1	Broad anti-coronaviral activity of FDA approved drugs against
2	SARS-CoV-2 in vitro and SARS-CoV in vivo
3	
4	Stuart Weston ¹ , Christopher M. Coleman ^{1†} , Rob Haupt ¹ , James Logue ¹ , Krystal Matthews ¹
5	and Matthew B. Frieman ^{*1}
6	1 - Department of Microbiology and Immunology, University of Maryland School of Medicine,
7	685 W. Baltimore St., Room 380, Baltimore, MD, 21201, USA
8	*Corresponding author. Email: MFrieman@som.umaryland.edu
9	† Current address: School of Life Sciences, University of Nottingham, Queen's Medical Centre,
10	Nottingham, NG7 2UH, United Kingdom
11	
12	Key words: SARS-CoV-2, nCoV-2019, COVID-19, coronavirus, drug repurposing, FDA
13	approved drugs, antiviral therapeutics, pandemic, chloroquine, hydroxychloroquine,
14	chlorpromazine
15	
16	Abstract
17	SARS-CoV-2 emerged in China at the end of 2019 and has rapidly become a pandemic with
18	roughly 2.7 million recorded COVID-19 cases and greater than 189,000 recorded deaths by
19	April 23rd, 2020 (www.WHO.org). There are no FDA approved antivirals or vaccines for any
20	coronavirus, including SARS-CoV-2. Current treatments for COVID-19 are limited to supportive
21	therapies and off-label use of FDA approved drugs. Rapid development and human testing of
22	potential antivirals is greatly needed. A quick way to test compounds with potential antiviral
23	activity is through drug repurposing. Numerous drugs are already approved for human use and
24	subsequently there is a good understanding of their safety profiles and potential side effects,

making them easier to fast-track to clinical studies in COVID-19 patients. Here, we present data on the antiviral activity of 20 FDA approved drugs against SARS-CoV-2 that also inhibit SARS-CoV and MERS-CoV. We found that 17 of these inhibit SARS-CoV-2 at a range of IC50 values at non-cytotoxic concentrations. We directly follow up with seven of these to demonstrate all are capable of inhibiting infectious SARS-CoV-2 production. Moreover, we have evaluated two of these, chloroquine and chlorpromazine, *in vivo* using a mouse-adapted SARS-CoV model and found both drugs protect mice from clinical disease.

32

33 Introduction

34 At the end of December 2019, reports started to emerge from China of patients suffering from 35 pneumonia of unknown etiology. By early January, a new coronavirus had been identified and 36 determined as the cause (1). Since then, the virus originally known as novel coronavirus 2019 37 (nCoV-2019), now severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2), has spread around the world. As of April 23th, 2020, there have been roughly 2.7 million confirmed cases of 38 39 COVID-19 (the disease caused by SARS-CoV-2 infection) with over to 189,000 recorded deaths 40 (www.WHO.org). Multiple countries have enacted social distancing and guarantine measures, 41 attempting to reduce person-to-person transmission of the virus. Healthcare providers lack 42 pharmaceutical countermeasures against SARS-CoV-2, beyond public health interventions, and 43 there remains a desperate need for rapid development of antiviral therapeutics. A potential route 44 to candidate antivirals is through repurposing of already approved drugs (for reviews; (2-4) and 45 for examples; (5-8). We have previously screened a library of FDA approved drugs for antiviral 46 activity against two other highly pathogenic human coronaviruses, SARS-CoV and Middle East 47 respiratory syndrome coronavirus (MERS-CoV)(6). We found 27 drugs that inhibited replication 48 of both of these coronaviruses, suggesting that they may have broad anti-coronaviral activity. 49 One of the hits from this work was imatinib with which we subsequently determined the

mechanism of action by demonstrating this drug inhibits fusion of coronaviruses with cellular
membranes, thus blocking entry (9, 10).

