Inhibition of PIKfyve kinase prevents infection by Zaire ebolavirus and SARS-CoV-2

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- **PIKfyve, pandemic**

41 Virus entry is a multistep process. It initiates when the virus attaches to the host 42 cell and ends when the viral contents reach the cytosol. Genetically unrelated viruses can subvert analogous subcellular mechanisms and use similar 43 trafficking pathways for successful entry. Antiviral strategies targeting early 44 steps of infection are therefore appealing, particularly when the probability for 45 successful interference through a common step is highest. We describe here 46 potent inhibitory effects on content release and infection by chimeric VSV 47 containing the envelope proteins of Zaire ebolavirus (VSV-ZEBOV) or SARS-CoV-48 49 2 (VSV-SARS-CoV-2) elicited by Apilimod and Vacuolin-1, small molecule inhibitors of the main endosomal Phosphatidvlinositol-3-50 Phosphate/Phosphatidylinositol 5-Kinase, PIKfyve. We also describe potent 51 inhibition of SARS-CoV-2 strain 2019-nCoV/USA-WA1/2020 by Apilimod. These 52 results define new tools for studying the intracellular trafficking of pathogens 53 54 elicited by inhibition of PIKfyve kinase and suggest the potential for targeting this kinase in developing small-molecule antivirals against SARS-CoV-2. 55

56 **INTRODUCTION**

Membrane-enveloped viruses deliver their contents to cells via envelope protein-57 catalyzed membrane fusion. Binding of virus to specific host cell receptor(s) triggers 58 59 membrane fusion, which can occur directly at the plasma membrane or following 60 endocytic uptake. Viruses that require endocytic uptake can use different initial 61 trafficking routes to reach the site of membrane fusion. In endosomes, acidic pH serves 62 to triggers conformational rearrangements in the viral envelope proteins that catalyze 63 membrane fusion, as seen for influenza A virus (IAV) and vesicular stomatitis virus 64 (VSV). For Zaire ebolavirus (ZEBOV), proteolytic processing of the envelope protein by host cell proteases (1) is necessary to expose the receptor binding domain prior to 65 engagement of Niemman-Pick disease type 1C (NPC1 or NPC Intracellular Cholesterol 66 67 Transporter 1) – the late endosomal-lysosomal receptor protein (2). Proteolytic processing is also required for severe acute respiratory syndrome coronavirus (SARS-68 69 CoV) (3, 4), and for the current pandemic SARS-CoV-2 (5). Lassa fever virus (LASV) 70 uses a different mechanism, binding alpha-dystroglycan at the plasma membrane (6), 71 for internalization with a subsequent pH-regulated switch that leads to engagement of 72 lysosomal associated membrane protein 1 (LAMP1) for membrane fusion (7). Lymphocytic choriomeningitis virus (LCMV) also uses alpha-dystroglycan (6) and is 73 74 internalized in a manner that depends on endosomal sorting complexes required for transport (ESCRT) proteins (8), although it remains unknown whether a second 75 76 receptor is required.

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78 A hallmark of the endolysosomal system is controlled dynamic trafficking of vesicular 79 carriers among its various sub-compartments. Phophoinositides are markers for defining the identity of these sub-compartments because they are restricted in their 80 81 distribution to specific intracellular membranes [reviewed in (9)]. Although it is one of the 82 least abundant of the phosphoinositides in cells, PI(3,5)P2 is particularly important for endomembrane homeostasis. It is produced by PIKfyve, which phosphorylates the D-5 83 84 position in phosphatidylinositol-3-phosphate (PI3P) to yield phosphatidylinositol 3,5-85 bisphosphate (PI(3,5)P2) (10). First cloned as mammalian p235 (11), PIKfyve is a 240 kDa class III lipid kinase, present on the cytosolic face of endosomal membranes (12, 86 13) as part of a ternary complex with the PI(3,5)P2 5-phosphatase Sac3 and ArPIKfyve 87 (14). 88

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Ablation of PIKfyve function by genetic (12, 15) or pharmacological means (16-20)
 causes endosomal swelling and vacuolation of late endosomes and endolysosomes. It

92 is thought that these changes result from decreased membrane fission and concomitant 93 interference in endosomal traffic (13, 21). Small-molecule inhibitors of PIKfyve, all of which have some structural resemblance to each other, have been studied as potential 94 drugs for treating cancer and autoimmune diseases. These inhibitors include Apilimod 95 96 (19), Vacuolin-1 (18), a series of 30 Vacuolin-related molecules (22), YM201636 (16), and WX8 chemical family members (20). Physiological effects of these compound in 97 98 cells include inhibition of autophagy (17, 22, 23), reduced generation of IL-12/IL-23 (24), 99 and reduced dendritic cell infiltration in psoriasis (25).

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Apilimod also inhibits infection by several viruses, including ZEBOV. Although it does 101 102 not alter the pH of endosomes nor inhibit cathepsin B or L (26), Apilimod blocks entry of 103 ZEBOV and other pathogenic filoviruses (27). Several groups reported that Apilimod prevents colocalization of VSV-ZEBOV pseudoviruses with the ZEBOV endosomal 104 105 receptor NPC1, but does not prevent colocalization with early endosomal antigen 1 106 (EEA1) (5, 27, 28). Apilimod also inhibits entry of pseudotyped viruses bearing the spike 107 proteins of MERS-CoV, SARS-CoV, and SARS-CoV-2, as well as of authentic mouse 108 hepatitis virus (MHV) particles (5).

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110 Here, we have studied the effects of Apilimod on infection of VSV-eGFP-SARS-CoV-2 111 and VSV-eGFP-ZEBOV chimeras and showed that Apilimod blocks infection of both, 112 with an IC50 of ~50 nM. Apilimod and Vacuolin-1 also prevented entry and infection of VSV-MeGFP-ZEBOV and many of the internalized VSV-MeGFP-ZEBOV virions 113 114 colocalized with NPC1 in the distended, vacuolated endosomes. This suggests that 115 blocking PIKfyve kinase has the same downstream effects on these viruses, even though VSV-eGFP-SARS-CoV-2 does not require interaction with NPC1 for membrane 116 117 fusion. Apilimod also inhibits infection by authentic SARS-CoV-2 strain 2019-118 nCoV/USA-WA1/2020 virus, with an IC50 slightly lower than its IC50 for the VSV-eGFP-119 SARS-CoV-2. We suggest that Apilimod, which has passed safety tests in previous 120 human clinical trials for non-viral indications (24, 25, 29, 30), is a potential starting point for developing small-molecule entry inhibitors of SARS-CoV-2 that could limit infection 121 122 and disease pathogenesis.

123 **RESULTS**

Apilimod inhibits infection of VSV-MeGFP-LCMV and VSV-ZEBOV. We inoculated 124 SVG-A cells with vesicular stomatitis virus (VSV) chimeras expressing the viral matrix 125 126 protein (M) fused to eGFP (MeGFP). The chimeras include VSV (VSV-MeGFP, which initiates fusion at pH<6.2), VSV-V269H GP (VSV-MeGFP-V269H, a variant of VSV GP 127 that initiates fusion at pH<5.8), rabies virus GP (VSV-MeGFP-RABV), Lassa virus GP 128 (VSV-MeGFP-LASV), lymphocytic choriomeningitis virus GP (VSV-MeGFP-LCMV) or 129 130 Zaire Ebola virus GP (VSV-MeGFP-ZEBOV). Following the incubation protocol 131 summarized in **Fig 1A**, we tested the effects on infection of Apilimod or Vacuolin-1; both compounds are small-molecule inhibitors of PIKfyve kinase, which generates PI(5)P 132 and PI(3,5)P2 in late endosomes and lysosomes. Using a flow cytometry based-assay 133 134 to monitor a single round of infection determined by expression of viral MeGFP (Fig. **1B**), we found that Apilimod and Vacuolin-1 potently inhibit VSV-MeGFP-ZEBOV 135 136 infection (Fig. 1C). These results agree with results obtained by others with Apilimod 137 (26, 31) in different cell types infected with MLV virus pseudotyped with ZEBOV GP or 138 with Ebola virus itself (26, 27, 32). Apilimod was a less effective inhibitor of VSV-MeGFP-LCMV infection, and Vacuolin-1 had no effect at the concentration used. In 139 contrast, Apilimod and Vacuolin-1 failed to prevent infection by VSV-MeGFP, VSV-140 141 MeGFP-V269H, VSV-MeGFP-RABV, or VSV-MeGFP-LASV (Fig. 1C). IN1 (33), an 142 inhibitor of the phosphoinositide kinase Vps34, the main endosomal generator of PI3P, also interfered with VSV-MeGFP-LCMV and VSV-MeGFP-ZEBOV infection (Fig. 1C). 143 144 All of these viruses require low pH to trigger viral membrane fusion with the endosomal 145 membranes, and as expected, infection was fully blocked by Bafilomycin A1, which 146 inhibits the vacuolar type H^+ -ATPase (V-ATPase) acidification activity (**Fig. 1C**).

