1 The effect of molnupiravir and nirmatrelvir on SARS-CoV-2

2 genome diversity in infected and immune suppressed mice.

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- Rebekah Penrice-Randal^{1*†}, Eleanor G. Bentley^{1*}, Parul Sharma¹, Adam Kirby¹, I'ah
 Donovan-Banfield^{1,2}, Anja Kipar^{1,3}, Daniele F. Mega¹ Chloe Bramwell^{1,4}, Joanne
 Sharp⁴, Andrew Owen^{4,5}, Julian A. Hiscox^{1,2,6} and James P. Stewart^{1,5}.
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- ¹Department of Infection Biology and Microbiomes, University of Liverpool, Liverpool,
 UK.
- 10 ²NIHR Health Protection Research Unit in Emerging and Zoonotic Infections,
- 11 Liverpool, UK.
- ¹² ³Laboratory for Animal Model Pathology, Institute of Veterinary Pathology, Vetsuisse
- 13 Faculty, University of Zurich, Switzerland.
- ⁴Department of Pharmacology and Therapeutics, University of Liverpool, UK.
- ⁵Centre of Excellence in Long-acting Therapeutics (CELT), University of Liverpool,
 UK.
- 17 ⁶A*STAR Infectious Diseases Laboratories (A*STAR ID Labs), Agency for Science,
- 18 Technology and Research (A*STAR), Singapore.
- 19
- 20 *These authors contributed equally.
- 21 [†] Corresponding author: <u>rebee@liverpool.ac.uk</u>
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- 23 Running title: Antiviral and SARS-CoV-2 genome diversity

24 Synopsis

Objectives. Immunocompromised individuals are susceptible to severe COVID-19 and potentially contribute to the emergence of variants with altered pathogenicity due to persistent infection. This study investigated the impact of immunosuppression on SARS-CoV-2 infection in k18-hACE2 mice and the effectiveness of antiviral treatments in this context.

Methods Mice were immunosuppressed using cyclophosphamide and infected with a
B lineage of SARS-CoV-2. Molnupiravir and nirmatrelvir, alone and in combination,
were administered and viral load and viral sequence diversity was assessed.

33 **Results** Treatment of infected but immune compromised mice with both compounds either singly or in combination resulted in decreased viral loads and pathological 34 35 changes compared to untreated animals. Treatment also abrogated infection of 36 neuronal tissue. However, no consistent changes in the viral consensus sequence 37 were observed, except for the emergence of the S:H655Y mutation. Molnupiravir, but 38 not nirmatrelvir or immunosuppression alone, increased the transition/transversion (Ts/Tv) ratio, representative of A>G and C>U mutations and this increase was not 39 altered by the co-administration of nirmatrelvir with molnupiravir. 40

41 Notably, immunosuppression itself did not appear to promote the emergence of
42 mutational characteristic of variants of concern (VOCs).

Conclusions Further investigations are warranted to fully understand the role of immunocompromised individuals in VOC development and to inform optimised public health strategies. It is more likely that immunodeficiency promotes viral persistence but does not necessarily lead to substantial consensus-level changes in the absence of antiviral selection pressure. Consistent with mechanisms of action, molnupiravir showed a stronger mutagenic effect than nirmatrelvir in this model.

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50 Keywords

SARS-CoV-2, COVID-19, immunocompromised, intra-host evolution, Molnupiravir,
Nirmatrelvir, Paxlovid.

53 Introduction

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Unsurprisingly, since the start of the SARS-CoV-2 pandemic and the first deposited genome sequences, and like other coronaviruses, SARS-CoV-2 has diverged through single nucleotide polymorphism, and homologous and heterologous recombination applications resulting in insertions and deletions ^{1, 2}. Over the course of the pandemic changes that have dominated have resulted in increased transmissibility such as the P323L/D614G changes in early 2020 ³⁻⁵, immune-evasion ⁶ and altered pathogenicity ⁷.

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Founder effects, population bottlenecks, selection pressures and behaviour have 63 contributed to the diversification of the SARS-CoV-2 genome but also to the apparent 64 65 waves of different variants. Several Variants of Concern (VoCs) have arisen that have 66 a transmission advantage and/or potential immune evasion. Some reports have suggested that such variants may have arisen in hosts with compromised immunity 67 68 and/or persistent infections, where infection leads to the generation of more diverse variants through longer viral evolution within an individual⁸. This includes a changing 69 landscape of dominant viral genome sequence and minor genomic variants in immune 70 compromised individuals e.g. in a patient with cancer ⁹. Changes within the individual 71 72 mapped to several different regions on the SARS-CoV-2 genome including the spike glycoprotein and orf8. 73

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Complicating the picture of potential rapid and dramatic genomic change in immune compromised hosts is that similar changes can be observed in immune competent patients. This can be either as part of the dominant genomic sequence ¹⁰ or minor variant genomes ¹. Indeed, genomic variants with deletions can be identified in the minor genomic variant population of Middle East respiratory syndrome coronavirus (MERS-CoV) from patients ¹¹ and as part of the dominant genomic sequence in camels ^{12, 13}.

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Parallels with other animal coronaviruses can be found where persistent infections are established, and this might be associated in pathogenicity; an example are feline coronavirus (FCoV) infections and feline infectious peritonitis (FIP) ¹⁴⁻¹⁷. Thus, one

concern with long term persistence of SARS-CoV-2 in immune compromised patients
 is that new transmissible variants could emerge ⁸.

