1 The replication of the human respiratory syncytial virus in a T cell line has

- 2 multiple ineffective steps.
- 3 Ricardo de Souza Cardoso^{1,3}, Ana Carolina Lunardello Coelho¹, Bruna Laís Santos de
- 4 Jesus¹, Brenda Cristina Vitti¹, Juliano de Paula Souza¹, Rosa Maria Mendes Viana¹,
- 5 Marjorie C. Pontelli¹, Tomoyuki Murakami³, Armando Moraes Ventura², Akira Ono^{3*},
- 6 Eurico Arruda^{1*}.
- 7 1 Department of Cell and Molecular Biology, School of Medicine of Ribeirao Preto,
- 8 University of Sao Paulo, Brazil
- 9 2 Institute of Biosciences, University of Sao Paulo, Brazil.
- 10 3 Department of Microbiology and Immunology, University of Michigan Medical
- 11 School, Ann Arbor, MI, USA.
- ¹² *Contributed equally to this study.

13 Abstract

14 Human respiratory syncytial virus is the most frequent cause of severe respiratory 15 disease in children. The main targets of HRSV infection are epithelial cells of the 16 respiratory tract and the great majority of the studies regarding HRSV infection are done 17 in respiratory cells. Recently, the interest on respiratory virus infection of lymphoid cells 18 has been growing, but details of the interaction of HRSV with lymphoid cells remain 19 unknown. Therefore, this study was done to assess the relationship of HRSV with A3.01 cells, a CD4⁺ T cell line. We found by flow cytometry and fluorescent focus assay that 20 A3.01 cells are susceptible but virtually not permissive to HRSV infection. De-quenching 21

22 experiments revealed that the fusion process of HRSV in A3.01 cells is reduced in comparison to HEp-2 cells, an epithelial cell lineage. Quantification of viral RNA by 23 gPCR determined that the replication of HRSV in A3.01 cells was modest. Western blot 24 and quantitative flow cytometry analyses demonstrated that the production of HRSV 25 proteins in A3.01 was significantly lower than in HEp-2 cells. Additionally, we found by 26 27 fluorescence in situ hybridization that the inclusion body-associated granules (IBAG's) are almost absent in HRSV inclusion bodies in A3.01 cells. We also assessed the 28 intracellular trafficking of HRSV proteins and found that HRSV proteins co-localized 29 30 partially with the secretory pathway in A3.01 cells, but these HRSV proteins and viral filaments are present only scarcely at the plasma membrane. HRSV infection of A3.01 31 CD4⁺ T cells is virtually unproductive as compared to HEp-2 cells, with virion production 32 hampered by low fusion, hypofunctional inclusion bodies, altered trafficking of viral 33 proteins to the plasma membrane. 34

35

36 Introduction

Human respiratory syncytial virus (HRSV), of the family *Pneumoviridae*, is a common 37 respiratory pathogen that circulates worldwide and a major cause of serious lower 38 respiratory tract disease, mainly bronchiolitis in children, but also severe disease in the 39 elderly [1]. HRSV infects mainly epithelial cells of the respiratory tract [1], but has also 40 been detected in non-respiratory tissues and cells [2][3]. In that regard, HRSV and other 41 respiratory viruses have been detected in tonsillar tissues and respiratory secretions 42 from children with tonsillar hypertrophy without symptoms of acute respiratory infection 43 [4], suggesting that HRSV may infect secondary lymphoid tissues. In addition, HRSV 44

45 antigen has been detected in circulating T lymphocytes, and thus it is conceivable that
46 it may affect immune function [5].

47 The HRSV genome is a single-stranded RNA with 10 genes that encode 11 proteins 48 [1]. Virus entry is mediated by the G protein that binds to the host cell [6], resulting in the fusion between viral envelope and cell membrane, mediated by the HRSV F protein 49 50 [7,8]. It is also known that HRSV enters cells by macropinocytosis, a process in which 51 nucleolin participates as virus receptor [9][10]. HRSV replication involves production of inclusion bodies [11], where viral proteins L, P, N, M, and M1,2 are present [12-14]. 52 53 However, the subsequent HRSV assembly process is not entirely understood. It has been shown that the viral glycoproteins follow the secretory pathway [15–18] to reach 54 the plasma membrane but also utilize Apical Recycling Endosome (ARE) machinery 55 56 during this process [19]. In contrast, the trafficking of HRSV non-glycosylated proteins during virus assembly has remained guite unclear. It is known that HRSV M protein 57 forms dimers prior to participating in the formation of viral progeny [20], and that this 58 protein has affinity to endomembrane system [21]. HRSV N protein coats the virus 59 genome and forms inclusion bodies, an essential role in which it is helped by the P 60 61 protein [22]. The inclusion bodies (IB's) of the HRSV are places where the viral replication and transcription occur; in addition, the IB's are responsible for stabilizing the 62 viral mRNAs that confers an efficient protein translation process [14]. HRSV N protein 63 64 was also observed to co-localize partially with the Golgi [23]. Consistent with these findings, recently we have shown that in HEp-2 cells the M and N engage partially with 65 secretory pathway and with the retromer complex [24]. Furthermore, HRSV P protein 66 67 was found partially co-localizing with endosomal vesicles [25]. Together, these findings

68	suggest the possibility that the non-glycosylated proteins of the HRSV reach the
69	assembly sites at least partially using secretory and/or endosomal pathways. The steps
70	of viral assembly and budding take place at the plasma membrane of the infected cells
71	[21,26,27]. It is known that HRSV F protein cytoplasmic tail is pivotal in the virus
72	budding process [7,28]. Finally, HRSV budding process results in a filamentous viral
73	particle, an event dependent on Rab11-FIP2 protein but not Vps4 [29].
74	Knowledge on HRSV-cell interactions has been accumulating through studies that use
75	models based on epithelial cell cultures susceptible and permissive to HRSV infection.
76	Nonetheless, virus-host cell interactions may be different between cells of epithelial
77	origin like HEp-2, in which virus progeny production results in cell death, and
78	lymphomononuclear cells, in which HRSV may cause long-term or persistent infection.
79	Previous studies have shown that HRSV infects $CD4^{+}T$ cells [5] and Breg cells of
80	neonates [30], which may enhance HRSV disease. Very little is known about HRSV
81	replication and progeny production in lymphoid cells, and hence the present study was
82	done to investigate intracellular HRSV assembly and replication in A3.01 cells, which
83	belong to the lymphoid cell lineage.

84

85 Material and Methods

86

87 Cells and viruses: A3.01 cells, obtained from the AIDS Research Reagent Program,

88 were maintained in Rosswell Park Memorial Institute (RPMI) culture medium,

supplemented with 10% of fetal bovine serum (FBS) and 1% of antimycotic/antibiotic.

HEp-2 and Vero cells were acquired from ATCC and maintained in MEM with 10% of
BFS in 1% of antimycotic/antibiotic. The virus used for this study was HRSV A long
strain (ATCC), propagated in HEp-2 cells and titrated in Vero cells, following routine
agar-based plaque assays. The multiplicity of infection (MOI) used for the infections are
specified in the experiment results.

95

Antibodies. The antibodies used were FITC-conjugated mouse monoclonal anti-RSV N 96 (MAB 858 3F, Millipore), mouse monoclonal anti-RSV F (MAB8262X, Millipore), mouse 97 anti-CD63 (Clone H5C6 RUO, BD Pharmingen), mouse polyclonal anti-RSV M [12], 98 rabbit polyclonal antibodies anti-TGN46 (ABT95, Millipore), anti-Giantin (PRB-114C-99 200, Covance), anti-SNX2 [31], anti-Lamp-1 D2D11 XPR (9091T, Cell Signaling); 100 mouse monoclonal anti-EEA1 (610457-BD), goat anti-RSV (Abcam ab20745); mouse 101 monoclonal anti-β-actin (sc 47778), Alexa Fluor 594-conjugated goat anti-rabbit 102 103 (ab150080, Abcam), Alexa Fluor 647-conjugated goat anti-mouse (ab150115, Abcam), HRP-conjugated goat anti-rabbit (656120 Invitrogen), HRP-conjugated rabbit anti-goat 104 (305 035 003, Jackson ImmunoResearch), and HRP-conjugated rabbit anti-mouse 105 106 (A9044, Sigma).