52

53 Here, we present our investigation of 20 priority compounds from our previous screening to test 54 if they can also inhibit SARS-CoV-2. Since these compounds are already approved for use in 55 humans, they make ideal candidates for drug repurposing and rapid development as antiviral 56 therapeutics. Our work found that 17 of the 20 of the drugs that inhibited SARS-CoV and 57 MERS-CoV could also inhibit SARS-CoV-2, with similar IC50 values. We further assessed a 58 subset of these drugs for their effects on SARS-CoV-2 RNA and infectious virus production and 59 found all to have inhibitory activity. Our screening based on cytopathic effect therefore appears 60 a favorable approach to find drugs capable of inhibiting production of infectious virus. Currently 61 there are no established small animal model systems for SARS-CoV-2. However, there is a 62 well-established mouse-adapted system for SARS-CoV (MA15 strain(11)) and we present data 63 here assessing the in vivo efficacy of chloroquine (CQ) and chlorpromazine (CPZ) against 64 SARS-CoV. We found that drug treatment does not inhibit virus replication in mouse lungs, but 65 significantly improves clinical outcome. Based on both of these drugs inhibiting SARS-CoV-2 66 infection in vitro and providing protection in vivo against SARS-CoV clinical disease we believe 67 they may be beneficial for SARS-CoV-2 therapy, but require further study in clinical contexts. 68

69 Materials and Methods

70 Cell lines and virus

Vero E6 cells (ATCC# CRL 1586) were cultured in DMEM (Quality Biological), supplemented
with 10% (v/v) fetal bovine serum (Sigma), 1% (v/v) penicillin/streptomycin (Gemini Bioproducts) and 1% (v/v) L-glutamine (2 mM final concentration, Gibco). Cells were maintained at
37°C and 5% CO₂. Samples of SARS-CoV-2 were obtained from the CDC following isolation

from a patient in Washington State (WA-1 strain - BEI #NR-52281). Stocks were prepared by
infection of Vero E6 cells for two days when CPE was starting to be visible. Media were
collected and clarified by centrifugation prior to being aliquoted for storage at -80°C. Titer of
stock was determined by plaque assay using Vero E6 cells as described previously (12). All
work with infectious virus was performed in a Biosafety Level 3 laboratory and approved by our
Institutional Biosafety Committee. SARS-CoV stock was prepared as previously described (13).
SARS-CoV spike (S) pseudotype viruses were produced as previously described (9).

82

83 Drug testing

84 All drug screens were performed with Vero E6 cells. Cells were plated in opaque 96 well plates 85 one day prior to infection. Drug stocks were made in either DMSO, water or methanol. Drugs 86 were diluted from stock to 50 µM and an 8-point 1:2 dilution series made. Cells were pre-treated 87 with drug for 2 hour (h) at 37°C/5% CO₂ prior to infection at MOI 0.01 or 0.004. Vehicle controls 88 were used on every plate, and all treatments were performed in triplicate for each screen. In 89 addition to plates that were infected, parallel plates were left uninfected to monitor cytotoxicity of 90 drug alone. Three independent screens with this set-up were performed. Cells were incubated 91 at 37°C/5% CO₂ for 3 days before performing CellTiter-Glo (CTG) assays as per the 92 manufacturer's instruction (Promega). Luminescence was read using a Molecular Devices 93 Spectramax L plate reader. Fluphenazine dihvdrochloride, benztropine mesvlate, amodiaguine 94 hydrochloride, amodiaquine dihydrochloride dihydrate, thiethylperazine maleate, mefloquine 95 hydrochloride, triparanol, terconazole vetranal, anisomycin, fluspirilene, clomipramine 96 hydrochloride, hydroxychloroquine sulfate, promethazine hydrochloride, emetine dihydrochloride 97 hydrate and chloroquine phosphate were all purchased from Sigma. Chlorpromazine 98 hydrochloride, toremifene citrate, tamoxifen citrate, gemcitabine hydrochloride and imatinib 99 mesylate were all purchased from Fisher Scientific.

101 Data analysis

102 Cytotoxicity (%TOX) data was normalized according to cell-only uninfected (cell only) controls
 103 and CTG-media-only (blank) controls:

%TOX =
$$\left(1 - \frac{(drug) - (blank)}{(cell only) - (blank)}\right) x \ 100$$

104 Inhibition (%Inhibit) data was normalized according to cell only and the activity of the vehicle105 controls:

$$\% Inhibit = \frac{(drug) - (vehicle)}{(cell only) - (vehicle)} x \ 100$$

106 Nonlinear regression analysis was performed on the normalized %inhibit and %TOX data and 107 IC50s and CC50s were calculated from fitted curves (log [agonist] versus response - variable 108 slope [four parameters]) (GraphPad Software, LaJolla, CA), as described previously (14). Drug 109 dilution points in a given run were excluded from IC50 analysis if the average cytotoxicity was 110 greater than 30% (arbitrary cutoff) across the 3 cytotoxicity replicates for that screen. IC50 or 111 CC50 values extrapolated outside the drug dilution range tested were reported as greater than 112 50µM or less than 0.39µM. Selectivity indexes (SI) were also calculated by dividing the CC50 by 113 the IC50.