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148 Apilimod and Vacuolin-1 prevent cytoplasmic entry of VSV-MeGFP-ZEBOV. 149 Productive infection requires delivery of the viral ribonucleoprotein core (RNP) into the cytosol. In these experiments, we deemed RNP delivery, as monitored by single cell 150 151 fluorescence microscopy imaging (experimental protocol summarized in Fig. 2A and 152 **3A**), to be successful when fluorescent MeGFP encapsulated in the incoming virus appeared at the nuclear margin of infected cells. The representative examples of VSV 153 infection and RNP core release shown in Fig. 2B were obtained in the absence or 154 presence of cycloheximide, which prevents viral protein expression. In the absence of 155 156 cycloheximide (*left panel*), large amounts of newly synthesized MeGFP are present throughout the cell. In the presence of cycloheximide (*right panel*), we observed MeGFP 157 in virions (fluorescent spots) as well as released MeGFP concentrated at the nuclear 158

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margin. We scored the effect of Apilimod, Vacuolin-1 or IN1 on RNP delivery by VSVMeGFP, VSV-MeGFP-V269H and VSV-MeGFP-ZEBOV by determining the appearance
of MeGFP at the nuclear margin in cycloheximide-treated cells. Consistent with the
infection results, Apilimod, Vacuolin-1 and IN1 prevented entry of VSV-MeGFP-ZEBOV
but not of VSV-MeGFP or VSV-MeGFP-V269H. As expected, Bafilomycin A1 blocked
entry of all viruses (images in Fig. 2C and quantification in Fig. 2D).

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166 Intracellular trafficking of virus particles in the presence of Apilimod or Vacuolin-167 1. Internalized virus particles traffic along the endocytic pathway to reach the endosomal compartment(s) from which membrane fusion and genome entry into the 168 cytosol occur. To establish the identity of the endosomal compartments, we used 169 genome-editing in SVG-A cells (Figs. 3C, G and 4B, D) to replace expression of a 170 subset of proteins enriched in different endosomal compartments (the small GTPases 171 172 Rab5c and Rab7a, EEA1, or NPC1) with their corresponding fluorescent chimeras 173 obtained by fusion with TagRFP, mScarlet, or Halo (Figs. 3B, E, F, I, and 4C, E). The 174 lack of fluorescently tagged filipin (a cholesterol binder) in the endolysosomal compartment in the absence but not in the presence of U18666A, a potent inhibitor of 175 176 NPC1 (Fig 4F), showed that NPC1-Halo remained active as a cholesterol transporter.

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178 Using live-cell spinning disk confocal microscopy (Fig. 3, 4), we monitored the presence 179 of virus particles in the fluorescently tagged endosomes by colocalization with the 180 fluorescent spots from the virus-incorporated MeGFP. We monitored entry by carrying 181 out the experiments in the presence of cycloheximide, thus ensuring that any MeGFP 182 fluorescent signal at the nuclear margin originated only from MeGFP molecules carried by incoming viral particles (Fig. 3B, F). All cells were maintained at 37°C throughout all 183 184 phases of the experiment to ensure normal and undisturbed intracellular trafficking. All 185 control experiments performed in the absence of inhibitors showed arrival of VSV-MeGFP, VSV-MeGFP-V269H, or VSV-MeGFP-ZEBOV virus particles to early (Rab5c 186 and EEA1) (Fig. 3E, 4E) or late endosomes and lysosomes (Rab7a or NPC1) (Fig. 3I. 187 4C, E). MeGFP released from all viruses appeared at the nuclear margin, showing 188 effective RNP release. NPC1, the receptor for VSV-MeGFP-ZEBOV entry is required for 189 fusion from endosomes (2). The successful VSV-MeGFP-ZEBOV infection observed in 190 the absence of drug in cells expressing NPC1-Halo alone or in combination with 191 192 mScarlet-EEA1 indicates that NPC1-Halo is capable of facilitating infection and that 193 VSV-MeGFP-ZEBOV trafficked to NPC1-Halo-containing endosomes.

194 Apilimod and Vacuolin-1 treatment of the SVG-A cells led to enlargement and vacuolization of their endosomes and lysosomes tagged with fluorescent EEA1, Rab5c, 195 Rab7a or NPC1 (Fig. 3-5), in agreement with earlier PIKfyve ablation studies (13, 21). 196 197 VSV-MeGFP and VSV-MeGFP-V269H (fluorescent dots, white) reached all tagged species of enlarged endolvsosomes and successfully penetrated into the cytosol, as 198 indicated by MeGFP at the nuclear margin (Fig. 3E. I). VSV-MeGFP-ZEBOV also 199 200 trafficked to all tagged species of enlarged endolysosomes (Fig. 3E, I), often reaching 201 one of the numerous NPC1-containing vacuoles enriched in EEA1 (Figs. 4E and 5B,C). 202 VSV-MeGFP-ZEBOV in EEA1-containing endosomes increased in the presence of Apilimod, as also reported for VLP ZEBOV (27). While able to reach NPC1-containing 203 functional endosomes in cells treated with Apilimod (Fig. 4C, E and 5B, C), VSV-204 205 MeGFP-ZEBOV failed to penetrate into the cytoplasm, as reflected by absence of MeGFP in the nuclear margin (Fig. 2C, 3E, I, 4C, E and 5B). 206

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208 Apilimod blocks infection of VSV SARS-CoV-2. Using a recombinant vesicular 209 stomatitis virus (VSV) expressing soluble eGFP (VSV-eGFP) where the glycoprotein (GP) was replaced with that of ZEBOV GP (VSV-eGFP-ZEBOV) or SARS-CoV-2 S 210 211 (VSV-eGFP-SARS-Cov2), we inoculated MA104 cells with these chimera viruses and 212 tested the effects of Apilimod on infection by flow cytometry (Fig. 6A). We found potent 213 inhibition of VSV-eGFP-SARS-CoV-2 infection by Apilimod and confirmed that the compound also inhibits VSV-eGFP-ZEBOV infection (Fig. 6B). The dose-response 214 curves indicated similar effects for VSV-eGFP-ZEBOV and VSV-eGFP-SARS-CoV-2 215 216 (IC50s ~ 50 nM), in contrast to the absence of any detectable inhibition of VSV-eGFP 217 infection, used here as a negative control.

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219 Apilimod blocks infection of SARS-CoV-2 virus.

To test the effect of Apilimod on bona fide SARS-CoV-2 infection, we exposed Vero E6 cells to fully infectious SARS-CoV-2 (strain 2019-nCoV/USA-WA1/2020); after 24 h incubation, supernatants were harvested and tittered by focus-forming assay on a separate set of Vero E6 cells (**Fig. 7A**). Apilimod strongly inhibited SARS-CoV-2 infection, and the dose-response curve was similar or more potent than those observed for VSV-eGFP-ZEBOV or VSV-eGFP-SARS-CoV-2 (IC50s ~ 10 nM) (**Fig. 7B**).