107

Flow Cytometry. HEp-2 or A3.01 cells were infected with HRSV, and the harvest occurred 24, 48, and 72 hours post infection, when cells were fixed with 4% PFA for 20 minutes. After that, cells were permeabilized and incubated with FITC-labeled mouse anti-RSV Nucleoprotein (Millipore), followed by washes with PBS containing 1% BSA. 112 The cells were analyzed in a BD LSR Fortessa flow cytometer: the results were analyzed in the Flow Jo software. The gating strategy used to perform the single cell 113 analysis for N-positive population to establish the mean fluorescence intensity was the 114 115 2D plot, in which the plans were fluorescence signal for N versus forward scatter. Experiments were done three times independently. 116 117 Biobond and Poly-Lysine coverslips treatment: Coverslips were treated with 0.1 N 118 HCl in 100% ethanol, then in 100% alcohol, air-dried, and then treated with 4% Biobond 119 120 (Koch Electron microscopy) in acetone for 4 minutes, followed by a wash in distilled water and air-drying. The same procedure was used for treatment with Poly-Lysine, 121 except for 20 µL of poly-lysine was used for 1 hour and then coverslips were air-dried. T 122 cells were deposited onto pre-treated coverslips and incubated stationary for 3 hours, 123 followed by fixation in 4% PFA for 20 minutes, washing, in PBB 1X and testing by 124 immunofluorescence or fluorescent in situ hybridization (FISH). 125 126 Immunofluorescence: After fixation, A3.01 or HEp-2 cells on coverslips were 127 permeabilized in 0.01% Triton X-100 in PBS for 15 minutes, and washed five times in 128

1X PBS. The cell preparations were incubated with the appropriate primary antibody for
1 h in a humidified chamber at 37°C. After that, preparations were washed five times in
PBS, incubated with the appropriate secondary antibody along with DAPI (Sigma) for 1

h, and finally washed in PBS. The coverslips were mounted on slides with

¹³³ Flouromounttm and analyzed by confocal microscopy in a Leica SP5 or a Zeiss 780

microscope. To perform analysis of quantity and size of the inclusion bodies in A3.01
and HEp-2 cells, the immunofluorescence images were subjected to Image J software
analysis. Using the tool Analyze>Analyze Particles, it was possible to determine the
area and number of the N-positive structures. The same procedure was done for
measuring the size of the A3.01 cells, but in this case, after the analysis, the mean of
the area of the A3.01 cells was established by conventional mathematic method.

Fluorescent in situ hybridization. HEp-2 or A3.01 cells were seeded on coverslips 140 treated or not with Poly-Lysine, and 24 and 48 hours post-infection were fixed with 4% 141 142 PFA and tested by FISH as per published protocol [14], slightly modified. Briefly, fixed cells were permeabilized with 0.1% Triton X-100 in 1% BSA-PBS and then treated with 143 free streptavidin for 1 h, re-fixed in 4% PFA, washed in PBS, incubated with the 144 hybridization mix (1µM biotinylated poly T probe, 35% formamide, 5% dextran sulfate, 145 herring sperm DNA in 2 X SSC2) at 37°C for 3 h, followed by serial washes in 2 X SSC, 146 SSC, and PBS. After that, coverslips were incubated with Alexa Fluor 594 streptavidin, 147 and then with FITC-conjugated mouse anti-RSV N (Millipore) for 1 h. Finally, the cells 148 were washed in PBS and mounted with a mounting solution containing DAPI prior to 149 150 observation using a fluorescent microscope. For counting the Inclusion bodies containing IBAGs, and the IBAGs within inclusion bodies in A3.01 and HEp-2 cells, the 151 immunofluorescence images from three independent experiments were subjected to a 152 153 manual counting. It was considered IBAGs all the clear distinct granular structures within the inclusion bodies. 154

155

156 Real-Time PCR. HEp-2 and A3.01 cells inoculated with HRSV for 1 h at 4°C, were centrifuged at 200 x g, and part of the cells and supernatant were placed into Trizol 157 reagent, which was considered as the time point zero. The remaining cells were 158 159 incubated in CO₂ at 37°C, and aliquots of cells and supernatant were collected in Trizol 160 reagent at 1, 3, 6, 12 and 24 hours post-infection. The total RNA was isolated following the Trizol protocol. Afterward, cDNAs from total RNAs were obtained by reverse 161 transcription with Superscript III Reverse Transcriptase (Invitrogen) primed with random 162 primers. The real-time PCR assays were performed using primers targeting the gene for 163 164 the N protein of HRSV. RNA quantification was extrapolated from a standard curve made with dilutions of plasmid containing the cloned RSV N gene segment. 165

166

R18 Fluorescence conjugation and dequenching assay. These protocols were the 167 same used by Evelyn M. Covés-Datson et al [32] with slight modifications. HRSV- or 168 169 culture supernants of HEp-2 cells were incubated with 30 µg/mL of octadecyl rhodamine 170 B chloride (R18) for 1 hour at room temperature protected from light. Then, fluorescently 171 labeled R18-HRSV or culture supernatants of HEp-2 were separated from excess R18 172 by a separation column (PD-10 Desalting Columns GE Healthcare). HRSV-R18 or culture supernatants of HEp-2 were incubated in suspension with A3.01 and HEp-2 173 cells at 4°C for 1 h to allow virus attachment, and then incubated in a CO₂ at 37°C, and 174 the dequenching of R18 was measured in a Synergy Photometer at the appropriate time 175 points. As positive control, 1% triton X-100 in PBS was used for total dequenching. The 176 177 amount of fluorescence emitted from R18 was measured at 560 nm excitation and 590 nm emission. 178

179 Fluorescent Focus Assay. A3.01 cells were infected with HRSV, and at 1, 6, 12, 24 and 48 hours post-infection, their supernatants were collected to investigate the 180 progeny production of the HRSV in these cells. Serial 10-fold dilutions of the 181 supernatants collected at each time point was made. Then, a HEp-2 cell monolayer was 182 infected with each of the HRSV dilutions. Three days after the infection, the cells were 183 permeabilized with 0,01% of 100X Triton and incubated with an antibody made in 184 mouse anti-HRSV F protein conjugated with Alexa Fluor 488. The fluorescent focus 185 were analyzed in a fluorescent microscope; the number of focus were counted and 186 187 plotted in a graph. 188 Results 189 190 A3.01 lymphocytes inoculated with HRSV are inefficient in progeny production. 191

The A3.01 cells were infected in suspension with HRSV (MOI=1) and analyzed by flow-192 193 cytometry at several times after infection (figure 1A). Infection was reproducibly 194 detected in three independent experiments at all times post infection, with peak at 48 hpi, when over 40% of the cells were infected. The highest numbers of cells positive for 195 196 HRSV N protein were found at 48 hpi (figure 1A). These results indicate that at least 197 under the condition used in these experiments, A3.01 cells are susceptible to HRSV infection and produce viral protein N. A3.01 cells were permissive for HRSV replication; 198 however, virus replication in this cell type was markedly reduced and delayed in 199 200 comparison with HEp-2 cells, as indicated by quantification of the HRSV genome

released into the culture supernatants (figure 1B). The fluorescent focus assay with
 mouse anti-HRSV F antibody indicated that infectious HRSV progeny production in
 A3.01 is also inefficient, with a replicative burst of less than one log₁₀ from 6 to 48 hours
 post-infection (figure 1C).