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Three small molecule direct acting anti-virals (DAAs) have received early use 89 authorisation for the treatment of COVID-19: remdesivir, molnupiravir (both nucleoside 90 91 analogues which target viral nucleic acid synthesis) and nirmatrelvir (which targets the main viral protease). Unlike remdesivir, molnupiravir and nirmatrelvir are orally 92 administered and thus more readily deployed for treatment in the community. 93 Nirmatrelvir is packaged with ritonavir (as Paxlovid), this later molecule acting as a 94 pharmacokinetic boosting agent to inhibit P450 (CYP) 3A4. However, adequate 95 nirmatrelvir plasma concentrations can be achieved in mice without the need for 96 ritonavir boosting. In cell culture single or combination treatment can result in 97 decreased viral replication ^{18, 19} and a natural extension is that such anti-virals may be 98 deployed as combination therapy to reduce the emergence of resistant genotypes ²⁰. 99 Resistant genotypes/phenotypes have been identified in vitro for remdesivir ²¹. 100 101 Molnupiravir has previously been shown to enhance viral transition/transversion mutations in a phase II clinical trial ²² and a molnupiravir associated signature has 102 103 been identified in circulating SARS-CoV-2 lineages since the introduction of molnupiravir in 2022²³. 104

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Immunocompromised patients with a SARS-CoV-2 infection are treated as a priority 106 107 with anti-virals, including those compounds that generically target virus replication by causing hyper-mutation or specifically preventing the function of a viral protein critical 108 109 to the life cycle of the virus. Such anti-virals may be deployed as combination therapy to reduce the emergence of resistant genotypes ²⁰ and may be particularly relevant 110 for patients with compromised immunity ²⁴. However, in the latter patients, anti-virals 111 may decrease viral loads but enhance genomic plasticity. To investigate this, the 112 genomic variation of SARS-CoV-2 was evaluated in an immune compromised host, in 113 the absence and presence of medical countermeasures. We have developed animal 114 models of COVID-19 to be able to assess pathogenicity of new variants and develop 115 interventions ²⁵⁻²⁷. An immune suppressed K18-hACE2 transgenic mouse model was 116 used to simulate patients with severe COVID-19^{28, 29}. Two anti-virals, molnupiravir 117 and nirmatrelvir, were evaluated either singly or in combination. 118

119 Methods

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121 Animal infection and treatment

A UK variant of SARS-CoV-2 (hCoV-2/human/Liverpool/REMRQ0001/2020), was
 used as described previously ^{30, 31}.

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Animal work was approved by the local University of Liverpool Animal Welfare and Ethical Review Body and performed under UK Home Office Project Licence PP4715265. Transgenic mice carrying the human ACE2 gene under the control of the keratin 18 promoter (K18-hACE2; formally B6.Cg-Tg(K18-ACE2)2Prlmn/J) were purchased from Jackson Laboratories (France) at 8 – 10 weeks of age. Mice were maintained under SPF barrier conditions in individually ventilated cages and underwent a week of acclimatisation in these conditions prior to experimental use.

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Experimental design is shown in Fig. 1 and treatment groups detailed in Table 1. 133 134 Animals were randomly assigned into multiple cohorts of four animals using a random number generator. For operational reasons at high containment the treatment groups 135 136 were not blinded during the experiment. Sample size was determined using prior 137 experience of similar experiments with SARS-CoV-2. For SARS-CoV-2 infection, mice 138 were anaesthetized lightly with isoflurane and inoculated intra-nasally with 50 µl containing 10⁴ PFU SARS-CoV-2 in PBS as described previously ²⁶. Some cohorts of 139 140 mice were immunosuppressed by treatment with cyclophosphamide (100 mg/kg) intraperitoneally (IP) at day -4 and -1 pre-infection. Molnupiravir was made up in 10% 141 142 PEG400 and 2.5% cremophor in water and used at 100 mg/kg. Nirmatrelvir was 143 dissolved in 2% Tween 80 in 98% (v/v) of 0.5% methyl cellulose and used at 500 mg/kg. Both drugs were administered via the oral route one hour prior to infection and 144 then twice daily up to 4 days post-infection via the oral (PO) route. Groups of animals 145 were kept in the same cages during the experiment and were always weighed and 146 treated in the same order. Mice were sacrificed at day 6 (vehicle and 147 cyclophosphamide treated group) or 7 (all others) after infection by an overdose of 148 pentobarbitone. Weights were recorded daily, and tissues were removed immediately 149 for downstream processing. The right lung and nasal turbinates were frozen at -80 °C 150 until further processing. The left lung and heads were fixed in 10% neutral buffered 151 formalin for 24-48 h and then stored in 70%. No data were excluded from the analyses. 152

153 Histology, immunohistology and morphometric analysis

The fixed left lung was routinely paraffin wax embedded. Heads were sawn 154 longitudinally in the midline using a diamond saw (Exakt 300; Exakt) and the brain left 155 in the skull. Heads were gently decalcified in RDF (Biosystems) for twice 5 days, at 156 room temperature and on a shaker, then both halves paraffin wax embedded. 157 158 Consecutive sections (3-5 µm) were either stained with hematoxylin and eosin (HE) or used for immunohistology (IH). IH was performed to detect viral antigen expression 159 using the horseradish peroxidase method and a rabbit anti-SARS-CoV nucleocapsid 160 protein (Rockland, 200-402-A50) as primary antibody, as previously described ^{26, 32,} 161 33 162

