Similar uptake but different trafficking and escape routes of reovirus virions and infectious subviral particles imaged in polarized Madin–Darby canine kidney cells

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ABSTRACT Polared epithelial cells that line the digestive, respiratory, and genitourinary tracts form a barrier that many viruses must breach to infect their hosts. Current understanding of cell entry by mammalian reovirus (MRV) virions and infectious subviral particles (ISVPs), generated from MRV virions by extracellular proteolysis in the digestive tract, are mostly derived from in vitro studies with nonpolarized cells. Recent live-cell imaging advances allow us for the first time to visualize events at the apical surface of polarized cells. In this study, we used spinning-disk confocal fluorescence microscopy with high temporal and spatial resolution to follow the uptake and trafficking dynamics of single MRV virions and ISVPs at the apical surface of live polarized Madin–Darby canine kidney cells. Both types of particles were internalized by clathrin-mediated endocytosis, but virions and ISVPs exhibited strikingly different trafficking after uptake. While virions reached early and late endosomes, ISVPs did not and instead escaped the endocytic pathway from an earlier location. This study highlights the broad advantages of using live-cell imaging combined with single-particle tracking for identifying key steps in cell entry by viruses.

INTRODUCTION During natural infections by many viruses, polarized epithelial cells that line the digestive, respiratory, and genitourinary tracts form a barrier that the viruses must breach to infect their hosts. In addition, viruses encounter similarly polarized cells in other settings, including endothelial cells in the circulatory system and ependymal cells in the CNS. Knowledge of the routes and mechanisms used by viruses to enter such polarized cells is of general interest, given the broad implications for understanding pathogenesis of viral diseases and for design of novel therapeutics and vaccines.

The nonfusogenic mammalian reoviruses (MRVs) constitute one of five approved species in genus Orthoreovirus, family Reoviridae. They are large (diameter \( \approx 85 \) nm, excluding their surface fibers) nonenveloped viruses, with a double-stranded RNA genome comprising 10 segments, and provide useful models for investigating the detailed routes and mechanisms of cell entry (Chandran and Nibert, 2003; Danthi et al., 2010) and host pathogenesis...
Experiments have demonstrated that JAM-A constitutes the primary receptor for productive infection of nonpolarized cells by ISVPs, as well as by virions (Barton et al., 2001b; Campbell et al., 2005), while β1 integrin appears to be important only for infection by virions (Maginnis et al., 2006). Together these observations suggest that ISVPs, as for virions described above, may exploit alternative pathways for productive entry, although those pathways have not been directly imaged.

Until recently, it was almost impossible to visualize entry of viruses from the apical surface of polarized cells, but a four-dimensional (4D) visualization approach with high temporal and spatial resolution is now available (Boulant et al., 2011; Kural et al., 2012), allowing us to follow the uptake and trafficking dynamics of single virus particles in live cells. In the current study, we took advantage of this new approach to compare the entry routes of MRV virions and ISVPs at the apical surface of polarized Madin–Darby canine kidney (MDCK) cells and found that both were internalized by clathrin-mediated endocytosis. We also found that, despite their similar uptake mechanism, virions and ISVPs exhibited strikingly different trafficking after internalization, with virions reaching early and late endosomes, while ISVPs escaped the endocytic pathway from a location reached before early endosomes.