226 **DISCUSSION**

227 Coronaviruses, filoviruses, and arenaviruses have different replication strategies and 228 unrelated surface glycoproteins that engage different receptor molecules during entry 229 (1, 2, 5-8). Coronavirus and filovirus surface glycoproteins share a requirement for 230 entry-associated proteolytic processing for activation as fusogens (1) Filoviruses require 231 passage through low pH compartments where cathepsins are active. Coronaviruses 232 may enter directly by fusion at the plasma membrane or following receptor mediated endocytosis. Cell entry of SARS-CoV and SARS-CoV-2 depends on the protease 233 234 TMPRSS2 in conjunction with ACE2 (34-37), and when TMPRSS2 is present, the entry pathway becomes insensitive to cathepsin inhibition (34, 37, 38). 235

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237 The common inhibition of viruses from all three groups by Apilimod is a consequence of perturbing their shared entry pathway. Moreover, it is not the cathepsin activity itself that 238 239 these compounds affect, judging from the outcome of the assays with Apilimod and Vacuolin-1 showing they inhibit VSV chimeras bearing the surface glycoproteins of 240 241 ZEBOV and LCMV and to a lesser extent LASV. Apilimod also inhibits infection of cells by VSV-SARS-CoV-2 as well as by authentic SARS-CoV-2; neither compound blocks 242 infection by wild-type VSV. For VSV-ZEBOV, we have shown that the virus reaches a 243 244 compartment enriched in NPC1, the ZEBOV co-receptor, and often also enriched in EEA1, but that it nonetheless fails to release internal proteins into the cytosol. Apilimod 245 246 does not inhibit cathepsin (26) but Apilimod (39) and Vacuolin-1 (17, 23) can interfere 247 with cathepsin maturation as evidenced by an increase in pro-cathepsin in treated cells; 248 they do not influence endosomal pH (18, 26, 40) although other studies report Apilimod 249 decreases cathepsin activity (41) and Vacuolin-1 increases pH (17, 23). Irrespective of this discrepancy, both Apilimod and Vacuolin-1 inhibit PI-3P-5-kinase (PIKfyve) (17, 19), 250 251 a three-subunit complex (14) with a PI-3P-binding FYVE domain (10, 11) that 252 recognizes the endosomal marker, PI-3-P. Functional ablation of this enzyme by 253 agenetic means (12, 15) gives rise to the same cellular phenotype as treatment with 254 either compound (17-19). The similar dose-response curves for Apilimod inhibition of the ZEBOV and SARS-CoV-2 chimeras (IC50 of ~ 50 nM) and of authentic SARS-CoV-255 2 virus (IC50 ~ 10 nM) are in good agreement with the IC50 of ~ 15 nM for Apilimod 256 inhibition of PIKfyve in vitro (19). Thus, perturbing normal endosomal trafficking by 257 258 inhibiting PIKfyve activity suggests it is the mechanism by which Apilimod and Vacuolin-259 1 block entry of such a diverse set of viral pathogens.

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261 One of the most striking consequence of PIKfyve inhibition, and hence of PI-3,5-P₂ 262 restriction in endosomal membranes, is the swelling of endosomes into small, spherical 263 vacuoles - the phenomenon that gave Vacuolin-1 its name (18). Our imaging data with 264 VSV-MeGFP-ZEBOV chimeras show that the virus particles accumulating in these 265 structures, many of which also contain the NPC1 co-receptor (2, 42), often appear to be 266 relatively immobile and adjacent to the endosomal limiting membrane. One possible 267 explanation is that when a virion reaches these distended endosomes, it can bind or 268 remain bound to the limiting membrane, but not fuse. Another is that virions may fuse 269 with smaller intraluminal vesicles in the endosomal lumen (43), but that PI-3,5-P2 270 depletion prevents back fusion of these vesicles with the endosomal limiting membrane 271 and inhibits release into the cytosol of the viral genome.

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273 Inhibition of infection by authentic SARS-CoV-2 shows that the blocked release of the 274 viral genome from a vacuolated endosome is independent of the shape, size, and 275 distribution of spike protein on the virion. The assay we used to determine effects on 276 infectivity of authentic virus measured release of virions after multiple rounds of 277 infection, rather than entry, which we monitored in the VSV-SARS-CoV-2 experiments 278 by detecting eGFP synthesis in the cytosol. Nevertheless, the IC50 of Apilimod in 279 experiments with authentic virus is remarkably similar (or even more potent) to that obtained with chimeric VSV-SARS-CoV-2. 280

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282 Although cathepsin L inhibitors block SARS-CoV and SARS-CoV-2 infection in cell 283 culture (4, 5), they have less pronounced effects when tested in animals (44). This may 284 because another protease, TMPRSS2 on the surface of cells in relevant tissues, appears to prime SARS-CoV (44) and SARS-CoV-2 (37) spike proteins for efficient 285 entry. As the effectiveness of Apilimod and Vacuolin-1 does not depend on cathepsin 286 287 inhibition, their capacity to block entry of several distinct families of viruses is likely to be 288 independent and downstream of the protease that primes their surface glycoprotein for 289 fusion. Phase I and phase II clinical trials have shown that Apilimod is safe and well-290 tolerated (24, 25, 29, 30). The trials were discontinued because of lack of effectiveness against the autoimmune condition for which the drug was tested. We suggest that one 291 292 of these compounds, or a potential derivative, could be a candidate broad-spectrum 293 therapeutic for several emerging human viral pathogens, including SARS-CoV-2.

294FIGURE LEGENDS

Figure 1. Apilimod and Vacuolin-1 inhibit VSV-MeGFP-ZEBOV infection.

- (A) Schematic of infectivity assay, where SVG-A cells were pretreated for 1 h with 5 μ M 296 297 Vacuolin, 5 µM Apilimod, 5 µM IN1, or 10 nM BAF A1 and subsequently infected with VSV-MeGFP (multiplicity of infection, MOI = 2), VSV-MeGFP-V269H (MOI = 1), VSV-298 MeGFP-RABV (MO I= 0.6), VSV-MeGFP-LASV (MOI = 0.6), VSV-MeGFP-LCMV (MOI 299 = 0.6) or VSV-MeGFP-ZEBOV (MOI = 0.6) for 1 h in the presence of drugs. The cells 300 were then washed to remove unbound virus and incubated for the indicated times in the 301 302 presence of drugs. The cells were then fixed and the percentage of cells expressing 303 viral MeGFP was measured by flow cytometry.
- 304 (B) Representative flow cytometry results of an infection assay using VSV-MeGFP-305 ZEBOV.
- 306 **(C)** Quantification of the infectivity is shown with averages from three independent 307 experiments per condition each determined as a duplicate measurement (error bars 308 show SEM). The statistical significance was determined using a one-way ANOVA and 309 Tukey *post-hoc* test (*, $P \le 0.05$; **, $P \le 0.01$; ***, $P \le 0.001$).
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311 Figure 2. Apilimod and Vacuolin-1 inhibit VSV-MeGFP-ZEBOV.

- (A) Schematic of entry assay where SVG-A cells were infected with VSV-MeGFP (MOI = 4), VSV-MeGFP-V269H (MOI = 4), or VSV-MeGFP-ZEBOV (MOI = 4). Experiments were performed in the presence of 5 μ g/mL cycloheximide (CHX) to prevent protein synthesis. Entry assay was based on the appearance of MeGFP fluorescence on the nuclear margin on a per cell basis, of fixed infected cells visualized by fluorescence microscopy. Staining the fixed cells with Alexa647 labeled wheat germ agglutinin identified the plasma membrane of each cell (dashed outlines in **C**).
- (B) Virus infection in the absence of CHX (left panel) resulted in the appearance of
 MeGFP fluorescence throughout the cell volume. The presence of CHX resulted in virus
 entry being observed by MeGFP fluorescence at the nuclear margin, which was
 released from incoming viral particles (right panel, white arrows). Scale bar indicates 10
 µm.
- (C) Representative examples of maximum-Z projections images from the whole cell
 volume obtained with optical sections separated by 0.3 µm using spinning disc confocal
 microscopy. Scale bar indicates 10 µm.
- 327 (D) Quantification of the number of cells with nuclear margin labeling from three328 independent experiments each determined from fields containing 59-90 cells (error bars

show SEM). The statistical significance of the entry data was analyzed for statistical significance by one-way ANOVA and Tukey *post-hoc* test (***, $P \le 0.001$).

