1 The P681H mutation in the Spike glycoprotein confers Type I interferon

2 resistance in the SARS-CoV-2 alpha (B.1.1.7) variant

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20 SUMMARY

21 Variants of concern (VOCs) of severe acute respiratory syndrome coronavirus 22 type-2 (SARS-CoV-2) threaten the global response to the COVID-19 pandemic. 23 The alpha (B.1.1.7) variant appeared in the UK became dominant in Europe and 24 North America in early 2021. The Spike glycoprotein of alpha has acquired a 25 number mutations including the P681H mutation in the polybasic cleavage site 26 that has been suggested to enhance Spike cleavage. Here, we show that the 27 alpha Spike protein confers a level of resistance to the effects of interferon- β (IFN β) in lung epithelial cells. This correlates with resistance to restriction 28 29 mediated by interferon-induced transmembrane protein-2 (IFITM2) and a 30 pronounced infection enhancement by IFITM3. Furthermore, the P681H 31 mutation is necessary for comparative resistance to IFN β in a molecularly 32 cloned SARS-CoV-2 encoding alpha Spike. Overall, we suggest that in 33 addition to adaptive immune escape, mutations associated with VOCs also 34 confer replication advantage through adaptation to resist innate immunity.

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38 INTRODUCTION

39 Both SARS-CoV-1 and SARS-CoV-2 enter target cells through the interaction of their 40 Spike proteins with the angiotensin converting enzyme 2 (ACE2) cell surface 41 receptor. Upon attachment and uptake, the Spike glycoprotein trimer is cleaved by 42 cellular proteases such as cathepsins and TMPRSS family members at two positions 43 - the S1/S2 junction and the S2' site - to facilitate the activation of the fusion 44 mechanism. Similar to more distantly related beta-CoVs, but so far unique in known 45 Sarbecoviruses, the SARS-CoV-2 glycoprotein contains a polybasic furin cleavage 46 site (FCS) with a (681-PRRAR-685) sequence at the S1/S2 junction. This allows the 47 Spike precursor to be processed to the S1 and S2 subunits by furin-like proteases 48 before viral release from the previously infected cell(Hoffmann et al., 2020). This 49 leads to a proportion of processed Spikes to be present on the virion before 50 engagement with the target cell, allowing for rapid activation and fusion at or near 51 the cell surface by TMPRSS2. The importance of the FCS is highlighted by the 52 observations that it enhances SARS-CoV-2 replication specifically in airway epithelial 53 cells and is essential for efficient transmission in animal models (Peacock et al., 54 2021a).

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The alpha variant of SARS-CoV-2 arose in the South-East of England in autumn 2020, and rapidly spread across the world in the first months of 2021. Various studies suggested that alpha had an increased transmissibility between individuals(Lindstrom et al., 2021; Mok et al., 2021; Tanaka et al., 2021). Alpha contains nine amino acid residue changes in Spike including a deletion of amino acid residues H and V in the N-terminal domain at position 69/70, thought to increase Spike incorporation into virions, a single amino acid deletion of Y144 (thought to

63 assist NTD antibody neutralization escape), and a N501Y mutation in the RBD which 64 enhances ACE2 binding affinity (Meng et al., 2021), (Chi et al., 2020). Together these 65 changes have been shown to reduce efficiency of neutralization by some 66 antibodies (Graham et al., 2021). Alpha also acquired a P681H change in the FCS 67 which has been shown to increase the accessibility of the site by furin leading to 68 enhanced cleavage(Mohammad et al., 2021),(Zhang et al., 2021). The alpha variant 69 Spike has also recently been reported to mediate more efficient cell-to-cell fusion 70 and syncytia formation (Michael Rajah et al., 2021), (Sanches et al., 2021).

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72 We and others have previously found that SARS-CoV-2 is variably sensitive to entry 73 inhibition by the interferon-regulated IFITM family (Winstone et al., 2021), (Shi et al., 74 2021). the three members of the family form multimeric complexes and have antiviral 75 activity against diverse enveloped viruses by blocking fusion of the viral and cellular 76 membranes (Bailey et al., 2014; Shi et al., 2017). While IFITM1 localizes primarily to 77 the plasma membrane, IFITM2 and IFITM3 are internalized via a conserved $Yxx\Phi$ endocytic motif to occupy distinct and overlapping endosomal compartments(Jia et 78 79 al., 2012; Jia et al., 2014). The sensitivity of a given virus to individual IFITM proteins 80 is largely determined by their route of cellular entry. We have previously shown that 81 for a prototypic Wuhan-like SARS-CoV-2 isolate from early 2020, IFITM2 reduced 82 viral entry and contributed to type I interferon (IFN-I)-induced inhibition in human 83 cells(Winstone et al., 2021). Sensitivity to IFITM2 could be markedly enhanced by 84 deletion of the FCS, suggesting that furin processing ameliorated SARS-CoV-2 85 sensitivity to IFITM2-restriction at least to some extent. We therefore postulated that 86 the altered cleavage site of alpha may have consequences for its sensitivity to IFN-I 87 and IFITMs. Here, we demonstrate that the Spike of the alpha variant is less

sensitive to restriction by IFN β and IFITMs in A549-ACE2 and Calu-3 cells. Furthermore, this resistance correlates with the enhanced polybasic site as reversion of this cleavage site increases the alpha variant's sensitivity to IFITM restriction. Finally, we demonstrate that the H681P reversion in the full-length virus confers IFN β sensitivity to alpha and suggest that part of this phenotype is driven by IFITMs.

93

94 **RESULTS**

95 The Spike proteins of currently circulating variants display differing 96 sensitivities to IFITMs in A549-ACE2 cells

97 Previously we have shown that viral entry mediated by the original Wuhan-1 Spike 98 pseudotyped lentiviral particle (PLV) or the England 02 isolate (hCoV-99 19/England/02/2020) was inhibited by IFITM2 in A549-ACE2 cells, and that this 100 effect correlated in part with the IFN β sensitivity of the virus(Winstone et al., 2021). 101 Over 2020 and 2021 several major variants of concern (VOCs) have arisen – alpha (B.1.1.7) in the UK, beta (B.1.351) in South Africa, gamma (P1) in Brazil, and delta 102 103 (B.1.617.2) in India. We wanted to compare the sensitivity of viral entry of the alpha, 104 beta, gamma and delta to the presence of IFITM proteins, given that these variants 105 have several changes in the Spike sequence (Figure 1A). Initially, PLVs bearing the 106 spike protein of each variant to test whether they were restricted by IFITMs 107 overexpressed on A549-ACE2 cells (Figure 1B-G). First, we confirmed that the 108 D614G mutation in the parental Wuhan-1 Spike that became prevalent in the first 109 wave of the pandemic displays a similar IFITM phenotype to the previously 110 characterised SARS-CoV-2 Spike (Wuhan-1) (Figure 1B, 1C)(Winstone et al., 2021). 111 The addition of D614G to the Wuhan-1 Spike had no effect on IFITM1 and IFITM2 112 sensitivity of PLV entry, while we observe a slight enhancement in the presence of