205

HRSV genome replication in A3.01 cells is inefficient.

Since we observed that A3.01 cells were inefficient in progeny production of HRSV, we 207 208 sought to investigate which step of the viral replicative cycle was compromised in these 209 cells. We set out to assess the virus genome production by quantitative real-time PCR 210 targeting the HRSV N gene in HRSV-infected A3.01 and HEp-2 cells, the latter of which 211 was cultured either as cells attached to plates (Att) on in suspension (Sus). Cells and viruses were incubated for absorption for 1 hour at 4° C, the inoculum was washed 212 away, and the cells were further incubated at 37°C. At several times post infection, cells 213 214 were collected and processed for RNA extraction in Trizol. The same amount of virus 215 inoculum was placed in virus-only wells, without cells, for quantification of the remaining virus inoculum after the incubation periods. The results revealed that the attachment of 216 HRSV to HEp-2 and A3.01 cells was only slightly different, but the virus replication was 217 almost 4 log₁₀ higher in HEp-2 than in A3.01 cells (figure 2). Furthermore, the amount of 218 cell-associated HRSV RNA decreased in A3.01 cells over 3 hours after inoculation, 219 220 suggesting a failure in a step after virus-cell attachment such as virus-cell fusion, which would result in degradation of internalized virions. 221

222

The fusion process of HRSV in A3.01 cells is defective.

224 The numbers of HRSV-positive A3.01 cells by flow cytometry were dependent on the 225 duration of the incubation period for virus adsorption to the cells (figure 3A). The fraction 226 of cells positive for HRSV N protein at 24 and 48 hpi was higher when the virus inoculum was not removed than when it was washed away after 3 hours. This is 227 228 consistent with the possibility that the entry of HRSV in A3.01 cells is not efficient. To test this possibility, both A3.01 and HEp-2 cell suspensions were inoculated with R18-229 labelled HRSV (R18-HRSV) or mixed with culture supernatants of mock-infected cells 230 231 containing an equivalent amount of R18, and the virus-cell fusion was evaluated based on dequenching of R18, which occurs when viral envelope fuses with cellular 232 membranes. We examined virus-cell fusion with $4x10^3$ cells per well. The virus or mock 233 inocula were incubated with cells on ice for 1 hour to allow adsorption but not entry, 234 after which the unbounded viruses were washed away. Subsequently, the virus-cell 235 suspensions were placed at 37°C, and the fluorescent emission by the R18 de-236 quenching process was measured. The results revealed that at 1, 3, 6, and 24 hpi, the 237 guantity of de-guenched R18 was significantly higher in HEp-2 than in A3.01 cells 238 239 (figure 3B). To test the possibility that the difference in the de-quenching of R18 between the two cell types is due to the difference in the amounts of attached virus, we 240 compared the attachment of R18-HRSV to HEp-2 and A3.01 cells using the same 241 242 approach. Both A3.01 and HEp-2 cells were treated with 1% Triton X-100 immediately after the washes with cold PBS, thereby completely de-guenching the attached viruses 243 244 (figure 3C). The results suggested that the amounts of HRSV attached in the A3.01 and

HEp-2 cells are similar (figure 3C). Altogether, we conclude that the process of HRSV
fusion to A3.01 cells is significantly compromised in comparison to HEp-2 cells.

247

248 HRSV protein production in A3.01 cells is small.

249 Even though the inefficient fusion process of the HRSV in A3.01 cells by itself could 250 explain the dramatic differences between genome replication in A3.01 and HEp-2, we 251 also hypothesized that other steps of the HRSV production could also be affected. 252 Since the replication of HRSV in A3.01 cells was inefficient, we examined whether the 253 production of N protein, a main component of virus factories, was also compromised in 254 A3.01 cells. Mean fluorescence intensity analysis of infected HEp-2 and A3.01 cells at 255 different time points using flow cytometry (figure 4A), showed that the N protein level in each HRSV-infected A3.01 cell was significantly lower than that in each HEp-2-infected 256 257 cell at all times. With the greatest difference at 48 hpi when the quantity of protein in 258 HEp-2 cells was approximately 42 times that found in A3.01 cells (figure 4B). Western 259 blot analysis of HRSV proteins produced in A3.01 or HEp-2 cells revealed that not only the N protein but also the other viral proteins were less abundant in infected A3.01 cells 260 at different times post-infection (figure 4C). Although the western blotting analysis is not 261 262 controlled for the number of infected cells, it revealed that the fold differences in M and P levels between HEp-2 and A3.01 are even greater than the difference in N. Therefore, 263 not only N, which was examined in the flow cytometry, but also other viral proteins are 264 present in A3.01 at lower levels than in HEp-2. In addition, it was not even possible to 265 266 see the bands corresponding to the HRSV G and F glycoproteins. These data indicate

that the HRSV protein levels at different times post-infection in A3.01 are broadly
 compromised.

269

270 HRSV inclusion body formation is compromised in A3.01.

271 The inclusion bodies (IB's) are important platforms for HRSV replication and assembly 272 [5,16,17]. Considering that the HRSV N, P and M are pivotal components of the IB 273 formation and that the levels of these proteins are very low in A3.01 cells, we expected 274 that the capacity of the HRSV to produce IBs in A3.01 cells is highly diminished. 275 Interestingly, immunofluorescence for HRSV N protein revealed that HRSV infection does induce inclusion body formation in A3.01 cells (figures 5B and C). However, by 276 277 measuring the perimeters of IBs in A3.01, we found that they were smaller than those in Hep-2 cells (figure 5G) and also present in significantly lower quantities in A3.01 278 compared to HEp-2 cells (figure 5H). Since the area of the HEp-2 cell is approximately 279 280 3.2 times larger than A3.01 cells, the difference in the area sizes can partially account for the differences found in the quantity of HRSV N inclusion bodies in HEp-2 and 281 A3.01. However, even when we normalized the quantity of HRSV N inclusions by the 282 area, the quantity of HRSV N-positive IBs was higher in HEp-2 than A3.01 cells (figure 283 5I). 284

285

The inclusion bodies of HRSV in A3.01 cells lack IBAGs.

287 Recently, Rincheval et al [14] showed that HRSV inclusion bodies are

compartmentalized and contain ultrastructural granules, which were called inclusion

289 body-associated granules (IBAGs). They also proposed that these structures are important for the IB's functionality, since this organization promotes stabilization of viral 290 transcripts and hence an efficient protein production process. We performed a FISH 291 292 analysis to examine the formation of IBAGs of HRSV-infected A3.01 cells. A biotinylated poly-T probe was used to reveal regions enriched for polyadenylated RNA. HRSV 293 produced IBAGs at both 24 and 48 hpi in HEp-2 cells (Figures 6I-L, and M-P), but in 294 A3.01 cells the majority of the HRSV-induced IBs did not appear to contain prominent 295 296 IBAG's and therefore can be considered hypofunctional (figures 6 Q-T and U-Z pointed by arrowheads). HRSV IBs in A3.01 cells did contain staining for polyA, but such polyA 297 signal was rarely found as clear distinct granules within the inclusion bodies as 298 observed in HEp-2 cells. A Z-stack and quantitative analyze was performed between 299 300 labelled HEp-2 (supplementary figure 1A) and A3.01 cells (supplementary figure 1B). The quantity of inclusion bodies of A3.01 and HEp-2 cells containing distinct IBAG's 301 within were counted (supplementary figure 1C). It showed that the quantity of inclusion 302 303 bodies of infected A3.01 cells containing IBAG's is significantly lower than those in infected HEp-2 cells (supplementary figure 1C). Further, the quantity of the IBAGs found 304 in inclusion bodies was significantly lower in A3.01 cells than in HEp-2 cells 305 (supplementary figure 1D). Therefore, we conclude that even when poly A-containing 306 mRNAs are present in the IBs of HRSV-infected A3.01 cells, the sequestration of such 307 308 mRNAs to subcompartments of IBs is inefficient in this cell type.