114

115 Viral infection

116 To further analyse candidate drugs, Vero E6 cells were grown in 24 well plate format for 24 h

prior to infection. As with the drug screens, cells were pre-treated with drug at a range of

- concentrations, or vehicle control for 2 h. Cells were then infected with SARS-CoV-2 at MOI 0.1
- 119 for 24 hour (h). Supernatant was collected, centrifuged in a table-top centrifuge for 3 minutes
- 120 (min) at max speed and stored at -80°C. After a wash in PBS, infected cells were collected in

TRIzol (Ambion) for RNA analysis (described below). Supernatant was used to titer viral
 production by TCID₅₀ assay (12).

123

124 RNA extraction and qRT-PCR

125 RNA was extracted from TRIzol samples using Direct-zol RNA miniprep kit (Zymo Research) as

126 per the manufacturer's instructions. RNA was converted to cDNA using RevertAid RT Kit

127 (Thermo Scientific), with 12 μ l of extracted RNA per reaction. For qRT-PCR, 2 μ l of cDNA

128 reaction product was mixed with PowerUp SYBR Green Master Mix (Applied Biosystems) and

129 WHO/Corman primers targeting N and RdRp: N FWD 5'-CACATTGGCACCCGCAATC-3', N

130 REV 5'-GAGGAACGAGAAGAGGCTTG-3', RdRp FWD 5'GTGARATGGTCATGTGGCGG-

131 3', RdRp REV 5'-CARATGTTAAASACACTATTAGCATA-3'. The qRT-PCR reactions were

132 performed with a QuantStudio 5 (Applied Biosystems). To normalize loading, 18S RNA was

used as a control, assessed with TaqMan Gene Expression Assays (Applied Biosystems) and

134 TaqMan Fast Advanced Master Mix. Fold change between drug treated and vehicle control was

135 determined by calculating $\Delta\Delta$ CT after normalization to the endogenous control of 18S.

136

137

138 **Pseudovirus fusion/entry assay**

The pseudovirion (PV) entry assay was performed as described (9, 15). Briefly, 2×10^4 BSC1 cells per well were in 96-well plates for 24 h, after which time cells were pre-treated with drug (1 h) and infected with PV (3 h). Media was removed and cells were washed with loading buffer (47 ml clear DMEM, 5 mM Probenecid, 2 mM L-glutamine, 25 mM HEPES, 200 nM bafilomycin, 5μ M E64D) and incubated for 1 h in CCF2 solution (LB, CCF2-AM, Solution B [CCF2-AM kit K1032] Thermo Fisher) in the dark. Cells were washed once with loading buffer and incubated from 6 h to overnight with 10% FBS in loading buffer. Percentage CCF2 cleavage was assessed

146	by flow cytomet	ry on the LSRI	(Beckton Dickinson)) in the flow cyte	ometry core facilit	v at the
140						y at the

- 147 University of Maryland, Baltimore. Data were analyzed using FlowJo.
- 148

149 Mouse infections.

- 150 All infections were performed in an animal biosafety level 3 facility at the University of Maryland,
- 151 Baltimore, using appropriate practices, including a HEPA-filtered bCON caging system, HEPA-
- 152 filtered powered air-purifying respirators (PAPRs), and Tyvek suiting. All animals were grown to
- 153 10 weeks of age prior to use in experiments. The animals were anesthetized using a mixture of
- 154 xylazine (0.38 mg/mouse) and ketamine (1.3 mg/mouse) in a 50 µl total volume by
- 155 intraperitoneal injection. The mice were inoculated intranasally with 50 µl of either PBS or 2.5 x
- 156 10³ PFU of rMA15 SARS-CoV (11) after which all animals were monitored daily for weight loss.
- 157 Mice were euthanized at day 4 post-infection, and lung tissue was harvested for further
- analysis. All animals were housed and used in accordance with the University of Maryland,
- 159 Baltimore, Institutional Animal Care and Use Committee guidelines.
- 160

