceivably direct) cellular transport machinery while gaining insights into viral trafficking. Deletion of US9 gene from the genomes of several alphaherpesviruses generates recombinant viruses with impairments in the ability to spread in the nervous system due to defects in anterograde transport [8–12,15,23]. Studies aimed at defining the intracellular localization of the viral protein showed a punctuate staining in cells expressing US9 fused to GFP, with a steady-state accumulation in the TGN region and vesicles scattered throughout the cell [8]. US9 was also found at the plasma membrane, from which it is recycled into cytoplasmic vesicles [20].

The key question we sought to address here is whether HSV US9 maintains its transport properties in the absence of other viral factors, and independently of its natural cargo, i.e. the virion. Our long-term goal is to use this information to be able to exploit US9 as tool for directed cellular/intercellular delivery, in the context of both mechanistic studies and drug-delivery purposes. In infected cells, besides being a constitutive component of transported virions, US9 is present on the membrane of transport vesicles. We reasoned that for a viral protein to be necessary for virus transport, it must be part of the molecular complex that connects the cellular transport machinery to its cargo. This implies that US9 (as any other protein with similar function) exerts two, possibly independent, activities. We may call these transport activity (i.e. the ability to engage cellular transport machinery and become part of vesicles that are exported along cytoskeleton structures) and cargo selection (i.e. the ability to direct virion load and consequent transport). Our study has been instrumental in determining the

Figure 6. Localization in neurons of US9 expressed from the viral genome or in virus-free context. RNs were either transfected with the GFP-US9 plasmid (g9: top panels) or infected (MOI = 10 pfu/cell) with the recombinant HSV-1(F) carrying the chimeric GFP-US9 gene (Fg9: bottom panels). Left first panels in both rows show the fluorescent pattern of transfected or infected cells, with the bright punctuate signal concentrated in the cell body and also dispersed along neurites. Two representative regions (indicated by small frames in left panels) from both transfected and infected cells were chosen and are shown at higher magnification in the right 2 panels (yellow frames: cell body; blue frames: neurites tips). Black & white rendering of the transfected or infected neurons shown on left panels were obtained by pseudo-coloring the microtubulin cytoskeleton revealed by immunofluorescence (shown in red in the right two panels), and are included to help locate the magnified regions in the complex neurons morphology. Both central and peripheral regions of transfected and infected cells show the microtubular architecture (red) with US9-vesicles (green). In both regions, localization of transfected US9 seems to mostly overlap that of US9 expressed in the viral context. Bar = 10 μm (in the left 2 panels); Bar = 5 μm (in the right 2 panels).

doi:10.1371/journal.pone.0104634.g006
extent to which these two functions are independent. This is a necessary condition to establish US9 as an autonomous transport protein for trafficking studies or specific delivery.

HSV-1 GFP-US9 chimera constructed here shows the distinctive fluorescent pattern in transfected cells, i.e. a bright punctuate signal indicating a vesicular expression, with labeled vesicles scattered throughout the cytoplasm and partially concentrated in a perinuclear region already defined as the steady-state site of accumulation of US9, and identified as Trans Golgi Network [8]. Additionally, US9 is detectable at the plasma membrane. GFP-US9 localization is almost identical in all tested cell types, suggesting that its behavior is an intrinsic property of the protein. Also interesting is the finding that US9-containing vesicles are always found in peripheral regions. In cells with elongated shape US9 is spotted at cellular tips, and this distal staining is not directly related to the intensity of the signal in the perinuclear region.

The key evidence for the US9 autonomous transport properties comes from real time experiments in which transfected cells are imaged in vivo, and GFP-US9 localization is recorded over an extended period of time. The resulting movie shows the actual highly dynamic behavior of US9 in the absence of other viral factors. Focusing on elongated regions of the cell (though the phenotype is not restricted to these areas), fluorescent vesicles can be seen traveling in both antero- and retrograde directions. Some vesicles invert the direction they move over the period the movement is recorded. This discontinuous activity is not surprising as often vesicles moving in one direction can apparently invert their motion by jumping on near microtubules tracks or through the coordination activity of a macromolecular complex [43–46]. As a consequence, the final transport may result from several short movements that can occur in both directions.