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Figure 3. Endolysosomal traffic of VSV-MeGFP-ZEBOV in cells expressing
 TagRFP-Rab5c or TagRFP-Rab7a in the presence of Apilimod or Vacuolin-1.
 (Associated Videos 1 and 2).

- (A) Schematic of live cell imaging experiment using SVG-A cells expressing
 fluorescently tagged TagRFP-Rab5c or TagRFP-Rab7a. Cells were infected with VSVMeGFP, VSV-MeGFP-V269H or VSV-MeGFP-ZEBOV (MOI = 4). Viruses trafficking
 (monitored with MeGFP) to the endo-lysosomal system (recognized by their labeling
 with TagRFP-Rab5c or TagRFP-Rab7a) and virus entry (established by MeGFP at the
 nuclear margin) were ascertained by live-cell florescence imaging using a spinning disc
 confocal microscope.
- (B) Visualization of VSV-MeGFP infection in TagRFP-Rab5c cells in the absence (left
 panel) or presence of CHX (right panel, white arrows) using live-cell imaging. Scale bar
 represents 10 μm.
- (C) Genomic PCR analysis of SVG-A cells showing biallelic integration of TagRFP into
 the *RAB5C* genomic locus by cotransfection of a plasmid coding for Cas9, a linear PCR
 product coding for the specific gRNAs targeting a region near the ATG codon of Rab5c
 under the control of the U6 promoter, and a template plasmid containing the RFP
 sequence flanked by 800 base pairs upstream and downstream of the targeted region
 (see materials and methods for more details) to generate a clonal gene-edited cell-line
 expressing TagRFP-Rab5c.
- 352 (**D**) Quantification of VSV-MeGFP and VSV-MeGFP-ZEBOV colocalization with Rab5c 353 containing endosomes in the presence of CHX together with absence or presence of 5 354 μ M Apilimod depicted in (E). Data shows number of viruses that colocalized with 355 endosomes containing or not Rab5c within the complete volume of the single cells 356 depicted in (e).
- 357 **(E)** Representative examples of maximum-Z projection images from four optical 358 sections spaced 0.35 μ m apart of virus entry without or with IN1, Vacuolin, or Apilimod 359 for VSV-MeGFP (top), VSV-Me-GFP-V269H (middle), and VSV-MeGFP-ZEBOV 360 (bottom). Each condition is in the presence of CHX. All viruses reach Rab5c-containing 361 endosomes but only VSV-MeGFP-ZEBOV fails to penetrate in the presence of IN1, 362 Vacuolin-1, or Apilimod. Scale bars are 10 μ m. Insets correspond to a single optical 363 section with scale bars of 3 μ m.

(F) Visualization of VSV infection in TagRFP-Rab7a cells in the absence of CHX (left
 panel) and entry in the presence of CHX (right panel, white arrows) with scale bar
 indicating 10 μm.

(G) Genomic PCR analysis showing biallelic integration of TagRFP into the *RAB7A* genomic locus to generate a clonal gene-edited cell-line expressing TagRFP-Rab7a,
 using the same approach as used for *RAB5C*.

- (H) Quantification of VSV-MeGFP and VSV-MeGFP-ZEBOV colocalization with Rab7a
 containing endosomes in the presence of CHX with or without 5 µM Apilimod within the
 complete cell volumes in the images depicted in (I).
- (I) Representative examples of maximum-Z projection images from four optical sections
 spaced 0.35 µm apart of virus entry without or with IN1, Vacuolin, or Apilimod for VSVMeGFP (top), VSV-Me-GFP-V269H (middle), and VSV-MeGFP-ZEBOV (bottom). All
 viruses reach Rab7a-containing endosomes but only VSV-MeGFP-ZEBOV fails to
 penetrate in the presence of IN1, Vacuolin-1 or Apilimod. Scale bars are 10 µm. Insets
- 378 correspond to a single optical section with scale bars of $3 \mu m$.
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Figure 4. Endolysosomal traffic of VSV-MeGFP-ZEBOV in cells expressing NPC1 Halo or coexpressing mScarlet-EEA1 and NPC1-Halo in the presence of Apilimod. (Associated Video 3).

- (A) Schematic of live cell imaging experiment with gene-edited SVG-A cells expressing
 NPC1-Halo or NPC1-Halo together with mScarlet-EEA1. Halo was labeled with either
 JF549 or JF647. Cells were infected with VSV-MeGFP-ZEBOV (MOI = 3).
- (B) Genomic PCR analysis showing biallelic integration of Halo into the *NPC1* genomic
 locus to generate a clonal gene-edited cell-line expressing NPC1-Halo, using the same
 approach as for *RAB5C* and *RAB7A*.
- (C) Representative examples of maximum-Z projection images from four optical
 sections spaced 0.25 μm apart in the absence (left) and presence of Apilimod (right)
 showing that VSV-MeGFP-ZEBOV reached NPC1-Halo-containing endosomes even in
 the presence of Apilimod, while failing to penetrate and infect. Scale bar indicates 10
 μm. Insets correspond to a single optical section with the scale bar indicating 3 μm.
- (D) SVG-A cells with genomic NPC1-Halo were further gene edited to contain EEA1
 tagged with mScarlet. Genomic PCR analysis shows biallelic integration into the *EEA1* locus of mScarlet-EEA1 (left) and into the *NPC1* locus of NPC1-Halo (right).
- (E) Representative examples of maximum-Z projection images in the absence (left) and
 presence of Apilimod (right) showing that VSV-MeGFP-ZEBOV reached endosomes
 containing mScarlet-EEA1 and endosomes containing both mScarlet-EEA1 and NPC1-

Halo in the presence of Apilimod, while failing to penetrate and infect. Scale bar
indicates 10 μm. Insets correspond to a single optical section with scale bar indicating 3
μm.

(F) Representative images of parental (top) and gene-edited SVG-A cells expressing
 NPC1-Halo (bottom) incubated with filipin III (naturally fluorescent polyene antibiotic
 that binds to cholesterol) in the absence (left) and presence of U18666A (right, NPC1
 inhibitor of cholesterol export) showing NPC1-Halo is a functional cholesterol
 transporter.

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Figure 5. Extent of VSV-MeGFP-ZEBOV traffic into endosomes enriched in in NPC1-Halo or NPC1-Halo and mScarlet-EEA1.

- (A) Schematic of imaging experiment of VSV-MeGFP-ZEBOV trafficking in NPC1-Halo
 or NPC1-Halo and mScarlet-EEA1 gene edited SVG-A cells.
- (B) Representative examples of maximum-Z projection images from four optical
 sections spaced 0.25 µm apart in the absence and presence of Apilimod after 2 or 4
 hours post infection. A large number of VSV-MeGFP-ZEBOV but not of VSV-MeGFP
 particles accumulated in the endosomes enlarged upon Apilimod treatment.
- 417 (C) Quantification of VSV-MeGFP-ZEBOV colocalization with mScarlet-EEA1 alone,
 418 both mScarlet-EEA1 and NPC1-Halo, or NPC1-Halo alone 2 and 4 hours post infection,
 419 in the absence or presence of 5 μM Apilimod. Data obtained from complete cell volumes
 420 are presented as numbers and corresponding % colocalizations of VSV-MeGFP-
- 421 ZEBOV particles associated with a given type of endosome.
- 422

423 Figure 6. Apilimod and Vacuolin-1 inhibit infection of VSV-eGFP-SARS-CoV-2

- (A) Schematic of infectivity assay of VSV-eGFP, VSV-eGFP-ZEBOV, and VSV-eGFPSARS-CoV-2 in MA104 cells. MA104 cells were pretreated for 1 h with the indicated
 concentration Apilimod. Pretreated cells were inoculated with the indicated virus (MOI =
 1) for 1 h at 37°C. At 6 hours post infection cells were harvested and the fraction of cell
 expressing eGFP cells quantified by flow cytometry.
- 429 **(B)** Quantification of the infectivity is shown with averages +/- SEM from three 430 independent experiments. Statistical significance was determined using a T-test (*, P \leq 431 0.05; **, P \leq 0.01).
- 432

433 Figure 7. Apilimod inhibits infection of SARS-CoV-2 virus

(A) Schematic of infectivity assay of fully infectious Sars-CoV-2 (strain 2019-nCoV/USA-