161 Plaque assay.

- 162 Vero cells were seeded in 35 mm dishes with 5×10^5 cells per dish 24 h prior to
- 163 infection. Supernatants from homogenized were serially diluted 10⁻¹ through 10⁻⁶ in serum-free
- 164 (SF) media. Cells were washed with SF media, 200 µl of diluted virus was added to each well
- and adsorption was allowed to proceed for 1 h at 37°C with gentle rocking every 10 min. 2X
- 166 DMEM and 1.6% agarose were mixed 1:1. Cells were washed with SF media, 2 ml DMEM-
- agarose was added to each well, and cells were incubated for 72 h at 37°C, after which time
 plagues were read.
- 169
- 170
- 171 <u>Results</u>

172 Screening FDA approved compounds for anti-SARS-CoV-2 activity

173 Previously, we performed a large-scale drug screen on 290 FDA approved compounds to 174 investigate which may have antiviral activity against SARS-CoV and MERS-CoV (6). With the 175 emergence of SARS-CoV-2, we prioritized testing 20 of the 27 hits that were determined to 176 inhibit both of the previously tested coronaviruses for antiviral activity against the novel virus. 177 The list of tested compounds is shown in Table 1. Our screening started at 50 μ M and used an 178 8-point, 1:2 dilution series with infections being performed at either MOI 0.01 or 0.004. CellTiter-179 Glo (CTG) assays were performed 3 days post-infection to determine relative cell viability 180 between drug and vehicle control treated cells. Uninfected samples were used to measure the 181 cytotoxicity of drug alone. From the relative luminescence data of the CTG assay, percent 182 inhibition (of cell death caused by viral infection) could be measured and plotted along with the 183 percent cytotoxicity of drug alone. Fig. 1 shows these plotted graphs from one representative of 184 three independent screens at MOI 0.01. For those drugs demonstrating a cell toxicity rate lower 185 than 30%, we were able to calculate IC50 values at both MOI from these graphs for 17 of the 20 186 drugs which is summarized in Table 1.

187

188 **Drug screen validation**

189 In order to validate our screening process as a means to identify compounds with antiviral effect 190 we decided to follow up with a subset of drugs. Chloroquine (CQ) has become the source of 191 much interest as a potential treatment for COVID19 (16), as such, we further investigated hydroxychloroquine (HCQ) and CQ as both were present in our screen (Table 1). Vero E6 cells 192 193 were plated and pre-treated with drug for 2 h prior to infection with SARS-CoV-2 at MOI 0.1. 194 Supernatant was collected 24 h post-infection to determine titer of virus by TCID₅₀ assay and 195 cells were collected in TRIzol to assess production of viral mRNA. Treatment with both drugs 196 caused a significant reduction in viral mRNA levels, especially at higher concentrations, without 197 drug induced cytotoxicity (Fig. 1 and Table 1). There was a significant decrease in relative

198 expression levels of both RdRp and N mRNA across the range of concentrations used (Fig. 2A-199 D). Along with causing a reduction in viral mRNA, treatment with both drugs caused a significant 200 reduction in viral replication (Fig. 2E and 2F). SARS-CoV-2 production was more sensitive to 201 HCQ than CQ with larger inhibition seen at the same concentration of treatment, which is in 202 agreement with HCQ having a lower IC50 in our cell viability assay (Table 1). We also 203 performed a time of addition assay with the highest concentration of HCQ to investigate whether 204 SARS-CoV-2 entry was the point of inhibition of this compound (Fig. 2G). Interestingly, while the 205 addition of HCQ at 2h post-infection did have some reduction in inhibitory activity there was not 206 a complete loss, suggesting that HCQ treatment may impact other stages of the viral life cycle 207 than just entry.

208

209 We have previously used a β -lactamase-Vpr chimeric protein (Vpr-BlaM) pseudotype system to 210 demonstrate that imatinib (a drug also seen to inhibit SARS-CoV-2 [Table 1 and Fig. 1]) inhibits 211 SARS-CoV and MERS-CoV spike-mediated entry (9). We used this system to more directly 212 investigate whether CQ could inhibit viral entry mediated by coronavirus spike, and additionally 213 included chlorpromazine (CPZ) as this is known to inhibit clathrin-mediated endocytosis (17) 214 and was also part of our drug screening (Table 1 and Fig. 2). In this assay, when the 215 pseudovirus fuses with a cellular membrane, BlaM is released into the cytoplasm of the infected 216 cell. BlaM cleaves cytoplasmic loaded CCF2 to change its emission spectrum from 520 nm 217 (green) to ~450 nm (blue), which can be quantified using flow cytometry.