For morphometric analysis, the immunostained sections were scanned (NanoZoomer-163 XR C12000; Hamamatsu, Hamamatsu City, Japan) and analysed using the software 164 165 program Visiopharm (Visiopharm 2020.08.1.8403; Visiopharm, Hoersholm, Denmark) 166 to quantify the area of viral antigen expression in relation to the total area (area occupied by lung parenchyma) in the sections. This was used to compare the extent 167 168 of viral antigen expression in the lungs between the different treatment groups. A first app was applied that outlined the entire lung tissue as ROI (total area). For this a 169 170 Decision Forest method was used and the software was trained to detect the lung 171 tissue section (total area). Once the lung section was outlined as ROI the lumen of 172 large bronchi and vessels was manually excluded from the ROI. Subsequently, a second app with Decision Forest method was trained to detect viral antigen expression 173 174 (as brown DAB precipitate) within the ROI.

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176 **qRT-PCR for viral load**

Viral loads were quantified using the GoTaq® Probe 1-Step RT-qPCR System
(Promega). For quantification of SARS-COV-2 the nCOV_N1 primer/probe mix from
the SARS-CoV-2 (2019-nCoV) CDC qPCR Probe Assay (IDT) were utilised and
murine 18S primers as described previously ^{25, 26}.

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182 Sequencing of SARS-CoV-2

Library preparation consisted of converting RNA to cDNA using LunaScript[™]
(Thermofisher), then amplified by reverse complement (RC)-PCR amplification
(EasySeq[™] SARS-CoV-2 Whole Genome Sequencing kit, Nimagen, Netherlands).
This kit barcodes and ligates Illumina adapters in a single PCR reaction, with two

187 separate pools of primers (pools 1 and 2). After amplification, each amplicon library 188 was pooled 1:1 before being cleaned with AmpliCleanTM beads and quantification. The 189 two pools were then added together and denatured. Finally, the denatured amplicon 190 library was loaded into the NovaSeq cartridge (2 x 150 bp run).

191

192 **Bioinformatics**

Supplementary Fig. S2 provides an overview of the workflow used in this study. In 193 short, raw paired end fast files were inputted into the EasySeg pipeline to generate 194 alignment files, vcf's and consensus sequences ³⁴.Consensus sequences were 195 inputted into Nextclade for lineage assignment and bam files were inputted into 196 (https://github.com/josephhughes/DiversiTools) to 197 **DiversiTools** assess minor variation. Sequencing data was analysed as previously described and statistical 198 analysis and visualisation was performed in R²². Raw fastg files are available under 199 200 SRA Project Accession: PRJNA886870. Code for analysis and figure generation is available at https://github.com/Hiscox-lab/viral-genomics-immunosupression-and-201 202 countermeasures.

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204 Statistics

Graphs were prepared and statistics performed using Prism 10 (Graphpad Inc). *P* values were set at 95% confidence interval. A repeated-measures two-way ANOVA (Bonferroni post-test) was used for time-courses of weight loss; log-rank (Mantel-Cox) test was used for survival curve and Mann-Whitney *U* test for side-by-side comparisons. All differences not specifically stated to be significant were not significant (p > 0.05). For all figures, *p < 0.05.

211 Results and Discussion

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Since the emergence of the Alpha VOC there has been discussion on the involvement 213 of the immunocompromised host and the generation of variants ^{8, 35-39}. There are many 214 the follow SARS-CoV-2 evolution 215 studies in literature that in case 216 immunocompromised hosts, however, little has been explored experimentally. In this study, mice were chemically immunocompromised with cyclophosphamide which is 217 known to efficiently remove adaptive immunity in the form of B and T cells ⁴⁰. 218 Additionally, therapeutic agents, molnupiravir and nirmatrelvir, were used 219 220 independently and in combination to determine the effectiveness of these compounds in an immunocompromised model, and the impact of these compounds on viral 221 222 sequence diversity.

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224 Modelling an immunocompromised state in animal models in the context of SARS-CoV-2 is important for the consideration of countermeasures that may be utilised for 225 226 humans who are considered vulnerable. Cyclophosphamide has been used previously to study the impact of immunosuppression in a hamster model ⁴¹⁻⁴³, where intranasally 227 228 infected hamsters with cyclophosphamide treatment before infection had prolonged 229 weight loss and an inadequate neutralising antibody response to SARS-CoV-2. 230 Distinct transcriptional profiles were identified between immunocompetent and immunosuppressed animals; however, the impact of antivirals or viral genome 231 232 diversity was not investigated.

233

234 To investigate the frequency of genomic changes that occur in SARS-CoV-2 in the 235 immune compromised or competent host in the presence or absence of antiviral drugs, K18-hACE2 transgenic mice were used as a model for severe SARS-CoV-2 infection 236 in humans ⁴⁴. We have found that the pathological changes in the lungs in this model 237 in many aspects resemble those in humans who have died of severe COVID-19^{26, 28,} 238 ^{29, 32, 33}. To mimic a host with compromised immunity, an experimental protocol was 239 developed in which mice were exposed to cyclophosphamide ⁴⁰ (Fig. 1, Table 1). 240 Several anti-viral regimes in humans were simulated in the mouse model by giving a 241 human equivalent dose of either molnupiravir (100 mg/kg), nirmatrelvir (500 mg/kg) or 242 both in combination. This included prophylactic followed by therapeutic treatment. 243 Mice were infected with 10⁴ PFU of SARS-CoV-2. 244