RESULTS

Fluorescent MRV virions and ISVPs used for imaging experiments

Purified virions of MRV strain type 1 Lang (T1L) were covalently labeled with fluorescent dye Alexa Fluor (AF) 647, using a labeling density per particle that did not reduce infectivity as measured by plaque assay ($3.7 \pm 0.9 \times 10^{11}$ plaque-forming units [PFU]/ml and $4.0 \pm 1.1 \times 10^{11}$ PFU/ml for AF647-labeled and nonlabeled virions, respectively). SDS–PAGE revealed that all three major surface-exposed proteins in the virions—outer-clamp protein μ3, T = 13 outer-shell protein μ1/μ1C, and core-turret protein λ2—were covalently labeled with the dye (Figure 1A, left). Fluorescent ISVPs were generated by standard chymotrypsin treatment of the AF647-labeled virions (Figure 1A), and plaque assays confirmed no reduction in infectivity ($6.2 \pm 1.3 \times 10^{11}$ and $6.5 \pm 1.5 \times 10^{11}$ PFU/ml for AF647-labeled and unlabeled ISVPs, respectively). SDS–PAGE confirmed the expected removal of dye-labeled μ3 from the ISVPs (Dryden et al., 1993), but the retention of labeled proteins μ1/μ1C (in the form of cleavage products μ1/μ1C and ϕ, as expected [Nibert and Fields, 1992]) and λ2 (Figure 1A, left). Imaging of the fluorescent particles by spinning-disk confocal microscopy revealed that they each belong to a single class of small, diffraction-limited objects (Figure 1A, right), consistent with presence of individual particles and absence of aggregates. Owing to the proteolytic removal of μ3, the average fluorescence intensity of AF647-labeled ISVPs was lower than that of AF647-labeled virions (Figure 1B). Comparable results were obtained for virions and ISVPs labeled with fluorescent dye AF568.

Virion and ISVP uptake from the apical surface of polarized MDCK cells

To examine the uptake and entry routes of MRV virions and ISVPs in polarized cells, we used MDCK cells, which are known to support MRV infection when nonpolarized (Kothdaraman et al., 1998). Polarization of these cells was confirmed by establishment of tight junctions between cells as determined by 1) immunostaining of ZO-1 in a typical “belt” pattern surrounding the cells (Figure 2, A and C); 2) accumulation of actin, reflecting development of apical microvilli (Figure 2A); and 3) increase of transepithelial resistance from ~300 to ~600 ohm/cm² after growth on Transwell supports for
Because clathrin-mediated endocytosis is dynamin dependent, we directly visualized the clathrin-mediated uptake of MRV particles and ISVPs at the apical surface of polarized MDCK cells. The imaging experiments allowed efficient visualization with high temporal resolution (1–2 s) of clathrin-mediated uptake of fluorescently labeled virions and ISVPs (bottom) imaged on coverslips were measured by use of Slidebook5 to identify particles by intensity thresholding and to record their intensities. Fluorescence intensity per pixel (in arbitrary units) is shown in bins on the x-axis, and percentage of total virions or ISVPs falling within each bin is shown on the y-axis. A Gaussian fit of the data for each type of particle is also shown. The legend shows the mean, SD, and total number of particles analyzed for each type.

To determine which uptake pathways are specifically important for MRV infection of polarized MDCK cells, we took advantage of the neutralizing activity of monoclonal antibody 5C6 (Virgin et al., 1991), directed against the MRV receptor-binding outer-fiber protein σ1, to assay for infection (protocol diagrammed in Figure 4A; see also Materials and Methods). We thereby found that dynasore or jasp reduced apical infection by virions by ∼70%, whereas amiloride alone or in combination with either dynasore or jasp had little or no effect (Figure 4B). The inhibitors of vacuolar acidification NH4Cl and bafilomycin A1, included as positive controls, each blocked infection by virions by >90% (Sturzenbecker et al., 1987; Martínez et al., 1996; Figure 4B). Effects on ISVP infection by the uptake inhibitors were only slightly less than those on virions (Figure 4B). These findings suggested to us that clathrin-mediated endocytosis is a major productive route of uptake for both virions and ISVPs at the apical surface of polarized MDCK cells and parallel the preceding findings obtained by imaging of internalized particles (see Figure 3B).