In PRV, US9 has been shown to interact with KIF1A, a microtubule-dependent kinesin-3 motor, and the interaction is dependent on the presence of other still unidentified viral factor(s) [17]. The same report demonstrates that HSV-1 US9 cannot be co-purified with KIF1A, launching the quest for a HSV US9 motor partner. Irrespective of the identity of that molecule, our data suggest that HSV US9 can engage the cellular transport machinery, and that its dynamic behavior occurs in the absence of other viral factors. It is possible that US9-dependent axonal

Figure 7. Localization of US9 mutated forms in RNs. A) Two major phosphorylation sites were mutated in the US9 sequence and the modified forms were expressed as fusions with GFP, in the same topological orientation as GFP-US9. r9 and g9 fusion proteins have a completely overlapping staining pattern, as shown in B (right panel is the merge of g9 and r9 images shown on the left). A perfect colocalization is obtained in cells cotransfected with r9-g9yyaa (C) and with r9-g9aea (D). In the figure, GFP- and RFP-chimeras are in green- and red-framed images, respectively, and merged images are framed in yellow in the right panels. Bar = 10 μm.
transport of HSV particles requires additional proteins, as suggested by the PRV data, with glycoproteins E and I being good, but not exclusive, candidates to play such a role. Nevertheless and not considering virion transport, U9-containing vesicles traffic occurs with no additional factors.

In primary neurons, the GFP-U9 localization could be clearly appreciated and may result extremely helpful to understand the molecular basis of U9 activity. In these cells (as on the other cell types analyzed), GFP-U9 fluorescent vesicles localize both at cell body and throughout neurites, frequently in close proximity to microtubules (figure 6, g9). Identical distribution and microtubules proximity (in distal regions) were seen in neurons infected with a recombinant virus expressing GFP-U9 (figure 6, Fg9), showing no differences with the behavior of the viral protein expressed in the absence of other viral factors. By using different fluorescent tags to distinguish stand alone- (transfected), and virus-encoded-U9, we were able to precisely assess the extent of co-localization. The two differently tagged versions of U9 co-expressed in the same cell mostly localize in the same vesicles, in both proximal and distal cellular regions. Few but constantly present individually tagged vesicles with no preferential sub-localization were detected in all the experiments, and this reproducible distribution can be interpreted with the equivalence between the two differently tagged U9. As well established, due to the peculiar way viruses replicate themselves, there is no difference between stand alone (transfected) RFP-U9 and virus-encoded GFP-U9, once they are co-expressed in the same infected cell. The presence of few green or red, and many yellow vesicles simply demonstrates U9 incorporation in these transport vesicles (not all necessarily carrying complete viral particles, as natural infection always produces empty and light particles).

In agreement with our hypothesis of an autonomous transport activity is the observation that ablations of two phosphorylation sites in the cytosolic tail of U9 do not affect U9 localization. PRV transneuronal spread depends (to different extents) on the presence of two tyrosines and two serines in a conserved acidic cluster [18,20]. Results presented here, showing co-localization of wild of two tyrosines and two serines in a conserved acidic cluster of the sequence encoding the 6 arginine stretch of US9 (figure 1A)

Materials and Methods

Ethics Statement

Animals were used as a source of brain tissue to prepare neuronal cultures, following the recommendations in the Guide for the Care and Use of laboratory Animals of the National Institute of Health. This protocol for harvesting brain tissue was approved by the Institutional Animal Care and Use Committee of Drexel University (PHS Animals Welfare Assurance #A-3222-01), approved on 10-18-2012 (permit #20098).

Cells, transfections, infections and treatments

MDA-MB-231 is a human breast cancer cell line [47]. HOS is a osteosarcoma cell line of human origin, with mixed, fibroblast and epithelial like morphology, (obtained through the NIH AIDS Reagent Program, Division of AIDS, NIAID, NIH: HOS- p.BABE-puro from Dr. Nathaniel Landau [48–49]). Saos-2 is a non-transformed cell line derived from the primary osteosarcoma of an 11-year-old Caucasian girl [50]. HEp2 cells [51] were derived via HeLa contamination. Vero cell line [51] was derived from kidney of a normal adult African green monkey. All cells lines were cultured in Dulbecco Modified Eagle Medium (DMEM) supplemented with 10% Fetal Calf Serum. Normal human astrocytes (HA) derived from neural precursor cells were purchased from ScienCell Research Laboratories (San Diego, CA), and cultured in Astrocyte Medium supplemented with fetal bovine serum (2%) and Astrocyte Growth Supplement provided by the vendor. Rat cortical neurons (RNs) were obtained from the brains of 17–18 day old rat embryos and cultured in Neurobasal medium containing B27 supplement as detailed by Sengupta et al. [52] and originally described by Brewer et al. [53].

For transfections, cells were seeded on 35 mm dishes or on glass coverslips in 35 mm dishes 16–24 hours prior to the experiment. RNs were plated on poly-L-lysine-coated coverslips. Transfections were performed with Lipofectamine 2000 (Invitrogen), using 2 μg of plasmid DNA. In all the experiments described, plasmids were introduced into the cells by transient transfection, and routinely ≥ 50% of transfected cells were obtained, as estimated by fluorescence microscopy. Nocodazole was obtained from Sigma, and used at 1 μg/ml concentration.