309

The HRSV F protein partially co-localizes with Golgi markers giantin and TGN46
 in A3.01 cells

312	In epithelial cells, F protein, a transmembrane protein, follows the conventional
313	anterograde pathway from ER through Golgi to the plasma membrane [36]. Also, the
314	Golgi participates in the intracellular traffic of the HRSV N protein to the plasma
315	membrane [23,24]. To examine whether the trafficking of HRSV F and N proteins in
316	A3.01 lymphocytes is similar to the one in HEp-2 cells, immunofluorescence staining
317	was done for a cis- and medial-Golgi marker, giantin, a trans-Golgi network marker,
318	TGN46, and the viral proteins F and N. A partial co-localization of HRSV F protein with
319	the Golgi markers was observed at 48 hours post-infection (Figure 7E-H). In HEp-2
320	cells, the HRSV F protein shows strong colocalization with Giantin and partial co-
321	localization with TGN46 (supplementary figures 2A-F). Even though the HRSV N protein
322	is a cytoplasmic and not transmembrane protein, a partial colocalization with TGN46
323	was also observed in HEp-2 cells (supplementary figures 3A-F). However, in A3.01
324	cells, the N protein did not co-localize with the Golgi markers (Figure 7 I-L). Therefore,
325	we conclude that the HRSV F and N proteins colocalized with the Golgi in A3.01 cells to
326	a lesser extent than in HEp-2 cells.

327

The HRSV F and N proteins partially co-localize with markers of endosomal
 pathway.

330

After trafficking through the Golgi stacks, the transmembrane proteins often associate with endosomal pathways [33–35]. In addition, it is already known that some of the HRSV proteins follow the endosomal system to reach the plasma membrane 334 [19,23,24,36]. To investigate whether the HRSV proteins also associate with endosomal machineries in A3.01 cells, immunofluorescence assays were performed for several 335 markers of this pathway. First, we aimed to investigate whether HRSV proteins were 336 337 preferentially targeted to late/lysosomal pathway in these cells. We assessed HRSV proteins co-localization with early endosome marker, the Early Endosome Antigen 1 338 (EEA1), and with a lysosome-specific marker, LAMP-1 (figure 8). Indeed, there was 339 strong co-localization of Lamp-1 signal with HRSV proteins in A3.01 cells (figures 8E-L) 340 when compared to HEp-2 (supplementary figures 2 and 3M-O, pointed by arrows), 341 suggesting an involvement of lysosomes in protein trafficking or degradation. However, 342 the imaging evidence is not enough to confirm specific involvement of lysosomal 343 machinery in HRSV protein degradation, since the co-localizations of HRSV proteins 344 with Lamp-1 and EEA1, an early endosome marker, were not statistically different in 345 A3.01 cells (figures 8M and N). We also did immunofluorescence microscopy for CD63 346 and Sorting Nexin 2 (SNX2). CD63 is a marker for late endosomes, while SNX2 is part 347 348 of the cellular retromer complex, responsible for carrying cargoes from early endosomes to the trans-Golgi [37,38]. In addition, recently, it was found that SNX2 is recruited to 349 HRSV N inclusion bodies and plays a role in the HRSV viral production in HEp-2 cells 350 351 [24]. It was possible to see that, similarly to what happens in HEp-2 cells, the HRSV F protein partially co-localized with SNX2 and CD63 in A3.01 (figures 9I to L, and 352 353 supplementary figures 2G to L). In addition, the HRSV N protein was found juxtaposed 354 to CD63 both in A3.01 and HEp-2 cells (figures 9I to L, and supplementary figures 3J to L, pointed by arrows). Interestingly, however, in contrast to HEp-2 cells, the HRSV N-355

positive structures failed to recruit SNX2 in A3.01 cells (figures 9E to H and

- 357 supplementary figures 3G to I).
- 358

The production of HRSV filaments at the plasma membrane of A3.01 cells is very
 low

HRSV assembly/budding takes place at the plasma membrane, with the appearance of
 viral protein-containing filament-shaped structures in infected epithelial cells [36,39].

363 Since the filaments emerging from the plasma membrane is one of the hallmarks of the

364 HRSV infection [1], we examined whether HRSV is capable of producing typical

365 filamentous structures in A3.01 cells. Immunofluorescence for viral proteins was done

with special attention to the plasma membrane. Although only scarcely, HRSV N and M

³⁶⁷ proteins could be seen in filament-shaped structures (figures 10A-C, arrowheads).

368 Nevertheless, the quantity of filament structures emerging from A3.01 cells is minimal

when compared to those found in HEp-2 cells (figures 10D-J). It is noteworthy that not

only the filament formation is reduced in A3.01 cells (figures 10A-C and H-I), the

quantity of the A3.01 cells displaying at least one filament is significantly lower than that

of HEp-2-infected cells (figure 10G). Therefore, while HRSV proteins were readily

detected in intracellular compartments of A3.01 cells, very little accumulation of HRSV

374 products was seen at the plasma membrane, which results in a fewer filament

375 production in these cells.

376

377 Discussion

378

379	Studies on the interaction of HRSV with host cells are usually conducted using
380	respiratory epithelial cells, which are the main targets of the natural infection that usually
381	results in cell death [1]. However, HRSV is also able to infect non-respiratory cells,
382	including CD4 ⁺ T lymphocytes [2,3,5], and may cause persistence in some cell types,
383	like murine macrophages [40]. The frequent detection of HRSV RNA in tonsillar tissues
384	from children without symptoms of acute HRSV infection [4] suggests that the agent
385	may cause prolonged infection in secondary lymphoid tissues. It is thus presumable that
386	HRSV causes patterns of infection that differ between lymphoid and epithelial cells,
387	which are likely to affect functions and survival of these cells. The present study was
388	undertaken to elucidate details of in vitro HRSV infection of the human $CD4^+$ T cell line
389	A3.01 in comparison to infection of HRSV in commonly used cell line, HEp-2. We
390	showed that HRSV infection in A3.01 cells is different from that in HEp-2 cells at
391	multiple steps of the virus replication cycle.
392	
393	While HRSV was able to infect A3.01 cells, the progeny production in these cells was
394	much lower than that of Hep-2 cells. This could be due to reduced susceptibility and/or
395	permissiveness of A3.01 cells to infection by HRSV. In that regard, we showed that
396	HRSV fusion with A3.01 cells was reduced as compared to the HEp-2 cells. This could
397	be due to some limitation of the HRSV fusion process itself or due to a difference in
398	HRSV-receptor engagement in A3.01 cells, creating subsequent hindrance to the fusion
399	process. It is noteworthy that HRSV can use different receptors to fuse to host cell
400	membranes [10,41,42]. Therefore, while binding of HRSV to HEp-2 and A3.01 cells

were comparable, differences in the levels of expression or affinity of individual
receptors between the two cell types may affect the efficiency of virus-cell fusion. It is
also worthwhile testing the activity of macropinocytosis in A3.01 cells versus HEp-2
cells, since this is the best known mechanism of HRSV internalization [9,43].