113 IFITM3. We next compared the IFITM sensitivities of the major global VOCs. The 114 alpha Spike appeared completely insensitive to IFITMs 1, 2 or 3 whilst beta, gamma 115 and delta still retain some sensitivity to IFITMs 1 and/or 2 (Figure 1B-G). None of the 116 VOCs were restricted by IFITM3 and, interestingly, we noted that IFITM3 appeared 117 to markedly enhance entry mediated by the alpha variant Spike. Next, we pre-treated 118 A549-ACE2-IFITM3 cells with cyclosporin H as this compound is known to drive 119 IFITM3 to degration(Petrillo et al., 2018), and showed that this led to a specific 120 abolishment of the enhanced infection by alpha PLVs (Supplemental Figure 1). 121 Henceforth, we selected alpha to further investigate and determine the mechanism 122 of IFITM resistance.

123 Next, to confirm whether IFITM sensitivity results seen with the alpha spike 124 PLVs could be recapitulated with the full-length virus (Figure 2A), we infected A549-125 ACE2 cells (stably expressing IFITMs or not) with either the Wuhan-like England/02 126 isolate or alpha at an MOI of 0.01 and measured infection by qPCR of E copies in 127 the infected cells 48 hours later (Figure 2B). As expected, replication of England/02 128 was significantly reduced in IFITM2 expressing cells over 48h. By contrast alpha 129 replicated as well in IFITM2-expressing cells and to a significantly higher level in 130 IFITM3-expressing cells in comparison to the control A549-ACE2 cells. This 131 confirmed that alpha SARS-CoV-2 is resistant to the effects of IFITMs 1 and 2 and 132 enhanced by IFITM3 on viral entry. Previous reports have suggested that the alpha 133 spike is more efficiently cleaved than Wuhan-like isolates due to the presence of the 134 P681H mutation optimising the accessibility of the FCS (Mohammad et al., 2021; 135 Zhang et al., 2021) We immunoblotted our viral stocks of England-02 and alpha and 136 confirmed that the alpha has more processed spike on the virions (Figure 2C, right 137 panel), although it was not as clearly discernible a difference in processing on PLVs

138 (Figure 2C, left panel). However, PLV infection in A549-ACE2 cells pre-treated with 139 E64D, a cathepsin inhibitor, that inhibits the entry of SARS-CoV-2 in endosomal 140 compartments because it cleaves S1/S2 junctions that have not been processed by 141 furin. This showed that the alpha variant Spike mediated entry of PLVs is markedly 142 less sensitive to this endosomal protease inhibition, when compared with the 143 Wuhan-1/D614G Spike, indicating differences in the site of virus entry mediated 144 consistent with enhanced Spike processing by furin in the producer cell 145 (Supplementary Figure 2).

146 The alpha variant is less sensitive to IFN β than a wave 1 isolate

147 While previous data has indicated that the original Wuhan-like SARS-CoV-2 virus 148 can delay pattern recognition of viral RNA in target cells, its replication highly 149 sensitive to exogenous interferon treatment in culture, in part determined by 150 IFITM2(Jouvenet, 2021). Having confirmed that the alpha variant is resistant to 151 IFITMs expression when ectopically expressed in cells, we then tested if alpha is 152 also resistant to the effects of IFN β , as suggested (Guo et al., 2021; Thorne et al., 153 2021). Indeed, we found from measuring supernatant viral RNA 48 hours after 154 infection of both A549-ACE2 cells and the human lung epithelial cell line Calu3, 155 which more faithfully represent target cells in the respiratory tract, that alpha is more 156 resistant than England/02 to pre-treatment with increasing doses of IFN β (Figure 3A, 157 3B). We have also confirmed this phenotype by measuring viral RNA in cell lysates, 158 and further extending these obervations to two clinical isolates of alpha isolated 159 (clinical isolates 10 and 28; Figure 3C). Thus in comparison to a representative 160 example of Wuhan-1-like SARS-CoV-2, the alpha variant has a marked resistance 161 to type I interferon.

162 The P681H mutation is necessary for conferring IFITM and IFNβ resistance in

163 **alpha**

164 Our previous data indicated IFITM sensitivity of SARS-CoV-2 Spike can be 165 increased by deleting the FCS. Given that alpha and delta Spikes have acquired 166 mutations at P681 to H or R respectively, and these enhance Spike cleavage to 167 S1/S2 (Supplementary figure 3), we hypothesized that P681H might be a 168 determinant of resistance to IFN and IFITM. First we confirmed that none of the other 169 individual alpha-Spike defining mutations were sufficient to confer IFITM resistance 170 to a Wuhan/D614G Spike in PLVs (Supplementary figure 4 A-F). By contrast a 171 P681H mutation in the Wuhan-1/D614G Spike was sufficient to abolish IFITM2-172 mediated inhibition PLVs entry, but not to fully confer the IFITM3-mediated 173 enhancement phenotype (Figure 4A). As expected, deletion of the HRRA cleavage 174 site in the alpha Spike conferred potent sensitivity to IFITM2 and reduced the level of 175 enhancement we showed with IFITM3. Finally, we reverted the mutation in alpha to 176 H681P and showed it gained IFITM2 sensitivity. Thus, the H681P mutation in the 177 alpha spike confers IFITM resistance consistent with its enhanced furin cleavage. 178 However delta bears a P681R mutation yet is not fully IFITM resistant (Figure 1G), 179 implying that the P681H phenotype may be context dependent. Indeed, we found 180 that combining the P681H mutation in the Wuhan spike with the deletion at position 181 69-70 in the NTD found in the alpha variant but not delta was sufficient to fully confer 182 an alpha spike phenotype to D614G (Supplementary figure 4G). This suggests that 183 the H681 confers IFITM resistance in the context of other adaptations in the alpha 184 Spike that are thought to affect the conformation of S1 and the interaction with 185 ACE2.