Live-cell imaging of clathrin-mediated endocytosis of virions and ISVPs at the apical surface of polarized MDCK cells

We directly visualized the clathrin-mediated uptake of MRV particles from the apical surface of polarized MDCK cells using our newly implemented approach for live-cell 4D imaging by spinning-disc confocal microscopy (Boullant et al., 2011). This imaging approach allows efficient visualization with high temporal resolution (1–2 s) of diffraction-limited fluorescently tagged objects (virions, ISVPs, and clathrin/AP2-coated pits and vesicles) located on or near the dome-shaped apical surface of polarized cells. The imaging experiments...
were carried out using a recently developed MDCK line expressing the α2 subunit of adaptor protein 2 fused to green fluorescent protein (AP2-GFP) (Boulant et al., 2011), which allows tracking of all AP2-containing endocytic carriers. Some experiments included transient expression of clathrin light chain A fused to fluorescent tomato protein (CLCa-TOM), to confirm that AP2 and clathrin always colocalized (Ehrlich et al., 2004; Cureton et al., 2009, 2010, 2012).

By using this approach, we obtained the first, real-time visualizations of capture and uptake of single MRV virions and ISVPs by the formation of clathrin/AP2-coated structures at the apical surface of polarized MDCK cells. Both virions and ISVPs colocalized extensively with clathrin/AP2 structures; this was followed by efficient internalization. We observed multiple such uptake events for single virions or ISVPs in all of the 4D series, as illustrated by the examples for a single virion tracked over time in Figure 5A and for virions and ISVPs in Supplemental Movies S1–S4. Typically, 30–40% of virions or ISVPs attached to the apical surface were internalized by clathrin-mediated endocytosis during the 3- to 5-min interval of the time series (Figure 5B and Movies S1–S4). Uptake by this route was divided into five phases: 1) apical cell surface attachment of an MRV virion or ISVP, and slow lateral diffusion; 2) cessation of lateral diffusion that always coincided with the site at which a clathrin/AP2 structure was forming; 3) growth of the coated pit (increase in AP2-GFP and CLCa-TOM fluorescence intensity); 4) separation of the virion- or ISVP-containing coated vesicular carrier from the apical membrane, followed by rapid release of the clathrin/AP2 coat (abrupt disappearance of AP2-GFP and CLCa-TOM fluorescence); and 5) rapid and directed intracellular movement of the virion- or ISVP-containing vesicular carrier (Figure 5A and Movies S1–S4). Also noteworthy was the distribution of clathrin-mediated uptake events across the entire apical surface of the polarized MDCK cells, and not concentrated near the edges where tight junctions are found (Movies S3 and S4), consistent with the apical distribution of some MRV receptor JAM-A (Figure 2C). In addition, the observed efficiency of virion and ISVP uptake by clathrin-mediated endocytosis in these cells was not specifically due to their polarization, as nonpolarized MDCK cells also exhibited efficient uptake of both particle types through this pathway (Figure 5B).

We compared the formation characteristics of clathrin/AP2 structures alone or in association with MRV particles by determining the lifetime of coated pits. Pit lifetime is the elapsed time between the
coated vesicles before completion of uncoating (loss of AP2-GFP signal) was 472 ± 83 nm, similar to the value obtained for coated vesicles not containing virions (451 ± 124 nm) (Figure 7B). The value for ISVP-containing coated vesicles was also similar (388 ± 83 nm). Thus the presence of an enclosed MRV particle, either virion or ISVP, did not significantly affect the distance traveled from the plasma membrane before release of the clathrin/AP2 coat.

Trafficking of virions and ISVPs after uptake from the apical surface of polarized MDCK cells

We next used our live-cell 4D imaging approach to examine the trafficking of virions and ISVPs following their uptake by clathrin-mediated endocytosis. To mark endosomal compartments, we used polarized MDCK cells transiently expressing low levels of Rab5-GFP (marking early endosomes) or Rab7-GFP (marking late endosomes). Additionally, in order to track the two types of MRV particles simultaneously, we coinfected the cells with AF568-labeled virions and AF647-labeled ISVPs (red and blue, respectively, in Figure 8A). This tracking strategy corrected for potential cell-to-cell variability in intracellular trafficking kinetics. Virions were readily found associated with both early and late endosomes (Figure 8A), in numbers that increased with time and reached a plateau at ∼20–25 min postinfection (Figure 8B). In striking contrast, substantially fewer ISVPs were found in either early or late endosomes (Figure 8A), and their numbers in these compartments did not increase with time (Figure 8A and B). Furthermore, we observed numerous, intracellular ISVPs that failed to colocalize with either Rab5 or Rab7 (Figure 8A).