Herpes Simplex Virus 1 (HSV-1) strain F, and the derived mutant strain F9 were amplified and titered in Vero cells, and inoculated in RNs at multiplicity of infection (MOI) = 10 plaques forming units per cell (pfu/cell).

Plasmids and recombinant viruses construction

The U9 sequence was amplified from HSV-1 genome and cloned into pEGFP-C1 (Clontech) to generate g9, with the insertion of the Hemoagglutinine (HA) epitope. The intercalary spacer between the two molecules results in the following peptide: KSGLRISSSSSEF-MAYPPYDVPDYASLGGMAMGMTS (EGFP - HA - U9).

To generate a red fluorescent version of tagged U9, the GFP sequence in g9 was substituted with the RFP sequence from pTagRFP-tubulin (Evrogen, Moscow, Russia).

g9-ATM was obtained by collapsing in g9 the trans-membrane sequence comprised between two Sall sites, present in the middle of the sequence encoding the 6 arginine stretch of U9 (figure 1A) and in the multiple cloning site of pEGFP-C1. g9-ATM has the same topological orientation as g9, with the U9 sequence lacking the trans-membrane coding portion.

GFP fused to the U9 trans-membrane domain was obtained by inserting the U9 sequence starting from Sall site in the middle of the 6 arginine stretch (figure 1A) into pEGFP-C1 XhoI site. The resulting plasmid was called g9-TM and lacks the HA epitope.
g9yyaa and g9aea are mutant versions of g9 in which U₉ g codons TAC-TAC (encoding two tyrosines at positions 32 and 33) and TCG-GAA-AGC (encoding serine-glutamic acid-serine at positions 34-36) have been mutagenized to obtain GCC-GCC (alanine-alanine) and GCC-GAA-GCG (alanine-glutamic acid-alanine), respectively.

To obtain the recombinant HSV-1 expressing the GFP-U₉ protein, the chimeric g9 coding sequence from g9 plasmid was cloned into an intermediate vector containing the BamHI X fragment from the HSV genome, with a short polylinker replacing the U₉ sequence downstream of the gene sixth codon. In the resulting plasmid, GFP-U₉ fusion gene substitutes the original U₉ sequence, with large enough viral genomic homologous regions on both sides of the insertion. This plasmid was co-transfected in Vero cells with the viral genomic DNA and recombinant viruses in fluorescent plaques were isolated from the monolayer. The recombinant virus, named Fg9, was plaque-purified for 5 times and finally titered in Vero cells. The resulting viral fusion protein expression is under U₉ natural promoter control and in the same genetic context, as GFP-U₉ translation begins after the first 6 U₉ amino acids.

All plasmids generated in the study were sequenced.

Protein electrophoretic analysis and western blotting

Protein extracts were separated on denaturing polyacrylamide gels (SDS Page) and transferred to a PVDF membrane for immunoblotting. Cell fractionation was performed as described by Sadoul et al. [33] and Fournier et al. [34]; post-nuclear fractions containing soluble (S) or crude membrane (M) proteins were analyzed by SDS Page. Primary antibodies used were the U₉ specific antibody [31], and anti-GFP antibody (sc-8334, SantaCruz Biotechnology). Secondary antibodies and detection reagents were from Pierce (SuperSignal West Femto kit).

Fluorescence microscopy analysis and acquisition

Except for real time experiments, cells were fixed in 4% paraformaldehyde at the indicated times post transfection/infection. For cytoskeleton immuno-staining of RNs, fixed cells were permeabilized in Phosphate Buffered Saline (PBS) containing 0.2% Triton, 1% Bovine Serum Albumine (BSA), 0.5% Newborn Calf Serum (NCS) for 30 minutes at Room Temperature. Permeabilized cells were first incubated with a monoclonal antibody against neuronal class III β-Tubulin, clone TUJ1 (Covance) and then with the anti-mouse Alexa Fluor 568 from Life Technologies. For MDA cells, antibodies used were anti β-tubulin (Sigma, T4026) and anti-mouse IgG (Fab specific)–TRITC (Sigma, T7782). Nuclear counter-staining was done via incubation of fixed cells with Hoechst 33342 (Molecular Probes - Invitrogen).

