The Length of Vesicular Stomatitis Virus Particles Dictates a Need for Actin Assembly during Clathrin-Dependent Endocytosis

The Harvard community has made this article openly available. Please share how this access benefits you. Your story matters

Citation
Cureton, David K., Ramiro H. Massol, Sean P. J. Whelan, and Tomas Kirchhausen. 2010. “The Length of Vesicular Stomatitis Virus Particles Dictates a Need for Actin Assembly during Clathrin-Dependent Endocytosis.” Edited by John A. T. Young. PLoS Pathogens 6 (9): e1001127. https://doi.org/10.1371/journal.ppat.1001127.

Citable link
http://nrs.harvard.edu/urn-3:HUL.InstRepos:41483486

Terms of Use
This article was downloaded from Harvard University’s DASH repository, and is made available under the terms and conditions applicable to Other Posted Material, as set forth at http://nrs.harvard.edu/urn-3:HUL.InstRepos:dash.current.terms-of-use#LAA
The Length of Vesicular Stomatitis Virus Particles Dictates a Need for Actin Assembly during Clathrin-Dependent Endocytosis

David K. Cureton1, Ramiro H. Massol2, Sean P. J. Whelan3, Tomas Kirchhausen1*

1 Department of Cell Biology, Harvard Medical School, and Immune Disease Institute at Children’s Hospital, Boston, Massachusetts, United States of America, 2 The Division of Gastroenterology and Nutrition, Children’s Hospital, Boston, Massachusetts, United States of America, 3 Department of Microbiology and Molecular Genetics, Harvard Medical School, Boston, Massachusetts, United States of America

Abstract

Microbial pathogens exploit the clathrin endocytic machinery to enter host cells. Vesicular stomatitis virus (VSV), an enveloped virus with bullet-shaped virions that measure 70 × 200 nm, enters cells by clathrin-dependent endocytosis. We showed previously that VSV particles exceed the capacity of typical clathrin-coated vesicles and instead enter through endocytic carriers that acquire a partial clathrin coat and require local actin filament assembly to complete vesicle budding and internalization. To understand why the actin system is required for VSV uptake, we compared the internalization mechanisms of VSV and its shorter (75 nm long) defective interfering particle, DI-T. By imaging the uptake of individual particles into live cells, we found that, as with parental virions, DI-T enters via the clathrin endocytic pathway. Unlike VSV, DI-T internalization occurs through complete clathrin-coated vesicles and does not require actin polymerization. Since VSV and DI-T particles display similar surface densities of the same attachment glycoprotein, we conclude that the physical properties of the particle dictate whether a virus-containing clathrin pit engages the actin system. We suggest that the elongated shape of a VSV particle prevents full enclosure by the clathrin coat and that stalling of coat assembly triggers recruitment of the actin machinery to finish the internalization process. Since some enveloped viruses have pleomorphic particle shapes and sizes, our work suggests that they may use altered modes of endocytic uptake. More generally, our findings show the importance of cargo geometry for specifying cellular entry modes, even when the receptor recognition properties of a ligand are maintained.

Coated pits incorporate and internalize soluble cargos of various sizes, such as transferrin (5 nm) [9,11] and low density lipoproteins (25 nm) [9,12]. Many viruses and intracellular bacteria are also internalized by the clathrin machinery [9,13–16]. We previously evaluated how cells internalize the 70 × 200 nm bullet-shaped vesicular stomatitis virus (VSV). We found that VSV internalization occurs through elongated, partially clathrin-coated structures that have longer lifetimes (~2 min.) than typical endocytic clathrin-coated vesicles and require local actin polymerization for uptake [15]. During VSV internalization, the clathrin coat first assembles as a partially closed dome at one end of the virion [15,17], and growth of the coat stalls when it encounters the long particle axis. Actin assembly then drives one or more late stage(s) of the internalization process, as recruitment of the actin machinery peaks during completion of clathrin assembly, and pharmacological inhibition of actin polymerization blocks VSV internalization without interfering with clathrin coat assembly [15]. Relatively small, spherical viruses like dengue virus (50 nm) [13] and some influenza A viruses (X-31 strain, ~120 nm) [14,18] also enter using a clathrin-dependent route, but it is unclear whether actin function is required for their uptake. Our observations with VSV led us to
Author Summary

We present a detailed comparison between the clathrin-dependent entry mechanisms of a parental virus (VSV) and its smaller defective interfering particle (DI-T). We used the difference in virion length to probe why actin assembly is required for the uptake of full-length VSV particles by nonpolarized mammalian cells. By imaging the entry of single particles in an unbiased manner, we resolved differences in the maturation kinetics, clathrin content, and actin dependency of clathrin endocytic structures internalizing VSV or DI-T virions. Our principal finding is that, unlike VSV uptake, DI-T internalization does not induce or require robust actin polymerization. We have also established, for the first time, that the geometry of an endocytic cargo can alter the mechanism of clathrin uptake. We propose that VSV-containing clathrin structures display characteristics of ‘frustrated’ endocytic intermediates that cells resolve by using the force of actin assembly to deform the plasma membrane into a complete endocytic vesicle.

Defective interfering (DI) particles arise spontaneously during virus replication. Such particles depend upon coinfecting helper virus to support their replication but contain all the essential acting regulatory elements for genome replication and assembly. One such well-characterized DI particle of VSV is termed DI-T, which lacks 82% of the viral genome [19,20]. Since the length of a VSV particle is dictated by the genome size [21], DI-T particles are 75 nm long and appear as truncated bullets by electron microscopy [22]. DI-T particles contain normal proportions of the viral structural proteins [23], including the viral surface glycoprotein (G), which mediates VSV attachment and entry into host cells.

Here we took advantage of significant differences in the physical dimensions of VSV and DI-T to investigate how the geometry of a viral cargo influences the actin-dependency of clathrin internalization. Using live cell fluorescence microscopic imaging, we compared the uptake mechanisms of VSV and DI-T at the single particle level. We report that in contrast to the clathrin- and actin-dependent uptake of VSV, the shorter DI-T particles enter cells through fully coated clathrin carriers that do not require actin dynamics for vesicle budding. These observations highlight the plasticity of the clathrin endocytic system, where clathrin coats serve as a scaffold to direct actin assembly when the clathrin machinery alone is not sufficient to mediate internalization.