186 Having established that P681H change is necessary for the resistance of the 187 alpha variant Spike to IFITMs, we next wanted to address if this was also a 188 determinant for the resistance to type-I IFN resistance of the virus itself. We constructed a recombinant molecular clone of SARS-CoV-2 Wuhan-1 encoding 189 190 Spike from the alpha variant. This virus essentially mimicked the resistance of the 191 alpha variant itself to IFN β in comparison to a representative Wuhan-1 like virus, 192 England-02, demonstrating that the alpha Spike alone is sufficient to confer a level 193 type I IFN resistance in A549-ACE2 cells (Figure 4B). We then took this recombinant 194 virus and reverted the amino acid residue H681 to a proline. Importantly, this single 195 point mutation was sufficient to confer a significant sensitivity to IFN β in Calu3 cells 196 (Figure 4C). Lastly, we wanted to confirm whether siRNA knockdown of IFITM2 was 197 sufficient to rescue the IFN sensitivity of the Wuhan(B.1.1.7 Spike H681P) virus. We 198 therefore knocked down IFITM2 expression using siRNA in A549-ACE2 cells and 199 treated the cells with IFN β as before. We found that the H681P reverted virus was 200 rescued from IFN_β restriction during IFITM2 knockdown, meanwhile the 201 Wuhan(B.1.1.7 Spike) virus was unaffected, consistent with this virus being resistant 202 to IFITM2 restriction. Thus, this confirmed that the Spike protein of the alpha variant 203 of SARS-CoV-2 is a determinant of type-I IFN resistance and that the P681H 204 mutation is necessary for this.

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206 **DISCUSSION**

Here we have shown that the Spike protein of the alpha variant of SARS-CoV-2 is resistant to IFN-I. Furthermore, we show that this maps to the histidine residue adjacent to the FCS that has been mutated from the parental proline, which has been shown to enhance Spike cleavage at the S1/S2 boundary (Peacock et al.,

211 2021b). This residue is necessary to confer resistance to IFITM2 and enhancement 212 by IFITM3, and as we demonstrated in our previous study(Winstone et al., 2021), 213 confirms that the FCS in Spike modulates IFITM entry restriction. Changes in the 214 FCS would be predicted to increase the efficiency of viral fusion and entry at or near 215 the plasma membrane, avoiding endosomal compartments where IFITMs 2 and 3 216 predominantly reside. Consistent with this, we show that the alpha Spike, as a PLV, 217 is less sensitive to the cathepsin inhibitor E64D. Thus we propose that these 218 changes in the alpha Spike have, in part, arisen to resist innate immunity.

219 At least two preprints suggest that variants of SARS-CoV-2 have begun to 220 evolve further resistance to interferon-induced innate immunity (Guo et al., 2021; 221 Thorne et al., 2021). In one, viral isolates over the pandemic show a reduced 222 sensitivity to type I interferons in culture(Guo et al., 2021); in a second the alpha 223 variant has a significantly reduced propensity to trigger pattern recognition in 224 epithelial cells(Thorne et al., 2021). In contrast, another study shows no difference in 225 IFN sensitivity of the new variants in African green monkey Vero-E6 cells(Michael 226 Rajah et al., 2021), although species-specificity in viral sensitivity to ISGs is a well 227 characterized trait. The SARS-CoV-2 genome contains multiple mechanisms to 228 counteract host innate immune responses, and much remains to be learned about 229 the mechanisms deployed by this virus and its relatives. While many reports on 230 SARS-CoV-2 evolution have naturally focussed on the pressing concern of potential 231 for vaccine escape, it is very unlikely that all selective adaptations that we see 232 arising in VOCs can be solely due to escape from adaptive immunity. The alpha 233 variant Spike, for example, only displays a minor reduction in sensitivity to 234 neutralizing antibodies (NAbs) (Graham et al., 2021; Mahase, 2021; Planas et al., 235 2021; Shen et al., 2021). However, this VOC had a considerable transmission

advantage, with suspicions that it may have arisen in an immunocompromised
individual with a persistent infection giving ample time for changes to be selected
that further evade innate immunity, including those that target viral entry(Corey et al.,

239 2021; Kemp et al., 2021).

240 In terms of IFITM resistance of VOC Spike proteins, so far we have only seen 241 marked change in phenotype for the alpha variant. This is despite the fact that delta, 242 which has superseded alpha in many places around the world in 2021, also shows 243 an adaptation for enhanced S1/S2 cleavage with an P681R change(Liu et al., 2021; 244 Peacock et al., 2021b). This would suggest that efficient cleavage of S1/S2 is 245 necessary but not sufficient for IFITM resistance, and indeed our data implicate a 246 context dependency of the NTD deletion at 69/70. While unique to the alpha VOC, 247 the 69/70 deletion has been observed in persistent infection of immunosuppressed 248 individuals and is thought to enhance viral fitness and Spike stability (Meng et al., 249 2021). While deletions in the NTD do affect NAb binding, it is primarily the 144 250 deletion in the alpha Spike that escapes neutralization by NTD-directed Nabs (Chi et 251 al., 2020) and we show that this has no impact on IFITM sensitivity. By contrast, the 252 more pronounced antibody evasion by the beta, gamma and delta variants is related 253 to mutations in the major neutralizing epitopes of the RBD, suggesting that they may 254 well have been driven by antibody escape(Planas et al., 2021; Zhou et al., 2021). 255 glycoproteins are dynamic structures that shift through large-scale Viral 256 conformational changes while interacting with their cognate receptors mediating viral 257 membrane fusion. Such context dependency is therefore likely to be complex and 258 will arise under competing selective pressures. Indeed, we have previously shown 259 that the HIV-1 envelope glycoprotein of transmitted viruses is IFITM insensitive and 260 this contributes to their overall type I IFN resistance (Foster et al., 2016). As HIV-1

infection progresses over the first 6 months in an infected person, the circulating
variants increase in IFN/IFITM sensitivity and this is determined by adaptive changes
in Env that resist the early neutralizing antibody response (Fenton-May et al., 2013).
Such escape has structural and functional implications for such dynamic proteins
that may impact upon receptor interactions and route of entry into the target cell.

266 While furin cleavage of the SARS-CoV-2 Spike reduces its IFITM sensitivity, 267 other interferon-induced proteins may contribute to this phenotype. The guanylate 268 binding protein family, and particularly GBP2 and GBP5, have been shown to have a 269 general antiviral activity against enveloped viruses by dysregulating furin processing 270 of diverse viral and cellular proteins (Braun et al., 2019). Similarly, IFITM 271 overexpression in HIV-infected cells can lead to their incorporation into virions and in 272 some cases promote defects in glycoprotein incorporation (Tartour et al., 2014). 273 Future studies will confirm whether either of these mechanisms are involved in the 274 IFN-resistance associated with the P681H mutation in alpha.