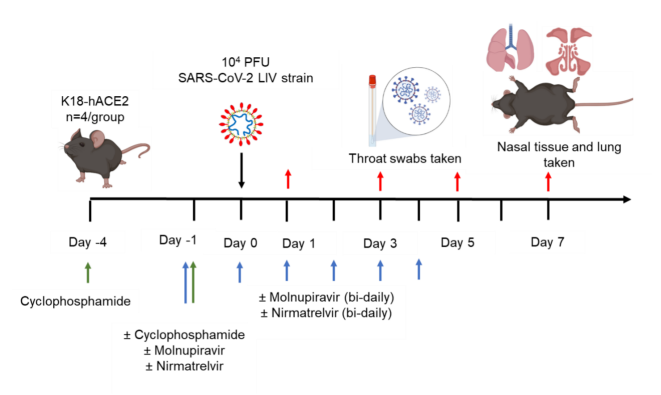


Figure 1. Schematic diagram of the experimental design for infection of immune compromised K18-hACE2 mice with SARS-CoV-2 and evaluation of two antiviral drugs given at a human equivalent dose; molnupiravir, a broad acting compound causing error catastrophe, or nirmatrelvir which specifically targets the viral 3C-like protease. Cyclophosphamide was used at 100 mg/kg via the intraperitoneal route to immunosuppress mice. Molnupiravir was used at 100 mg/kg and nirmatrelvir at 500 mg/kg both via the oral route. Effects of infection and treatment were evaluated by measuring the weight of the mice daily, determining viral loads in sequential oral/throat swabs and at day 7 post-infection, and examining nose, brain and lung at day 7 post infection for any histological changes and the expression of SARS-CoV-2 nucleoprotein.

245

246 Table 1. Treatment groups for in vivo analysis

Group Treatment

- **1** Control (vehicle)
- 2 Cyclophosphamide
- 3 Molnupiravir
- 4 Cyclophosphamide + molnupiravir
- **5** Cyclophosphamide + nirmatrelvir
- 6 Cyclophosphamide + molnupiravir + nirmatrelvir
- 247
- 248

Treatment with Molnupiravir or Nirmatrelvir either individually or in combination provides recovery in immune compromised mice infected with SARS-CoV-2.

Cyclophosphamide treatment prior to SARS-CoV-2 infection of hACE2 mice led to a 251 more pronounced early weight loss in comparison to immunocompetent mice, a 252 253 phenomenon previously reported in hamsters ⁴³. This was not associated with earlier mortality than in vehicle treated immunocompetent mice, although in human, a 254 delayed adaptive immune response has been shown to be associated with fatality in 255 256 COVID-19 patients, which may have been observed over longer timeframes ⁴⁵. Daily weighing of the animals indicated that all groups lost body weight after day 1 (Fig. 2). 257 We attribute this to aversion to eating as all therapies were applied by gavage. 258 However, starting at day 3 all groups, except for mice exposed to cyclophosphamide, 259 260 or mice exposed to cyclophosphamide and treated with molnupiravir, started to gain, or stabilise weight. By days 5 and 6 a clear pattern had emerged where all groups 261 262 treated with molnupiravir or nirmatrelvir either individually or in combination had regained their starting weight. The exception to this were mice exposed to vehicle only 263 264 (controls) or cyclophosphamide; these reached a humane end point on day 6 (Fig. 2). Comparison of survival curves again indicated that immune compromised animals 265 266 treated either singly or in combination with each therapeutic went on to survive (Fig. 267 3).

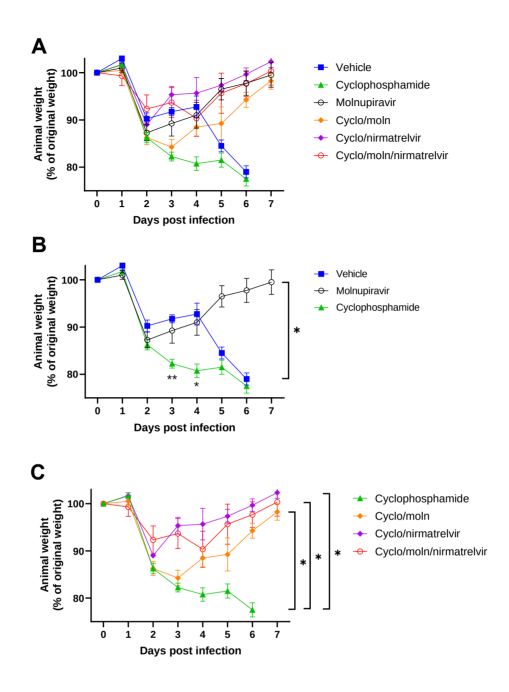


Figure 2: Treatment of SARS-CoV-2-infected immunocompromised mice leads 269 to decreased weight loss. K18-hACE2 mice were challenged intranasally with 10⁴ 270 271 PFU SARS-CoV-2 and their body weight monitored at indicated time-points (n = 4). Data represent the mean residual weight \pm SEM. Comparisons were made using a 272 repeated-measures two-way ANOVA (Bonferroni post-test). * Represents P < 0.05. 273 Data from the same experiment were presented differently grouped in three separate 274 graphs for clarity. (A) Curves for all groups. (B) Curves for vehicle, cyclophosphamide 275 and molnupiravir groups. Asterisks below the curves represent * P < 0.05 and ** P < 276 0.01 between the cyclophosphamide and vehicle groups. (C) Curves for the groups 277 278 treated with cyclophosphamide. Panels B and C were plotted using data shown in A 279 but for added clarity.