Results

Biological properties of DI-T particles

To generate a clonal population of VSV DI-T particles, we recovered DI-T from cDNA (Figure 1A) [24] and amplified the particles by co-infection of cells with VSV [25]. We separated DI-T particles from VSV by rate zonal centrifugation in a sucrose density gradient. Electron microscopic analysis (Figure 1B, C) confirmed that VSV virions measure 70+/−8 nm by 204+/−14 nm (n = 114) [21], while the shorter DI-T particles have a length of 76+/−8 nm (n = 81) [22]. DI-T particles, like VSV, are covered with spike-like projections that correspond to homotrimers of G protein (Figure 1B), and SDS-PAGE analysis of purified particles confirmed that VSV and DI-T particles contain similar ratios of G protein to core virion components (Figure 1D) [23]. The purified stocks of DI-T lacked full-length virions (Figure 1B), with only a single VSV virion observed amongst more than 3,000 DI-T particles. Limited dilutions of the purified DI-T stock contained ∼1x10⁶ plaque forming units of virus per microgram of total viral protein, or 10³ times fewer infectious particles than for an equivalent protein quantity of VSV particles (not shown). Thus, we have successfully purified relatively homogeneous populations of VSV and DI-T particles, which differ only in their physical dimensions.

DI-T particles enter cells by clathrin-dependent endocytosis

To visualize VSV and DI-T by fluorescence microscopy, we covalently labeled the G proteins with spectrally separable fluorescent dye molecules (Alexa Fluor 568 and 647, respectively) using conditions that do not reduce viral infectivity [15]. Spinning disk confocal images of labeled particles adsorbed onto glass coverslips showed diffraction-limited objects with single-peaked distributions of fluorescence intensity values (Figure 1E), indicating that the DI-T and VSV populations primarily consist of individual particles [15]. We tracked the entry of DI-T particles into BSC1 cells stably expressing an eGFP-tagged σ2 subunit of the AP-2 adaptor complex (σ2-eGFP), which incorporates into all clathrin-coated structures that form on the plasma membrane [9,15,26]. Single DI-T particles readily attached to cells and progressed through the following set of defined events (see Figures 2A, B; Video S1 for examples): (1) membrane-bound DI-T particles diffused slowly (D = 5x10⁻¹¹–5x10⁻¹² cm² s⁻¹) and with the random directionality characteristic of Brownian motion; (2) shortly after DI-T attachment, a dim spot of AP-2 signal arose and remained colocalized with the particle, signifying incorporation of DI-T into an assembling clathrin-coated pit; (3) the AP-2 signal steadily increased over time until it peaked as coat assembly completed, and the DI-T particle then underwent an abrupt movement into the cell, after which the AP-2 signal disappeared due to clathrin uncoating. This sequence of events is identical to what we previously observed for VSV entering cells by clathrin-dependent endocytosis [15]. Moreover, the efficiency of DI-T uptake via the clathrin pathway is similar to that of the full-length VSV particles, as 89% (55/62) of DI-T particles that attached to 3 individual cells during imaging entered by clathrin-dependent endocytosis. These data show that DI-T efficiently enters cells through the clathrin pathway and validate the use of DI-T and VSV as comparative endocytic cargos.

Clathrin structures capture VSV and DI-T particles with similar kinetics

To directly compare how DI-T and VSV particles engage the clathrin machinery, we simultaneously inoculated BSC1 cells with the two spectrally distinct particle forms and then analyzed their mode of incorporation into AP-2 containing clathrin structures on a single cell basis (Figure 2C). For each complete virus uptake event, we quantified the kinetics of particle capture by measuring the elapsed time between virion attachment and the appearance of an AP-2 signal that colocalized with the bound particle. The capture time for DI-T particles was 110+/−80 s (n = 121), which is statistically indistinguishable (Student’s t-test, p = 0.2) to that measured for VSV (130+/−125 s (n = 87)) and agrees well with our prior measurements (Figure 2D) [15]. The AP-2 structures that captured DI-T or VSV initiated within a ~250 nm zone (the resolution limit of the optical system) of the attached particle. We therefore conclude that DI-T and VSV particles engage the clathrin system in an indistinguishable manner, which likely reflects a shared mechanism triggered by the same viral glycoprotein-receptor interactions.
Cells internalize DI-T particles using conventional clathrin-coated vesicles

To investigate the characteristics of the clathrin coat responsible for DI-T internalization, we imaged the uptake of both particle forms by the same BSC1 cell. We found that DI-T particles are internalized through AP-2 containing structures significantly faster than full-length virions (Figure 3A, B; Video S2). Quantitative analysis of data compiled from 4 cells showed that pits incorporating DI-T (n = 36) form in 43 ± 14 s, which is similar to the assembly kinetics of pits that lack virus particles (n = 212, 35 ± 10 s) (Figure 3C, Table 1). As expected, AP-2 structures that capture VSV (n = 29) require longer (75 ± 22 s) to complete (Figure 3C, Table 1). A similar analysis conducted in cells transiently expressing eGFP-tagged clathrin light chain A1 (eGFP-LCa) yielded analogous results (Figure 3D, E, Table 1, Videos S3, S4). The interaction of full-length VSV with a cell had no impact on the uptake kinetics of DI-T into the same cell (Table 1).

The different kinetics of DI-T versus VSV internalization suggests that DI-T enters cells through conventional fully coated clathrin structures and not the partially coated vesicles responsible for VSV uptake. To further investigate this possibility, we measured the maximum fluorescent signal of eGFP-tagged AP-2 or clathrin molecules, as this peak signal is known to be proportional to the overall size of a clathrin coat [9,15,26]. We compared this value for structures that contained DI-T with those that contained VSV or lacked either particle. Similar quantities of coat components were present in structures associated with DI-T to those lacking viral particles (Figure 3C, E, Table 1). As expected, VSV-containing structures accumulate more AP-2 and LCa molecules than structures lacking virus (Figure 3C, E, Table 1). Taken together, the above experiments suggest that DI-T enters cells through pits that acquire a full clathrin coat.

Consistent with this, electron micrographs of DI-T particles captured during cell entry show particles present in circular pits entirely surrounded by a clathrin coat (Figure 4A). This is in marked contrast to the partial clathrin coat found at one end of the endocytic carriers that internalize VSV (Figure 4B) [15].

Clathrin structures containing VSV recruit more cortactin that pits that internalize DI-T

During the final phase of coat assembly, endocytic clathrin structures associated with VSV show a strong recruitment of cortactin, an F-actin and dynamin binding protein that activates Arp2/3-mediated assembly of branched actin filaments [15,27,28]. To determine whether DI-T uptake is associated with an acute recruitment of cortactin, we monitored internalization...
into BSC1 cells transiently co-expressing monomeric Cherry-LCa (mCherry-LCa) and low levels of cortactin-eGFP. As previously shown [15, 26, 29], conventional clathrin-coated pits exhibit minimal cortactin recruitment that typically peaks just before completion of clathrin assembly (Figure 5A, Video S5). Cortactin recruitment is similarly sparse during the uptake of DI-T particles (Figure 5B, Video S6). In marked contrast, and as expected [15], large bursts of cortactin accompany the internalization of VSV (Figure 5C, Video S7). Quantitative analysis revealed that the peak fluorescence intensity of cortactin detected in the late phase of VSV uptake averaged 3-fold higher than the signal associated with pits containing DI-T or pits that did not capture a virus particle (Figure 5D, Table 1). These data suggest that while formation of short-branched actin filaments is required during the late stages of clathrin-mediated VSV entry, this need is obviated during clathrin-dependent DI-T entry.

