Reorganization of F-actin nanostructures is required for the late phases of SARS-CoV-2 replication in pulmonary cells.

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17 Abstract

The severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) is worldwide the 18 main cause of the COVID-19 pandemic. After infection of human pulmonary cells, 19 intracellular viral replication take place in different cellular compartments resulting in the 20 destruction of the host cells and causing severe respiratory diseases. Although cellular 21 trafficking of SARS-CoV-2 have been explored, little is known about the role of the 22 23 cytoskeleton during viral replication in pulmonary cells. Here we show that SARS-CoV-2 infection induces dramatic changes of F-actin nanostructures overtime. Ring-like actin 24 nanostructures are surrounding viral intracellular organelles, suggesting a functional 25 interplay between F-actin and viral M clusters during particle assembly. Filopodia-like 26 structures loaded with viruses to neighbour cells suggest these structures as mechanism for 27 cell-to-cell virus transmission. Strikingly, gene expression profile analysis and PKN 28 inhibitor treatments of infected pulmonary cells reveal a major role of alpha-actinins 29 superfamily proteins in SARS-CoV-2 replication. Overall, our results highlight cell actors 30 required for SARS-CoV2 replication that are promises for antiviral targets. 31

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Teaser : Impairing regulation of actin filaments inhibits SARS-CoV-2 particle production in human pulmonary cells.

Keywords: SARS-CoV-2; F-actin nanostructures; alpha-actinins; Transcriptomic; STED microscopy

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39 Introduction

40 SARS-CoV-2 is worldwide a major public health burden as the main cause of the still 41 ongoing COVID-19 pandemic with almost 244 million confirmed cases in 190 countries 42 and more than 5 million deaths until now (1). SARS-CoV-2 infects mainly human 43 pulmonary cells destroying the target cells and causing severe respiratory diseases with 44 excessive inflammation capable of inducing respiratory failure, multi-organ failure and 45 death (2). SARS-CoV-2 is an enveloped virus with a positive sense, single-stranded RNA 46 genome and belongs to the Beta coronavirus family (3,4,5). After SARS-CoV-2 infection

of target cells, intracellular viral replication take place consisting of a series of complex 47 processes (e.g., viral RNA translation, particle packaging, assembly and release) that are 48 tightly orchestrated to one another and often mutually exclusive (reviewed in 6). Viral 49 translation often takes place first in order to create a stock of viral proteins that will serve 50 to assemble the newly made viral particles. The SARS-CoV-2 transcriptome consists of a 51 long unspliced genomic RNA and 9 sub-genomic RNAs that are generated by alternative 52 splicing. After viral RNA translation, once the structural nucleocapsid protein N is 53 produced in the cytosol of infected cells, SARS-CoV-2 assembly continues with the 54 interactions of the N proteins with the unspliced genomic RNA. These interactions lead to 55 a ribonucleoprotein complex that will assemble at the membrane of the Endoplasmic 56 Reticulum – Golgi intermediate Compartment (ERGIC) with the structural proteins 57 transmembrane (M), envelop protein (E) and spike protein (S) (7,8,9). The viral particles, 58 ranging between 90 and 200 nm as recently described (7,8,10,11), will bud from the 59 ERGIC and egress through the secretory pathway. A number of studies over the years 60 have shown that most viruses hijack the cytoskeletal network to fulfill their own 61 replication cycle, which motivated us to perform a detail study on the role of the 62 cytoskeleton during SARS-CoV-2 infection in human host pulmonary cells (12,13,14). 63

The cytoskeleton is a complex and dynamic network of protein filaments in the cytoplasm 64 of the cells, extending from the cell nucleus to the cell membrane. Its primary function is 65 to give the cell its morphology and mechanical resistance to deformation (15). In addition, 66 the cytoskeleton has been related to many different cellular processes including cell 67 migration, cell signalling, endocytosis, cell division, chromosome segregation, 68 intracellular transport, etc (12, 16,17, 18). It can also build specialized structures, such as 69 flagella, cilia, lamellipodia and podosomes. In eukaryotic cells, the cytoskeleton consists 70 of three main components: microfilaments, intermediate filaments and microtubules, and 71 all these components are rapidly growing or disassembling depending on the requirements 72 of the cell. Microfilaments are mainly composed of linear polymers of G-actin proteins. 73 The G-actin monomer combines to form a polymer which continues to form the actin 74 filament. These actin filament subunits assemble into two chains that intertwine into 75 nanostructures called F-actin chains or fibers (15). F-actin fibers generate force when the 76 growing end of the actin filaments push against a barrier, such as the cell membrane. They 77 also act as tracks for the movement of myosin molecules that affix to the microfilament 78 and "move" along them. Interestingly, the network of F-actin fibers under the plasma 79 membrane can be a carrier for virus entry or transfer from one cell to other (12,17). 80 Although mechanisms of trafficking implicated in SARS-CoV-2 infection have been 81 explored mostly in simian Vero cell lines (19,20), little is known about the participation of 82 F-actin nanostructures during SARS-CoV-2 replication in human pulmonary cells. Here, 83 we implemented confocal and super-resolution 2D and 3D STED microscopy (21) to 84 study the kinetics of M cluster formation during SARS-CoV-2 particle assembly and 85 release, as well as the effects of SARS-CoV-2 infection on the morphology of human 86 pulmonary cells as consequence of intracellular rearrangements of F-actin nanostructures. 87 The human pulmonary A549-hACE2 cells implemented here are a well-established 88 experimental model for SARS-CoV-2 research due to their high susceptibility to SARS-89 CoV-2 infection, which can be explained by the stably overexpression of the host receptor 90 protein for the viral S protein (human angiotensin-converting enzyme 2, hACE2) and the 91 92 presence of the the co-receptor (human transmembrane protease serine 2, TMPRSS2) (22). Our results demonstrate that the kinetics of M cluster formation during SARS-CoV-2 93 particle assembly and release correlate with rearrangements of intracellular F-actin fibers 94 and morphological changes of SARS-CoV-2-infected human pulmonary A549-hACE2 95

- cells. Moreover, we show that the reorganization of F-actin nanostructures is required for 96 97
 - SARS-CoV-2 replication in human pulmonary A549-hACE2 cells.