218

Cells were treated with CQ or CPZ for 1 h before infection with BlaM-containing SARS-S
pseudovirions (PV). Cells were then analyzed by flow cytometry to quantify the cleavage of
CCF2. In mock treated cells infected with SARS-S PV there was a shift in the CCF2 emission
spectrum indicating release of BlaM to the cytosol, and that spike-mediated fusion with cellular
had occurred. Upon treatment with CQ or CPZ there was a greater than 90% reduction in CCF2

cleavage caused by SARS-S PV (Fig. 2H). These data demonstrate that both drugs inhibit
SARS-CoV spike-mediated fusion with cellular membranes. These pseudotype assays suggest
that the inhibition of coronavirus replication caused by CQ and CPZ is at the stage of entry to
cells but combined with the time of addition assays (Fig. 2G), there is a suggestion that later
stages may also be impacted.

229

230 HCQ and CQ are used as anti-malarial drugs and are in the class of aminoquinolines which are 231 hemozoin inhibitors, similarly to 4-methanologinolines. Interestingly, from our drug screening. 232 three other hemozoin inhibitors were identified: amodiaquine dihydrochloride dihydrate, 233 amodiaguine hydrochloride and mefloguine. We therefore decided to directly test these drugs 234 for antiviral activity against SARS-CoV-2. We directly tested CPZ against SARS-CoV-2 having 235 seen that it could inhibit SARS-CoV S-mediated entry to cells (Fig. 2H). We also included 236 imatinib since we have previously shown that this can inhibit entry of both SARS-CoV and 237 MERS-CoV (9) and was a hit against SARS-CoV-2 (Fig. 1 and Table 1). Again, cells were pre-238 treated with drugs at the indicated concentrations and infected with SARS-CoV-2 at MOI 0.1 for 239 24h, after which supernatant samples were collected. As can be seen in Fig. 3, at the highest 240 concentrations of all drugs there is significant inhibition of SARS-CoV-2 infection. All five drugs 241 showed very strong inhibition at 20 µM.

242

Overall, the data from Fig. 2 and Fig. 3 indicate that there are various FDA approved drugs that have broad-spectrum anti-coronavirus activity *in vitro* and that our initial screening based on cytopathic effect is a good method to identify compounds with antiviral activity.

246

247 Chloroquine and chlorpromazine do not inhibit SARS-CoV (MA15) replication in mouse

248 lungs, but significantly reduces weight loss and clinical signs

249 CQ and CPZ treatment displayed significant inhibition of coronavirus replication in vitro, with our data suggesting entry is inhibited. We therefore decided to investigate whether these drugs 250 251 were efficacious in vivo using SARS-CoV strain, MA15 in BALB/c mice. This model displays 252 ~15-20% weight loss by 4 days post infection (dpi), occasionally resulting in death. We tested 253 whether prophylactically administered CQ or CPZ could protect mice from severe MA15 254 infection. Mice were injected intraperitoneally with either water, 0.8 mg CQ, 1.6 mg CQ, 20 µg 255 CPZ, 100 µg CPZ or 200 µg CPZ at day -1 of infection and then were dosed every day through 256 the 4 days of infection. On day 0, mice were intranasally infected with 2.5 x 10³ pfu of SARS-257 CoV (MA15) or PBS as control. Weight loss was measured as a correlate of disease and mice 258 were euthanized at 4 dpi for analysis.

259

260 PBS inoculated mice showed no weight loss or clinical signs of disease when treated with either 261 water, CQ or CPZ over the experiment time course indicating drug treatment did not adversely 262 affect morbidity (Fig. 4A and 4D). Mice that were infected with MA15 and treated with water lost 263 ~15% of their starting body weight over 4 days and had significant clinical signs of disease 264 including ruffled fur, labored breathing and lethargy (Fig. 4A and 4D). Mice that were treated 265 with 0.8 mg of CQ each day, displayed similar weight loss as the water control through the first 266 3 days of infection, however by 4 dpi the weight loss was halted in the drug treated mice (Fig. 267 4A). Mice that were treated with 1.6 mg CQ per day showed markedly reduced weight loss 268 compared to the water control (Fig. 4A). Pathological analysis was also performed on H&E 269 stained sections. Mice infected with MA15 and treated with water displayed significant 270 inflammation and denuding bronchiolitis suggesting severe disease (Fig. 4B). By contrast, 0.8 271 mg CQ dose group had moderate inflammation that was reduced compared to control and the 272 1.6 mg dose group had minimal lung pathology (Fig. 4B). Interestingly, even though CQ 273 treatment appeared to protect against weight loss and inflammation in the lungs, the viral titer 274 was equivalent between drug treated and vehicle control mice (Fig. 4C).