Actin polymerization is not required for DI-T internalization

To directly test whether actin assembly is required for DI-T entry, we treated BSC1 cells with latrunculin B (latB), a chemical inhibitor of actin filament assembly [30], and tracked the endocytic fate of DI-T and VSV in the same cells. Treatment of cells with 6.3 μM latB did not change the efficiency of DI-T entry (Figure 6A–C, Video S8), but it reduced the internalization of VSV by 0.75% (Figure 6A–C). As expected [15], latB treatment did not affect the capture efficiency of either particle type by clathrin (Figure 6C). The lifetimes and AP-2 content of pits lacking particles or containing DI-T was similarly unaffected by latB (Figure 6D). We conclude that the shorter DI-T particles bypass the actin requirement displayed by the larger VSV for efficient clathrin-based uptake.
Discussion

The major conclusion of our study is that the physical properties of a virus particle dictate the need for engagement of the actin system during its clathrin-dependent uptake into cells. We formulate this conclusion based on tracking the clathrin-dependent internalization of VSV and its shorter DI-T particle into live cells. Internalization of VSV is accompanied by the recruitment of AP-2, which is involved in the formation of coated vesicles. The kinetics and AP-2 content of endocytic structures were analyzed to determine the relative lifetime and maximum fluorescence intensity of AP-2 during the uptake of coated pits lacking virus or structures that internalized DI-T or VSV. The values were expressed as percentages to facilitate comparison of viral and nonviral uptake events across multiple cells. Approximately 50 pits lacking virus were analyzed in each cell, and the mean of the measured values was calculated for each parameter. Each open circle represents a single uptake event, and horizontal red lines demark the mean of the compiled population. Numerical values and statistical analyses are provided in Table 1. doi:10.1371/journal.ppat.1001127.g003
The importance of particle length for the clathrin-dependent endocytosis of VSV is illustrated by the differences in the physical properties of the virus particle captured by a clathrin-coated pit. We found that DI-T particles do not acquire a full clathrin coat [15], which agreed with earlier electron micrographs depicting virus particles in tubular invaginations with clathrin at the cytosolic tip [17]. The morphology of those structures suggests that cells initially capture one tip of the virus particle, and clathrin assembly stalls when the constricting coat encounters the long axis of the virion (Figure 7). Here we found that DI-T particles do not alter the process of clathrin-coated vesicle formation (Figures 3, 4). This finding implies that clathrin can fully enclose cargos displaying VSV G provided that the particle shape does not physically prohibit clathrin assembly or closure of the plasma membrane (Figure 7). Consequently, the physical properties of the VSV particle captured by a clathrin-coated pit are what dictate the altered mode of actin-dependent uptake. We therefore propose the model (Figure 7) that it is the incomplete clathrin structure formed during VSV uptake that elicits the actin-based response to rescue the endocytic process and leads to the successful engulfment of the trapped virus particle.

We previously showed that endocytic structures containing VSV do not acquire a full clathrin coat [15], which agreed with earlier electron micrographs depicting virus particles in tubular invaginations with clathrin at the cytosolic tip [17]. The morphology of those structures suggests that cells initially capture one tip of the virus particle, and clathrin assembly stalls when the constricting coat encounters the long axis of the virion (Figure 7). Here we found that DI-T particles do not alter the process of clathrin-coated vesicle formation (Figures 3, 4). This finding implies that clathrin can fully enclose cargos displaying VSV G provided that the particle shape does not physically prohibit clathrin assembly or closure of the plasma membrane (Figure 7). Consequently, the physical properties of the VSV particle captured by a clathrin-coated pit are what dictate the altered mode of actin-dependent uptake. We therefore propose the model (Figure 7) that it is the incomplete clathrin structure formed during VSV uptake that elicits the actin-based response to rescue the endocytic process and leads to the successful engulfment of the trapped virus particle.
plaques and not conventional coated pits in several types of nonpolarized mammalian cells [26,41]. Thus, while actin dynamics play an evolutionarily conserved role in clathrin-dependent endocytosis, mammalian cells regulate the interplay between the clathrin and actin systems.

Our analyses of VSV and coated plaque internalization reveal two properties that correlate with their actin-dependent uptake mechanisms. First, plaques and VSV-containing structures remain on the plasma membrane for >2-fold longer than standard coated pits [15,26] (Figure 3C, E). The prolonged presence of a clathrin lattice might promote interactions between clathrin-associated proteins and regulators of actin polymerization. Second, clathrin plaques and VSV-containing structures physically differ from standard coated pits. Plaques fail to constrict their outer boundaries during the final phase of clathrin assembly [26], and pits containing VSV lack a complete clathrin coat [15,17]. Such unusual structural features might attract proteins that associate with exposed lipids, such as dynamin. Indeed, significantly more dynamin molecules accumulate during the final stages of VSV uptake [13]. This localized increase in dynamin may enhance the recruitment of dynamin-interacting proteins with the capacity to bind lipids and activate the Arp2/3 complex through N-WASP, including endophilin, syndapin, and SNX9 [42–46]. Although these proteins may link the clathrin and actin systems and facilitate localized membrane remodeling during endocytosis (reviewed in [47]), further studies are needed to determine whether they play a role in clathrin-dependent VSV endocytosis. Comparative studies of VSV and DI-T uptake may provide a useful tool to dissect the mechanisms that regulate actin assembly during clathrin-mediated endocytosis.

Implications for the internalization mechanisms of other viruses

The dimensions of the actin-dependent VSV particle and the actin-independent DI-T particle fall within the range of shapes present in many pleomorphic viruses. Our work suggests that this may lead to important distinctions in their mode of uptake. For example, some influenza A virus strains, such as X-31, produce spherical particles that measure 80–120 nm in diameter [18]. By contrast, the Udorn strain forms filamentous particles that measure 80–100 nm wide and up to 30 microns in length [48,49]. Although influenza A virus can enter cells by clathrin-dependent and -independent mechanisms [14], the impact of particle shape on the entry pathway remains unknown. It seems likely that remodeling of the cortical actin cytoskeleton will be important for uptake of filamentous influenza particles, and clathrin may facilitate local membrane deformation during the endocytic process. The Arenaviridae generate roughly spherical particles that range in diameter from 40–300 nm [50,51], and it is known that clathrin function is important for efficient infection of cells by some New World arenaviruses [52]. It will now be of interest to determine whether spherical particles of different diameter employ altered modes of clathrin-based endocytosis.