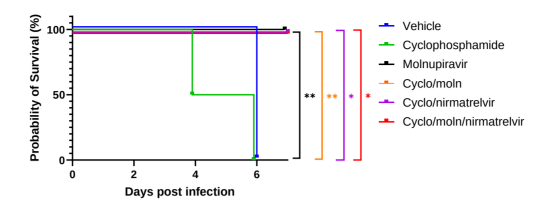




Figure 3: Treatment of SARS-CoV-2-infected mice leads to enhanced survival. K18-hACE2 mice were challenged intranasally with 10⁴ PFU SARS-CoV-2. Survival was assessed at indicated time points and significance determined using log rank (Mantel-Cox) test (n = 4).

Viral load decreases in immune compromised mice treated with Molnupiravir or Nirmatrelvir either individually or in combination.

Viral load in terms of copy numbers of the SARS-CoV-2 genome were calculated for 288 289 throat swabs during infection and compared to nasal tissue and lung tissue at the end 290 of the experiment. The data indicated that for throat swabs on days 1 and 3 post-291 infection there was a significant decrease in viral load in animals treated with molnupiravir or nirmatrelvir either individually or in combination compared to untreated 292 controls (Figure 4A). At day 3 there was a significant difference between both 293 compounds used in combination and molnupiravir only (Figure 4A). No significant 294 295 differences were observed between vehicle control and cyclophosphamide only 296 groups.

297

Comparison of viral loads and titres in nasal and lung tissue respectively (Figure 4B 298 and 4C, respectively) at day 7 post-infection reflected that there was a significantly 299 lower viral load in animals treated with molnupiravir or nirmatrelvir either individually 300 301 or in combination compared to untreated mice. However, nirmatrelvir treatment resulted in a greater decrease in viral load compared to molnupiravir. The 302 molnupiravir/nirmatrelvir combination was also more effective at decreasing viral load 303 than either drug alone, but this was only statistically significant in the case of 304 molnupiravir vs the drug combination. 305

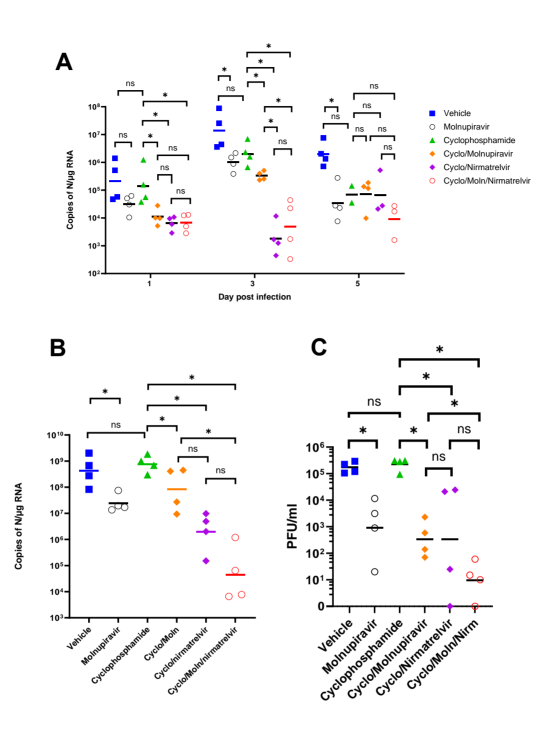
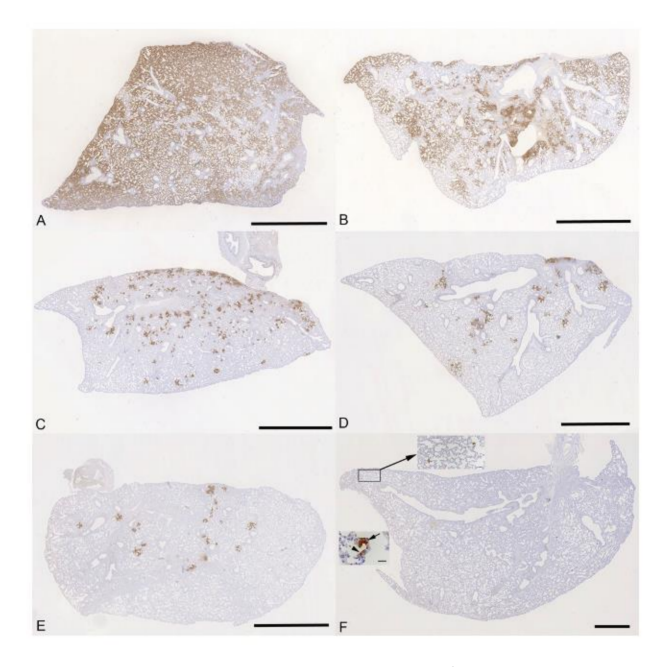


Figure 4. Viral loads in swabs and tissues. K18-hACE2 mice were challenged 307 intranasally with 10^4 PFU SARS-CoV-2 and treated as indicated (n = 4 per group). 308 309 RNA extracted from oral/throat swabs and nasal tissue was analysed for virus RNA load using gRT-PCR and primers specific for the SARS-CoV-2 N gene. Assays were 310 normalised relative to levels of 18S RNA. Lung tissue was analysed for live virus by 311 plaque assay. Data for individual animals are shown with the median value 312 represented by a black line. (A) Throat swabs; (B) nasal tissue; (C) lung tissue. 313 Comparisons were made using two-way ANOVA (Bonferroni post-test) in panel A and 314 315 Mann-Whitney U test (Panels B and C). * Represents p < 0.05. 316