276	Similar to the CQ results, CPZ treatment reduced weight loss in mice infected with MA15 at 100
277	μg and 200 $\mu g,$ but the 20 μg treatment group were equivalent to vehicle control (Fig. 4D) and
278	the H&E sections showed protection against inflammation and denuding bronchiolitis at the
279	higher doses (Fig. 4E). Again, as with CQ treatment, even though there were reduced signs of
280	infection with CPZ treatment, there was no difference in MA15 titer in the mouse lungs (Fig. 4F).
281	Overall these data indicate that even though CQ and CPZ treatment do not inhibit viral
282	replication in the lungs, both can protect mice from signs of disease following SARS-CoV
283	(MA15) infection.

284

285 Discussion

286 The SARS-CoV-2 pandemic has demonstrated the desperate need for antiviral drugs. Since the 287 emergence of SARS-CoV in 2002, research has uncovered many details of coronavirus biology 288 and pathogenesis, however there are currently no approved therapeutics against this emerging 289 virus family. Whether being used for treating SARS-CoV-2 in this current pandemic or the next 290 unknown viral pathogen in the future, we must attempt to develop and validated antiviral drugs 291 that are ready to be used at the first signs of an outbreak. Many FDA approved drugs have been 292 found to have antiviral activity in addition to their approved use (e.g; (5–8)), and since these are 293 extensively used in humans for other conditions, they could be streamlined for rapid approval 294 and repositioning as antivirals. In our previous work, 290 FDA approved drugs were screened 295 for antiviral activity and 27 were found to inhibit both SARS-CoV and MERS-CoV (6). We 296 prioritized testing these for antiviral activity against SARS-CoV-2 since they displayed broad-297 spectrum antiviral activity. From multiple independent screens performed with two MOI, we 298 found that 17 of our 20 tested priority compounds display significant antiviral activity at non-299 cytotoxic concentrations. Many of the compounds have IC50 values under 10 µM and these will 300 be the source of follow up testing on additional cell lines and in mouse models of SARS-CoV-2.

302 We further investigated seven of the hits to directly test if they inhibited SARS-CoV-2 replication. 303 We performed follow-up experiments with HCQ, CQ, amodiaguine and mefloguine because 304 chloroquine has garnered much interest as a potential treatment for COVID19 (16) and the 305 others are similarly used as anti-malarial compounds (18). In addition, we have previously 306 demonstrated that imatinib is an inhibitor of SARS-CoV, MERS-CoV and infectious bronchitis 307 virus entry to cells (9, 10) so included that here as the mechanism of coronavirus inhibition is 308 understood. Finally, CPZ inhibits clathrin function in cells (17) so can disrupt infection by many 309 viruses that require clathrin-mediated endocytosis and was therefore also chosen for further 310 analysis. Treatment of cells with all these drugs showed inhibition of infectious viral particle 311 production (measured by TCID₅₀ assay) at non-cytotoxic levels. 312 313 Having demonstrated that HCQ. CQ and CPZ can inhibit cytopathic effect, mRNA synthesis and 314 infectious viral particle production of SARS-CoV-2, we used a previously published system of 315 SARS-CoV pseudotype viruses carrying Vpr-BlaM to investigate whether CQ and CPZ inhibit 316 coronavirus spike-mediated entry to better define mechanism of action. We have previously 317 used this system to define imatinib as an entry inhibitor of these viruses (9) and found similar 318 results for CQ and CPZ, thus better defining their mechanism of antiviral activity. 319 320 Finally, we investigated the efficacy of CQ and CPZ with an *in vivo* model using SARS-CoV 321 MA15. There is currently a lack of an established mouse model for SARS-CoV-2 so we used the