Pseudotyping is often used to study the entry pathway of highly pathogenic viruses, including the long filamentous filoviruses, Ebola and Marburg, as well as several arenaviruses. Such pseudotypes are frequently based on VSV or retroviral virions in which the endogenous entry proteins have been replaced with the surface glycoproteins of the pathogenic virus [53–56]. Although viral pseudotypes are useful for studying the entry process, VSV and the spherical virions of retroviruses (~100 nm in diameter) do not accurately recapitulate the sizes or shapes of the pleomorphic viruses. Our studies of VSV and DI-T clearly show that virion geometry can fundamentally alter aspects of the viral internalization process. Therefore, it is critically important to study viral endocytosis using pseudotyped or virus-like particles that closely approximate the physical properties of a virus in question.

Materials and Methods

Cells and viruses

African green monkey kidney BS-C-1 cells (herein BSC1, American Type Culture Collection (ATCC) CCL-26; Manassas, VA) and Vero cells (ATCC) were maintained at 37°C and 5% CO2 in Dulbecco’s Modified Eagle Medium (DMEM, Invitrogen Corporation; Carlsbad, CA) supplemented with 10% fetal bovine serum (Tissue Culture Biologicals; Tulare, CA). BSC1 cells stably expressing rat σ2 adaptin-eGFP (σ2-eGFP) [9] were maintained as above in the presence of 0.4 mg mL⁻¹ genetin (G418, Invitrogen).
Recombinant VSV (rVSV) [25] was amplified and purified as before [15]. Defective interfering T (DI-T) particles of VSV were recovered from a cDNA clone of the DI-T genome [24]. The DI-T particles were amplified by co-infecting baby hamster kidney cells (BHK-21, ATCC C-13) with rVSV (multiplicity of infection (MOI) 50). A subsequent passage was performed by inoculating cells with filtered, undiluted supernatant from the primary amplification and rVSV (MOI of 50). Viruses were concentrated by centrifugation at 44,000 g, and the virus pellet was resuspended in NTE (10 mM Tris pH 7.4, 100 mM NaCl, 1 mM EDTA). The two particle forms were separated on a 15–45% sucrose gradient prepared in NTE by centrifugation at 77,000 g for 5 h. The DI-T particles were extracted from the upper virus band, concentrated as before, and resuspended to 1 mg mL−1 of total protein in PBS.

Dye conjugation to virus particles

Purified DI-T and VSV particles were labeled with Alexa Fluor dye molecules (Molecular Probes, Invitrogen) as previously described [15] except that the final dye concentration in the labeling reaction was reduced to 25 μg ml−1. Plaque assays of virus preps before and after labeling showed that dye conjugation did not affect the infectivity of VSV particles or the capacity of DI-T virions to inhibit plaque formation by VSV. The surface density of G protein on VSV or DI-T particles was estimated by measuring the ratio of G protein to N protein in each particle population. To separate and visualize the viral proteins, purified virions were subjected to SDS-PAGE using 10% polyacrylamide gel and stained with Coomassie blue. The relative amounts of N or G protein were established using ImageJ (U. S. National Institutes of Health, Bethesda, Maryland; http://rsb.info.nih.gov/ij/).

Nucleic acid transfection

BSC1 cells were seeded into 6-well plates at ~60,000 cells per well 16–20 h prior to transfection. Plasmid DNA was introduced into the cells using FuGENE 6 (Roche Diagnostics; Indianapolis,

Figure 5. Clathrin structures containing VSV recruit more cortactin that pits that internalize DI-T. (A) Cortactin recruitment during coated pit formation. Left, snapshot showing the surface of a BSC1 cell transiently expressing cortactin-eGFP (green) and mCherry-LCa (red) at 18 h post-transfection. Time-lapse images were acquired at 3 s intervals, and frame 83 is shown. Middle, split channel kymographs of coated pit formation in the cell at left. White arrowheads highlight pits in which cortactin recruitment is clearly visible above the local background. Right, example plot of cortactin and clathrin fluorescence intensity over time during the formation of a single clathrin-coated pit in the cell shown at left (Video S5). (B and C) Examples of cortactin recruitment during DI-T (B) and VSV (C) uptake. BSC1 cells transiently expressing mCherry-LCa (red) and cortactin-eGFP (green) were inoculated with Alexa Fluor 647-labeled DI-T or VSV, and images were acquired as in A. Left, split-channel kymographs views of protein and virion (blue) fluorescence intensity over time (Videos S6, S7). Images in the right-hand panels show a snapshot of the maximal cortactin or clathrin fluorescence, and white arrowheads highlight the peak cortactin signal. Right, plots of the cortactin and clathrin fluorescence intensity over time for each internalization event. (D) Relative peak fluorescence intensity of cortactin in cells co-expressing mCherry-LCa and cortactin-eGFP. At 18 h post-transfection, samples were separately inoculated with DI-T or VSV particles, and images were acquired as in A. For each cell that was imaged, the maximum cortactin fluorescence associated with ~50 pits lacking virus particles (pits, black) and all pits that internalized a DI-T (left, blue, 3 cells) or VSV (right, red, 5 cells) particle was measured. The data are plotted as described in the legend of Figure 3C, and the number of events is shown above each plot. Numerical values and statistical analyses are provided in Table 1.

doi:10.1371/journal.ppat.1001127.g005
IN) according to the manufacturer’s instructions. Prior to transfection, media on the cells was replaced with 1 ml OPTIMEM (Invitrogen). After addition of the transfection mixture, cells were incubated at 37°C for 5 h, and the existing media was supplemented with 2 ml DMEM containing 10% FBS. To ensure optimal replacement of endogenous clathrin light chain molecules with rat eGFP-clathrin light chain A1 (eGFP-LCa) [9], cells were transfected with 0.75 μg of plasmid DNA encoding eGFP-LCa. The cells were cultured for ~36 h and seeded onto glass coverslips ~14 h prior to image acquisition. Co-expression of mCherry-LCa (constructed as described for tomato-LCa [8]) and mouse cortactin-eGFP [57, 58] was achieved by transfection of cells on glass coverslips with 1 μg of each plasmid, and the cells were imaged ~18 h later.