98 99 **Results**

M cluster formation during SARS-CoV-2 assembly correlates with changes of F-100 actin nanostructures and cell morphology 101

To investigate the kinetics of SARS-CoV-2 assembly, we monitored viral M clusters 102 formation at different time points upon SARS-CoV-2 infection (6 to 77 h post-infection, 103 pi) of human pulmonary A549-hACE2 cells by immunofluorescence confocal microscopy 104 using an anti-COV-2 membrane protein (α M) antibody (Fig. 1A-B, Supplementary Fig. 105 1A-B and 2). Indeed, human pulmonary A549-hACE2 cells were highly susceptible to 106 SARS-CoV-2 infection (Supplementary Fig. 1 and 2). Further, size quantification of the 107 intracellular M clusters per cell at different time points pi (Fig. 1C, left, and 108 Supplementary Fig. 1B) showed an increase of intracellular M cluster size from a mean of 109 0.6 μ m² (interquartile range, IQR -0.1 -3 μ m²) at 24h pi to a median of 1.46 μ m² 110 $(IQR=0.4-15 \ \mu m^2)$ at 48h pi. Interestingly, the size of intracellular M clusters per cell 111 significantly decreased after 48h pi to 0.77 μ m2 (IQR=0.1-9 μ m²) at 54h pi, 0.72 μ m2 112 (IQR=0.1-8 μ m²) at 72h pi and 0.64 μ m2 (IQR=0.1-7 μ m²) at 77h pi. Remarkably, the 113 maximal area occupied by intracellular M clusters per cell was 15 μ m² at 48h pi, whereas 114 it ranged between 3 and 9 µm2 at the other time points analysed. Quantification of the 115 total intensity of viral M clusters per cell (Figure 1C, right) showed also a significant peak 116 at 48h pi (Mean= 8.06×10^7) when compared to the other time points analysed. Supporting 117 these results, RNA-sequencing (RNA-seq) based expression analysis in SARS-CoV-2-118 infected A549-hACE2 cells showed significant increase of all viral transcripts from 24h pi 119 to 48h pi (Supplementary Fig. 1C), suggesting an increase of unsliced viral genomic RNA 120 during SARS-CoV-2 assembly. Interestingly, we detected intracellular increase of the 121 viral structural proteins N, M and S from 24h pi to 72h pi (Supplementary Fig. 1D), as 122 well as a peak of viral particle release in the cell culture medium at 72h pi (Fig. 1D and 123 Supplementary Fig. 1E). Our results support that during the kinetic of SARS-CoV-2 124 infection of human pulmonary cells, intracellular SARS-CoV-2 assembly peaks at 48h pi, 125 whereas the release of SARS-CoV-2 particles peaks at 72h pi. 126

- We also monitored the effect of SARS-CoV-2 infection on the cytoskeleton of A549-127 hACE2 cells by confocal microscopy at different time points pi after labelling F-actin by 128 phalloidin stain (Figure 1E and Supplementary Fig. 2). We observed changes in the 129 distribution of F-actin in A549-hACE2 cells at different time points SARS-CoV-2 pi. E.g., 130 F-actin stress fibers were visible in non-infected control cells and disappeared 24h pi, 131 whereas other F-actin structures, such as elongated filopodia, appeared 24h pi. Further, 132 quantification of F-actin intensity implementing Z-stack projection images (Fig. 1F) 133 revealed significant increase of F-actin intensity from 999 a.u. at 0h pi to 1572 a.u. at 24h 134 135 pi and 1857 a.u. at 48h pi, whereas F-actin intensity decreased to 1184 a.u. at 72h pi. Interestingly, we did not detect significant changes in the total actin content of the cell by 136 western blot analysis (WB) of proteins extracts (Figure 1G). Our results indicate that 137 SARS-CoV-2 infection induced extensive reorganization of intracellular actin fibers 138 without significantly affecting total actin levels. 139
- Further analysis of the morphological changes that we observed in A549-hACE2 cells 140 upon SARS-CoV-2 infection (Fig. 2A-B, Supplementary Movies 1 and 2) revealed 141

significant increase of cell height from 2.96 µm (IOR=2.2 - 3.9 µm) at 0h pi to 3.75 µm 142 $(IQR=2.2-4.72 \ \mu m)$ at 24h pi and 4.89 μm (IQR= 3.4-6 μm) at 48h pi, whereas cell height 143 decreased to 3.29 µm (IOR=2.4-4.8 µm) at 72h pi. In contrast to cell height, cell surface 144 and cell volume showed the opposite effects upon SARS-CoV-2 infection with decreases 145 values above 24h pi (Fig. 2C-D), suggesting a contraction of the A549-hACE2 cells at this 146 time point. Summarizing, our results indicate that the kinetics of M cluster formation 147 during SARS-CoV-2 particle assembly and release correlate with rearrangements of 148 149 intracellular F-actin fibers and morphological changes of SARS-CoV-2-infected human pulmonary A549-hACE2 cells. 150

151 **Reorganization of F-actin nanostructures in SARS-CoV-2 infected pulmonary cells**

Since we detected a maximum of M cluster formation during SARS-CoV-2 assembly at 152 48h pi, we focused our further analysis on this specific time point and optimized 153 experimental conditions to achieve a higher resolution for imaging of F-actin structures 154 using super-resolution 2D STED microscopy in non-infected and SARS-CoV-2 infected 155 A549-hACE2 cells (Figure 3A), thereby increasing resolution below 70nm by 10 times as 156 compared to confocal microscopy and being able to observe dual colour structures. To 157 quantify F-actin rearrangements, we have analyzed the orientation angle of actin fibers 158 from STED images (as in 18). We clearly observed the parallel orientation of actin stress 159 fibers in non-infected cells, whereas actin stress fibers were not visible in SARS-CoV-2 160 infected cells at 48h pi (Fig. 3A). The color map of F-actin orientation and data for 161 distribution of orientation angle, suggested a significant rearrangement of F-actin network 162 with protrusion of filaments at the cell plasma membrane (Fig. 3A-B). The major possible 163 orientation angle of F-actin fibers in non-infected cells was significantly lower (around 90 164 and -90 degree) than in SARS-CoV-2 infected cells at 48h pi (Figure 2 B). We also 165 detected an increase in the random orientation angle after infection, probably due to F-166 actin rearrangement, distortion or reorganization. To investigate the potential formation of 167 intracellular self-organizing F-actin structures in SARS-CoV-2 infected A549-hACE2 168 cells, we monitored the organization of the F-actin cytoskeleton at 48h pi using super-169 resolution 2D microscopy and detected in the infected cells F-actin structures resembling 170 intracellular "actin rings" (Fig. 3C). The diameter of the observed "actin rings" ranged 171 between 0.5 and 2.5 µm with a mean of 1.03 µm and a standard deviation (STD) of 0.35 172 um (Fig. 3D, top). However, detection of viral M clusters around intracellular "actin 173 rings" by 2D STED microscopy was limited by signal saturation in Z direction. To bypass 174 these limitations, super-resolution 3D STED microscopy with 185 nm slice in Z direction 175 was implemented (Fig. 3E) and surprisingly revealed that viral M clusters formed similar 176 "ring-like" structures in close proximity to the "actin rings". Transmission electron 177 microscopy (TEM) slices of the infected cells also show vesicular structures, of 0.6 to 1 178 um size, full of viruses with particle budding events at the cell membranes (Fig.3E, TEM). 179 By STED 3D, the diameter of the "viral rings" ranged between 0.5 and 2 µm with a mean 180 of 0.95 µm and a STD of 0.27 µm (Fig. 3D, bottom). By superposing both F-actin and 181 virus STED images, it appears that M labelled intracellular organelles ("viral rings") are 182 surrounded by F-actin nanostructures ("actin rings") (Fig. 3E). Our results allow the 183 hypothesis of a spatial and functional interplay between F-actin nanostructures and M 184 cluster formation during assembly of SARS-CoV-2 particles, suggesting a stabilization of 185 the viral assembly platforms by F-actin or the need of F-actin for the transport of virus 186 loaded vesicles towards the cell plasma membrane. 187