Failure to detect substantial numbers of ISVPs in early or late endosomes of polarized MDCK cells could be the result of ISVPs

appearance and sudden disappearance of the coat constituents (Figure 6A). The lifetimes of coated pits containing or lacking MRV particles were measured in the same polarized MDCK cells (Figure 6A). Coated pits containing virions had significantly longer lifetimes (78 ± 21 s) than those lacking virions (49 ± 8 s) (Figure 6B). Similarly, coated pits containing ISVPs had significantly longer lifetimes (61 ± 13 s) than those without, but significantly shorter lifetimes than those with virions (Figure 6B). These increased coated-pit lifetimes with virions or ISVPs are not easily explained by the time it would take to assemble a larger coat, however, because pits containing or lacking MRV particles recruited similar amounts of AP2, an indicator of coat size (Ehrlich et al., 2004; Figure 6C). Thus, though not increasing coat size in polarized MDCK cells, both particle types (virions to a greater extent than ISVPs) slowed the progression of pit formation.

Our live-cell imaging approach further allowed us to track MRV-containing clathrin-coated vesicles immediately after they budded from the plasma membrane. Soon after recruitment of AP2-GFP reached its peak (Figure 7A, green circles), we observed a small displacement of the MRV particle away from the plasma membrane into the cell interior (Figure 7A, red circles). This displacement corresponds to inward movement of the virion-containing coated vesicle just after budding, but before complete release of the clathrin/AP2 coat, as depicted in the schematic (Figure 7A, top panel). At about the same time that uncoating was completed, we observed a sudden but brief, high-velocity displacement of the MRV particle (Figure 7A, blue line), presumably still within its vesicular carrier. This movement is similar to one previously described during the clathrin-dependent uptake of vesicular stomatitis virus (Cureton et al., 2010). The average Z-displacement from the apical membrane of virion-containing coated vesicles before completion of uncoating (loss of AP2-GFP signal) was 472 ± 83 nm, similar to the value obtained for coated vesicles not containing virions (451 ± 124 nm) (Figure 7B). The value for ISVP-containing coated vesicles was also similar (388 ± 83 nm). Thus the presence of an enclosed MRV particle, either virion or ISVP, did not significantly affect the distance traveled from the plasma membrane before release of the clathrin/AP2 coat.

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μ1(N42A)-ISVPs, the latter of which are unable to escape to the MDCK cells with AF568-labeled ISVPs and AF647-labeled between these two main possibilities, we coinfected polarized they were unable to reach those endosomes. To try to distinguish 1) having escaped the endocytic pathway from a location reached before early endosomes or 2) having been redirected such that 1) having escaped the endocytic pathway from a location reached before early endosomes. Important implications of these conclusions are discussed below.

**DISCUSSION**

The above results lead us to the following main conclusions. 1) MRV virions and ISVPs underwent efficient uptake at the apical surface of polarized MDCK cells by clathrin-mediated endocytosis, with little or no evidence for uptake by macropinocytosis. 2) Clathrin-mediated endocytosis at the apical surface of these cells led efficiently in turn to productive infection by both virions and ISVPs. 3) Clathrin-mediated endocytosis of virions and ISVPs was widely distributed at the apical surface of these cells and not concentrated at regions near tight junctions, consistent with our evidence that the MRV receptor JAM-A was found throughout the apical surface in these cells. Additional conclusions pertain to real-time studies by live-cell 4D imaging of individual MRV particles undergoing clathrin-mediated endocytosis at the apical surface of polarized MDCK cells. 4) Virions and ISVPs both lengthened the lifetime of coated-pit formation, and virions did so to a greater extent than ISVPs. 5) Virions and ISVPs did not affect coat size or intracellular displacement before uncoating. Finally, 5) following internalization, virions underwent efficient trafficking to early and late endosomes, whereas ISVPs did not and instead escaped the endocytic pathway from a location reached before early endosomes. Important implications of these conclusions are discussed below.