435 WA1/2020). Vero E6 cell monolayers were pretreated with medium containing DMSO

436 or serial dilutions of Apilimod at the indicated concentrations. SARS-CoV-2 was diluted

- 437 (MOI = 0.01) in Apilimod-containing medium and added to Vero E6 cells for 1 h at 37° C.
- After adsorption, the viral inocula were removed, and medium containing the respective
 concentration of Apilimod was reapplied. After 24 h incubation, supernatants were
 harvested and titrated by focus-forming assay on a separate set of Vero E6 cells.
- (B) Quantification of the infectivity is shown with averages +/- SEM from three
 independent experiments per condition and expressed as the percent infection relative
 to mock-treated SARS-CoV-2 infected cells.
- 444
- Table I. Primer sequences used to generate the sgRNAs and corresponding
 genomic fragments.
- 447

448 **Table II. Primer sequences used for screening.**

449

450Video 1. Apilimod doesn't inhibit VSV-MeGFP entry. Maximal Z-projection from four451optical sections separated 0.25 μm apart of SVG-A cells gene-edited to express452TagRFP-Rab5c imaged by spinning disc confocal microscopy every 3 seconds for 3453min. Cells were infected with VSV-MeGFP (MOI = 4) in the presence of CHX with or454without 5 μM Apilimod and imaged ~ 3-4 h post-infection.

455

Video 2. Apilimod inhibits VSV-MeGFP-ZEBOV entry. Maximal Z-projection from four
optical sections separated 0.25 μM apart of SVG-A cells gene-edited to express
TagRFP-Rab5c imaged by spinning disc confocal microscopy every 3 seconds for 3
min. Cells were infected with VSV-MeGFP-ZEBOV (MOI = 3) in the presence of CHX
with or without 5 μM Apilimod and imaged ~ 6-7 h post infection.

Video 3. Apilimod inhibits VSV-MeGFP-ZEBOV entry. Maximal Z-projection from four optical sections separated 0.25 μM apart of SVG-A cells gene-edited to express NPC1-Halo imaged by spinning disc confocal microscopy every 3 seconds for 3 min. Cells were infected with VSV-MeGFP-ZEBOV (MOI = 3) with or without 5 μM Apilimod and imaged ~ 5 h post infection.

467 **MATERIAL AND METHODS**

468 Cell culture

Human astroglial SVG-A derived cells (a kind gift from Walter J. Atwood) were grown at 469 470 37° C and 5% CO₂ in Minimum Essential Medium (MEM) (10-010-CV; Corning) 471 supplemented with 10% heat inactivated fetal bovine serum (S11150; Atlanta Biologicals), penicillin and streptomycin (1406-05-9; VWR International). African Green 472 473 Monkey kidney epithelial MA104 cells (a kind gift from Siyuan Ding, WUSM) were grown 474 at 37°C and 5% CO₂ in Medium 199 supplemented with 10% heat inactivated fetal 475 bovine serum. Vero C1008 [Vero 76, clone E6, Vero E6] (ATCC CRL-1586) cells were cultured in Dulbecco's Modified Eagle Medium (DMEM) supplemented with 10% fetal 476 477 bovine serum, and penicillin and streptomycin. Vero CCL-81 (ATCC CCL-81) cells were 478 maintained in DMEM supplemented with 10% FBS, 10mM HEPES pH 7.4, 1% 479 Glutamax, and penicillin/streptomycin.

480

481 **Reagents**

Vacuolin-1 (18) was custom synthesized; Apilimod (HY-14644) was from MedChem
Express, IN1 was a kind gift from Dr. N. Gray (33), U-18666A (10009085), and Filipin III
(70440) were from Cayman Chemical, Bafilomycin A1 (B1793-2UG) was from SigmaAldrich, Cycloheximide (239764) was from Calbiochem, and wheat germ agglutinin
conjugated with Alexa Fluor®-647 (W32466) was from ThermoFisher.

487

488 Viruses

- Recombinant VSV (Indiana serotype) expressing MeGFP alone which initiates fusion at
 pH<6.2 (VSV-MeGFP) (45) (or in combination with V269H GP, VSV-MeGFP-V269H),
 RABV GP (VSV-MeGFP-RABV) (46), LASV GP (VSV-MeGFP-LASV) (7), LCMV GP
 (VSV-MeGFP-LCMV), Zaire EBOV GP (VSV-MeGFP-ZEBOV) (47) or SARS-CoV-2 S
 Wuhan-Hu-1 strain (VSV-eGFP-SARS-CoV-2 description to be published elsewhere)
 were used for infection, entry and live cell imaging assays. All viruses were generated
 and recovered according to (48).
- 496

497 SARS-CoV-2 strain 2019-nCoV/USA-WA1/2020 was obtained from the Centers for
498 Disease Control and Prevention (gift of Natalie Thornburg). Virus was passaged once in
499 Vero CCL81 cells (ATCC) and titrated by focus-forming assay also on Vero E6 cells.

- 500
- 501 Genome-editing

Individual cell lines of SVG-A were gene edited in both alleles using the CRISPR/Cas9
system to incorporate fluorescent tags into the N-terminus of Rab5c (TagRFP), Rab7a
(TagRFP), EEA1 (mScarlet) or the C-terminus of NPC1 (Halo). The NPC1-Halo
expressing cells were further gene edited to incorporate mScarlet-EEA1 creating SVG-A
cells simultaneously expressing mScarlet-EEA1 and NPC1-Halo.

507

508 A free PCR strategy (49, 50) was used to generate small guide RNAs (sgRNA) with 509 target sequences for either Rab5c, Rab7a, NPC1, or EEA1.

510

The genomic DNA fragment of Rab5c, Rab7a, NPC1, or EEA1 genes fused with either 511 512 TagRFP, Halo, or mScarlet were cloned into the pUC19 vector (donor constructs) which 513 then served as homologous recombination repair templates for the Cas9 enzymecleaved genomic DNA. Donor constructs were obtained by a ligation of PCR 514 515 amplification products from the genomic DNA fragments, TagRFP, Halo, and mScarlet 516 sequences. Primers F1-R1 and F3-R3 amplified approximately 800 base pairs of 517 genomic sequences upstream and downstream of the start codon of Rab5c, Rab7a or EEA1, or the stop codon of NPC1, respectively. Primers F1 and R3 contain sequences 518 519 complementary to the pUC19 vector linearized using the Smal restriction enzyme (lower 520 case in the primer sequences). The TagRFP sequence containing the GGS peptide linker was amplified using primers F2-R2 from a TagRFP mammalian expression 521 522 plasmid used as a template. The F2 primer contains complementary sequences to the 523 3' end of the F1-R1 fragment, while the F3 primer contains complementary sequences 524 to the 3' end of the TagRFP sequences. Primer sequences used to generate the 525 sgRNAs and corresponding genomic fragments are listed in Table I; primers used for 526 screening are listed in Table II.

527

528 PCR products (fragments F1-R1, F2-R2, and F3-R3) were subjected to electrophoresis 529 in 1% agarose and gel purified using a purification kit from Zymogen. The PCR 530 fragments were cloned into the linearized pUC19 vector using the Gibson Assembly 531 Cloning Kit (E5510S; New England Biolabs).

532

533 SVG-A cells (1.5 x 10^5 cells) were co-transfected with 0.8 µg of *Streptococcus* 534 *pyogenes* Cas9, 0.8 µg free PCR product coding for the target sgRNA, and 0.8 µg 535 pUC19 vector using Lipofectamine 2000 reagent (Invitrogen) according to the 536 manufacturer's instructions. Transfected cells were grown for 7 to 10 days and sorted 537 for TagRFP, Halo, or mScarlet expression using fluorescence-activated cell sorting (FACS) (SH-800S; Sony). Prior to FACS, NPC1-Halo cells were labeled for 15 minutes
with Janelia FluorTM 647 (JF647). Single cells expressing the desired chimera were
isolated, clonally expanded, and then screened by genomic PCR for TagRFP, Halo, or
mScarlet insertion into both alleles.