Confocal images were taken under a Zeiss Axio Imager.Z1m microscope equipped with Plan-apochromat 63x/1.4 oil DIC objective. For all the other experiments, cells were observed and recorded under the Nikon Eclipse E600 fluorescence microscope (filters UV2A, B2A, G2A) equipped with 100× objective. Acquisition software was the provided Nikon ACT-2U. For real time imaging, the output generated by ACT-2U was captured by the freeware Debut. All images were analyzed and assembled using ImageJ. Graphical elements were added using Gimp2 and Inkscape. Images are representative of the original data and comply with the journal image manipulation policy.

Supporting Information

Figure S1 U₉ localization is dependent on the integrity of cytoskeleton. Intracellular trafficking is organized along cytoskeletal structures. Cells treated with Nocodazole, an antimitotic agent commonly used to disrupt the cellular cytoskeleton, undergo a complete rearrangement of cytoskeleton, with tubulin molecules becoming redistributed throughout the cytoplasm. Nocodazole effect is reversible, as incubation of treated cells in medium lacking the drug leads to a complete recovery of normal cellular cytoskeletal structures. MDA cells transfected with g9 plasmid were treated for the indicated time (0, 30, 60, and 120 minutes) with Nocodazole, immunostained with an anti-tubulin antibody, and imaged under a fluorescence microscope. Effect of treatment on U₉ localization (in green) is compared to the disruption of cytoskeleton (immunostained in red with an anti-tubulin antibody) induced by the presence of the drug. The complete rearrangement of U₉-associated vesicular staining (g9 t30 through g9 t120) increases in a time dependent manner and perfectly matches tubulin disorganization. 30 minutes after removal of Nocodazole from culturing medium (g9 t120 rec) cytoskeleton structures begin to appear, and corresponding U₉-labeled normal vesicular pattern becomes detectable (white arrows in panel ‘g9 t120 rec’). 2 hours nocodazole treatment of cells expressing g9-TM truncated form does not seem to have a major impact on the chimera localization. Bar = 10 μm. (TIF)

Figure S2 Localization in neurons of U₉ expressed from the viral genome or in virus-free context. RNs were either transfected with the GFP-U₉ plasmid (g9: top panels) or infected (MOI = 10 pfu/cell) with the recombinant HSV-1(F) carrying the chimeric GFP-U₉ gene (Fg9: bottom panels). Left first panels in both rows show the bright fluorescent punctuate pattern in transfected or infected cells. Central panels show the immunostained microtubular cytoskeleton. In right panels, the two images on the left have been merged. Localization of transfected U₉ seems to mostly overlap that of U₉ expressed in the viral context. This supplemental figure shows a different magnification of the same cells in figure 6. Bar = 10 μm. (TIF)

Movie S1 U₉ trafficking does not depend on other viral functions. Real time imaging of GFP-U₉ vesicles trafficking inside a MDA transfected cell. Only a portion of a g9-transfected MDA cell is shown in the movie. (AVI)

Movie S2 Directional transport of U₉ vesicles. Movie showing a portion of a GFP-U₉-transfected MDA cell imaged in real time. Frames from Movie S1 have been chosen and labeled with green and red arrows to track the movements of arbitrary chosen vesicles. Green arrows = anterograde direction; red arrows = retrograde direction. (AVI)

Acknowledgments

The authors would like to thank Alessandro Fatatis (Drexel University College of Medicine) for critical discussion.

Author Contributions

Conceived and designed the experiments: OM RB. Performed the experiments: MP RB. Analyzed the data: OM RB. Contributed reagents/materials/analysis tools: BN. Wrote the paper: RB OM.
References

1. Hirokawa N, Noda Y, Okada Y (1998) Kinesin and dynein superfamily proteins in organellar transport and cell division. Curr Opin Cell Biol 10: 60–73.

2. Hirokawa N (1996) Organellar transport along microtubules — the role of KIFs. Trends in Cell Biology 6: 135–141.

3. Mitchison T, Kirschner M (1984) Dynamic instability of microtubule growth. Nature 312: 237–241.

4. Nascimento R, Costa H, Parkhouse RME (2012) Virus manipulation of cell cycle. Protoplasma 249: 519–528.

5. Hill A, Juguve P, York I, Russ G, Bennek J, et al. (1995) Herpes simplex virus turns off the TAP to evade host immunity. Nature 375: 411–415.

6. Fruh K, Ahs K, Djaballah H, Sempe P, van Endert PM, et al. (1995) A viral inhibitor of peptide transporters for antigen presentation. Nature 375: 415–418.

7. Brideau AD, Carancho, D, Enquist LW (1999) Intracellular trafficking and localization of the pseudorabies virus US9 type II envelope protein to host and viral membranes. J Virol 73: 4372–4380.