188 SARS-CoV-2 infection induced actin filament rearrangement forming virus loaded 189 protrusions

Production of SARS-CoV-2 particles is a multistep process that occurs in different 190 compartments of an infected cell and culminates in the assembly of virus components at 191 the plasma membrane followed by budding and release of infectious virus particles. To 192 investigate the role of F-actin during virus particle formation and release, we used super-193 194 resolution 2D STED microscopy for quantitative analysis (Fig. 4A-C). The number of filopodia-like structures significantly increased from 2 per 10 µm of infected cell plasma 195 membrane (IQR=1-3) in non-infected A549-hACE2 cells to 10 per 10 µm of infected cell 196 plasma membrane (IQR=6-18) in SARS-CoV-2-infected cells 48h pi (Fig. 4B). We also 197 detected a significant increase in the maximum length of filopodia-like structures from 2-4 198 µM in non-infected cells to 10-12 µm in SARS-CoV-2-infected cells 48h pi (Fig. 4C). 199 200 Interestingly, we observed these structures being loaded with viruses. A quantitative analysis of the viral M clusters and individuals particles using 2D/3D STED images 201 revealed three different population of viral M clusters in 3 different cellular regions: 202 intracellular (see Fig. 3 D, E), particle release sites at the cell plasma membrane (Fig.4D, 203 zone 1) and viral particles at the filopodia of infected cells (Fig. 4D, zone 2). The size of 204 viral particles ranged in zone 2 from 70 nm to 350 nm with a mean of 209 nm and a STD 205 of 87 nm (Fig. 4D), in zone 1 from 150 nm to 1000 nm (Mean=478 nm; STD=216 nm; 206 Fig. 4D). These quantitative analyses suggested that viral particles are released in package 207 at the cell membrane and then particles migrate (possibly one or two together) on 208 filopodia-like structures. Strikingly, in some infected cells at close proximity, we also 209 observed an inter-connection between cells via filopodia-like structures loaded with 210 viruses, suggesting that virus-containing filopodia could be one of the possible 211 mechanisms for cell-to-cell SARS-CoV-2 infection spread (Fig. 4D). 212

213 SARS-CoV-2 infection enhances expression of actin cytoskeleton regulating genes

Our results suggest that F-actin rearrangements during viral particle assembly are 214 important for viral replication during SARS-CoV-2 infection progression. To gain further 215 insights, we performed a RNA-seq-based transcriptome analysis in non-infected and 216 SARS-CoV-2 infected A549-hACE2 cells at 48h pi (Fig. 5A and Supplementary Fig. 3A). 217 We detected increased levels after infection in 9.91% of the transcripts mapped to the 218 human genome (4376 transcripts with FC \geq 2), whereas the levels of 7.10% of the 219 transcripts mapped to the human genome were reduced (3136 transcripts with FC ≤ 0.3). 220 Gene set enrichment analysis (GSEA) based on Reactome [23] from the top 9.91% of the 221 genes with increased levels 48h pi (4376transcripts; Fig. 5B) revealed significant 222 enrichment of genes related to RHO GTPases activate PKNs (P=0.368) as the top item of 223 the ranked list. In addition, graphical representation of the enrichment profile (Fig. 5C) 224 showed a high enrichment score (ES) of 0.559 for RHO GTPases activate PKNs. Since 225 RHO GTPases are best known for their roles in regulating cytoskeletal rearrangements, we 226 227 monitored the transcript levels of various proteins related to the cytoskeleton (Fig. 5D) and detected increased transcript levels of proteins that are known to be regulated by RHO 228 GTPases, thereby being Alpha-actinin superfamily, and in particular Alpha-Actinins 2 and 229 3 (ACTN2, ACTN3), the ones with the most prominent increase. Type II myosins are also 230 reported (MHY7 and MHY6), suggesting together with ACTN2 and ACTN3 a possible 231 contractile activity of the infected cells (24), as previously suggested by our cell 232 233 morphology analysis (Fig. 2). In addition, 2D STED microscopy images and actin orientation analysis (Fig. 3A) revealed the formation of large actin fibers near the cell 234

plasma membrane. Furthermore, WB of protein extracts showed a 2-fold increase of
 alpha-actinins (ACTN) in SARS-CoV-2 infected cells 48h pi when compared to non infected cells (Fig. 5E).

238 PKN inhibitor reduced SARS-CoV-2 replication in human pulmonary cells

Following the line of ideas from our previous results (Fig. 5, 2 and 3), we investigated the 239 240 effect of the 2 inhibitors Rho/SFR and PKN inhibitors on SARS-CoV-2 replication, since they regulate actin fibers formation and alpha-actinins regulation, respectively. Data 241 reveals a reduction of SARS-CoV-2 replication in human pulmonary cells overtime and in 242 a dose-dependent manner with an IC50 of 1.36 µM and 0.65 µM for Rho/SFR and PKN 243 inhibitors, respectively (Fig. 6A, Supplementary Fig. 4). The LD50 being determined as 244 7.02 μ M for Rho/SFR inhibitor and 37.7 μ M for PKN inhibitor, indicating a selectivity 245 index > 10, thereby supporting that these are potent antiviral inhibitors for the 246 development of therapeutic strategies against SARS-CoV-2. Moreover, confocal 247 immunofluorescence images of human pulmonary A549-hACE2 cells infected with 248 SARS-CoV-2 in the presence of 0.5µM PKN inhibitor showed a restoration of cell 249 morphology and of F-actin structural pattern (Fig. 6B). Strikingly, PKN inhibitor 250 treatment of SARS-CoV-2-infected pulmonary cells reduced the size of intracellular viral 251 M clusters (Fig. 6C) and decreased the levels of viral particle release (Fig. 6D). We used 252 for these experiments remdesivir (IC50 equal 1 µM) as positive control [25], since it is an 253 antiviral drug that targets the virus replication complex reducing the number and size of 254 viral assembly M clusters (Fig. 6C) and decreasing the levels of viral particle release (Fig. 255 6D). Interestingly, PKN inhibition blocked M clusters in the ER (Supplementary Fig. 5), 256 thus probably at the level of virus assembly or virus egress from the ER, as shown by 257 immunofluorescence staining for M and grp78 ER marker (Supplementary Fig. 5) and 258 suggesting a role for the alpha-actinins superfamily proteins in SARS-CoV-2 assembly 259 and particle egress. 260