**Figure 6. Actin polymerization is not required for DI-T internalization.** (A) The endocytic fate of virus particles after inhibition of actin polymerization. BSC1 cells stably expressing eGFP (green) were treated with 6.3 μM latB for 12 min. and inoculated with DI-T and VSV particles in the continued presence of latB. Time-lapse images of a single cell were acquired at 4 s intervals for 692 s, and an 8.8 × 5.0 μm² area of the cell surface is shown. The upper panels show the complete internalization of a DI-T particle (blue, arrowheads), where +0 s indicates the first frame of the time-lapse series. The lower panels show the subsequent capture but failed internalization of 2 VSV (red, arrowheads) particles on the same area of cell membrane (time scale continued from above) (Video S8). (B) AP-2 fluorescence intensity for the events shown in A. Note that the adaptor fluorescence associated with the DI-T particle (blue) and a conventional coated pit (black) that formed within the same membrane area peak and disappear normally, while the adaptor signal associated with the upper-most VSV particle (red) does not, signifying failed internalization. (C) Effect of latB on the efficiency of virus capture and internalization. BSC1 cells stably expressing eGFP were treated and imaged as described in A. Left, the percentage of virus particles that were captured by a clathrin structure after attachment. Right, the percentage of captured virus particles that were successfully internalized within 300 s of capture (see D. for details). Cumulative data are from 5 cells. (D) Effect of latB on the lifetime and peak fluorescence intensity of AP-2 in clathrin structures. Data were acquired as described in A. and displayed as in the legend of Figure 3C. Left, relative lifetime of AP-2 in structures that lack (pits, black) or capture a virus particle. Inset shows a rescaled distribution of the pit and DI-T internalization events. Right, maximum fluorescence intensity of AP-2 in the events at left. Data are from 4 of the 5 cells analyzed in C, as thermal drift during imaging prevented accurate fluorescence intensity measurements in one cell. The number of events in each category is shown above the corresponding plots at right. DI-T (blue) data consists only of productive internalization events. VSV events are categorized as productive internalizations (VSV entry, red) or non-productive captures (trapped VSV, red). A non-productive capture is defined as a stable colocalization between a spot of AP-2 and a VSV particle that began at least 300 s before the last captured image and did not result in virus uptake before cessation of imaging. The 300 s cutoff was chosen because a majority (22/24) of productive internalizations occurred within 300 s of capture. Captures in which the final image frame was acquired before 300 s elapsed were excluded from the analysis, as the eventual endocytic fate of the particle cannot be predicted.

doi:10.1371/journal.ppat.1001127.g006
Virus Length Alters Clathrin-Dependent Endocytosis

Live cell imaging

Cells on 25 mm coverslips (No. 1.5, Electron Microscopy Sciences; Hatfield, PA) were placed into a perfusion chamber and overlaid with z-MEM lacking phenol red (Invitrogen) and supplemented with 20 mM HEPES pH 7.4 and 2% FBS. The chamber was placed in a heated sample holder (20/20 Technology Inc.; Wilmington, NC) mounted on the stage of a Mariana imaging system (Intelligent Imaging Innovations, Denver, CO) based on an Axiovert 200M inverted microscope (Carl Zeiss, Inc.; Thornwood, NY). The microscope stage and objective lenses were maintained at 37°C within an environmental chamber, and the air above the cells was supplied with 5% CO₂. The position of the sample holder with respect to the objective lens was manipulated using a PZ-2000 automated stage (Applied Scientific Instrumentation; Eugene, OR). Samples were illuminated using 40–50 mW solid state lasers (λ = 488, Coherent, Inc.; Santa Clara, CA, λ = 561, Cobolt AB; Solna, Sweden, λ = 640, 40 mW; Coherent) directed through an acousto-optic tunable filter (AOTF; Gooch and Housego L.L.C.; Melbourne, FL). Laser illumination was imparted on the sample through a CSU-X1 spinning disk confocal unit (Yokogawa Electric Corporation; Tokyo, Japan) and a 63X objective lens (Plan-Apochromat, NA 1.4, Carl Zeiss). Emission spectra were selected using single band width emission filters (LF405/488/561/635-A, Semrock; Rochester, NY), and after transmission through a spherical aberration correction device (Infinity Photo-Optical; Boulder, CO), the emission photons were collected using a Cascade II:512B back-illuminated EMCCD camera (Roper Scientific, Photometrics; Tuscon, AZ). Under this configuration, a single pixel corresponds to 0.07 μm². Image analysis was performed as previously described [15] with the following modifications. Slidebook 4.2.13 (III) was used to view, scale, and export images for publication. Movies were generated by exporting a series of continuous TIFF files from a Slidebook time-lapse acquisition and compiling the images into a single AVI file using ImageJ (NIH). SigmaPlot 8.0 (SYSTAT; Point Richmond, CA) was used to plot data. Microsoft Excel was used to determine whether data from 2 categories differ in a statistically significant manner using two-tailed Student’s t-tests. An automated image analysis application (IMAB) [8] developed within MATLAB (Mathworks; Natick, MA) was used to track the formation of clathrin-coated structures and the internalization of single virus particles. Images were processed for analysis as previously described, and established criteria were used to exclude incomplete endocytic events and events in which pixels assigned to one clathrin structure merged with or split from an adjacent structure [8]. For each cell of interest, the first ~50 pits lacking virus particles were analyzed in detail to measure the coat lifetime and protein composition (see below). All complete virus uptake events that occurred in these cells were also analyzed in a similar manner. With the exception of internalization events blocked by latB, incomplete internalization events or events truncated by the start/end of image acquisition were not analyzed. Aggregates of virus were identified as objects with fluorescence intensities greater than that of single particles and were excluded from all analyses. IMAB was used to measure the fluorescence intensity of coat components or virus particles in the following manner. A roughly spherical mask encompassing 69 pixels was centered on the peak fluorescence intensity of the object in all frames that the object was detectable above the local background fluorescence. The contribution of the local background fluorescence was estimated by measuring the average intensity of pixels within a ring of single pixel width that extended from the outer boundary of the object mask. The intensity of the pixels within the object mask was then summed, and the average intensity value of pixels in the local background was subtracted from each pixel within the object mask to estimate the fluorescence contributed by proteins (or dye molecules) within the object of interest.

Electron microscopy

Purified virus particles were deposited onto carbon-coated copper grids and stained with 2% phosphotungstic acid (PTA) in H₂O₂ (pH 7.5). To visualize DI-T particles in clathrin-coated pits, BSC1 cells were inoculated with unlabeled DI-T particles using a dose that yielded ~1000 attached particles per cell after 10 min. Samples were then processed for ultra-thin sectioning as previously described [15,59]. Electron micrographs of VSV particles in clathrin endocytic structures were obtained from cells infected with VSV for 6 h, where the entry of newly released particles could readily be visualized. Virus particles and ultra-thin sections of cells were viewed using a Tecnai G² Spirit BioTWIN transmission electron microscope (FEI, Hillsboro, OR).