**Details of clathrin-mediated endocytosis of MRV revealed by live-cell 4D imaging**

Incorporation of MRV virions into clathrin-coated pits at the apical surface of polarized MDCK cells significantly prolonged the pits’ mean lifetime (78 vs. 49 s for pits lacking an MRV virion in the same cells) but did not alter the number of clathrin and adaptor molecules that ultimately assembled into the endocytic coat. Virion incorporation therefore altered the rate of one or more steps in the pit-maturation process, but not the eventual size of the endocytic vesicle. Analyses of individual traces revealed that coated pits containing or lacking an MRV virion recruited clathrin and adaptor at similar rates during the earlier stages of pit formation, but virion-containing pits took longer to complete the later stages of pit maturation. These findings suggest that assembly of the coat around the first one-half to two-thirds of the virion does not constitute a limitation, but that completion of coat assembly and pit constriction into a vesicle likely take longer due to steric hindrance created by the virion. Some perturbation to coated-pit formation is in fact expected, because the diameter of the MRV virion (85 nm, excluding the σ1 fibers, which would nearly double the particle diameter if fully extended [Furlong et al., 1988; Dryden et al., 1993]) approaches the upper limit for the internal diameter of a canonical coated vesicle (Fotin et al., 2006; Kirchhausen, 2009). Because the σ1 fiber can bend...
formation may therefore have been slowed less by ISVPs (mean pit lifetime: 61 s) than by virions, because the effective diameter of the ISVP is smaller, due to the absence of outer-clamp protein σ3 (Dryden et al., 1993).

at several points along its length, including near its base (Furlong et al., 1988; Fraser et al., 1990), we suspect it may contribute little to the effective diameter of virions and ISVPs as far as coated-pit formation is concerned. Completion of coated-pit formation may therefore have been slowed less by ISVPs (mean pit lifetime: 61 s) than by virions, because the effective diameter of the ISVP is smaller, due to the absence of outer-clamp protein σ3 (Dryden et al., 1993).
One question that naturally arises from the current results is whether different, productive uptake routes may be used by MRV virions to infect nonpolarized versus polarized cells, and to infect polarized cells from their apical versus basolateral surfaces. This study was not designed to address those questions, but during the course of our experiments, we found that MDCK cells before polarization appeared to be as efficient at internalizing MRV virions by clathrin-mediated endocytosis as after polarization. Thus the large difference in high efficiency of clathrin-mediated endocytosis identified here with polarized MDCK cells compared with the lower efficiency described earlier in nonpolarized BSC-1 cells (Ehrlich et al., 2004) is likely to reflect a difference between these two cell types unrelated to polarization, perhaps due to the lower level of JAM-A expression in BSC-1 cells. We suspect that clathrin-mediated endocytosis is the main productive uptake route for MRV virions in most types of cells, unless a deficiency in JAM-A, and/or perhaps β1 integrins (Maginnis et al., 2006, 2008), allows alternative uptake routes, such as macropinocytosis, to gain greater prominence.

Uptake, trafficking, and escape of ISVPs

We previously reported that the lifetime of coated pits engaged in the capture of MRV virions averaged 400 s in nonpolarized BSC-1 cells (Ehrlich et al., 2004), much longer than the 78-s mean lifetime in polarized MDCK cells reported here. Our previous study indicated, however, that only a small fraction of MRV virions was internalized by clathrin-mediated endocytosis. In contrast, we now find evidence for very efficient clathrin-mediated capture of virions at the apical surface of polarized MDCK cells. We believe that this difference may have been due in part to a relative deficiency of one or more MRV cell surface receptors in the BSC-1 cells. Indeed, BSC-1 cells are greatly deficient in JAM-A (the MRV protein receptor [Barton et al., 2001a; Campbell et al., 2005]) relative to MDCK cells, and transient expression of JAM-A in BSC-1 cells substantially increases the capture of virions by clathrin/AP2 carriers (S. Boulant, unpublished data).