261 Discussion

SARS-CoV-2 is a recently discovered virus. Despite the fact that it is in the spotlight of 262 the scientific community worldwide for being the main cause of the COVID-19 pandemic. 263 the role of the cytoskeleton during SARS-CoV-2 replication has remained elusive. Here, 264 we implemented confocal and super-resolution 2D and 3D STED microscopy to analyse 265 the correlation between that host cell F-actin content and the kinetic of SARS-CoV-2 266 infection in human pulmonary cells: intracellular SARS-CoV-2 assembly and F-actin 267 content peak at 48h pi, accompanied with a cell morphology deformation and SARS-CoV-268 2 particle release peaking at 72h pi. The RNA-seq-based analysis of viral transcripts in 269 these infected human pulmonary cells (Supplementary Fig. 1C) correlated with the peak of 270 SARS-CoV-2 assembly at 48h pi. The apparent discrepancies between our SARS-CoV-2 271 transcriptome and the recently published SARS-CoV-2 transcriptome (26) could be 272 explained by differences in the experimental design, such as the implementation of 273 different cell lines (Calu-3, Caco-2 and Vero cells versus A549-hACE2) and higher viral 274 titters (MOI 1 and 0.1 versus 0.01), among others. The higher susceptibility to SARS-275 CoV-2 infection of the here implemented human pulmonary A549-hACE2 cells allowed 276 us to use lower viral titers during our RNA-seq experiment (MOI 0.01). More relevant 277 from the present study, the observed kinetics of M cluster formation during SARS-CoV-2 278 particle assembly and release correlated with rearrangements of cytoskeletal F-actin 279 nanostructures and morphological changes of SARS-CoV-2-infected A549-hACE2 cells. 280 Our results agree with previous reports demonstrating that actin rearrangements are 281

involved during the replication of various viruses targeting the respiratory tract, including 282 the respiratory syncytial virus (RSV) and influenza virus [27,28,29,30]. Further, the strong 283 cytopathic effect observed after SARS-CoV-2 infection could be related with the dramatic 284 changes in F-actin nanostructures and cell morphology. However, the expression of 285 caspases as apoptosis markers were reduced in SARS-CoV-2 infected cells at 48h pi, 286 supporting the viability of the cells analyzed here (Supplementary Fig. 3). It is well known 287 that actin polymerization has a role in replication of influenza viruses (31). For RSV, actin 288 289 was not completely proven to be involved in assembly, rather actin was playing a role in virus spread driven filopodia induction through Arp2/3 complexes [28]. In our study, 290 increase in F-actin content during infection, with a global actin content remaining quite 291 constant (Figure 1 G), included a redistribution of F-actin polymerization into new 292 nanostructures that appears quite crucial for the late phases of viral replication. 293 Remarkably, we found that SARS-CoV-2 infection induces ring-like F-actin 294 295 nanostructures surrounding also ring-like viral M-containing structures, suggesting the formation of large intracellular viral organelles, in which SARS-CoV-2 particle assembly 296 take place. This interpretation of our results is in agreement with earlier discoveries, 297 showing that large intracellular structures at the ERGIC containing SARS-CoV-2 298 structural proteins (M, N, E and S proteins) together with viral genomic RNA and driving 299 the assembly of new viral particles (7,8). Strikingly, we also found that SARS-CoV-2 300 infection promote filopodia-like structures loaded with viruses to neighbour cells, 301 suggesting these structures as mechanism for cell-to-cell SARS-CoV-2 infection spread. 302 Consistent with these findings, viral cell-to-cell transmission has been reported for other 303 RNA enveloped viruses, such as the human immunodeficiency virus 1 (HIV-1) (32). 304 Summarizing, all these results support a spatial and functional interplay between F-actin 305 nanostructures and M cluster formation during assembly of SARS-CoV-2 particles, which 306 needs to be further investigated. 307

- Actin polymerization starts with the formation of a small aggregate consisting of three 308 actin monomers. Actin filaments are then able to grow by the reversible addition of 309 monomers to both ends. However, one end (the plus end) growth up to ten times faster 310 than the minus end. The polymerisation of actin and the reorganization of actin filaments 311 are complex processes regulated by different factors. One of these factors is the protein 312 kinase N (PKN), which is a fatty acid-activated serine/threonine kinase, whose catalytic 313 domain exhibits homology with that of the protein kinase C family. It has been reported 314 that the interaction of PKN with alpha-actinins is promoted by phosphatidylinositol 4,5-315 bisphosphate (33), suggesting that PKN/alpha-actinin complexes locate at the cell plasma 316 membrane to promote cortical actin fiber rearrangement (34). Alpha-actinins belong to the 317 spectrin gene superfamily which represents a diverse group of cytoskeletal proteins. 318 Alpha-actinins are F-actin-crosslinking proteins found in various subcellular localizations 319 both in muscle and non-muscle cells (35), and being involved in diverse cellular 320 processes. Besides the involvement of alpha-actinins regulating cortical actin dynamics 321 during HIV-1 entry (36), the role of alpha-actinins during replication of virus has 322 remained largely unknown. Here, we showed that SARS-CoV-2 infection increased levels 323 of ACTN2 and ACTN3 transcripts of ACTN proteins. Furthermore, interfering with alpha-324 actinins function through PKN inhibitor treatment in SARS-CoV-2 infected cells restored 325 F-actin structures and reduced SARS-CoV-2 replication. Overall, our results reveal that F-326 actin nanostrucutres and F-actin rearrangement are required for SARS-CoV-2 replication 327 in pulmonary host cells and support the idea of using PKN inhibitors for the development 328 of therapeutic approaches against SARS-CoV-2 infection. 329
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331 Materials and Methods

Cell culture and infection

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Human pulmonary Alveolar A549-hACE2 cells were obtained from original A549 334 (ECACC) transduced with a lentiviral vector expressing human ACE2 receptor 335 (manufactured by FlashTherapeutics company, Toulouse, France) and sorted by cytometry 336 for having more than 80% hACE2 on their surface. The sorted A549-hACE2 cells were 337 maintained in RPMI supplemented with 10% heat inactivated fetal bovine serum (FBS), 338 339 1% sodium Pyruvate, 0.5% HEPES and antibiotics (penicillin/Streptavidin) and cultivated at 37°C with 5% CO₂. For virus production, VeroE6 cells were obtained from (ECACC) 340 and maintained in Dulbecco's minimal essential medium (DMEM) supplemented with 341 10% heat inactivated fetal bovine serum (FBS) at 37°C with 5% CO₂. 342

The strain BetaCoV/France/IDF0372/2020, was supplied by the National Reference 343 Center for Respiratory Viruses hosted by Institut Pasteur (Paris, France) and headed by Pr. 344 345 **Sylvie** van der Werf. The human sample from which strain BetaCoV/France/IDF0372/2020 was isolated has been provided by Dr. X. Lescure and Pr. 346 Yazdanpanah from the Bichat Hospital, Paris, France. Y. Moreover, 347 the BetaCoV/France/IDF0372/2020 strain was supplied through the European Virus Archive 348 goes Global (EVAg) platform, a project that has received funding from the European 349 Union's Horizon 2020 research and innovation program under the grant agreement No 350 653316. COV-2 Virus was propagated in VeroE6 cells with DMEM containing 2.5% FBS 351 at 37°C with 5% CO₂ and harvested 72 hours post inoculation. Virus stocks were stored at 352 -80°C and titer using plaque assays as previously described (11). 353

Quantitative reverse transcription polymerase chain reaction (qRT-PCR)

RNAs from mock infected or infected (MOI=0.01) A549-hACE2 cell culture supernatant were extracted using the Nucleospin Dx Virus RNA purification kit (Macherey-Nagel). Then qRT-PCR was performed in triplicate as described²⁰, using primers targeting the E gene of SARS-CoV-2 (E_Sarbeco-Forward ACAGGTACGTTAATAGTTAATAGCGT; E_Sarbeco-Reverse ATATTGCAGCAGTACGCACAA) and Luna Universal One-Step qRT-PCR Kit (New England Biolabs) on a Roche Light Cycler 480. The calibration of the assay was performed with a nCoV-E-Sarbeco-Control Plasmid (Eurofins Genomics).