In summary, the spike protein of SARS-CoV-2 alpha increases resistance to IFN-I and this correlates with the P681H mutation. Furthermore, this correlates with resistance to IFITM-mediated entry restriction. This suggests that in addition to adaptive immune escape, fixed mutations associated with VOCs may well also confer replication and/or transmission advantage through adaptation to resist innate immune mechanisms.

281

283 MATERIALS AND METHODS

284 Cells and plasmids

HEK293T-17 (ATCC, CRL-11268TM), Calu-3 (ATCC, HTB-55TM), A549-ACE2, VeroE6, Vero-E6-TMPRSS2 and A549-ACE2 expressing the individual IFITM proteins
were cultured in DMEM (Gibco) with 10% FBS (Invitrogen) and 200µg/ml Gentamicin
(Sigma), and incubated at 37°C, 5% CO₂. ACE2, TMPRSS2, and IFITM stable
overexpression cells were generated as previously described (Winstone et al.,
2021).

291 Codon optimised SARS-CoV-2 Wuhan Spike and ACE2 were kindly given by Dr. 292 Nigel Temperton. Codon optimised variant Spikes (B.1.1.7, B.1.351) were kindly 293 given by Dr. Katie Doores. Codon optimised variant Spikes (P1, B.617.2) were kindly 294 given by Professor Wendy Barclay. Plasmid containing TMPRSS2 gene was kindly 295 given by Dr. Caroline Goujon. Spike mutants were generated with Q5® Site-Directed 296 Mutagenesis Kit (E0554) following the manufacturer's instructions, and using the 297 following forward and reverse primers:

298 D614G (GCTGTACCAGGGCGTGAATTGCA, ACGGCCACCTGATTGCTG)

299 B.1.351. Δ242-244 (ATTTCATATCTTACACCAGGC, ATGCAGGGTCTGGAATCTG)

300 D614G P681H (GACCAATAGCcacAGAAGAGCCAGAAGC,

301 TGGGTCTGGTAGCTGGCG)

B117 △HRRA (AGAAGCGTGGCCAGCCAG, GCTATTGGTCTGGGTCTGGTAG)

303 B117 H681P (GACCAATAGCcccAGAAGAGCCAG, TGGGTCTGGTAGCTGGCG)

 $304 \quad \Delta 144$ (CATAAGAACAACAAGAGC, ATAAACACCCAGGAAAGG), N501Y

305 (CCAGCCTACCtacGGCGTGGGCT, AAGCCGTAGCTCTGCAGAG), E484K

306 (TAATGGCGTGAACGGCTTCAATTGCTACTT, CACGGTGTGCTGCCGGCC).

307 A549 stable cell lines expressing ACE2 (pMIGR1-puro), and IFITMs (pLHCX) were

308 generated and selected as described previously (Winstone et al., 2021).

309 Production of Pseudotyped Lentiviral Vectors (PLVs) and infection

- 310 HEK293T-17 were transfected with firefly luciferase expressing vector (CSXW), HIV
- 311 gag-pol (8.91) and Spike plasmid with PEI-max as previously described (Winstone et
- al., 2021). Viral supernatant was then used to transduce each cell line of interest and
- 313 readout measured by Luciferase activity 48 hours later (Promega Steady-Glo®
- 314 (E2550)).

315 Cyclosporin H assay

Cells were pre-treated with 30µM of Cyclosporin H (Sigma, SML1575) for 18 hours.

Cells were then infected with PLVs and viral entry quantified by Luciferase activity 48
hours later.

319 Passage and titration of SARS-CoV-2

320 PHE England strain 02/2020 was propagated in Vero-E6-TMPRSS2 cells and titre 321 was determined by plaque assay (Winstone et al., 2021). Vero-E6-TMPRSS2 were 322 infected with serial dilutions of SARS-CoV-2 for 1h. Subsequently, 2X overlay media 323 (DMEM + 2% FBS + 0.1% agarose) was added, and infected cells were fixed 72 324 hours after infection and stained with Crystal Violet. Plaques were counted and 325 multiplicity of infection calculated for subsequent experiments. A replication-326 competent alpha variant was kindly provided by Professor Wendy Barclay (Imperial 327 College London)(Brown et al., 2021). All virus stocks were sequence confirmed in 328 the Spike gene at each passage to ensure no loss of the FCS.

329 Generation of recombinant full-length viruses

330 We used the previously described Transformation-Associated Recombination 331 (TAR) in yeast method(Thi Nhu Thao et al., 2020), with some modifications, to 332 generate the mutant viruses described in this study. Briefly, a set of overlapping 333 cDNA fragments representing the entire genomes of SARS-CoV-2 Wuhan isolate 334 (GenBank: MN908947.3) and the B.1.1.7 alpha variant were chemically synthesized 335 and cloned into pUC57-Kan (Bio Basic Canada Inc and Genewiz, respectively). The 336 cDNA fragment representing the 5' terminus of the viral genome contained the 337 bacteriophage T7 RNA polymerase promoter preceded by a short sequence stretch 338 homologous to the Xhol-cut end of the TAR in yeast vector pEB2(Gaida et al., 2011). 339 The fragment representing the 3' terminus contained the T7 RNA polymerase 340 termination sequences followed by a short segment homologous to the BamHI-cut 341 end of pEB2.

342 To generate Wuhan virus carrying the alpha variant spike, a mixture of the 343 relevant synthetic cDNA fragments of the Wuhan and alpha variants was co-344 transformed with Xhol-BamHI-cut pEB2 into the Saccharomyces cerevisiae strain 345 TYC1 (MATa, ura3-52, leu2 Δ 1, cyh2^r, containing a knockout of DNA Ligase 4) 346 (Gaida et al., 2011) that had been made competent for DNA uptake using the LiCl₂-347 based Yeast transformation kit (YEAST1-1KT, Merck). The transformed cells were 348 plated on minimal synthetic defined (SD) agar medium lacking uracil (Ura) but 349 containing 0.002% (w/v) cycloheximide to prevent selection of cells carrying the 350 empty vector. Following incubation at 30[°]C for 4 to 5 days, colonies of the yeast 351 transformants were screened by PCR using specific primers to identify those 352 carrying plasmid with fully assembled genomes. Selected positive colonies were 353 then expanded to grow in 200 ml SD-Ura dropout medium and the plasmid 354 extracted. Approximately 4 µg of the extracted material was then used as template to

355 in vitro synthesized viral genomic RNA transcripts using the Ribomax T7 RNA 356 transcription Kit (Promega) and Ribo m7G Cap Analogue (Promega) as per the 357 manufacturer's protocol. Approximately 2.5 µg of the in vitro synthesized RNA was used to transfect ~6 x10⁵ BHK-hACE2-N cells stably expressing the SARS-CoV-2 N 358 359 and the human ACE2 genes(Rihn et al., 2021) using the MessengerMax lipofection 360 kit (Thermo Scientific) as per the manufacturer's instructions. Cells were then 361 incubated until signs of viral replication (syncytia formation) became visible (usually 362 after 2-3 days), at which time the medium was collected (P0 stock) and used further 363 as a source of rescued virus to infect VERO E6 cells to generate P1 and P2 stocks. 364 Full genome sequences of viruses collected from from P0 and P1 stocks were 365 obtained in order to confirm the presence of the desired mutations and exclude the 366 presence of other spurious mutations. Viruses were sequenced using Oxford 367 Nanopore as previously described(da Silva Filipe et al., 2021).