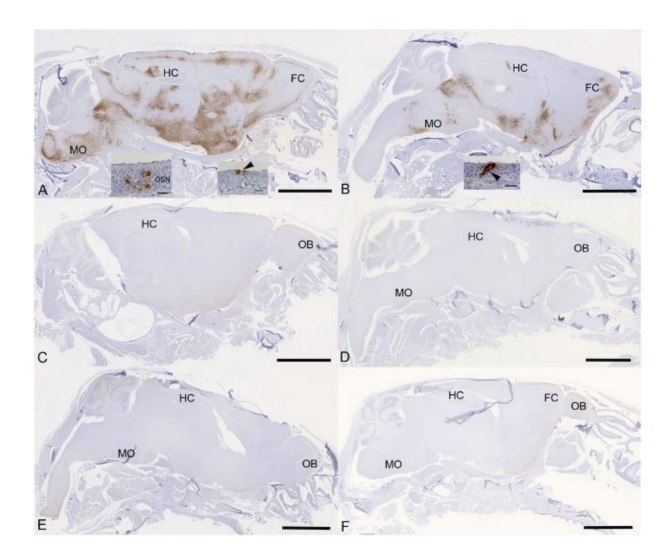
317 Treatment with molnupiravir or nirmatrelvir or both in combination results in 318 marked reduction of pulmonary infection and inhibits viral spread to the brain.

The lung, nose and brain of all animals were examined for any histopathological 319 changes and the expression of viral antigen by immunohistology, to determine 320 whether treatment of the animals with molnupiravir and/or nirmatrelvir influenced the 321 322 outcome of infection. The lungs of vehicle treated, immunocompetent animals showed the typical changes previously reported in K18-hACE2 mice infected with this virus 323 324 strain ²⁶, i.e. multifocal areas with pneumocyte degeneration, type II pneumocyte activation, mild neutrophil infiltration, and mild vasculitis, with a diffuse increase in 325 interstitial cellularity and widespread SARS-CoV-2 antigen expression in alveolar 326 epithelial cells (Fig. 5A). In mice that had received cyclophosphamide alone, the 327 changes were very similar, but slightly less widespread, with some unaltered 328 parenchyma and less extensive viral antigen expression (Fig. 5B). With molnupiravir 329 330 treatment, both inflammatory processes and viral antigen expression were markedly decreased; indeed, SARS-CoV-2 antigen was only found in disseminated patches of 331 332 alveoli with positive pneumocytes (Fig. 5C). With cyclophosphamide and molnupiravir treatment, the lung parenchyma was widely unaltered, and there were only small 333 334 patches of inflammation and alveoli with viral antigen expression, respectively (Fig. 335 5D). These were further reduced in number and size in animals that had received 336 cyclophosphamide and nirmatrelvir (Fig. 5E). Treatment with all three compounds, cyclophosphamide, molnupiravir and nirmatrelvir, resulted in widely unaltered lung 337 338 parenchyma with no or minimal viral antigen expression (Fig. 5F). The morphometric 339 analysis to quantify the extent of viral antigen expression in the lungs in the different 340 groups of animals confirmed that the various antiviral treatment regimens significantly 341 reduced the extent of lung infection (Figure S1).



343 Figure 5: K18-hACE2 mice were challenged intranasally with 10⁴ PFU SARS-CoV-2 and treated as indicated below (n = 4 per group). Immunohistology for the detection of viral antigen 344 345 in the lung at day 6 or 7 post infection. Sections from the formalin-fixed, paraffin embedded 346 left lung lobe were stained using anti-SARS-CoV nucleoprotein and counterstained with 347 hematoxylin. Representative images from the individual treatment groups are shown as follows: A. vehicle; B. cyclophosphamide; C. molnupiravir; D. cyclophosphamide and 348 349 molnupiravir; E. cyclophosphamide and nirmatrelvir; F. cyclophosphamide, molnupiravir and 350 nirmatrelvir. Viral antigen expression is restricted to pneumocytes in a few individual alveoli 351 (higher magnifications in insets). Bars represent 2.5 mm (A-E), 1 mm (F) and 20 µm (F, insets).

Examination of the heads using longitudinal sections (midline) revealed consistent and 352 widespread infection of the brain in animals treated with the vehicle or with 353 cyclophosphamide alone (Fig. 6A, B); this was associated with mild perivascular 354 mononuclear infiltration in particular in the brain stem, as described before in K18-355 hACE2 mice infected with this virus strain ³³. In both groups of animals, 356 357 immunohistology confirmed viral antigen expression in the respiratory and/or olfactory epithelium, in the latter with evidence of infection in olfactory sensory neurons (Fig. 358 6A, B). In the other groups, there was no evidence of viral infection of the brain (Fig. 359 6C-F), and viral antigen expression in the nasal mucosa was not seen or restricted to 360 361 scattered individual epithelial cells. In vehicle control and cyclophosphamide mice, the 362 nasal mucosa harboured viral antigen at this stage, in the respiratory epithelium and in the olfactory epithelium; in the latter it also appeared to be present in sensory 363 364 neurons. Consequently, the virus had reached and spread widely in the brain where it 365 was detected in neurons; the infection was associated with mild inflammatory response in particular in the brain stem, as described before in K18-hACE2 mice 366 367 infected with this virus strain ^{26, 33}. After treatment with all three compounds, cyclophosphamide, molnupiravir and nirmatrelvir, the lung parenchyma was basically 368 369 unaltered, with no or minimal viral antigen expression. In all groups of mice, viral 370 antigen expression in the nasal mucosa was not seen or restricted to scattered 371 individual epithelial cells and there was no evidence of viral infection of the brain, suggesting that the antiviral treatment blocked infection of the brain. Whether the latter 372 373 is purely a consequence of reduced viral replication in the upper respiratory tract cannot be assessed in the present study; it does, however, appear likely. 374