542

543 Infection assays

SVG-A cells were plated at about 30-40% confluency into 24-well plates and incubated 544 545 for 1 day at 37°C and 5% CO₂. At the start of the experiment, cells were incubated with 546 the indicated drug or DMSO at 37°C for one hour. Following this, cells were incubated for 1 h at 37°C with VSV, VSV-MeGFP-V269H, VSV-MeGFP-RABV, VSV-MeGFP-547 548 LASV, VSV-MeGFP-LCMV or VSV-MeGFP-ZEBOV in drug or DMSO-containing 549 infection medium (α -MEM, 50mM HEPES, 2% FBS). Cells were then washed to remove non-adsorbed viruses and further incubated at 37°C in medium containing the drug or 550 551 DMSO with experiments ending at the indicated times by fixation with 3.7% formaldehyde in PBS. Fluorescent intensity from 20,000 single cells from a single round 552 of infection was determined by flow cytometry using a BD FACSCanto[™] II equipped 553 554 with DIVA software package.

555

556 MA104 cells were pretreated for 1 h with the indicated concentration Apilimod or DMSO. 557 Pretreated cells were inoculated with VSV-eGFP, VSV-eGFP-ZEBOV or VSV-eGFP-558 SARS-CoV-2 at an MOI = 1 (based on titers in MA104 cells) in the presence of Apilimod 559 or DMSO for 1 h at 37°C. Six to 8 h post infection, cells were collected and fixed in 2% 560 PFA and then subjected to flow cytometry. The percentage of GFP cells was 561 determined using FlowJo software (Tree Star Industries, Ashland, OR).

562

Vero E6 cell monolayers were pretreated for 1 h at 37°C with serial dilutions of Apilimod at the indicated concentrations. Next, SARS-CoV-2 was diluted to an MOI of 0.01 focusforming units (FFU)/cell in Apilimod-containing medium and added to Vero E6 cells for 1 h at 37°C. After adsorption, cells were washed once with PBS, and medium containing the respective concentration of Apilimod was added. Cell were incubated for a 24 h at 37°C, and at which time cell culture supernatants were removed and used for determination of viral titer by focus forming assay.

570

571 SARS-CoV-2 focus forming assay

572 Cell culture supernatants from virus-infected cells were diluted serially 10-fold and 573 added to Vero E6 cell monolayers in 96-well plates and incubated at 37°C for 1 h.

574 Subsequently, cells were overlaid with 1 % (w/v) methylcellulose in MEM supplemented with 2% FBS. Plates were harvested 30 h later by removing overlays and fixed with 4% 575 576 paraformaldehdye in PBS for 20 min at room temperature. Plates were washed and sequentially incubated with 1 µg/mL of CR3022 anti-spike antibody (51) and HRP-577 578 conjugated goat anti-human IgG in PBS supplemented with 0.1% saponin and 0.1% 579 BSA. SARS-CoV-2-infected cell foci were visualized using TrueBlue peroxidase 580 substrate (KPL) and quantitated on an ImmunoSpot microanalyzer (Cellular 581 Technologies). Data were processed using Prism software (GraphPad Prism 8.0) and 582 viral titers are reported as percent inhibition relative to mock-treated SARS-CoV-2 infected cells. 583

584

585 Entry assay and intracellular traffic

SVG-A cells plated on glass #1.5 coverslips at about 30-40% confluency one day prior 586 587 to experiment were treated with drug or DMSO for 1 h at 37°C. Following this, cells were incubated at 37°C with VSV, VSV-MeGFP-V269H, VSV-MeGFP-RABV. VSV-588 MeGFP-LASV, VSV-MeGFP-LCMV or VSV-MeGFP-ZEBOV in drug or DMSO 589 containing infection medium. After this, cells were washed then further incubated in 590 medium containing the drug or DMSO at 37°C with the experiment ending at the 591 indicated time by fixation for 20 min at room temperature with 3.7% formaldehyde in 592 PBS. This was followed with a 10-min incubation of 5 µg/mL of Alexa647-labeled wheat 593 594 germ agglutinin in PBS to label the outline of the cells.

595

596 Cells were imaged using a spinning disk confocal microscope with optical planes 597 spaced 0.3 µm apart (52). The entry assay scored the presence of MeGFP at the 598 nuclear margin in each cell. Trafficking of viruses to endosomal compartments was observed using live-cell imaging using the spinning disc confocal microscope. Chemical 599 600 fixation tends to eliminate the large endolysosomal vacuoles generated by Vacuolin-1 or 601 Applied and reduces the colocalization with viral particles contained within. Time series 602 with images taken every 3 seconds for 3 min in a single optical plane with the 603 appropriate fluorescent channels (52) were acquired from non-fixed samples imaged at the end of the experimental period. For experiments containing NPC1-Halo, the Halo-604 tagged cells were labeled with either 250 nM JF549 or JF647 dye in media for 30 min at 605 37°C. Following labeling, cells were washed three times with media. The microscope 606 607 was operated using the Slidebook 6.4 software package (31) and images were displayed 608 also using this software.

609

610 Statistical tests

To compare the means from cells with different treatments, one-way ANOVA and the

612 *post-hoc* Tukey test analysis were used to take into account unequal sample sizes as 613 indicated in figure legends.

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624

625 AUTHOR CONTRIBUTIONS (NAMES GIVEN AS INITIALS)

T.K., S.P.W., and M.S.D. were responsible for the overall design of the study; Y.K. 626 627 carried out virus infection, entry and imaging experiments and prepared figures in the lab of T.K. (Fig. 1-5). P.W.R. designed, generated and characterized VSV-eGFP-628 629 SARS-CoV-2 and Z.L. carried out VSV-chimera infection experiments in the lab of S.P.W. (Fig. 6). J.B.C. and R.E.C. carried out the experiments with authentic SARS-630 CoV-2 under BSL3 conditions in the lab of M.S.D. (Fig. 7). Recombinant viruses were 631 632 generated and characterized by D.K.C., S.P., M.R and T.S. in the lab of S.P.W; T.K. 633 drafted the manuscript and editorially reviewed it in close association with SP.W. and 634 M.S.D; the authors commented on the manuscript.