Image analysis

Image analysis was performed as previously described [15] with the following modifications. Slidebook 4.2.13 (III) was used to view, scale, and export images for publication. Movies were generated by exporting a series of continuous TIFF files from a Slidebook time-lapse acquisition and compiling the images into a single AVI file using ImageJ (NIH). SigmaPlot 8.0 (SYSTAT; Point Richmond, CA) was used to plot data. Microsoft Excel was used to determine whether data from 2 categories differ in a statistically significant manner using two-tailed Student’s t-tests. An automated image analysis application (IMAB) [8] developed within MATLAB (Mathworks; Natick, MA) was used to track the formation of clathrin-coated structures and the internalization of single virus particles. Images were processed for analysis as previously described, and established criteria were used to exclude incomplete endocytic events and events in which pixels assigned to one clathrin structure merged with or split from an adjacent structure [8]. For each cell of interest, the first ~50 pits lacking virus particles were analyzed in detail to measure the coat lifetime and protein composition (see below). All complete virus uptake events that occurred in these cells were also analyzed in a similar manner. With the exception of internalization events blocked by latB, incomplete internalization events or events truncated by the start/end of image acquisition were not analyzed. Aggregates of virus were identified as objects with fluorescence intensities greater than that of single particles and were excluded from all analyses. IMAB was used to measure the fluorescence intensity of coat components or virus particles in the following manner. A roughly spherical mask encompassing 69 pixels was centered on the peak fluorescence intensity of the object in all frames that the object was detectable above the local background fluorescence. The contribution of the local background fluorescence was estimated by measuring the average intensity of pixels within a ring of single pixel width that extended from the outer boundary of the object mask. The intensity of the pixels within the object mask was then summed, and the average intensity value of pixels in the local background was subtracted from each pixel within the object mask to estimate the fluorescence contributed by proteins (or dye molecules) within the object of interest.

Electron microscopy

Purified virus particles were deposited onto carbon-coated copper grids and stained with 2% phosphotungstic acid (PTA) in H₂O₂ (pH 7.5). To visualize DI-T particles in clathrin-coated pits, BSC1 cells were inoculated with unlabeled DI-T particles using a dose that yielded ~1000 attached particles per cell after 10 min. Samples were then processed for ultra-thin sectioning as previously described [15,59]. Electron micrographs of VSV particles in clathrin endocytic structures were obtained from cells infected with VSV for 6 h, where the entry of newly released particles could readily be visualized. Virus particles and ultra-thin sections of cells were viewed using a Tecnai G² Spirit BioTWIN transmission electron microscope (FEI, Hillsboro, OR).
Virus Length Alters Clathrin-Dependent Endocytosis

Supporting Information

Video S1 Clathrin-dependent DI-T internalization. BSC1 cells stably expressing ε2-eGFP (green) were inoculated with DI-T particles, and time-lapse images were acquired at 4 s intervals. A 4.2×4.2 μm² area of the cell surface is shown as in Figure 2B. The video depicts a single DI-T particle (red) that attaches to the plasma membrane and diffuses slowly. A clathrin-coated pit captures the virus particle when a dim adaptor spot colocalizes with the virus signal. The adaptor fluorescence intensity increases as coated pit assembly proceeds, and particle internalization occurs shortly after the adaptor signal peaks. Disappearance of the adaptor signal signifies clathrin uncoating, and the virus-containing vesicle is then transported toward the cell interior. Found at: doi:10.1371/journal.ppat.1001127.s005 (3.01 MB AVI)

Video S2 Sequential internalization of single VSV and DI-T particles by the same cell. BSC1 cells stably expressing ε2-eGFP (green) were simultaneously inoculated with wild-type VSV and DI-T particles, and time-lapse images were acquired at 4 s intervals. A 3.5×3.5 μm² area of the cell surface is shown as in Figure 3A. At the outset of imaging (t = 0), one VSV (red) particle and one DI-T particle (blue) are visible on the cell surface. The VSV particle enters first, followed by the DI-T particle. Found at: doi:10.1371/journal.ppat.1001127.s002 (0.39 MB AVI)

Video S3 Uptake of a single DI-T particle by a cell expressing eGFP-LCa. BSC1 cells transiently expressing eGFP-LCa were inoculated with DI-T and full-length VSV particles, and images were captured as for Video S2. At the onset of imaging, a DI-T particle is present within a 3.5×3.5 μm² area of the cell surface. Shortly thereafter, the particle briefly colocalizes with a spot of eGFP-LCa but does not enter. As visualized in Figure 2B, a clathrin-coated pit subsequently initiates near the virion, and the particle disappears into the cell after the clathrin signal peaks. Found at: doi:10.1371/journal.ppat.1001127.s003 (0.77 MB AVI)

Video S4 Uptake of a single VSV particle by a cell expressing eGFP-LCa. The movie depicts a separate area (of equal size) of the plasma membrane from the same cell that internalized the particle shown in Video S3. A VSV particle (red) attaches to the cell surface, and a clathrin endocytic structure subsequently internalizes the particle (Figure 3D). Found at: doi:10.1371/journal.ppat.1001127.s004 (0.77 MB AVI)

Video S5 Cortactin recruitment during conventional clathrin-coated pit formation. Time-lapse images were acquired at 3 s intervals from a cell transiently co-expressing mCherry-LCa (red) and cortactin-eGFP (green) (Figure 5A). A 7.0×7.0 μm² area of the cell surface is shown. Note that the cortactin signal is nearly indistinguishable from the local background during most clathrin endocytic events. Found at: doi:10.1371/journal.ppat.1001127.s005 (3.01 MB AVI)

Video S6 Cortactin recruitment during the internalization of a single DI-T particle. BSC1 cells transiently co-expressing mCherry-LCa (red) and cortactin-eGFP (green) for 18 h were inoculated with DI-T particles (blue), and time-lapse images were acquired at 3 s intervals. The video shows the internalization of a single DI-T particle by a clathrin-coated vesicle. The internalization event is centered within a 3.5×3.5 μm² area of the cellular plasma membrane (Figure 5B). The left panel shows an overlay of the 3 channels, and the right panel shows only the cortactin channel displayed in monochrome. Note that the cortactin fluorescence intensity during DI-T entry is nearly indistinguishable from the local background. Found at: doi:10.1371/journal.ppat.1001127.s006 (1.42 MB AVI)