Alternative uptake pathways for MRV virions

Several studies preceding this one have provided evidence that MRV virions can enter nonpolarized cells by clathrin-mediated endocytosis (Sturzenbecker et al., 1987; Rubin et al., 1992; Ehrlich et al., 2004; Maginnis et al., 2008; Schulz et al., 2012); however, in some of those same studies, inhibitors of clathrin-mediated endocytosis did not completely block virion uptake and infection, suggesting that virions can productively enter cells via one or more alternative routes as well. In our current study, even though we obtained evidence that clathrin-mediated endocytosis is a major uptake route for productive infection by MRV virions at the apical surface of polarized MDCK cells, 20–30% of virion uptake and infection occurred, despite inhibition of the clathrin-dependent route. Thus one or more clathrin-independent routes for productive infection of MRV virions must have also been available for use, though those routes remain to be elucidated in polarized MDCK cells.

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more alternative uptake routes in these cells, as also recently suggested for other, nonpolarized cells (Schulz et al., 2012).

Another key conclusion from our study is that ISVPs escaped the endocytic pathway of polarized MDCK cells after clathrin/AP2-vesicle budding and uncoating but before reaching early endosomes. These findings likely explain why previous studies in nonpolarized cells have detected few infecting ISVPs in endosomes and have furthermore detected ISVPs free in the cytosol near the plasma membrane (Borsa et al., 1979). They are also consistent with our “sequestration” model, whereby the ISVP undergoes structural changes associated with membrane penetration (Chandran et al., 2002, 2003; Odegard et al., 2004) only after its surrounding volume is restricted inside an endocytic vesicle or similar structure (Agosto et al., 2008). Whether infecting ISVPs escaped directly from uncoated vesicles in this study remains to be demonstrated. Another question of interest concerns ISVPs generated from virions only after proteolysis inside late endosomes or lysosomes (Silverstein and Dales, 1968; Borsa et al., 1979; Sturzenbecker et al., 1987; Baer and Dermody, 1997; Mainou and Dermody, 2012; Kothandaraman et al., 1998; Ebert et al., 2002; Chandran et al., 2003). Can ISVPs escape

**FIGURE 8:** Endosomal association of MRV virion and ISVP particles after cell entry. Cells transiently expressing Rab5-GFP or Rab7-GFP for 16 h were inoculated with fluorescent virus particles, and Z-series images were acquired from individual cells at different times postinoculation by 4D spinning-disk confocal microscopy, as described in Materials and Methods. (A) Maximum-intensity projection images of MDCK cells expressing early endosome marker Rab5-GFP (left panels) or late endosome marker Rab7-GFP (right panels) at 30 min postinoculation with both AF563-labeled virions (red) and AF647-labeled ISVPs (blue). White boxes correspond to the magnified insets. (B) Quantification of the fraction of virions and ISVPs (same particles as shown in A) colocalizing with Rab5-GFP (left panel) and Rab7-GFP (right panel). The number of virions or ISVPs in these endosomal compartments at different times postinoculation was normalized to the total number of virions or ISVPs in those cells at each respective time. The histogram represents ± SD for 10 cells analyzed. (C) Similar experiment as described in (B), but this time inoculating the cells with AF647-labeled ISVPs (blue) and AF563-labeled μ1(N42A)-ISVPs (magenta). The histogram represents the mean ± SD for 10 cells analyzed.
directly from either of those compartments, or must they first be trafficked elsewhere?