405

406 The replication of the HRSV genome occurs in IBs [1]. Recently Rincheval et al. demonstrated that the HRSV mRNAs are sequestered within the IBs, more specifically 407 in organized structures that they called inclusion bodies-associated granules or IBAGs 408 409 [14]. Using IF and FISH approaches in the present study, we found that HRSV IBs in A3.01 cells are significantly less abundant, smaller, and morphologically different than 410 those seen in Hep-2 cells and that most of the IBs that are formed in A3.01 cells lack 411 IBAGs. It is thought that in IBAGs, the M2-1 protein of the HRSV binds to viral mRNAs 412 and thereby makes them more stable, which consequently ensures a better protein 413 414 production [14]. Therefore, it is likely that the lack of discrete HRSV IBAGs in A3.01 cells at least partially accounts for the reduction in HRSV proteins N, M and P in A3.01 415 cells in comparison with HEp-2 cells at all times post infection (figure 4). HRSV proteins 416 N and P are integral parts of the RNA transcription complex and IBs [44–46]. Therefore, 417 lower levels of these proteins could help to explain significantly reduced rates of HRSV 418 genome production in A3.01 cells compared to the HEp-2 cells. Together, these findings 419 lead us to speculate that the low permissiveness to the HRSV genome replication in 420 A3.01 cells is a sum of defects in formation of IBs and IBAGs and protein translation, 421 422 where inefficiencies in each one of these steps exacerbate the other.

Since HRSV uses the secretory pathway to deliver viral proteins to the assembly sites 423 at the plasma membrane [1,15,17,19,23,24], we examined the presence of virus 424 proteins along the main components of the secretory and endosomal pathways by 425 immunofluorescence. Our results and previous literature have shown that HRSV F and 426 N proteins partially co-localized with markers of the secretory pathway [1,23,24]. 427 However, it was noteworthy that in contrast to HEp-2 cells, A3.01 cells did not display 428 an evident accumulation of SNX2 at places where the HRSV N protein is. In HEp-2 429 cells, SNX2 was found recruited to N-positive structures, and the knockdown of SNX2 430 431 impacted negatively in the HRSV production [24]. The absence of the recruitment of SNX2 to N-positive structures in A3.01 cells could contribute to the inefficiency in the 432 viral production in A3.01 cells. The co-localization of CD63 and Lamp-1 with HRSV 433 proteins is consistent with viral protein degradation, which could explain the lower 434 amounts of the HRSV proteins in A3.01 cells. However, more studies should be 435 performed to specifically assess this question, since it is currently unknown whether the 436 viral protein synthesis is impaired or whether they are degraded in this cell type. 437 Interestingly, even though the HRSV proteins partially co-localized with secretory 438 439 pathway markers in A3.01 cells, the number of the A3.01-infected cells displaying filaments at its surface was dramatically low. These results suggest that the trafficking 440 of viral components to the virus assembly sites is defective in A3.01 relative to HEp-2 441 442 cells. We do not rule out at this time that the absence of viral filaments at the cell surface is due to a defect in the assembly process, which may be caused by low viral 443

444 protein levels or by yet another block on the process at the plasma membrane.

Overall, the present results showed that HRSV infection of A3.01 CD4⁺ T cells is 445 virtually unproductive, with the meager virion production as compared to HEp-2 cells, 446 and that this is due to multiple defects during HRSV replication in A3.01 cells, namely, 447 low virus-cell fusion, formation of hypo-functional inclusion bodies lacking IBAGs, failure 448 to achieve high viral protein levels, and possibly altered trafficking of viral proteins and 449 genome to virus assembly sites at the plasma membrane. 450 451 References 452 453 454 1. Collins PL and RAK. Respiratory Syncytial Virus and Metapneumovirus. 6th ed. In: Knipe MD, Howley, MP, editors. Fields Virology, 6th Edition. 6th ed. 2013. pp. 455 1086-1123. 456 2. Rezaee F, Gibson LF, Piktel D, Othumpangat S, Piedimonte G. Respiratory 457 syncytial virus infection in human bone marrow stromal cells. Am J Respir Cell 458 459 Mol Biol. 2011. doi:10.1165/rcmb.2010-0121OC 460 3. Mills BG, Singer FR, Weiner LP, Holst PA. Immunohistological demonstration of 461 respiratory syncytial virus antigens in Paget disease of bone. Proc Natl Acad Sci U S A. 1981. doi:10.1073/pnas.78.2.1209 462 4. Proenca-Modena JL, Pereira Valera FC, Jacob MG, Buzatto GP, Saturno TH, 463 Lopes L, et al. High rates of detection of respiratory viruses in tonsillar tissues 464 from children with chronic adenotonsillar disease. PLoS One. 2012:7. 465 doi:10.1371/journal.pone.0042136 466 Raiden S, Sananez I, Remes-Lenicov F, Pandolfi J, Romero C, De Lillo L, et al. 467 5. Respiratory syncytial virus (RSV) infects CD4+ T cells: Frequency of circulating 468 CD4+ RSV+ T cells as a marker of disease severity in young children. Journal of 469 Infectious Diseases. 2017. doi:10.1093/infdis/jix070 470

- 6. Levine S, Kaliaber-Franco R, Paradiso PR. Demonstration that glycoprotein G is
- the attachment protein of respiratory syncytial virus. J Gen Virol. 1987.
- 473 doi:10.1099/0022-1317-68-9-2521
- 474 7. Feldman SA, Audet S, Beeler JA. The fusion glycoprotein of human respiratory
- syncytial virus facilitates virus attachment and infectivity via an interaction with
- 476 cellular heparan sulfate. J Virol. 2000;74: 6442–7. doi:10.1128/JVI.74.14.6442477 6447.2000.Updated
- Matthews JM, Young TF, Tucker SP, Mackay JP. The core of the respiratory
 syncytial virus fusion protein is a trimeric coiled coil. J Virol. 2000;74: 5911–5920.
 doi:10.1128/JVI.74.13.5911-5920.2000
- Krzyzaniak MA, Zumstein MT, Gerez JA, Picotti P, Helenius A. Host cell entry of
 respiratory syncytial virus involves macropinocytosis followed by proteolytic
 activation of the F protein. PLoS Pathog. 2013;9: e1003309.
- 484 doi:10.1371/journal.ppat.1003309
- Tayyari F, Marchant D, Moraes TJ, Duan W, Mastrangelo P, Hegele RG.
 Identification of nucleolin as a cellular receptor for human respiratory syncytial
 virus. Nat Med. 2011;17: 1132–1135. doi:10.1038/nm.2444
- 488 11. García J, García-Barreno B, Vivo A, Melero JA. Cytoplasmic inclusions of
 489 respiratory syncytial virus-infected cells: formation of inclusion bodies in
 490 transfected cells that coexpress the nucleoprotein, the phosphoprotein, and the
 491 22K protein. Virology. 1993;195: 243–7. doi:10.1006/viro.1993.1366
- 492 12. Carromeu C, Simabuco FM, Tamura RE, Farinha Arcieri LE, Ventura a M.
 493 Intracellular localization of human respiratory syncytial virus L protein. Arch Virol.
 494 2007;152: 2259–63. doi:10.1007/s00705-007-1048-4
- Lifland AW, Jung J, Alonas E, Zurla C, Crowe JE, Santangelo PJ. Human
 respiratory syncytial virus nucleoprotein and inclusion bodies antagonize the
 innate immune response mediated by MDA5 and MAVS. J Virol. 2012;86: 8245–
 58. doi:10.1128/JVI.00215-12

14. Rincheval V, Lelek M, Gault E, Bouillier C, Sitterlin D, Blouquit-Laye S, et al.

- 500 Functional organization of cytoplasmic inclusion bodies in cells infected by
- respiratory syncytial virus. Nat Commun. 2017;8. doi:10.1038/s41467-017-00655-
- 502