RNA sequencing and data analysis.

RNA was sequenced as previously described [PMID 31110176; 29867223]. Briefly, total 365 RNA from non-infected (Ctrl) or SARS-CoV-2 infected A549-hACE2 and hPCLS was 366 isolated using Trizol (Invitrogen). RNA was treated with DNase (DNase-Free DNase Set, 367 Qiagen) and repurified using the miRNeasy micro plus Kit (Qiagen). Total RNA and 368 library integrity were verified on LabChip Gx Touch 24 (Perkin Elmer). One ug of total 369 RNA was used as input for SMARTer Stranded Total RNA Sample Prep Kit-HI 370 Mammalian (Clontech). Sequencing was performed on the NextSeq500 instrument 371 (Illumina) using v2 chemistry with 1x75bp single end setup. Raw reads were visualized by 372 FastQC to determine the quality of the sequencing. Trimming was performed using 373 the trimmomatic with following parameters LEADING:3 374 TRAILING:3 SLIDINGWINDOW:4:15 HEADCROP:4, MINLEN:4. High quality reads were mapped 375 using with HISAT2 v2.1.0 with reads corresponding to the transcript with default 376 parameters. RNA-seq reads were mapped to human genome hg19. After mapping, Tag 377 libraries were obtained with MakeTaglibrary from HOMER (default setting). Samples 378 were quantified by using analyzeRepeats.pl with the parameters (hg19 -count genes – 379 rpkm; reads per kilobase per millions mapped). 380

RNA-seq based expression analysis of viral transcripts.

Fastq files from infected A549-hACE2 cells after 24h, and 48h were used as input; for 383 each timepoint we used 2 replicates. Read trimming was performed using trimmomatic (v 384 0.39) with the following parameters "ILLUMINACLIP: all adapters v0.38.fa:2:30:10 385 AVGQUAL:30 LEADING:0 TRAILING:0 SLIDINGWINDOW:6:30 MINLEN:38". 386 Trimmed reads where then aligned to the SARS-CoV-2 reference genome 387 388 NC 045512.2.fasta (downloaded May 2021 from https://www.ncbi.nlm.nih.gov/nuccore/NC 045512) using the STAR (v 2.7.9a) aligner; 389 **STAR** parameters where the following "--outFilterType BySJout 390 -outFilterMultimapNmax 20 --alignSJoverhangMin 8 --outSJfilterOverhangMin 12 12 12 391 12 --outSJfilterCountUniqueMin 1 1 1 1 --outSJfilterCountTotalMin 1 1 1 1 --392 0 outSJfilterDistToOtherSJmin 0 0 0 --outFilterMismatchNmax 999 393 394 outFilterMismatchNoverReadLmax 0.04 --scoreGapNoncan -4 --scoreGapATAC -4 -chimOutType WithinBAM HardClip --chimScoreJunctionNonGTAG 0 --alignIntronMin 395 --alignIntronMax 1000000 --alignMatesGapMax 20 1000000 396 alignSJstitchMismatchNmax -1 -1 -1 -1". Samtools (v 1.12) was used to handle the 397 coverage was used to count reads in each viral feature alignments, and bedtools 398 (gene) genomic coordinates using the from 399 GCF_009858895.2_ASM985889v3_genomic.gff (downloaded May 2021 from 400 https://ftp.ncbi.nlm.nih.gov/genomes/all/GCF/009/858/895/GCF_009858895.2_ASM9858 401 89v3/GCF_009858895.2_ASM985889v3_genomic.gff.gz) only for protein 402 coding features. Feature counts were transformed to reads per kilobase million (RPKM) we 403 calculated mean RPKM from the duplicates for each feature and then calculated a Fold 404 Change as mean RPKM at 48h / mean RPKM at 24h; this data handling and plotting was 405 performed using R. 406

Western Blot analysis

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A549-hACE2 cells were infected for 2 hours with SARS-CoV-2 (MOI = 0.01). At 409 different time point (6h, 24h, 48h, 54h, 72h and 77h) post infection (pi), cells were washed 410 twice in PBS, detached with Versen (0.1M EDTA), pelleted at 250g for 6min and lysed in 411 RIPA buffer for western blot analysis. Total protein concentration was calculated using a 412 Bradford protein assay kit (ThermoFisher). 20ug of total cell lysates were diluted in 413 Laemmli buffer and proteins were separated by SDS-PAGE on 8% (for COV-2 S- and N 414 proteins) and 12% (for COV-2 M protein) acrylamide gels. Gels were transferred to PVDF 415 membrane using wet transfer with Tris-glycine-methanol buffer. Membranes were washed 416 in TBS, blocked with 5% milk in TBS-Tween 0.1% for 30min and incubated overnight at 417 4°C with primary antibodies against the spike S protein (Gentex, cat# GTX632604), N-418 protein (Gentex, cat# GTX632269) or M-protein (Tebu, cat# 039100-401-A55), all three 419 diluted at 1:1000 in TBS-T. After washing with 5% milk in TBS-Tween, the membranes 420 were incubated with HRP conjugated anti-mouse antibodies for N and S protein, and with 421 HRP conjugated anti-rabbit antibody for M protein and alpha-actinins for 2h at room 422 temperature, then washed in TBS-Tween buffer, incubated with ECL reagent (Amersham 423 cat#RPN2236) and imaged using a Chemidoc Imager (Biorad). 424

426Immuno-fluorescence confocal and 2D/3D STED super-resolution microscopy

A549-hACE2 cells seeded on glass coverslips were infected with SARS-CoV-2 at a MOI=0.1 or MOI=0.01 (low multiplicity of infection). At different time interval from 6h to 77h post-infection cells were washed with PBS and fixed in 4% paraformaldehyde in PBS for 15 minutes at room temperature, followed by permeabilization with 0.2% Triton