375

376 Figure 6: K18-hACE2 mice were challenged intranasally with 10⁴ PFU SARS-CoV-2 and 377 treated as indicated below (n = 4 per group). Immunohistology for the detection of viral antigen 378 in the brain and nose at day 6 or 7 post infection. Sections from formalin-fixed, decalcified and 379 paraffin embedded heads after longitudinal sawing in the midline were stained using anti-380 SARS-CoV nucleoprotein, and counterstained with hematoxylin. Only small fragments of 381 nasal mucosa were available for the examination, as the nasal turbinates had been sampled 382 for PCR. Representative images from the individual treatment groups are shown as follows: 383 A. Vehicle. There is widespread infection of the brain. The insets show infection of individual cells with the morphology of olfactory sensory neurons and epithelial cells in the olfactory 384 385 epithelial layer (left inset) and individual respiratory epithelial cells in the nasal mucosa (arrowhead; right inset); B. Cyclophosphamide. There is widespread infection of the brain. 386 387 The inset shows a group of positive epithelial cells/sensory neurons in the olfactory epithelial 388 layer (arrowhead); C. Molnupiravir. There is no evidence of brain infection. D. 389 Cyclophosphamide and molnupiravir. There is no evidence of brain infection. E. 390 Cyclophosphamide and nirmatrelvir. There is no evidence of brain infection. **F**. Cyclophosphamide, molnupiravir and nirmatrelvir. There is no evidence of brain infection. Bars 391 392 represent 2.5 mm (A-F) and 20 µm (A, B insets). FC – frontal cortex, HC – hippocampus, MO 393 - medulla oblongata, OB - olfactory bulb, OSN - olfactory sensory neurons.

394 Evaluation of dominant and minor variants in SARS-CoV-2

To determine the impact of immunosuppression on viral diversity, 116 RNA samples 395 from swabs and tissue were sequenced and analysed using the EasySeg WGS 396 397 protocol by Nimagen. alignment files and associated index files were inputted into DiversiTools to provide mutation data and outputs were analysed in R. Samples with 398 399 less than 90% breadth of coverage were discarded for mutational analysis (n=12), as 400 well as samples that returned bad or mediocre quality scores in nextclade (n=13). The samples that were excluded were associated with higher Ct values and later time 401 points belonging in the nirmatrelvir treatment groups. Sequencing data from 89 402 403 samples were taken forward in the analysis (swab, n=50, tissue n=39, Supplementary 404 Table S1).

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406 The input virus contained 5 substitutions and 3 amino acid substitutions in comparison 407 to the reference sequence and were thus not considered as changes during the analysis (Supplementary Table S2). The S: H655Y mutation was present in 76% of 408 409 the genomes that passed QC at the dominant level and observed as a minor variant across all samples (Supplementary Fig. S3). This mutation has been reported 410 previously as a spike adaptation to other species such as cats, hamsters, and mink ⁴⁶⁻ 411 ⁴⁸ and of course has independently arisen in human lineages such as Omicron ⁴⁹. As 412 413 this mutation was clearly associated with a species adaptation, it was disregarded for the evaluation of treatment and immune status driven mutations. The other mutations 414 415 appear to be novel at the time of writing; however, no distinct group was associated with driving these mutations, and can be overall interpreted as a rare event. The 416 417 sequences showing the highest number of mutations were sequences derived from 418 tissue samples. Species specific adaptations were more frequently reported in the dataset than the immunocompromised and antiviral environments, putting the 419 420 evolutionary pressures into perspective.

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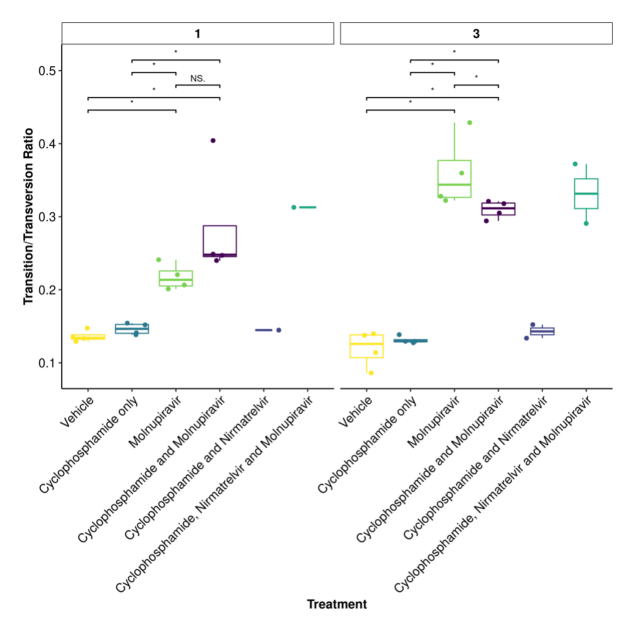
422 Molnupiravir increases the Ts/Tv ratio at the minor variant level in genomes 423 derived from swabs