To generate Wuhan virus carrying alpha spike gene with the H681P mutation, we first introduced this mutation into the relevant alpha variant cDNA fragment by site-directed mutagenesis. This fragment was combined with those described above and the mixture was then used to generate plasmid pEB2 carrying the cDNA genome of Wuhan encoding the alpha spike H681P by the TAR in yeast procedure. The virus rescue and subsequent characterisation were performed as described above.

375 Generation of Clinical Viral Isolates

Viruses were isolated on Vero.E6 cells (ATCC CRL 1586[™]) from combined nasooropharyngeal swabs submitted for routine diagnostic testing by real-time RT-PCR
and shown to be from the B.1.1.7 (alpha) variant by on-site whole-genome
sequencing (Oxford Nanopore Technologies, Oxford, UK) (Pickering et al., 2021).

Infected cells were cultured at 37°C and 5% CO2, in Dulbecco's modified Eagle's
medium (DMEM, Gibco[™], Thermo Fisher, UK) supplemented with 2% foetal bovine
serum (FBS, Merck, Germany), pen/strep and amphotericin B.

All work performed with full-length SARS-CoV-2 preparations, as well as isolation and propagation of viral isolates from swabs, was conducted inside a class II microbiological safety cabinet in a biosafety level 3 (BSL3) facility at King's College London.

387 Infection with replication competent SARS-CoV-2

1.5x10⁵ A549-ACE2 cells were infected for 1 hour at 37°C with SARS-CoV-2 replication competent viruses at MOI 0.01 or 500 E gene mRNA copies/cell. 2x10⁵ Calu-3 cells were infected for 1h at 37°C with SARS-CoV-2 replication competent viruses at 5000 copies/cell. Media was replaced and cells were incubated for 48 hours at 37°C, after which cells or supernatant were harvested for RNA extraction or protein analysis.

394 Interferon assays

Cells were treated with different doses of IFN β (PBL Assay Science, 11415-1) for 18 hours prior infection. The following day media was replaced, and the infection performed as described above. Viral RNA levels in cells or supernatants were measured 48 hours after infection by RT-qPCR.

399 siRNA knockdown of IFITM2

400 A549-ACE2 cells were reverse transfected using 20pmol of Non-targeting siRNA (D-

401 001206-13-20) or IFITM2 siRNA (M-020103-02-0010) and 1μL of RNAi max

402 (Invitrogen). Cells were incubated for 24h prior to a second round of reverse

transfection. 8h later, cells were treated with different doses of IFN β . Following 18h

404 of IFN treatment cells were infected with full-length viruses as previously described.

405 **RT-qPCR**

RNA from infected cells was extracted using QIAGEN RNeasy (QIAGEN RNeasy
Mini Kit, 74106) following the manufacturer's instructions. 1µL of each extracted
RNA was used to performed one step RT-qPCR using TaqMan Fast Virus 1-Step
Master Mix (Invitrogen). The relative quantities of envelope (E) gene were measured
using SARS-CoV-2 (2019-nCoV) CDC qPCR Probe Assay (IDT DNA technologies).
Relative quantities of E gene were normalised to GAPDH mRNA levels (Applied
Bioscience, Hs9999905_m1).

Supernatant RNA was extracted using RNAdvance Viral XP (Beckman) following the
manufacturer's instructions. 5µL of each RNA was used for one-step RT-qPCR
(TaqMan[™] Fast Virus 1-Step Master Mix) to measured relative quantities of E and
calibrated to a standard curve of E kindly provided by Professor Wendy Barclay.

417 SDS-PAGE and Western blotting

418 Cellular samples were lysed in reducing Laemmli buffer at 95°C for 10 minutes. 419 Supernatant or viral stock samples were centrifuged at 18,000 RCF through a 20% 420 sucrose cushion for 1 hour at 4°C prior to lysis in reducing Laemmli buffer. Samples 421 were separated on 8–16 % Mini-PROTEAN® TGX[™] Precast gels (Bio-Rad) and 422 transferred onto nitrocellulose membrane. Membranes were blocked in milk prior to 423 detection with specific antibodies: 1:1000 ACE2 rabbit (Abcam, Ab108209),1:5000 424 GAPDH rabbit (Abcam, Ab9485), 1:5000 HSP90 mouse (Genetex, Gtx109753), 1:50 425 HIV-1 p24Gag mouse (48 ref before) 1:1000 Spike mouse (Genetex, Gtx632604),

426 1:1000 anti-SARS-CoV-2 N rabbit (GeneTex, GTX135357). Proteins were detected
427 using LI-COR and ImageQuant LAS 4000 cameras.

428 Ethics

Clinical samples were retrieved by the direct care team in the Directorate of Infection, at St Thomas Hospital, London, UK, and anonymised before sending to the King's College London laboratories for virus isolation and propagation. Sample collection and studies were performed in accordance with the UK Policy Framework for Health and Social Care Research and with specific Research Ethics Committee approval (REC 20/SC/0310).

435 ACKNOWLEDGEMENTS

We are grateful to Nigel Temperton, Caroline Goujon, Katie Doores, Wendy Barclay and Public Health England for reagents. We acknowledge the G2P-UK National Virology consortium funded by MRC/UKRI (grant ref: MR/W005611/1) and the Barclay Lab at Imperial College London for providing the alpha variant. We thank E. J. Louis, University of Leicester for generously providing the TAR in yeast system.