Video S7 Cortactin recruitment during the internalization of a wild-type VSV particle. BSC1 cells transiently co-expressing mCherry-LCa (red) and cortactin-eGFP (green) for 18 h were inoculated with VSV particles (blue), and time-lapse images were acquired at 3 s intervals. The video is displayed as described for Video S3 and shows the clathrin-dependent uptake of a single VSV particle in a 3.5×3.5 μm² area of the plasma membrane (Figure 5C). Note the visible burst of cortactin that occurs in the final stages of VSV internalization. Found at: doi:10.1371/journal.ppat.1001127.s007 (1.51 MB AVI)

Video S8 BSC1 cells stably expressing ε2-eGFP were treated with 6.3 μM latB for 12 min. Cells were inoculated with DI-T (blue) and VSV (red) particles, and images were acquired at 4 s intervals. The movie shows an 8.8×5.0 μm² area of the plasma membrane. The 2 DI-T particles (the lower particle is already in a coated pit at the onset of imaging) enter the presence of latB, while the 2 VSV particles are subsequently captured by coated pits but fail to enter the cell (Figure 6A, B). Found at: doi:10.1371/journal.ppat.1001127.s008 (4.65 MB AVI)

Acknowledgments

We express gratitude to Eric Marino for maintaining the imaging resource used in this study. We also gratefully acknowledge Maria Ericsson and Irene Kim for preparation of samples for electron microscopic analysis and members of the Kirchhausen and Whelan labs for thought-provoking discussions.

Author Contributions

Conceived and designed the experiments: DKC RHM SPJW TK. Performed the experiments: DKC. Analyzed the data: DKC SPJW TK. Contributed reagents/materials/analysis tools: DKC RHM SPJW TK. Wrote the paper: DKC SPJW TK.

References

1. Conner SD, Schmid SL (2003) Regulated portals of entry into the cell. Nature 422: 37–44.
2. Kirchhausen T (2000) Clathrin. Annu Rev Biochem 69: 699–727.
3. Kirchhausen T (2009) Imaging endocytic clathrin structures in living cells. Trends Cell Biol 19: 596–605.
4. Owen DJ, Evans PR (1998) A structural explanation for the recognition of tyrosine-based endocytotic signals. Science 229: 1327–1332.
5. Hossing S, Ricotta D, Krauss M, Spate K, Spolaore B, et al. (2005) Phosphatidylinositol-(4,5)-bisphosphate regulates sorting signal recognition by the clathrin-associated adaptor complex AP2. Mol Cell 18: 519–531.
6. Danke H, Baba T, Warnock DE, Schmid SL (1994) Induction of mutant dynamin specifically blocks endocytic coated vesicle formation. J Cell Biol 127: 913–934.
7. Lee DW, Zhao X, Zhang F, Eisenberg E, Greene LE (2005) Depletion of GAK/auxillin 2 inhibits receptor-mediated endocytosis and recruitment of both clathrin and clathrin adaptors. J Cell Sci 118: 4311–4321.
8. Massoul RH, Boll W, Grünig AM, Kirchhausen T (2006) A burst of auxillin recruitment determines the onset of clathrin-coated vesicle uncoating. Proc Natl Acad Sci U S A 103: 10265–10270.
9. Ehrlich M, Boll W, Van Oijen A, Hanifarian R, Chandran K, et al. (2004) Endocytosis by random initiation and stabilization of clathrin-coated pits. Cell 118: 591–605.
10. Loerke D, Mettlen M, Yarar D, Jaqaman K, Jaqaman H, et al. (2009) Cargo and dynamin regulate clathrin-coated pit maturation. PLoS Biol 7: e257.
11. Hanover JA, Beguinot L, Willingham MC, Pastan IH (1985) Transit of receptors for epidermal growth factor and transferrin through clathrin-coated pits. Proc Natl Acad Sci U S A 82: 3827–3831.
12. Anderson RG, Brown MS, Goldstein JL (1977) Role of the coated endocytic vesicle in the uptake of receptor-bound low density lipoprotein in human fibroblasts. Cell 10: 351–364.
13. van der Schaar HM, Rust MJ, Chen C, van der Eede-Metselaar H, Wilchut J, et al. (2008) Dissecting the cell entry pathway of dengue virus by single-particle tracking in living cells. PLoS Pathog 4: e100024.

14. Rust MJ, Lakadamyali M, Zhang F, Zhaou N (2004) Assembly of endocytic machinery around individual influenza viruses during viral entry. Nat Struct Mol Biol 11: 567–573.

15. Cureton DK, Massol RH, Saffarian S, Kirchhausen TL, Whelan SP (2009) Vesicular stomatitis virus enters cells through vesicles incompletely coated with clathrin that depend upon actin for internalization. PLoS Pathog 5: e1000394.

16. Veiga E, Guttman JA, Bonazzi M, Bocourt E, Todeo-Arau A, et al. (2007) Invasive and adverse bacterial pathogens co-Opt host clathrin for infection. Cell Host Microbe 2: 340–351.

17. Simpson RW, Hauser RE, Davila S (1969) Viropexis of vesicular stomatitis virus by I cells. Virology 37: 253–290.

18. Harris A, Cardone G, Winkler DC, Heymann JB, Brecher M, et al. (2006) Influenza virus pleiomorphism characterized by cryo-electron tomography. Proc Natl Acad Sci U S A 103: 19123–19127.

19. Meier E, Harmin G, Geine JD, Schubert M (1984) Sites of copy choice replication involved in generation of vesicular stomatitis virus defective-interfering particle RNAs. J Virol 51: 511–521.

20. Huang AS, Wagner RR (1966) Defective T particles of vesicular stomatitis virus. II. Biologic role in homologous interference. Virology 30: 173–181.

21. Peng G, Tsoo J, Schimm S, Green TJ, Luo M, et al. (2010) Cryo-EM model of the bullet-shaped vesicular stomatitis virus. Science 327: 689–693.

22. Huang AS, Greenwald JW, Wagner RR (1966) Defective T particles of vesicular stomatitis virus. I. Preparation, morphology, and some biologic properties. Virology 30: 161–172.

23. Wagner RR, Schimm TM, Snyder RM (1969) Structural proteins of vesicular stomatitis viruses. J Virol 3: 393–403.

24. Pattnaik AK, Ball LA, LeGronne AW, Wertz GW (1992) Infectious defective interfering particles of VSV from transcripts of a cDNA clone. Cell 69: 1011–1020.

25. Whelan SP, Ball LA, Barr J, Wertz GT (1995) Efficient recovery of infectious defective interfering particle RNAs. J Virol 51: 515–521.

26. Saffarian S, Cocucci E, Kirchhausen T (2009) Distinct dynamics of endocytic machinery around individual influenza viruses at the plasma membrane. J Cell Biol 180: 1219–1232.