635

636 COMPETING FINANCIAL INTEREST STATEMENT

M.S.D. is a consultant for Inbios, Vir Biotechnology, NGM Biopharmaceuticals, and on
the Scientific Advisory Board of Moderna. The Diamond laboratory at Washington
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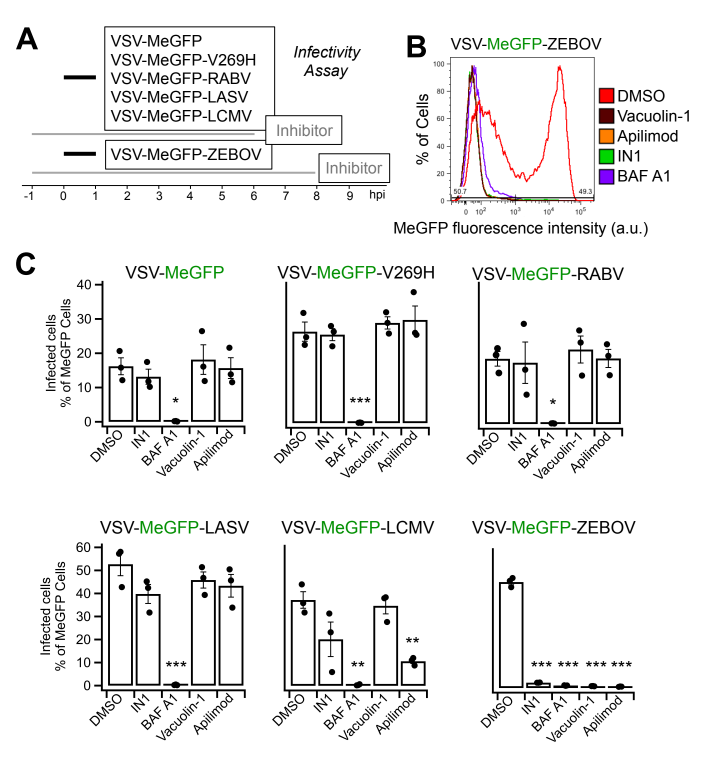
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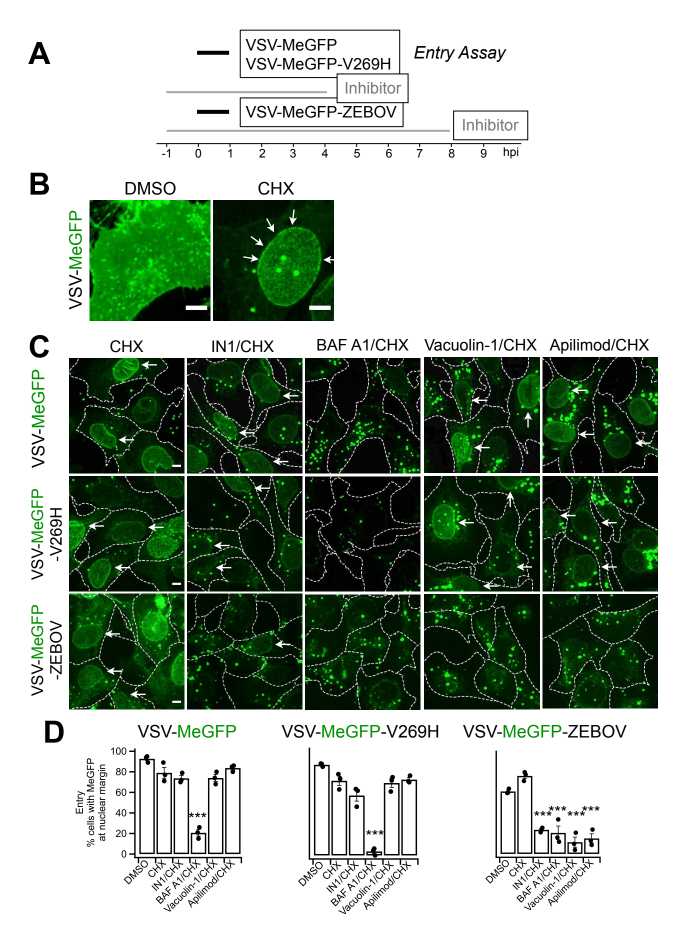
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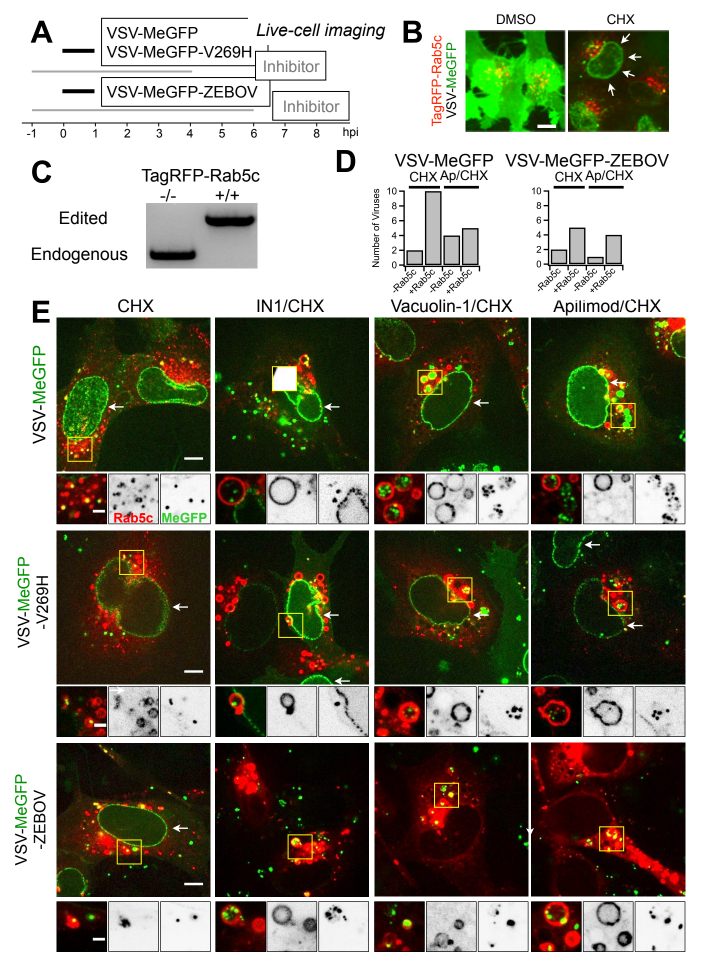
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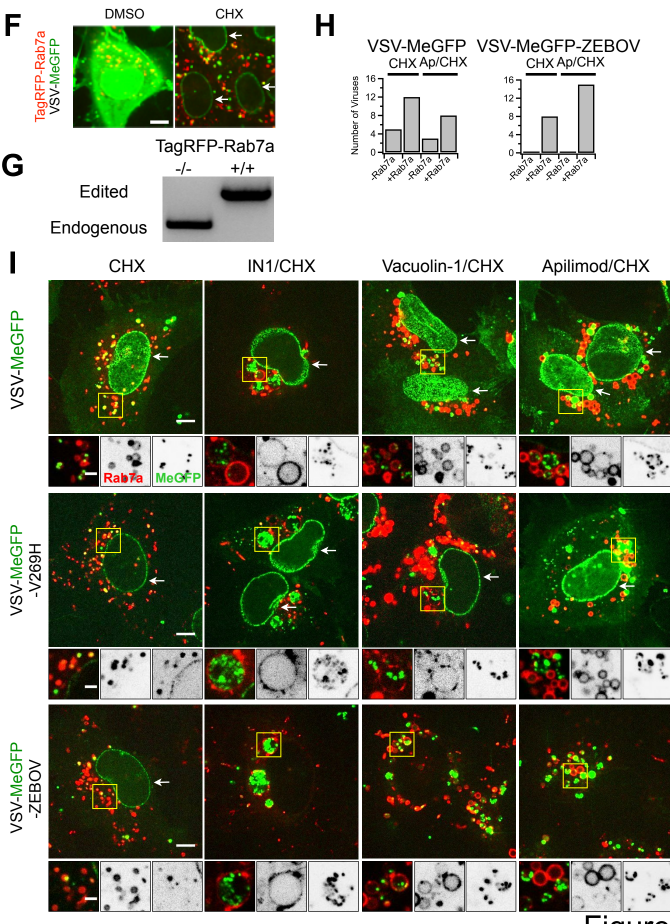
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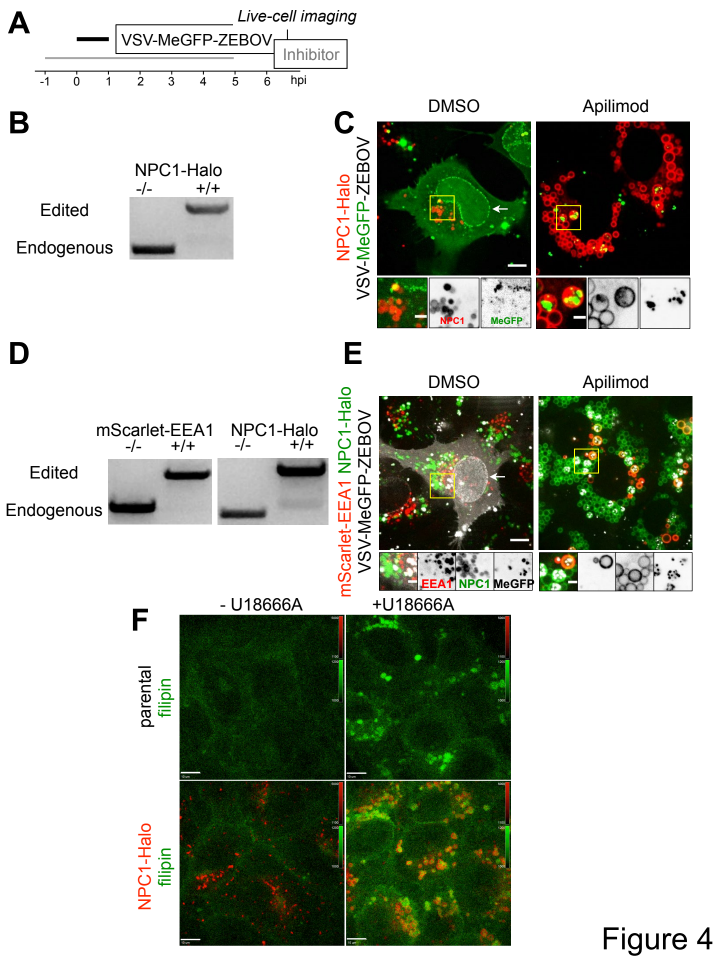
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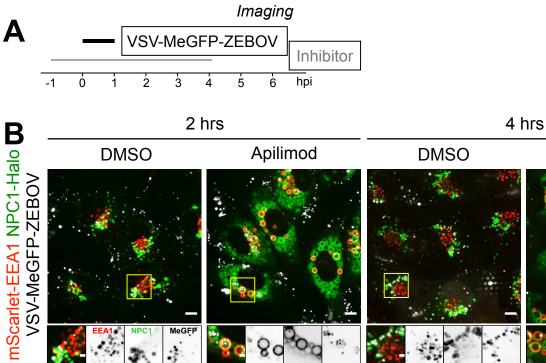