441 FUNDING

442 This work was funded by Wellcome Trust Senior Research Fellowship 443 WT098049AIA to SJDN, MRC Project Grant MR/S000844/1 to SJDN and CMS, and 444 funding from the Huo Family Foundation jointly to SJDN, Katie Doores, Michael 445 Malim and Rocio Martinez Nunez. MR/S000844/1 is part of the EDCTP2 programme 446 supported by the European Union. HW is supported by the UK Medical Research 447 Council (MR/N013700/1) and is a King's College London member of the MRC 448 Doctoral Training Partnership in Biomedical Sciences. This work is supported by the 449 UKRI SARS-CoV-2 Genotype-2-Phenotype consortium. We also benefit from 450 infrastructure support from the KCL Biomedical Research Centre, King's Health

- 451 Partners. Work at the CVR was also supported by the MRC MC_UU12014/2 and the
- 452 Wellcome Trust (206369/Z/17/Z).

453 AUTHORS CONTRIBUTION

- 454 Experiments were performed by MJL, HW, HDW and AD. SP, RPG, LS and GN
- 455 collected, sequenced and isolated clinical viral isolates. MP, AHP, GDL, VMC, WF,
- 456 NS, and RO generated reverse genetics-derived viruses. MJL, HW, HDW and AD
- 457 analysed data. CMS provided reagents, funding support and advice. HW, MJL and
- 458 SJDN analysed the data and wrote the manuscript. All authors edited the manuscript
- and provided comments.

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596 **FIGURE LEGENDS**

597 Figure 1. IFITM sensitivity of SARS-CoV-2 variants of concern. A) Schematic of 598 Spike protein domains of the different variants of concern relative to the original 599 Wuhan Spike sequence: alpha, beta, gamma and delta. The different mutations 600 between the variants are represented in red. B-G) IFITM sensitivity of Wuhan, 601 D614G, alpha, beta, gamma and delta PLVs in A549-ACE2 cells stably expressing 602 the individual IFITMs. PLV entry was guantified by Luciferase activity 48 hours after 603 infection and normalized to control cells. Data shown are mean ± SEM, n=3. 604 Statistics were calculated in Prism using *t*-test, stars indiciate significance between 605 control cell and individual IFITM (*P=0.05).

606

607 Figure 2. The alpha variant of SARS-CoV-2 is resistant to IFITMs. A) D614G and 608 Alpha PLVs infection of A549-ACE2 cells stably expressing the individual IFITMs. 609 Infection was guantified by Luciferase activity 48 hours later and normalized to 610 control cells. Data shown are mean ± SEM, n=3. Statistics were calculated in Prism 611 using t-test, stars indiciate significance between control cell and individual IFITM 612 (*P=<0.05). B) Infection of A549-ACE2 stably expressing the individual IFITMs with 613 England 02 and alpha full-length viruses at MOI 0.01. Infection was guantified by RT-614 qPCR of E gene relative to GAPDH 48 hours later; graph represents E mRNA levels 615 relative to GAPDH. Data shown are mean ± SEM, n=3. Statistics were calculated in 616 Prism using t-test, stars indiciate significance between control cell and individual 617 IFITM (*P=<0.05). C) Western blot from representative D614G and alpha PLVs 618 produced in HEK293T/17 cells, and virions from full-length England-02 and alpha 619 viruses. Virions were purified through a 20% sucrose gradient.

620

621 Figure 3. The alpha variant is resistant to IFN β . A) England 02 and alpha full-622 length virus infection in A549-ACE2 cells pre-treated with IFNβ. Cells were pre-623 treated with increasing concentrations of IFN β for 18 hours prior to infection with 624 either virus at 500 E mRNA copies/cell. Infection was quantified by RT-qPCR of E 625 mRNA from the supernatant 48 hours later and normalised to the un-treated control. 626 Data shown are mean \pm SEM, n=3. Statistics were calculated in Prism using *t*-test, 627 stars indicate signifincance between the different viruses at individual IFN 628 concentrations (*P=<0.05). B) England 02 and alpha full-length virus infection in 629 Calu-3 cells pre-treated with IFNB. Cells were pre-treated with increasing 630 concentrations of IFN_β for 18 hours prior infection with either virus at 5000 E 631 copies/cell. Infection was quantified by RT-qPCR of E mRNA from the supernatant 632 48 hours later and normalised to the un-treated control. Data shown are mean ± 633 SEM, n=3. Statistics were calculated in Prism using t-test, stars indicate 634 signifincance between the different viruses at individual IFN concentrations 635 (*P=<0.05). C) England 02 and clinical isolates of alpha full-length virus infection in 636 Calu-3 cells pre-treated with IFN^β and harvested as in A and B. Cells were pre-637 treated with increasing concentrations of IFN β for 18 hours prior to infection with 638 either virus at 5000 E copies/cell. Infection was quantified by RT-qPCR of cellular E 639 mRNA relative to GAPDH 48 hours later and normalised to the un-treated control. 640 Data shown are mean \pm SEM, n=3. Statistics were calculated in Prism using *t*-test, 641 stars indicate signifincance between the different viruses at individual IFN 642 concentrations (*P = < 0.05).

643

Figure 4. The P681H mutation is necessary but not sufficient for IFITM
 resistance, and necessary and sufficient for IFNβ resistance. A) D614G,

646 D614G- P681H, alpha, alpha-ΔHRRA, and alpha-H681P PLVs infection in A549-647 ACE2 cells stably expressing the individual IFITMs. PLVs entry was quantified by 648 Luciferase activity 48 hours later and normalized to control cells. Data shown are 649 mean ± SEM, n=3. Statistics were calculated in Prism using t-test, black stars 650 indicate significance relative to the control cells, red stars indicate significance 651 between alpha and alpha-H681P in IFITM2 cells (*P=<0.05). B) England 02, alpha, 652 and Wuhan-alpha Spike full-length virus infection in A549-ACE2 cells pre-treated 653 with IFN_β. Cells were pre-treated with increasing concentrations of IFN_β for 18 hours 654 prior to infection with either virus at 500 E copies/cell. Infection was quantified by RT-655 qPCR of E mRNA in the supernatant 48 hours later and normalised to the un-treated 656 control. Data shown are mean \pm SEM, n=3. Statistics were calculated in Prism using 657 t-test, stars indicate signifincance between the different viruses at individual IFN 658 concentrations (*P=<0.05). C) Wuhan(B.1.1.7 spike) and Wuhan(B1.1.7 spike 659 H681P) Spike full-length virus infection in Calu-3 cells pre-treated with IFNβ. Cells 660 were pre-treated with increasing concentrations of IFN^β for 18 hours prior to infection with either virus at 5000 E copies/cell. Infection was quantified by RT-qPCR 661 662 of E mRNA in the supernatant 48 hours later and normalised to the un-treated 663 control. Data shown are mean \pm SEM, n=3. Statistics were calculated in Prism using 664 t-test, stars indicate signifincance between the different viruses at individual IFN 665 concentrations (*P=<0.05). D) A549-ACE2 cells were transfected with siRNAs 666 against non-targeting control or IFITM2 for 24 hours and then treated with IFNß for 667 18 hours prior to infection with Wuhan(B.1.1.7 spike) or Wuhan(B.1.1.7 spike 668 H681P) at 500 copies/cell. Infection was quantified by RT-qPCR of E gene relative to 669 GAPDH 48 hours later; graph represents E mRNA levels relative to GAPDH. Data 670 shown are mean \pm SEM, n=3. Statistics were calculated in Prism using *t*-test, stars

671 indicate signifincance between the different viruses at individual IFN concentrations

672 (*P=<0.05).