27. Riezman H, et al. (2008) Distinct acto/myosin-I structures associate with clathrin that depend upon actin for internalization. PLoS Pathog 5: e1000394.

28. Meier E, Harmison GG, Keene JD, Schubert M (1984) Sites of copy choice replication involved in generation of vesicular stomatitis virus defective-interfering particle RNAs. J Virol 51: 511–521.

29. Merrifield CJ, Perrais D, Zenisek D (2005) Coupling between clathrin-coated-pit adaptors, and actin during endocytic internalization. Cell 115: 475–487.

30. Coue M, Brenner SL, Spector I, Korn ED (1987) Inhibition of actin polymerization by cortactin. J Biol Chem 262: 119–131.

31. Yang D, Rust MJ, Chen C, van der Ende-Metselaar H, Wilschut J, et al. (2008) Distinct acto/myosin-I structures associate with clathrin that depend upon actin for internalization. PLoS Pathog 5: e1000394.

32. Coyne CB, Kim KS, Bergelson JM (2007) Poliovirus entry into human brain microvascular cells. Cell Host Microbe 2: 340–351.

33. Coyne CB, Bergelson JM (2006) Virus-induced Abl and Fyn kinase signals permit coxsackievirus entry through epithelial tight junctions. Cell 124: 301–313.

34. Kaksonen M, Peng HB, Rauvala H (2000) Association of cortactin with dynamic endocytic profiles at the plasma membrane. J Cell Biol 151: 187–198.

35. Dawson JC, Legg JA, Machesky LM (2006) Bar domain proteins: a role in assembly and is required for efficient clathrin-mediated endocytosis. Mol Cell Biol 16: 2058–2067.

36. Yarar D, Waiseman-Sorer CM, Schmid SL (2007) SNX9 couples assembly to phosphoinositide signals and is required for membrane remodeling during endocytosis. Dev Cell 13: 43–56.

37. Kingtad NL, Nempto Y, De Camilli P (1997) The SH3p4/Sh3p8/Sh3p13 protein family: binding partners for synaptojanin and dynamin via a Grb2-like Src homology 3 domain. Proc Natl Acad Sci U S A 94: 8569–8574.

38. Okus M, Itoh T, Takenawa T (2003) Neural Wiskott-Aldrich syndrome protein is recruited to rafts and associates with endophilin A in response to epidermal growth factor. J Biol Chem 278: 6461–6469.

39. Dawson JC, Legg JA, Machensky LM (2006) Bar domain proteins: a role in tubulation, scission and actin assembly in clathrin-mediated endocytosis. Trends Cell Biol 16: 493–498.

40. Roberts PC, Companis RW (1998) Host cell dependence of viral morphology. Proc Natl Acad Sci U S A 95: 5746–5751.

41. Roberts PC, Lamb RA, Companis RW (1998) The M1 and M2 proteins of influenza A virus are important determinants in filamentous particle formation. Virology 240: 127–137.

42. Neumaen BW, Adair BD, Barra JW, Milligan RA, Buchmeier MJ, et al. (2005) Complemenntarity in the supramolecular design of arenaviruses and retroviruses revealed by electron cryomicroscopy and image analysis. J Virol 79: 3022–3030.

43. Murphy FA, Webb PA, Johnson KM, Whittfield SG, Chappell WA (1970) Arenaviruses in Vero cells: ultrastructural studies. J Virol 6: 507–518.

44. Cole JH, Warren JM, Kenton C, Robinson J, Parton RG, et al. (2005) PheRIR2 is the entry receptor for Nipah virus, an emergent deadly paramyxovirus. Nature 436: 401–405.

45. Clemente R, da la Torre JC (2009) Cell entry of Borna disease virus follows a lipoprotein receptor-mediated pathway. J Virol 83: 549–557.

46. Neuman BW, Adair BD, Burd JW, Milligan RA, Buchmeier MJ, et al. (2005) Complemenntarity in the supramolecular design of arenaviruses and retroviruses revealed by electron cryomicroscopy and image analysis. J Virol 79: 3022–3030.

47. Kedou F, Kurz A, Takanawa T (2003) Neural Wiskott-Aldrich syndrome protein is recruited to rafts and associates with endophilin A in response to epidermal growth factor. J Biol Chem 278: 6461–6469.

48. Dawson JC, Legg JA, Machensky LM (2006) Bar domain proteins: a role in tubulation, scission and actin assembly in clathrin-mediated endocytosis. Trends Cell Biol 16: 493–498.

49. Roberts PC, Companis RW (1998) Host cell dependence of viral morphology. Proc Natl Acad Sci U S A 95: 5746–5751.

50. Roberts PC, Lamb RA, Companis RW (1998) The M1 and M2 proteins of influenza A virus are important determinants in filamentous particle formation. Virology 240: 127–137.

51. Neumaen BW, Adair BD, Burd JW, Milligan RA, Buchmeier MJ, et al. (2005) Complemenntarity in the supramolecular design of arenaviruses and retroviruses revealed by electron cryomicroscopy and image analysis. J Virol 79: 3022–3030.

52. Murphy FA, Webb PA, Johnson KM, Whittfield SG, Chappell WA (1970) Arenaviruses in Vero cells: ultrastructural studies. J Virol 6: 507–518.

53. Koek J, Sanchez AB, Nguyen NT, de la Torre JC, Kunz S (2000) Different mechanisms of cell entry by human-pathogenic Old World and New World arenaviruses. J Virol 74: 7677–7687.

54. Chandraan K, Sullivan NJ, Fulher U, Whelan SP, Cunningham JM (2005) Endosomal proteolysis of the Ebolavirus glycoprotein is necessary for infection. Science 308: 1643–1645.

55. Nogte C, Avera EL, Aguilar HC, Bertoloti-Caulet A, Nazarian R, et al. (2005) E6PurinB2 is the entry receptor for Nipah virus, an emergent deadly paramyxovirus. Nature 436: 401–405.

56. Clemente R, da la Torre JC (2009) Cell entry of Borna disease virus follows a lipoprotein receptor-mediated pathway that requires Rab5 and microtubules. J Virol 83: 10406–10416.

57. Bhattacharya S, Warfield KL, Ruthig G, Bavari S, Aman MJ, et al. (2005) E6PurinB2 is the entry receptor for Nipah virus, an emergent deadly paramyxovirus. Nature 436: 401–405.

58. Moctezuma B, Avera EL, Aguilar HC, Bertoloti-Caulet A, Nazarian R, et al. (2005) E6PurinB2 is the entry receptor for Nipah virus, an emergent deadly paramyxovirus. Nature 436: 401–405.