9

- 15. Brock SC, Heck JM, McGraw PA, Crowe JE. The transmembrane domain of the
 respiratory syncytial virus F protein is an orientation-independent apical plasma
 membrane sorting sequence. J Virol. 2005;79: 12528–35.
 doi:10.1128/JVI.79.19.12528-12535.2005
- Shaikh FY, Utley TJ, Craven RE, Rogers MC, Lapierre L a, Goldenring JR, et al.
 Respiratory syncytial virus assembles into structured filamentous virion particles
- independently of host cytoskeleton and related proteins. PLoS One. 2012;7:
- 510 e40826. doi:10.1371/journal.pone.0040826
- Anderson K, Stott EJ, Wertz GW. Intracellular processing of the human
 respiratory syncytial virus fusion glycoprotein: Amino acid substitutions affecting
 folding, transport and cleavage. J Gen Virol. 1992. doi:10.1099/0022-1317-73-5 1177
- Kwilas S, Liesman RM, Zhang L, Walsh E, Pickles RJ, Peeples ME. Respiratory
 Syncytial Virus Grown in Vero Cells Contains a Truncated Attachment Protein
 That Alters Its Infectivity and Dependence on Glycosaminoglycans. J Virol. 2009.
 doi:10.1128/jvi.00986-09
- Brock SC, Goldenring JR, Crowe JE. Apical recycling systems regulate directional
 budding of respiratory syncytial virus from polarized epithelial cells. Proc Natl
 Acad Sci U S A. 2003;100: 15143–8. doi:10.1073/pnas.2434327100
- 522 20. Förster A, Maertens GN, Farrell PJ, Bajorek M. Dimerization of Matrix protein is
 523 required for budding of Respiratory Syncytial Virus. J Virol. 2015;89: JVI.03500524 14. doi:10.1128/JVI.03500-14
- Henderson G, Murray J, Yeo RP. Sorting of the Respiratory Syncytial Virus Matrix
 Protein into Detergent-Resistant Structures Is Dependent on Cell-Surface

527 Expression of the Glycoproteins. Virology. 2002;300: 244–254.

- 528 doi:10.1006/viro.2002.1540
- 529 22. Galloux M, Gabiane G, Sourimant J, Richard C-A, England P, Moudjou M, et al.
- Identification and Characterization of the Binding Site of the Respiratory Syncytial
- 531 Virus Phosphoprotein to RNA-Free Nucleoprotein. J Virol. 2015.
- 532 doi:10.1128/jvi.03666-14
- 23. Céspedes PF, Bueno SM, Ramírez B a, Gomez RS, Riquelme S a, Palavecino
 CE, et al. Surface expression of the hRSV nucleoprotein impairs immunological
 synapse formation with T cells. Proc Natl Acad Sci U S A. 2014; 1–10.
 doi:10.1073/pnas.1400760111
- 537 24. Cardoso RS, Tavares LA, Jesus BLS, Criado MF, de Carvalho AN, Souza J de P,
 538 et al. Host retromer protein sorting nexin 2 interacts with human respiratory
 539 syncytial virus structural proteins and is required for efficient viral production.
 540 MBio. 2020. doi:10.1128/mBio.01869-20
- 541 25. Vanover D, Smith D V., Blanchard EL, Alonas E, Kirschman JL, Lifland AW, et al.
 542 RSV glycoprotein and genomic RNA dynamics reveal filament assembly prior to
 543 the plasma membrane. Nat Commun. 2017;8. doi:10.1038/s41467-017-00732-z
- San-Juan-Vergara H, Sampayo-Escobar V, Reyes N, Cha B, Pacheco-Lugo L,
 Wong T, et al. Cholesterol-Rich Microdomains as Docking Platforms for
 Respiratory Syncytial Virus in Normal Human Bronchial Epithelial Cells. J Virol.
 2012. doi:10.1128/jvi.06274-11
- Shaikh FY, Crowe JE. Molecular mechanisms driving respiratory syncytial virus
 assembly. Future Microbiol. 2013;8: 123–31. doi:10.2217/fmb.12.132
- Shaikh FY, Cox RG, Lifland AW. A Critical Phenylalanine Residue in the
 Respiratory Syncytial Virus Fusion Protein Cytoplasmic Tail Mediates Assembly
 of Internal Viral. 2012. doi:10.1128/mBio.00270-11.Editor
- Utley TJ, Ducharme NA, Varthakavi V, Shepherd BE, Santangelo PJ, Lindquist
 ME, et al. Respiratory syncytial virus uses a Vps4-independent budding

555 mechanism controlled by Rab11-FIP2. 2008;105: 10209–10214.

- 30. Zhivaki D, Lemoine S, Lim A, Morva A, Vidalain PO, Schandene L, et al.
- 557 Respiratory Syncytial Virus Infects Regulatory B Cells in Human Neonates via
- 558 Chemokine Receptor CX3CR1 and Promotes Lung Disease Severity. Immunity.
- 559 2017;46: 301–314. doi:10.1016/j.immuni.2017.01.010
- Amorim NA, Da Silva EML, De Castro RO, Da Silva-Januário ME, Mendonça LM,
 Bonifacino JS, et al. Interaction of HIV-1 nef protein with the host protein Alix
 promotes lysosomal targeting of cd4 receptor. J Biol Chem. 2014.
- 563 doi:10.1074/jbc.M114.560193
- S64 32. Covés-Datson EM, King SR, Legendre M, Gupta A, Chan SM, Gitlin E, et al. A
 molecularly engineered antiviral banana lectin inhibits fusion and is efficacious
 against influenza virus infection in vivo. Proc Natl Acad Sci U S A. 2020.
- 567 doi:10.1073/pnas.1915152117
- 33. Banting G, Maile R, Roquemore EP. The steady state distribution of humTGN46
 is not significantly altered in cells defective in clathrin-mediated endocytosis. J
 Cell Sci. 1998.
- S71 34. Chia PZC, Gunn P, Gleeson PA. Cargo trafficking between endosomes and the
 trans-Golgi network. Histochemistry and Cell Biology. 2013. pp. 307–315.
 doi:10.1007/s00418-013-1125-6
- 57435.Progida C, Bakke O. Bidirectional traffic between the Golgi and the endosomes -575machineries and regulation. Journal of Cell Science. 2016.
- 576 doi:10.1242/jcs.185702
- 577 36. Santangelo PJ, Bao G. Dynamics of filamentous viral RNPs prior to egress.
- 578 Nucleic Acids Res. 2007;35: 3602–11. doi:10.1093/nar/gkm246
- 579 37. Rojas R, Kametaka S, Haft CR, Bonifacino JS. Interchangeable but essential
- 580 functions of SNX1 and SNX2 in the association of retromer with endosomes and
- the trafficking of mannose 6-phosphate receptors. Mol Cell Biol. 2007;27: 1112–
- 582 24. doi:10.1128/MCB.00156-06

38. 583 Haft CR, Sierra M d. I. L. Bafford R, Lesniak MA, Barr VA, Taylor SI. Human Orthologs of Yeast Vacuolar Protein Sorting Proteins Vps26, 29, and 35: 584 585 Assembly into Multimeric Complexes. Mol Biol Cell. 2000. doi:10.1091/mbc.11.12.4105 586 Brown G, Rixon HWM, Sugrue RJ. Respiratory syncytial virus assembly occurs in 587 39. GM1-rich regions of the host-cell membrane and alters the cellular distribution of 588 tyrosine phosphorylated caveolin-1. J Gen Virol. 2002;83: 1841–50. Available: 589 http://www.ncbi.nlm.nih.gov/pubmed/12124448 590 Rivera-Toledo E, Gómez B. Respiratory syncytial virus persistence in 591 40. 592 macrophages alters the profile of cellular gene expression. Viruses, 2012. 593 doi:10.3390/v4123270 594 41. Kurt-Jones EA, Popova L, Kwinn L, Haynes LM, Jones LP, Tripp RA, et al. 595 Pattern recognition receptors TLR4 and CD14 mediate response to respiratory syncytial virus. Nat Immunol. 2000. doi:10.1038/80833 596 597 42. Johnson SM, McNally BA, Ioannidis I, Flano E, Teng MN, Oomens AG, et al. Respiratory Syncytial Virus Uses CX3CR1 as a Receptor on Primary Human 598 599 Airway Epithelial Cultures. PLoS Pathog. 2015. doi:10.1371/journal.ppat.1005318 600 43. M., Lingemann., T. M, X., Liu., U.J., Buchholz., S., Surman., S.E., Martin., P.L., 601 Collins., S., Munir. The alpha-1 subunit of the Na+,K+-ATPase (ATP1A1) is required for macropinocytic entry of respiratory syncytial virus (RSV) in human 602 603 respiratory epithelial cells. PLoS Pathog. 2019. doi:10.1371/journal.ppat.1007963 LK -604 http://sfx.library.uu.nl/utrecht?sid=EMBASE&issn=15537374&id=doi:10.1371%2Fj 605 ournal.ppat.1007963&atitle=The+alpha-1+subunit+of+the+Na%2B%2CK%2B-606 ATPase+%28ATP1A1%29+is+required+for+macropinocytic+entry+of+respiratory 607 +syncytial+virus+%28RSV%29+in+human+respiratory+epithelial+cells&stitle=PLo 608 S+Pathog.&title=PLoS+pathogens&volume=15&issue=8&spage=e1007963&epag 609 e=&aulast=Lingemann&aufirst=Matthias&auinit=M.&aufull=Lingemann+M.&coden 610 =&isbn=&pages=e1007963-&date=2019&au 611