To further assess the impact of immunocompromising mice by cyclophosphamide, and the therapeutic agents molnupiravir and the nirmatrelvir, a minor variant analysis was conducted on samples derived from throat swabs. The average transition/transversion (Ts/Tv) ratio for SARS-CoV-2 genomes from each mouse and

the mean of each group was compared across cohorts. On day 1, an increase in Ts/Tv 428 ration was observed in the molnupiravir cohort and the cyclophosphamide and 429 molnupiravir cohort and had a p value < 0.05 when compared to the vehicle control 430 and cyclophosphamide only groups (Fig. 7). The number of samples analysed for 431 cyclophosphamide and nirmatrelvir only was too small for statistical analysis, however, 432 433 the trend resembles that of vehicle and cyclophosphamide only. Likewise, the combined cyclophosphamide and molnupiravir and nirmatrelvir cohort only resembles 434 one genome, however, the trend resembles that of other genomes with exposure to 435 molnupiravir. The same is observed at day 3 of sampling, however, there is a 436 437 significant difference between the mean Ts/Tv ratio between the molnupiravir only and cyclophosphamide and molnupiravir groups. Importantly, the Ts/Tv ratios between the 438 vehicle control and cyclophosphamide only groups resemble each other. The 439 440 proportion of base changes were also observed, with particular interest in the C to U and G to A transitions as previously seen in a phase II clinical trial ²² (Figure 8). 441

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443 Further investigations are warranted to understand completely the role of 444 immunocompromised individuals in the development of SARS-CoV-2 variants. It is 445 more likely that immunodeficiency promotes viral persistence providing the virus 446 more opportunity to replicate and introduce mutations. Molnupiravir, compared to 447 nirmatrelvir, shows a stronger mutagenic effect in this model at the minor variant level, however, data is insufficient to make conclusions regards consensus level 448 449 changes over the timeframes used in this study. When these therapies are used 450 individually or in combination, there is successful depletion in viral load and animals 451 recover from infection, whilst preventing infiltration into brain tissue. Given the concern of molnupiravir associated lineages in circulation ²³, combination therapy 452 may reduce this through more effective clearance of the virus ²⁰, although this would 453 need to be evaluated over time in a real-world setting as the mutational signatures 454 455 were observed in the combined therapy group.



457 Figure 7: The mean Ts/Tv ratio per genome plotted as boxplots. The plot is facetted by day 458 post infection. Less genomes were recovered for cyclophosphamide and nirmatrelvir and 459 cyclophosphamide, nirmatrelvir and molnupiravir, therefore statistical analysis returns the 460 differences as non-significant. Trends can be concluded with caution. * Represents a P value 461 <0.05 (Mann Whitney U test).</p>

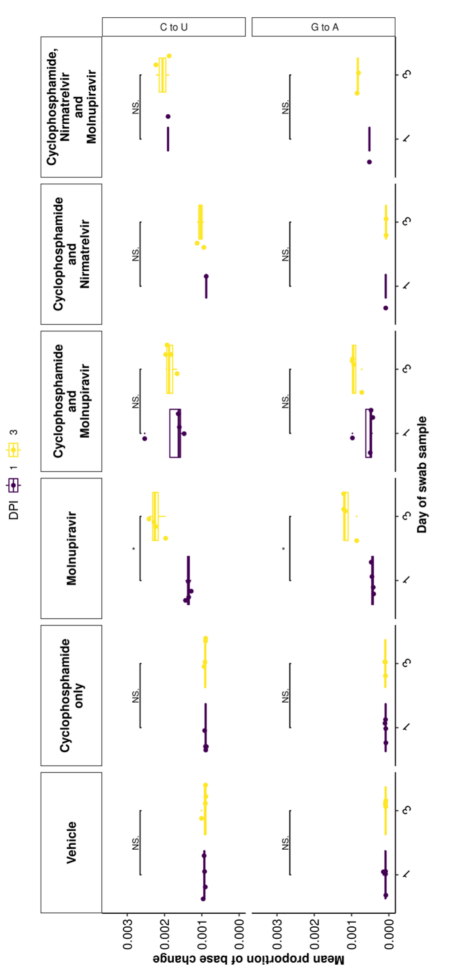


Figure 8: C to U and G to A minor variation changes significantly increased between day 1 and day 3 post infection in the molnupiravir only group. A similar trend is observed between other groups including molnupiravir treatment, however, the change is not reported as significant. * represents a P value <0.05 (Mann Whitney U test).

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488 Transparency Declaration

A.O. is a director of Tandem Nano Ltd and co-inventor of patents relating to drug 489 delivery. A.O. has been co-investigator on funding received by the University of 490 Liverpool from ViiV Healthcare and Gilead Sciences in the past 3 years unrelated to 491 COVID-19. A.O. has received personal fees from Gilead and Assembly Biosciences 492 in the past 3 years, also unrelated to COVID-19. JPS has received funding from ENA 493 494 respiratory Pty Ltd, Bicycle Tx Ltd, and Infex Therapeutics Ltd unrelated to this study. R.P.R. is an employee at TopMD Precision Medicine Ltd. No other conflicts are 495 496 declared by the authors.

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