**MATERIAL AND METHODS**

**Virus particles**

Nonlabeled virions of MRV strain T1L, derived from stocks originally obtained from Bernard N. Fields (Harvard Medical School), were grown and purified by standard protocols (Furlong et al., 1988). Nonlabeled ISVPs were generated from nonlabeled virions (1 × 10^13 particles/ml) by treatment with chymotrypsin (200 μg/ml) in virion buffer (150 mM NaCl, 10 mM MgCl₂, 10 mM Tris, pH 7.5) at 32°C for 10 min, after which the protease was inactivated with 1 mM phenylmethylsulfonyl fluoride. Before being fluorescently labeled in vitro, virions were dialyzed overnight against phosphate-buffered saline (PBS). Amine-reactive dyes AF647 or AF568 (Molecular Probes) were solubilized in dimethyl sulfoxide at 10 mg/ml and diluted to a final concentration of 62.5 μg/ml in PBS, along with purified virions at a final concentration of 1 mg/ml. After incubation at room temperature for 90 min, labeled virions were separated from free dye using a PD-10 desalting column (GE Healthcare, Waukesha, WI) and dialyzed against virion buffer. Fluorescent ISVPs were obtained by treatment of labeled virions, as described above for nonlabeled particles. Recoated μ1(N42A)-virions (Odegard et al., 2004) were generated from T1L cores and proteins and purified using standard protocols (Chandran et al., 1999). They were then fluorescently labeled with AF647 or AF568, as described above for native virions. Fluorescent μ1(N42A)-ISVPs were prepared from μ1(N42A)-virions, as described above. All particles were titrated by plaque assay, as described previously (Middleton et al., 2002).

**Antibodies and inhibitors**

The following primary antibodies were used: rabbit polyclonal antibodies against virions of MRV strain T1L (Tyler et al., 1993), used at 1/1000 dilution for immunofluorescence and immunoblotting; rabbit polyclonal antibodies against MRV nonstructural protein μNS (Broering et al., 2000), used at 1/1000 dilution for immunofluorescence and immunoblotting; mouse monoclonal antibodies against ZO-1 (Invitrogen, Carlsbad, CA) and JAM-A (J10.4; Santa Cruz Biotechnology, Santa Cruz, CA) used at 1/1000 and 1/200 dilutions for immunofluorescence; and mouse monoclonal antibody 5C6 against MRV attachment protein α1 (Virgin et al., 1991), used at 1–2 μg/ml in PBS for neutralization. Secondary antibodies were conjugated with AF488, AF-568 or AF-647 and directed against the corresponding animal source of primary antibodies (Molecular Probes). The following pharmacological inhibitors were used: 1 μM jasplakinolide (Alexis Biochemical, Farmingdale, NY), 80 μM dynasore (Maca et al., 2006), 20 mM 5-((N-ethyl-N-isopropyl)amiloride (Sigma-Aldrich, St. Louis, MO), 20 mM NH₄Cl (Sigma-Aldrich), and 1 μM bafilomycin A (Sigma-Aldrich).

**Cell culture and transfection**

MDCK-2 cells stably expressing AP2-EGFP (Boulant et al., 2011) were grown in DMEM containing 10% fetal calf serum (FCS), penicillin, and streptomycin (Invitrogen). Cells were seeded on coverslips at 5 d before infection to allow formation of polarized monolayers. Transfected plasmids were ones encoding 1) CLCa-TOM (Cureton et al., 2009, 2010), 2) EGFP fused to early endosome marker Rab5 (Rab5-GFP; Addgene, Cambridge, MA), and 3) GFP fused to late endosome marker Rab7 (Rab7-GFP; Addgene). Transfection was carried out using TransIT-LT1 (Mirus Bio, Madison, WI), and cells were analyzed 12–16 h after transfection unless otherwise indicated.

**Spinning-disk confocal microscopy**

Imaging of MRV entry was performed as described for vesicular stomatitis virus and canine parvovirus (Cureton et al., 2009, 2010, 2012), and the microscopy setup was identical to that described for another recent study of polarized MDCK cells (Boulant et al., 2011). Briefly, imaging was performed in a temperature-controlled, 5% CO₂, humidified chamber (20/20 Technologies, Wilmington, NC) mounted on a Marianas system platform (Intelligent Imaging Innovations, Denver, CO) based on a Zeiss AxioVert 200M inverted microscope equipped with a CSU-X1 spinning-disk confocal unit (Yokogawa Electric, Tokyo, Japan) and a spherical-aberration correction device (Infinity Photo-Optical, Boulder, CO). The correction device was needed because of the severe spherical aberration on the apical surface of polarized MDCK cells located 8–10 μm away from the coverslip.