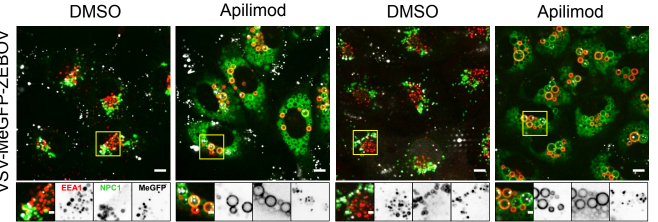




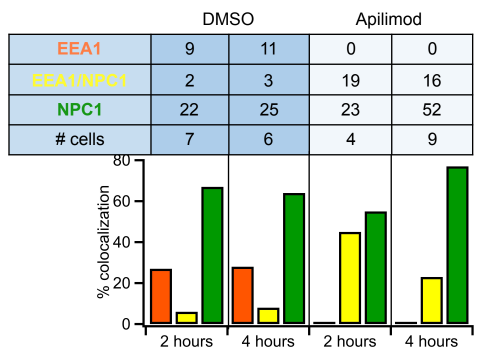


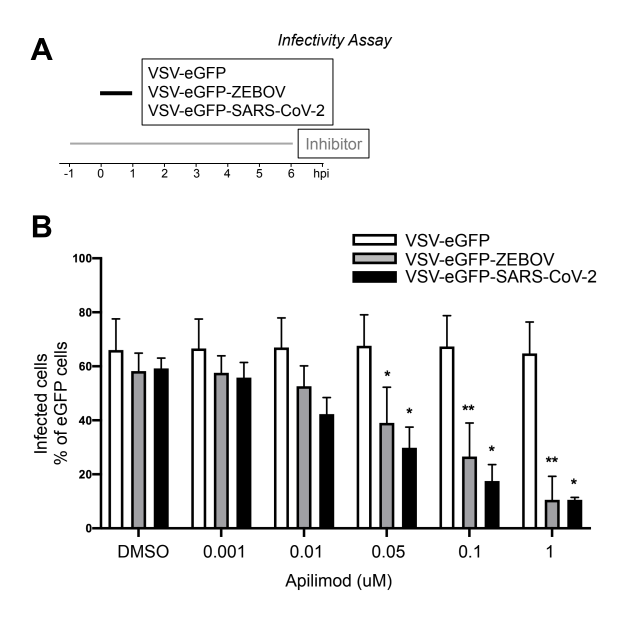


С



of endosomes containing VSV-MeGFP-ZEBOV





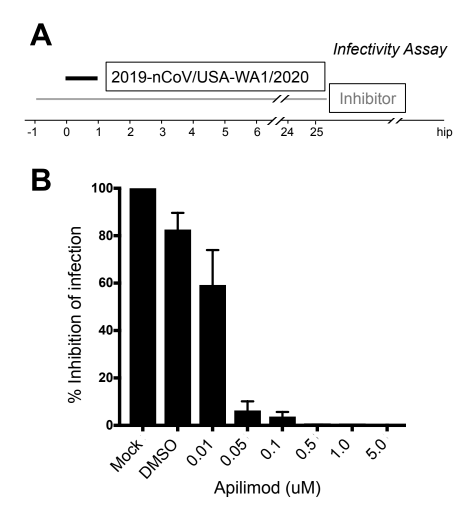


Table I

Primer Des	cription	Sequence		
	forward primer	5' GCCGGTACCTGAGGGCCTATTTCCC 3'		
U6 promoter		5'ACCTCTAGAAAAAAGCACCGACTCGGTGCCACTTTTTCAAGTTGATAACGGACTAGCCTTATTTTAACTTGCTATTTCTAGCTCTAAAACN		
	reverse primer	NNNNNNNNNNNNNNNNCGGTGTTTCGTCCTTTCCACAAG 3		
	Rab5c	GACCCGCCATTGCCCGTCCA		
target sequence for	Rab7a	TCAAACTAAAGGGGGAAAAG		
reverse primer	NPC1	TAAATTTCTAGCCCTCTCGC		
	EEA1	GGTGGTGGTTGAACCATG		
	F1	gaattcgagctcggtacccGAGAGAACTAGGGAAGAAGAAGAACGA		
	R1	TGCCCGTCCAGCTGTAGTG		
Rab5c	F2	CCACTACAGCTGGACGGGCAatggtgtctaagggcgaagagc		
Raboc	R2	GGAACCACCAGAACCACCAGAA		
	F3	GGTTCTGGTGGTTCTGGTGGTTCCCTGGCGGGGCGGGGGG		
	R3	gtcgactctagaggatccccCCTCCTACCAAGAGAGTAGAGAAAG		
	F1	gaattcgagctcggtacccACTGCTGTCAGCCTTGCCTTCA		
	R1	CCTTCAAACTAAAGGGGGAAAAGG		
Rab7a	F2	CCTTTTCCCCCCTTTAGTTTGAAGGatggtgtctaagggcgaag		
Kab/a	R2	GGAACCACCAGAACCACCAGAA		
	F3	TTCTGGTGGTTCTGGTGGTTCCACCTCTAGGAAGAAGTGTTGCTG		
	R3	gtcgactctagaggatccccCCTCACCCAACCTACCACAGAAT		
	F1	gaattcgagctcggtacccCCACTGAGATGAAGGAGTCCAT		
	R1	GAAATTTAGAAGCCGTTCGCGC		
NPC1	F2	CGCGAACGGCTTCTAAATTTCggaggttctggtggttctggtggttccGCAGAAATCGGTACTGGCTTTCCA		
NPCI	R2	GCCGGAAATCTCGAGCGTCGACAG		
	F3	CTGTCGACGCTCGAGATTTCCGGCtagccctctcgcagggcatcc		
	R3	gtcgactctagaggatcccccGCTGTCTAATGAAACTTCTAGGTC		
	F1	gaattcgagctcggtacccCTCTTTGGCTGAAATTAGAAGCAGG		
	R1	CATGGTTTAACCACCGGCG		
5544	F2	CGCCGGGTGGTGGTAAACCATGgtgagcaaggcgaggcgagtgat		
EEA1	R2	cttgtacagctcgtccatgccgc		
	F3	GCGGCATGGACGAGCTGTACAAGggaggttctggtggttctggtggttccTTAAGGAGGATTTTACAGAGGGTAAGAG		
	R3	gtcgactctagaggatccccGCTCTAATCTTTCTATCCTCAAGGTTTTC		

Table II

escription	Sequence
ing forward	GAGCCTGAAGTTGGGAGACC
ing reverse	CATGCCCACTCACCTCCAAT
ing forward	GCGGTCACTTCTTTGAGAAAGT
ing reverse	AAGTGGCAGCACGGACAGTGT
ing forward	TCTCCAAAAGAGAGGGAGAGAGAGAT
ing reverse	AAGTTTAGTGTCCTGTGGTTGCCT
ing forward	CATCTGTCAGTTACGGGGGCTG
ing reverse	CGGCACCACACCCTCCAGCTC
	ing forward ing reverse ing forward ing reverse ing forward ing reverse ing forward ing reverse