673

674 Supplementary figure 1. Cyclosporin H treatment abolishes IFITM3 675 enhancement of alpha PLVs. A) D614G PLVs pre-treated with Cyclosporin H. 676 A549-ACE2s stably expressing the individual IFITMs were pre-treated with 30 µM of 677 Cyclosporin H for 18 hours prior to infection with D614G PLVs. Infection was 678 quantified by Luciferase activity 48 hours after infection and normalized to control 679 cells. Data shown are mean ± SEM, n=3. Statistics were calculated in Prism using t-680 test (*P=<0.05). B) A549-ACE2s stably expressing the individual IFITMs were pre-681 treated with 30 μ M of Cyclosporin H for 18 hours prior to infection with alpha PLVs. 682 Infection was quantified by Luciferase activity 48 hours after infection and normalized 683 to control cells. Data shown are mean ± SEM, n=3. Statistics were calculated in 684 Prism using t-test, stars indiciate significance between IFITM3 mock and IFITM3 685 CsH (*P=<0.05).

686

Supplementary figure 2. alpha PLVs are less sensitive to inhibition by E64D than D614G PLVs. A549-ACE2 cells were pre-treated with increasing concentrations of E64D for 1 hour prior infection with D614G or alpha PLVs. PLV entry was quantified by Luciferase activity 48 hours after infection and normalized to control cells. Data shown are mean \pm SEM, n=3.

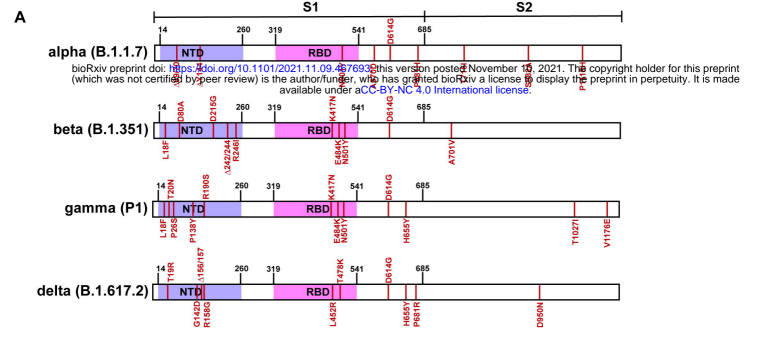
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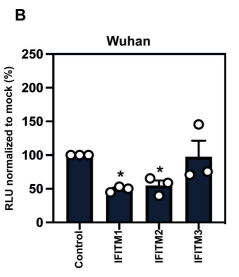
Supplementary figure 3. Spike processing of full-length virus and cleavage site
PLVs mutants. A) England-02, alpha, Wuhan(B.1.1.7 Spike), Wuhan(B.1.1.7 Spike)
H681P) were purified through 20% sucrose and immunoblotted for Spike and N

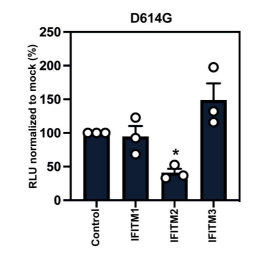
- 696 proteins. B) PLVs expressing different Spike mutants were produced in HEK293T-17
- 697 cells and cell lysates and supernatant immunoblotted for gag and Spike. Supernatant
- 698 was purified through 20% sucrose.
- 699

Supplementary figure 4. IFITM sensitivity of individual alpha Spike mutations.

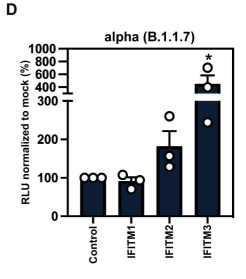
- 701 A-F) PLVs with individual alpha mutations were used to infect A549-ACE2 cells
- stably expressing the individual IFITMs. Infection was quantified by Luciferase
- activity 48 hours after infection and normalized to control cells. Data shown are
- mean ± SEM, n=3. Statistics were calculated in Prism using *t*-test, stars indiciate
- significance between each PLVs control cell and individual IFITM (*P=<0.05).

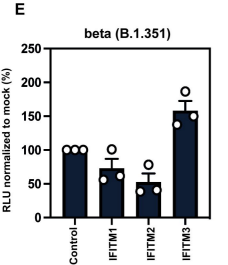


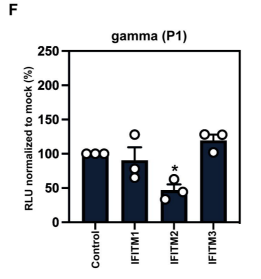


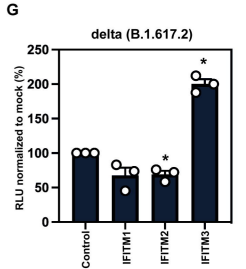


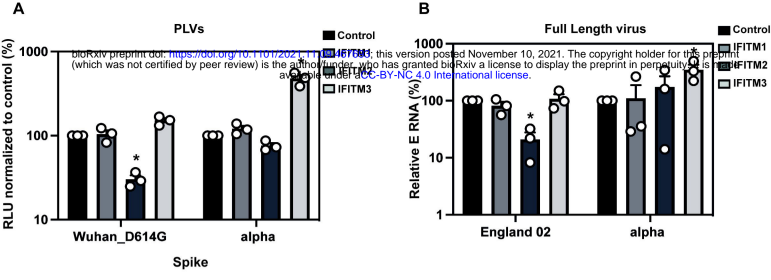
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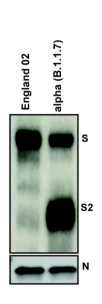