X-100 in PBS for 4-5 minutes and blocking in 2% BSA in PBS for 15 min. Incubation 431 with primary antibodies anti-SARS-CoV2 rabbit membrane (M) protein (1:100) was 432 performed for 2 hours at room temperature. After washing with PBS, cells were incubated 433 with secondary antibodies AF568-labeled goat-anti-rabbit (1:200) and Star orange for high 134 resolution STED imaging (1:100) as well as AF488-labeled Phalloidin and Star red 435 phalloidin (1:100) (for high resolution STED microscopy) for 2 hours at room 436 temperature. We have used mounting media prolong gold antifade reagent with DAPI and 437 prolong gold antifade reagent without DAPI for confocal and STED microscopy 438 respectively. Confocal fluorescence images were generated using a LSM800 confocal 439 laser-scanning microscope (Zeiss) equipped with a 63X, 1.4 NA oil objective and STED 440 2D and 3D measurements were performed on the Abberior Instrument Expert Line STED 441 super-resolution microscope (Abberior Instruments GmbH, Göttingen, Germany) using 442 Star orange 580 and Star red pulsed excitation laser sources with a pulsed STED laser 443 144 operating at 775 nm. For STED 2D (25% laser) lateral resolution was 67nm and for STED 3D (30% laser) resolution was 185nm in Z. All the images processed with ImageJ/Fiji. For 445 3D-reconstruction of confocal images, cells were fixed and stained as indicated and 446 imaged as z stack with 0.3 µm sections. Z stack was processed using ImageJ/Fiji, Imaris 147 viewer. 448

Electron Microcopy

A549-hACE2 pulmonary cells infected with SARS-CoV-2 were fixed with 2.5% (v/v) 451 glutaraldehyde in PHEM buffer and post fixed in osmium tetroxide 1% / K₄Fe(CN)₆ 0,8%, 452 at room temperature for 1h for each treatment. The samples were then dehydrated in 453 successive ethanol bathes (50/70/90/100%) and infiltrated with propylene oxide/ 154 EMbed812 mixes before embedding. 70 nm ultrathin cuts were made on a PTXL 455 ultramicrotome (RMC,France), stained with OTE/lead citrate and observed on a Tecnai 456 G2 F20 (200kV, FEG) TEM at the Electron Microscopy Facility COMET, INM, Platform 457 Montpellier RIO Imaging, Biocampus, Montpellier. 458

Statistical analysis

Statistical tests were performed using Origin 2021 software. Statistically significant 461 analysis was evaluated using one \Box way ANOVA tests. **p < 0.05. Cell area and cell 462 volume and height were calculated using 3D viewer plugin from Fiji image J. The 463 orientation angle properties of a given region of interest in an image were computed based 464 on the evaluation of the structure tensor in a local neighborhood using the Java plug-in for 465 ImageJ/Fiji (http://imagej.nih.gov/) 'OrientationJ'. The Materials and Methods section 466 should provide sufficient information to allow replication of the results. Begin with a 467 section titled Experimental Design describing the objectives and design of the study as 468 well as prespecified components. 469

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Author contributions: PM, DB performed cell culture, BSL3 infection, viral stock amplification and titer, viral RNA extraction and qRT-PCR, immunoblots. DB participated to the TEM. JS performed immunofluorescence sample preparation, confocal and STED 2D and 3D Microscopy and quantitative analysis. KR performed RNA extraction and sequencing. KR, IA, SG and GB performed RNA sequencing and analyzed RNA-seq data. DM, JS, GB and KR were involved in manuscript writing. JS, GB and DM conceptualized the study, edited the figures and wrote the manuscript. DM supervised the study. DM and GB raised funding for the study.

- **Competing interests:** Authors declare that they have no competing interests.
- **Data and materials availability:**
- All data are available in the main text or the supplementary materials.

- **Figures and Tables**

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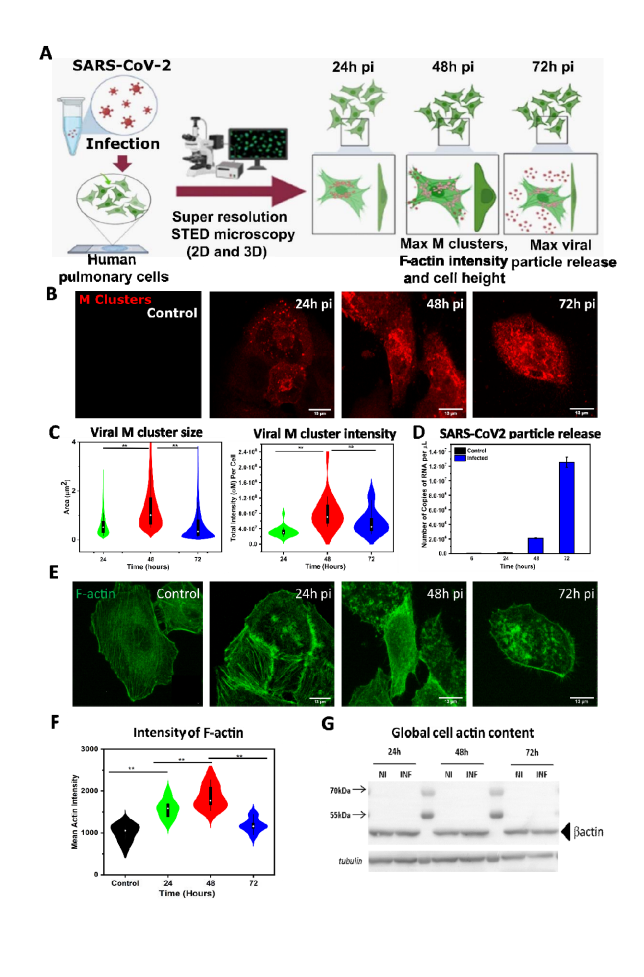
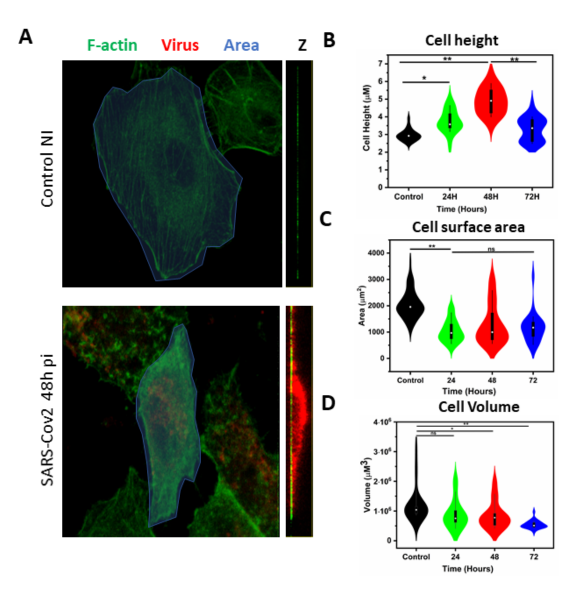




Fig 1. Increase in F-actin content correlated with M clusters of human pulmonary cells upon SARS-CoV-2 infection