8. Chowdhury SI, Onderci M, Bhattacharjee PS, Al-Mubarak A, Weiss, et al. (2002) Bovine herpesvirus 5 (BHV-5) US9 is essential for BHV-5 neuropathogenesis. J Virol 76: 3039–3051.

9. LaVail JH, Tauscher AN, Sucher A, Harrabi O, Brandimarti R (2007) Viral regulation of the long distance axonal transport of herpes simplex virus nucleocapsid. Neuroscience 146: 974–985.

10. Poliozova K, Bwasa PS, Banerjee K, Wimen WT, Rouse BT, et al. (2005) Herpesvirus kentosis in the anterior transport of virus from sensory ganglia to the central nervous system. Proc Natl Acad Sci USA 102: 11462–11467.

11. Tomishima MJ, Enquist LW (2001) A conserved alpha-herpesvirus protein necessary for axonal localization of viral membrane proteins. J Cell Biol 154: 711–725.

12. Enquist LW, Tomishima MJ, Gross S, Smith GA (2002) Directional spread of an alpha-herpesvirus in the nervous system. Vet Microbiol 86: 5–16.

13. Feierbach B, Bisher M, Goodhouse J, Enquist LW (2007) In vitro analysis of transneuronal spread of an alphaherpesvirus infection in peripheral nervous system neurons. J Virol 81: 6846–6857.

14. Lyman MG, Curanovic D, Brideau AD, Enquist LW (2008) Targeting of pseudorabies virus US9 protein with homologs from other veterinary and human alphaherpesviruses. J Virol 82: e58776.

15. Lyman MG, Curanovic D, Brideau AD, Enquist LW (2008) Role of pseudorabies virus US9, a type II membrane protein, in infection of tissue culture cells and the rat nervous system. J Virol 74: 934–945.

16. Lyman MG, Curanovic D, Brideau AD, Enquist LW (2008) Fusion of Enhanced Green Fluorescent Protein to the Pseudorabies Virus A2126II Membrane Protein. J Virol 82: 10308–10311.

17. Kato A, Tanaka M, Yamamoto M, Asai R, Sata T, et al. (2005) Identification of a phylogenetic phylogenetic analysis of the herpes simplex virus 1-encoded protein kinase US5 which regulates its optimal catalytic activity in vitro and influences its function in infected cells. J Virol 82: 6172–6189.

18. Roizman B, Knipe DM (2001) Herpes simplex viruses and their replication. In: Roizman B, Knipe DM (eds) Herpes simplex virus type 1. Lippincott Williams & Wilkins, Philadelphia, PA, pp 741–752.

19. Chandra-Sakker M, Armiti P, Boadle RA, Holland DJ, Cunningham AL (2000) Antergastric transport of herpes simplex virus type 1 in cultured, dissociated human and rat dorsal root ganglion neurons. J Virol 74: 1827–1839.

20. Kennedy SW, Serpinskaya AS, Vaughan PS, Lopez Fanarraga M, Vernos I, et al. (2003) Dynactin is required for bidirectional organelle transport. J Cell Biol 160: 297–305.

21. Gross SP, Welte MA, Block SM, Wieschaus EF (2002) Coordination of opposite polarity Motor protein activities to drive bidirectional transport. Cell 113: 527–538.

22. Cailleau R, Young R, Olivere M, Reeves WJ Jr (1974) Breast tumor cell lines from human mammary carcinomas. J Natl Cancer Inst 52: 754–763.

23. Polcicova K, Biswas PS, Banerjee K, Wisner TW, Rouse BT, et al. (2005) Visualization of an alphaherpesvirus membrane protein that is essential for fast axonal transport. Mol Biol Cell 16: 741–752.

24. Lyman MG, Curanovic D, Enquist LW (2007) Pseudorabies virus US9 directs axonal sorting of viral capsids. J Virol 81: 11363–11371.

25. Butcher D, Banfield BW, Enquist LW (1998) The US9 gene product of pseudorabies virus, an alphaherpesvirus, is a phosphorylated, tail-anchored type II membrane protein. J Virol 72: 4560–4570.

26. Butcher D, Kemp CD, Taylor MP, Enquist LW (2009) Comparison of the pseudorabies virus US9 protein with homologs from other veterinary and human alphaherpesviruses. J Virol 83: 6978–6986.

27. Chowdhury SI, Brum MCS, Coats C, Doster A, Wei H, et al. (2011) The bovine herpesvirus type II envelope protein US9 acidic domain is crucial for anterograde axonal transport. Vet Microbiol 152: 270–279.

28. Collier KE, Smith GA (2000) Two viral kinases are required for sustained long distance axon transport of a neuroinvasive herpesvirus. Traffic 9: 1458–1470.