**Immunostaining**

Cells grown on coverslips were fixed at room temperature for 15 min with 4% wt/vol paraformaldehyde dissolved in PBS and then processed by sequential incubation with the appropriate primary and secondary antibodies dissolved in PBS containing 0.05% wt/vol Triton X-100 and 2% FCS. Sequential optical planes spaced 200 nm in the z-axis were acquired using spinning-disk confocal microscopy, and representative maximum-intensity projection images are shown.

**Internalization assay**

Polarized monolayers of MDCK cells on coverslips were pretreated with respective pharmacological inhibitors for 15 min at 37°C in serum-free DMEM. Fluorescent virions or ISVPs (MOI = 100) were then added on top of the cells in the presence of the same respective inhibitors and incubated at 37°C for indicated times up to 30 min. Cells were then washed with ice-cold PBS and fixed with 0.5–1% paraformaldehyde for 5 min on ice. Cells were washed three times with ice-cold PBS and immunostained, without being permeabilized, using anti-virion polyclonal antibodies. The fraction of internalized particles determined by spinning-disk confocal microscopy was obtained using the intensity-thresholding routine in Slidebook 5 (Intelligent Imaging Innovations) to determine the number of fluorescent virus particles colocalizing (“outside”) or not (“inside”) with differently fluorescent secondary antibodies specific for the anti-virion primary antibodies. The virus particles and secondary antibodies were respectively labeled with AF568 and AF647, or with AF647 and AF568. AF647-labeled human transferrin and fluorescein-labeled dextran (Molecular Probes) were used at 200 μg/ml, respectively.

**Infection assay**

Polarized monolayers of MDCK cells were washed with PBS, and serum-free DMEM with respective pharmacological inhibitors was added for 30 min at 37°C. Virions or ISVPs were then added on top of the cells (MOI = 3) and incubated for 2 h at 37°C to allow attachment and uptake in the presence of the same respective inhibitors. Cells were washed three times with ice-cold PBS, and anti-α1 monoclonal antibody 5C6 was then added for 2 h at 4°C to neutralize any MRV particles that had not yet been internalized. Cells were washed three times with ice-cold PBS, and fresh complete medium was added for 8–12h at 37°C to allow MRV gene expression and replication. Cells were then fixed with 4% paraformaldehyde and immunostained using anti-μNS polyclonal antibodies and fluorescent secondary antibodies. The fraction of infected cells (i.e., cells expressing μNS) was determined by fluorescence microscopy. The neutralizing
activity of 5C6 was validated beforehand in comparable experiments in nonpolarized cells.

**Live-cell imaging of virus uptake**

MRV particles (MOI = 100) were added atop polarized monolayers of MDCK cells at 2–3 min before imaging. Virus uptake was then monitored by 4D spinning-disk confocal microscopy. Three-dimensional (3D) movies comprising Z-stacks of three to five consecutive optical planes spaced 0.5 μm apart were acquired at a frequency of 0.3 Hz using a 100-ms exposure time for a total of 50–100 frames. Two-dimensional movies were generated from these 3D movies by merging the maximum-intensity Z-projection for a particular virus particle from each frame. AP2-coated fluorescent structures were identified and tracked by using a 3D-tracking MATLAB routine (Kural et al., 2012).

**Live-cell imaging of virus trafficking**

At 16 h before infection, MDCK cells were transfected according to the manufacturer’s instructions with plasmids encoding Rab5-GFP or Rab7-GFP. Fluorescent virions, ISVPs, or μ1(N42A)-ISVPs (MOI = 100) were then added on top of the transfected cells, and at different times postinfection, a Z-series encompassing the full volume of at least 10 cells was acquired by 4D spinning-disk confocal microscopy. Virus particles were identified using an intensity-thresholding routine in Slidebook 5 (Intelligent Imaging Innovations), and their colocalization with the endosomal markers was ascertained. The fraction of virus particles colocalizing with either marker was normalized to the total number of particles detected in the respective compartments at 30 min postinfection. A minimum of 75 particles was counted for each histogram value shown in Figure 8.

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