612	44.	Mason SW, Aberg E, Lawetz C, DeLong R, Whitehead P, Liuzzi M. Interaction
613		between human respiratory syncytial virus (RSV) M2-1 and P proteins is required
614		for reconstitution of M2-1-dependent RSV minigenome activity. J Virol. 2003;77:
615		10670–6. doi:10.1128/JVI.77.19.10670-10676.2003
616	45.	Bailly B, Richard CA, Sharma G, Wang L, Johansen L, Cao J, et al. Targeting
617		human respiratory syncytial virus transcription anti-termination factor M2-1 to
618		inhibit in vivo viral replication. Sci Rep. 2016. doi:10.1038/srep25806
619	46.	Blondot ML, Dubosclard V, Fix J, Lassoued S, Aumont-Nicaise M, Bontems F, et
620		al. Structure and functional analysis of the RNA- and viral phosphoprotein-binding
621		domain of respiratory syncytial virus M2-1 protein. PLoS Pathog. 2012.
622		doi:10.1371/journal.ppat.1002734

623

624 Acknowledgements

625

We thank the expert technical support of Maria Dolores Seabra Ferreira and Jose 626 Augusto Maulin (FMRP-USP Electron Microscopy Facility); Elizabete Rosa Milani and 627 Roberta Ribeiro Costa Rosales (FMRP-USP Bioimaging Facility); and Maria Lucia Silva. 628 This work was made possible by a research grant from the State of Sao Paulo 629 630 Research Foundation (FAPESP, Grant No. 2013/16349-2). We also thank the funding agencies CNPq and CAPES (Brazilian Ministry of Education) for scholarships. We also 631 would like to thank the Department of Microbiology and Immunology of the University of 632 633 Michigan, and all the Ono's Lab members for their precious suggestions. In addition, we thank all of the Eurico's lab members for their pivotal ideas, experimental and discusses 634 for the paper. Finally, we thank the Lukacss lab, especially Susan Morris for all the 635 support and help. 636

637

638 Legends for figures

639

Infection of HRSV in A3.01 cells. (A) flow cytometry analysis of mock and HRSV-infected cells
 showing the percentage of the infected cells by detection of HRSV N protein. (B) qPCR of HRSV
 genome in supernatant of infected HEp-2 and A3.01 cells over time post-infection. (C) HRSV
 progeny production in A3.01 cells determined by fluorescent focus assay. All results are from
 three independent experiments.

646 2. Intracellular accumulation of HRSV genome in A3.01 and HEp-2 cells. (A) A3.01 and HEp-2
647 cells attached (Att) or in suspension (Sus) were inoculated with HRSV or mock-inoculated, and
648 kept at 4°C for 1 hour for attachment, then cells were centrifuged, and collected for qPCR
649 analysis at time zero and at different times thereafter. Genome quantification was plotted in Y
650 axis. The "virus only" well received only virus, in the absence of cells. This graph is a
651 representation of three independent experiments.

652

3. The fusion process in A3.01 is inefficient. (A) differences between HRSV-infected cells with
and without inoculum removal at different times post-infection. (B) comparison of the fusion
process in A3.01 and HEp-2 cells over time, in less populated, 40,000 cells per well. (C)
comparison of the HRSV attachment in A3.01 and HEp-2 cells. The graphs in A, B and C
represent at least three independent experiments; the statistical method used was Student's Ttest, p*<0.05, p**< 0.01, and p***< 0.001. The intensity of fluorescence emitted by R18 was
measured by a SynergyTM Multi-Mode Microplate Reader.

660

4. HRSV protein production in A3.01 cells is discrete. (A) histogram of mean intensities of 661 fluorescence of cells by a flow cytometry. (B) graph plotted from three different experiments 662 comparing the production of HRSV N protein in A3.01 and HEp-2 attached (Att) cells at 24, 48 663 664 and 72 hours post-infection. The graph in B represents at least three independent experiments; (C) western blot of HEp-2 and A3.01 cells infected or not (MOCK) by HRSV, the graphs represent 665 the analysis of the proteins bands of three independent experiments. The statistical method 666 used in figure B was Two-Way ANOVA, and the statistical method used in C was Student's T-667 test, p*<0.05, p**< 0.01, and p***< 0.001. 668

669 670

671 5. HRSV inclusion bodies in HRSV-infected A3.01 and HEp-2 cells. (A-C) A3.01 cells stained for HRSV N protein at 48 hpi (green fluorescence). (D-F) staining for HRSV N protein in HEp-2 cells 672 at 48hpi. (G) comparative analysis of inclusion body sizes in A3.01 and HEp-2 cells. (H) 673 674 comparative analysis of numbers of vesicular structures stained for HRSV N in A3.01 and HEp-2 675 cells. (I) comparative analysis of the structures stained for HRSV N in A3.01 and HEp-2 cells normalized by cell area. The immunofluorescence images shown in figures A-F represent a 676 677 single focal plane of at least three independent experiments. The images were taken in a Zeiss 780 confocal microscope. Magnification 63x. The scale bars represent = $10\mu m$. The graphs in G 678

and H represent at least three independent experiments; the statistical method used was
Student's T-test, p*<0.05, p**< 0.01, and p***< 0.001.

681

682 **6. The HRSV inclusion bodies in A3.01 are IBAG's absent. (**A-D) and (E-H) A3.01 and HEp-2 683 mock-infected cells. (I-L) and (M-P) HRSV infected HEp-2 cells at 24 and 48 hpi as shown by IBs 684 in J and N, respectively. Within IBs in J and N it is possible to see FISH-signal for IBAGs, as shown 685 in K, L, O and P (arrows). (Q-T) and (U-Z) represent A3.01-infected cells at 24 and 48 hpi. No 686 IBAGs were seen within HRSV inclusion bodies (R and W) in these cells, as shown in S, T, Y and 687 Z. This set of figures represents a single focal plane of three independent experiments taken in 688 a Zeiss 780 Confocal. Magnification 63x. Scale bars = 10µm.