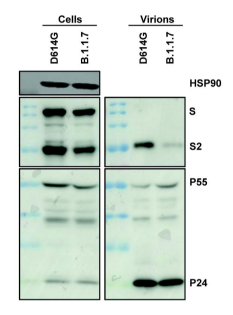


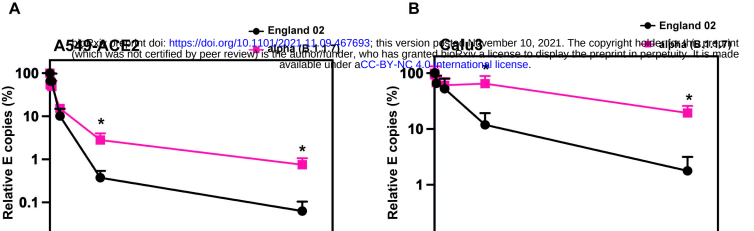
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Full Length virus

PLVs







0.01

С

100

Relative E mRNA (%)

10+0

0

Calu3

40 IFNß (U/mL)

60

alpha (B.1.1.7) clinical isolate 10

✓ alpha (B.1.1.7) clinical isolate 28 ★

- England 02 - alpha (B.1.1.7)

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-

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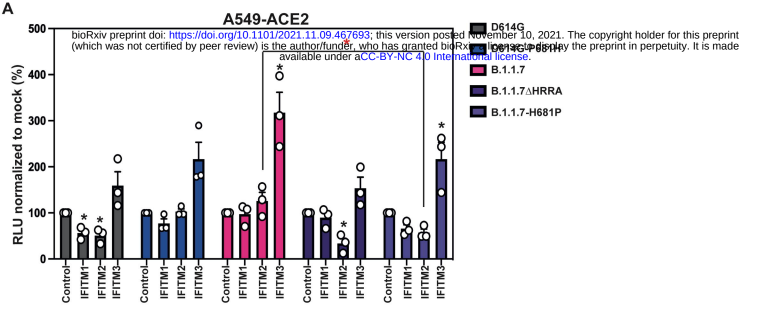
0.1

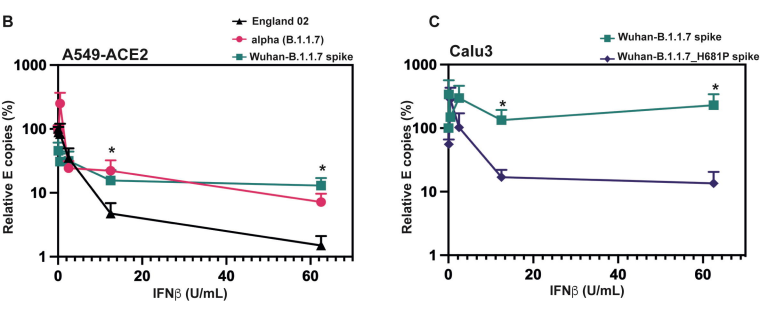
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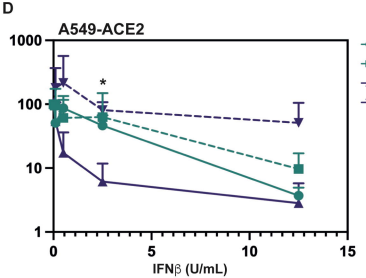
40 IFNß (U/mL) 60

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IFNß (U/mL)

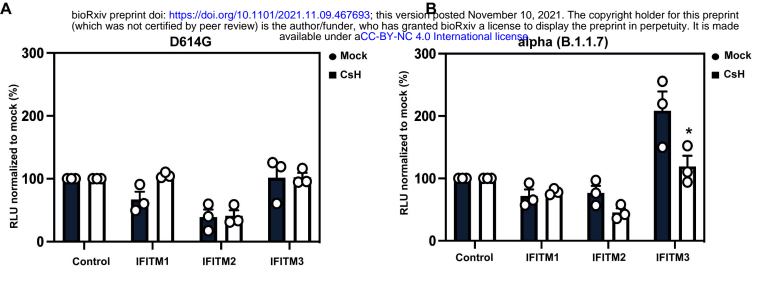






- Wuhan-B.1.1.7 spike/Control siRNA

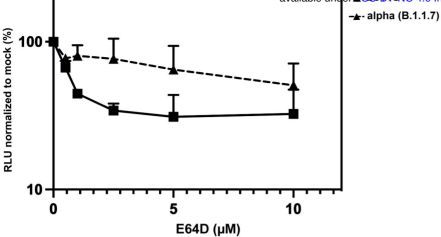
Figure 4



Supplementary Fig1

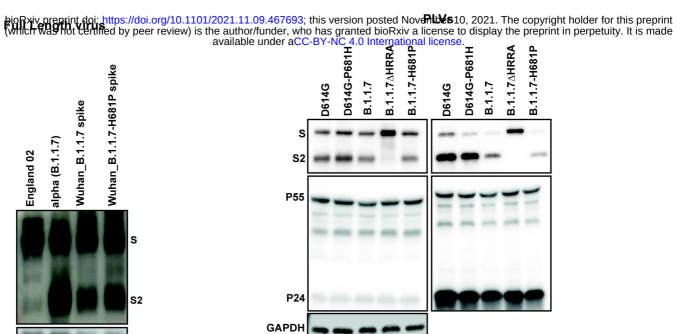






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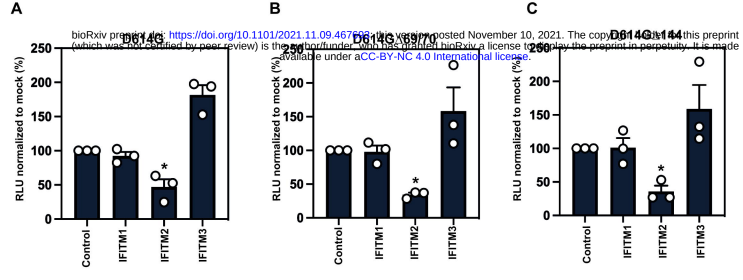
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CELLS

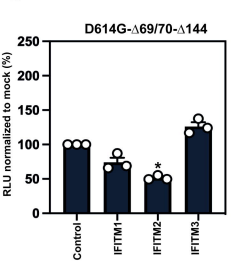
VIRIONS

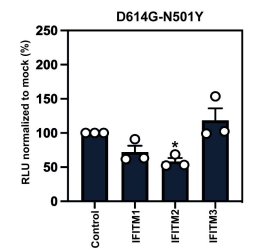
Supplementary Fig3

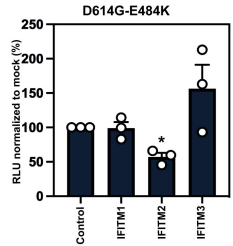


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