Imaging and quantitative analysis of time course changes in viral M clusters area, 535 morphology and mean F-actin intensity of SARS-CoV-2 infected A549-hACE2 cells. 536 A549-hACE2 cells were fixed at 0h, 24h, 48h, and 72h post infection and processed for 537 immunofluorescence. SARS membrane protein anti-M rabbit antibody and a secondary 538 antibody Alexa Fluor 568 (in red) and for F-actin Phalloidin Alexa Fluor 488 were used 539 for confocal microscopy. (A) Schematic representation and confocal images of viral 540 clusters and F-actin with different time post infection 0h to 72h. (B) Images for changes in 541 viral M clusters size at different time post infection. (C) Plot for viral M clusters size and 542 viral M clusters intensity at different time post infection. (D) Plot for Number of copies of 543 RNA/µL in the supernatant of infected cells with different time post infection. (E) Images 544 for changes in F-actin intensity per cell at different time post infection. (F) Plot for mean 545 F-actin intensity with or without infection at different time of post infection. (G) Western 546 blot data for global actin content of non-infected and infected cell at different time point of 547 infection. All F-actin intensity of infected and non-infected cells are calculated from Z-548 projection images. A number of 20 < n < 50 cells were analyzed from at least 3 549 independent experiments. Statistical significant analysis were evaluated using one way 550 ANOVA tests. ***p* < 0.05. 551

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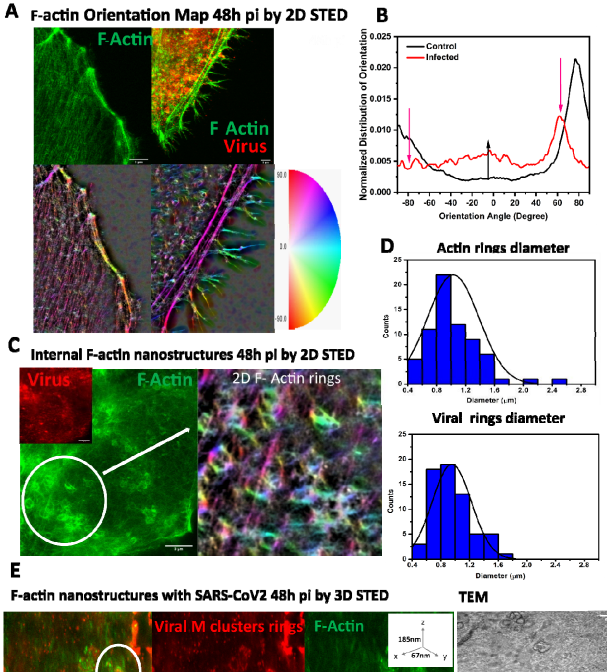
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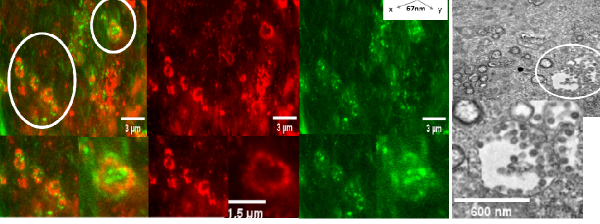
554 Fig 2. Morphological changes of human pulmonary cells upon SARS-CoV-2 infection

Imaging and quantitative analysis of time course changes in morphology of SARS-CoV-2 555 infected A549-hACE2 cells. A549-hACE2 cells were fixed at 0h, 24h, 48h, and 72h post 556 infection and processed for immunofluorescence. SARS membrane protein anti-M rabbit 557 antibody and a secondary antibody Alexa Fluor 568 (in red) and for F-actin Phalloidin 558 Alexa Fluor 488 were used for confocal microscopy. (A) Confocal images of Control and 559 Infected A549-hACE2 cells (48h post infection. (B) Plot for cell height. (C) Plot for 560 surface area of the cell. (D) plot for cell volume. All cell surface area, height, volume of 561 infected and non-infected cells are calculated from actin and M protein spreading at XY 562 and Z direction at different time of post infection. A number of 20 < n < 50 cells were 563 analyzed from at least 3 independent experiments. Statistically significant analysis was 564 evaluated using one \Box way ANOVA tests. **p < 0.05. 565

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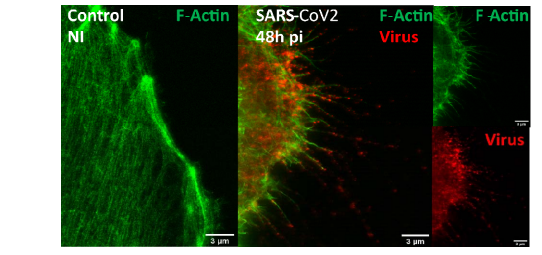
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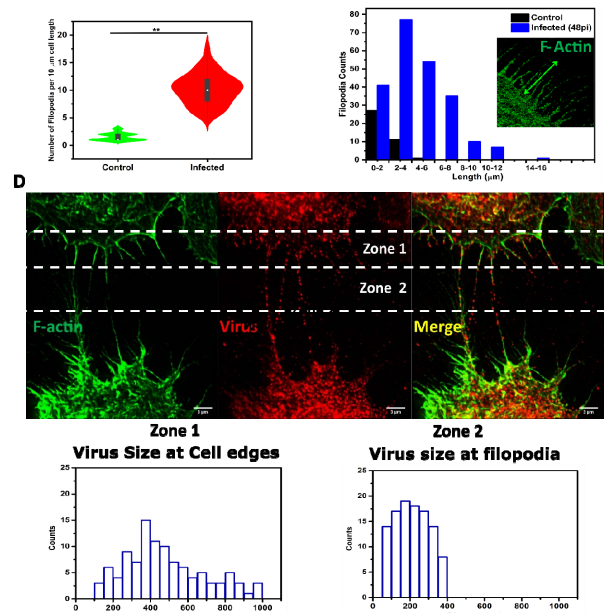
Fig 3. Reorganization of F-actin nanostructures and intracellular actin ring formation in SARS-CoV-2 infected pulmonary cells

STED 2D and 3D images of changes in F-actin and viral clusters of SARS-CoV-2 infected 574 pulmonary cells. A549-hACE2 cells were fixed for control and 48h post infection 575 processed for immunofluorescence coupled to STED microscopy. For imaging virus 576 SARS membrane protein anti-M rabbit antibody and then secondary antibody Star orange 577 (green) and for F-actin Phalloidin Star red (red) were used. (A) 2D STED images and 578 color representation of orientation angle of F-actin network with (48h post infection) or 579 without infection. (B) Plot for distribution of orientation angle, with (48h post infection) 580 or without infection. (C) STED 2D images of actin ring with color representation of rings. 581 (D) Plot for distribution of F-Actin rings and viral ring diameters. (E) STED 3D images of 582 intracellular F-actin rings and viral M clusters in infected pulmonary cells. A number of 583 25 < n < 50 cells were analyzed from at least 3 independent experiments. Statistical 584 significant analysis were evaluated using one \Box way ANOVA tests. **p < 0.05. 585 Transmission electron microscopy (TEM) images of intracellular structures filled with 586 budding viruses in SARS-CoV-2 infected pulmonary cells. Scale bars are 500 to 600nm 587 for the zoom image, as indicated. 588

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Diameter (nm)

Diameter (nm)