689

7. Co-localization analysis of HRSV proteins at the Golgi in A3.01 cells. (A-D) mock-infected 690 691 cells stained for cis and medial-Golgi (Giantin) in red (B) trans-Golgi (TGN46) in magenta (C) 692 merge in (D). (E-H) HRSV-infected A3.01 cells at 48 hpi stained for HRSV F (green) in €, giantin in 693 (F) and TGN46 in (G). Merge is depicted in (H). The arrowheads point to a region where the F protein is located in the plasma membrane. (I-L) HRSV-infected A3.01 cells at 48 hpi stained for 694 HRSV N in (I), giantin in (J) and TGN46 in (K). The merge of the figure is in (L). All the figures 695 represent a single focal plane of at least three independent experiments taken in a Zeiss 780 696 Confocal microscope. Magnification 63x. Scale bars = $10\mu m$. 697

- 698
- 699

8. Co-localization of EEA1 and Lamp-1 with HRSV proteins in A3.01 cells. A-D, represent A3.01
mock-infected cells. In B is showed Lamp-1 (red), in C EEA1 (magenta) and in D the merge of the
set of the figures. E-H, correspond to A3.01-infected cells, stained by HRSV F, shown in E, Lamp1 in F and EEA1 in G. The merge to this set of figures is depicted in H. I-L, correspond to A3.01infected cells, stained by HRSV N, shown in I, CD63 in J and SNX2 in K. The merge of this figure
set is depicted in L. All the images were taken in a Zeiss 780 Confocal and are a representation
of a single focal plane. Magnification 63x. Scale bars = 10µm.

707

9. CD63 co-localization with HRSV proteins. (A-D), represent A3.01 mock-infected cells. In B is 708 709 showed CD63 (red), in C SNX2 (magenta) and in D the merge. E-H, correspond to A3.01-infected cells, stained by HRSV F, shown in E, CD63 in F and SNX2 in G. The merge to this set of figures is 710 depicted in H. The arrows point to the places where the cells are not infected, and there were 711 712 not CD63 accumulation. The arrowhead point to the places where the HRSV F protein is found 713 and the CD63 is accumulated. I-L, correspond to A3.01-infected cells, stained by HRSV N, shown 714 in I, CD63 in J and SNX2 in K. The merge to this set of figures is depicted in L. The arrows point to the places where the cells are not infected, and there were not CD63 accumulation. The 715 716 arrowhead points to the place where the HRSV N protein is found, and the CD63 is 717 accumulated. M-O represent the surface plot of the figures in I, J and K respectively showing 718 the heating map of the CD63 where HRSV N is located. P, graph of co-localization between CD63 and SNX2 with HRSV proteins, showing significant co-localization of CD63. The figure P 719 720 represents more than three independent experiments and was done in at least five cells per field, the statistical method used was Student's T-test, p*<0.05, p**< 0.01, and p***< 0.001. All 721

the images were taken in a Zeiss 780 Confocal and are a representation of a single focal plane.
Magnification 63x. Scale bars = 10μm.

724

725

726

727 10. Filament formation of HRSV in A3.01 is rare. (A-C) immunofluorescence for HRSV N and M 728 in A3.01-infected cells at 48 hpi depicting some filaments, pointed by arrowheads. (D-F) 729 immunofluorescence for HRSV N and M in HEp-2-infected cells at 48 hpi depicting filaments 730 pointed by arrowheads. Figures A to F represent a single focal plane of at least three 731 independent experiments taken in a Leica SP5 Confocal. Magnification 63x. This experiment was repeated at least three independent times. The scale bar of figure C = $10\mu m$. (H) graph of 732 the percentage of A3.01 and HEp-2-infected cells displaying at least one filament emerging 733 734 from the plasma membrane per field. (I) graph of the quantity of the filaments in A3.01 and 735 HEp-2-infected cells. (J) graph of the quantity of the filaments in A3.01 and HEp-2-infected cells 736 normalized by the cell area. The graphs depicted in H, I and J are representative of more than 5 737 independent experiments. The statistical method used was Student's T-test, p*<0.05, p**< 738 0.01, and p***< 0.001. All the images were taken in a Zeiss 780 or Leica SP5 Confocal and are a representation of a single focal plane. Magnification 63x. Scale bars = $10\mu m$. 739

740

741 Supplementary Figure 1. Sectional slices from HEp-2 and A3.01-infected cells

742 (supplementary). A, a Z stack was performed, and it is possible to observe |BAG's within HRSV

- in HEp-2 IB's along with the stacks, highlighted in the crops. B, sequential sectional slices from
- A3.01 cells, showing |B and the absence of clear |BAG's within, highlighted in the crops. (C)
- graph of the quantity of the Inclusion Bodies containing IBAG's found in A3.01 and HEp-2-
- infected cells at 48 hpi. The images were acquired in a Zeiss 780 Confocal, and are
- 747 representative of three independent experiments. (D) graph of the quantity of IBAG's counted 748 in inclusion bodies in A3.01 and HEp-2-infected cells. Magnification 63x. Scale bars = 10μ m. To 749 the (C) and (D) 0graphs was counted at least 10 fields of three independent experiments from
- single focal plane of Z-stacks imaging acquisition. The statistical method used was Student's T test, p*<0.05, p**< 0.01, and p***< 0.001.
- 752

753 754 Supplementary Figure 2. Colocalization of the HRSV Fusion protein with secretory pathway

markers. (A, B and C) immunofluorescence of HRSV F, Giantin and the merge respectively, the
 arrow points to colocalization. (D, E and F) immunofluorescence of HRSV F, TGN46 and the
 merge respectively, the arrow points to colocalization. (G, H and I) immunofluorescence of
 HRSV F, SNX2 and the merge respectively. (J, K and L) immunofluorescence of HRSV F, CD63 and

HRSV F, SNX2 and the merge respectively. (J, K and L) immunofluorescence of HRSV F, CD63 an
 the merge respectively. (M, N and O) immunofluorescence of HRSV F, Lamp1-1 and the merge

- respectively, the arrows point the areas of colocalization. All figures represent a single plane of
- 761 Z-stack or not experiments. The images were acquired in a Zeiss 780 or Leica SP5Confocal.
- 762 Magnification 63x. Scale Bars = $10\mu m$.
- 763

764 Supplementary Figure 3. Colocalization of the HRSV Nucleoprotein protein with secretory

765 pathway markers. (A, B and C) immunofluorescence of HRSV N, Giantin and the merge

- respectively, the arrows point to colocalization area. (D, E and F) immunofluorescence of HRSV
- N, TGN46 and the merge respectively, the arrow points to colocalization. (G, H and I)
- immunofluorescence of HRSV N, SNX2 and the merge respectively, the arrows point to
- 769 colocalization areas. (J, K and L) immunofluorescence of HRSV N, CD63 and the merge
- respectively, arrows point to colocalization areas. (M, N and O) immunofluorescence of HRSV N,
- Lamp1-1 and the merge respectively, the arrows point the areas of colocalization. All figures
- represent a single plane of Z-stack or not experiments. The images were acquired in a Zeiss 780
- or Leica SP5Confocal. Magnification63x. Scale Bars = $10\mu m$.
- 774
- 775
- 776
- 777
- 778
- 779





5.0×10⁵¬

bioRxiv preprint doi: https://doi.org/10.1101/2020.12.28.424605; this version posted December 28, 2020. The copyright holder for this preprint (which was not certified by peer review) is the author/funder, who has granted bioRxiv a license to display the preprint in perpetuity. It is made available under aCC-BY-NC-ND 4.0 International license.

 3.0×10^{4} 3.0×10^{4} 3.0

bioRxiv preprint doi: https://doi.org/10.1101/2020.12.28.424605; this version posted December 28, 2020. The copyright holder for this preprint (which was not certified by peer review) is the author/funder, who has granted bioRxiv a license to display the preprint in perpetuity. It is made available under aCC-BY-NC-ND 4.0 International license.

HRSV-infected cells

P.S.

bioRxiv preprint doi: https://doi.org/10.1101/2020.12.28.424605; this version posted December 28, 2020. The copyright holder for this preprint (which was not certified by peer review) is the author/funder, who has granted bioRxiv a license to display the preprint in perpetuity. It is made available under aCC-BY-NC-ND 4.0 International license.

N Y

L L D S

A3.01 24 bS/SV bj

HED-Z HED-Z

HRSVN

Non-infected cells HRSV N

A

B

48 hpi

C

HRSV N

Merge