Α

591 Fig 4. Reorganization of actin fibers into virus loaded filopodia-like protrusions at the cell surface of SARS-CoV-2 infected pulmonary cells. 592

STED 2D images and Quantitative data of changes in F-actin nanostructures in SARS-593 CoV-2 infected A549-hACE2 cells. A549-hACE2 cells were fixed for control and 48h 594 post infection processed for immunostaining and STED microscopy. For imaging viral 595 particles and clustered SARS membrane protein anti-M rabbit antibody and then 596 secondary antibody Star orange (green) and for F- actin Phalloidin Star red (red) were 597 used. (A) Merge STED 2D Images of control and 48h post infected pulmonary cells. (B) 598 Plot for distribution of number of filopodia per 10 micrometer lengths of each cell infected 599 and control. (C) Plot for distribution of length of individual filopodia in control and 700 infected cells. (D) 3D projection Image and plots showing viral M cluster size at cell edge 701 (Zone 1), at filopodia-like structures (Zone 2), and at cell-to-cell connections. A number 702 of 15 < n < 20 cells were analyzed from at least 3 independent experiments. Statistical 703 704 significant analysis were evaluated using one \Box way ANOVA tests. **p < 0.05.





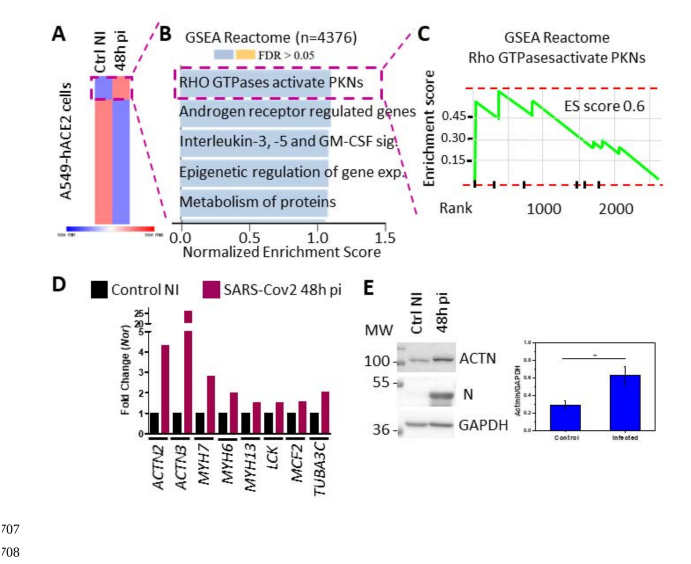
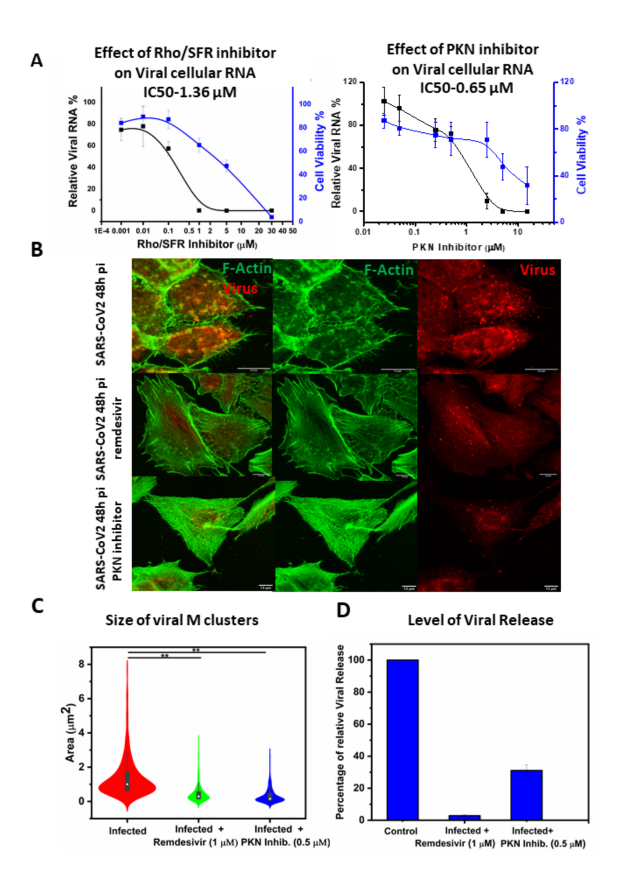


Fig 5. Cellular gene expression analysis of SARS-CoV-2 infected human pulmonary cells using RNAseq reveals an upregulation of alpha-Actinins

(A) Heat map showing RNA-seq-based expression analysis of differentially expressed 712 transcripts in non-infected (Ctrl) and A549hACE2 cells infected with SARS-CoV-2 for 713 48h (48h pi). *n*=44141 differentially expressed transcripts; 2 individual cell replicates per 714 715 condition. (B) Top. Reactome-based Gene Set Enrichment Analysis (GSEA) of candidates with FC ≥ 2 (*n*=4376 upregulated transcripts) using WebGestalt (WEB-based Gene SeT 716 AnaLysis Toolkit, 2019). Bottom. Panther-based Gene Set Enrichment Analysis (GSEA) 717 of candidates with FC ≤ 0.3 (*n*=3136 downregulated transcripts) using WebGestalt (WEB-718 based Gene SeT AnaLysis Toolkit, 2019). FDR: False Discovery Rate. (C) Reactome-719 based Gene Set Enrichment Analysis (GSEA) for Rho GTPAses pathway of candidates 720 with FC≥2. (D) Histogram plots representing the basal transcription activity (48h pi 721 normalized to Control) of components of the Rho-GTPase pathway that are differentially 722 expressed in Control and SARS-CoV-2 infected A549hACE2 cells. 723



726	Fig 6. Reduction of SARS-CoV-2 replication in pulmonary cells upon PKN and Rho/SFR
727	inhibitors treatment accompanied with cellular F-actin and cell shape restoration
728	(A) Dose effect of Rho/SFR and PKN inhibitors on SARS-CoV-2 replication in pulmonary
729	cells using qRT-PCR and cell viability. (B) Confocal images of changes SARS-CoV-2
730	infected A549-hACE2 cells with the treatment of Remdesivir (1µM) or PKN Inhibitor
731	(0.5 µM). (C) Plot for changes in viral clustered size with or without (infected) treatment
732	of Remdesivir (1µM) or PKN Inhibitor (0.5 µM). (D) Plot for Number of copies of
733	RNA/µL in the supernatant of infected cells at 48h post infection. A549-hACE2 cells were
734	fixed at 48h post infection and processed for immunofluorescence and laser confocal light
735	microscopy using a SARS-CoV-2 membrane protein anti-M rabbit antibody and a
736	secondary antibody Alexa Fluor 568 (in red) and for F-actin imaging Phalloidin Alexa
737	flour 488 (Green) were used. A number of $15 < n < 20$ cells were analyzed from at least 3
738	independent experiments. Statistical significant analysis were evaluated using one way
739	ANOVA tests. ** $p < 0.05$.
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743	Supplementary Materials

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- see Supplementary Materials document.
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