322 mouse adapted SARS-CoV (MA15) strain as a surrogate to assess the *in vivo* efficacy of these

drugs against a closely related coronavirus. We are of the opinion that this is a good model

324 since both viruses use ACE2 as a receptor (19–22) and therefore have a similar cellular tropism

- 325 which is important since both of these compounds appear to inhibit viral entry. Prophylactic
- dosing in MA15 infection experiments demonstrated that, in contrast to the *in vitro* antiviral

327 activity, CQ and CPZ did not inhibit viral replication in mouse lungs based on viral titer 328 recovered from lungs at 4 dpi. However, both drugs resulted in reduced weight loss and 329 improved clinical outcome, with the higher dose giving greater protection. Along with being an 330 anti-malarial, CQ is used in humans for the treatment of systemic lupus erythematosus and 331 rheumatoid arthritis because of anti-inflammatory properties and has effects on antigen 332 presentation (23–25). We speculate that these properties may have a role in the protection we 333 observe in vivo since much of the pathology from SARS-CoV is a consequence of 334 immunopathology during infection (in mice; (26), in non-human primates (27) and for a detail 335 review (28)). These results suggest that CQ alone may not be a viable therapeutic but may be 336 beneficial for treatment of SARS-CoV-2 in combination with more directly acting antivirals such 337 as remdesivir (29-31).

338

339 The development of antiviral drugs for emerging coronaviruses is a global priority. In the middle 340 of the COVID19 pandemic, we must identify rapidly accessible therapeutics that are validated in 341 both in vitro and in vivo models. FDA approved drugs being assessed for repurposing and other 342 experimental drugs in development must be properly validated in animal studies to best assess 343 their potential utility in people. We have presented here a list of FDA approved drugs that are 344 effective in vitro against SARS-CoV-2 as well as being effective against SARS-CoV and MERS-345 CoV (6). Moreover, we have demonstrated that two of these, CQ and CPZ, can protect mice 346 from severe clinical disease from SARS-CoV. Future research will be aimed at testing these 347 compounds in SARS-CoV-2 animal models to further assess their potential utility for human 348 treatment.

349

350 Acknowledgments

We kindly thank Emergent BioSolutions for financial support to perform these experiments. Wealso kindly thank Julie Dyall for helpful discussions regarding data analysis.

354 <u>References</u>

- 355 1. Zhu N, Zhang D, Wang W, Li X, Yang B, Song J, Zhao X, Huang B, Shi W, Lu R, Niu P,
- 356 Zhan F, Ma X, Wang D, Xu W, Wu G, Gao GF, Tan W. 2020. A novel coronavirus from
- 357 patients with pneumonia in China, 2019. N Engl J Med 382:727–733.
- 358 2. Sisk JM, Frieman MB. 2016. Screening of FDA-Approved Drugs for Treatment of

359 Emerging Pathogens. ACS Infect Dis. American Chemical Society.

- 360 3. Pushpakom S, Iorio F, Eyers PA, Escott KJ, Hopper S, Wells A, Doig A, Guilliams T,
- 361 Latimer J, McNamee C, Norris A, Sanseau P, Cavalla D, Pirmohamed M. 2018. Drug
- 362 repurposing: Progress, challenges and recommendations. Nat Rev Drug Discov.
- Mercorelli B, Palù G, Loregian A. 2018. Drug Repurposing for Viral Infectious Diseases:
 How Far Are We? Trends Microbiol 26:865–876.
- 365 5. Madrid PB, Chopra S, Manger ID, Gilfillan L, Keepers TR, Shurtleff AC, Green CE, Iyer L
- 366 V., Dilks HH, Davey RA, Kolokoltsov AA, Carrion R, Patterson JL, Bavari S, Panchal RG,
- 367 Warren TK, Wells JB, Moos WH, Burke RLL, Tanga MJ. 2013. A Systematic Screen of

368 FDA-Approved Drugs for Inhibitors of Biological Threat Agents. PLoS One 8.

- 369 6. Dyall J, Coleman CM, Hart BJ, Venkataraman T, Holbrook MR, Kindrachuk J, Johnson
- 370 RF, Olinger GG, Jahrling PB, Laidlaw M, Johansen LM, Lear-rooney CM, Glass PJ,
- 371 Hensley LE, Frieman B. 2014. Repurposing of Clinically Developed Drugs for Treatment
- 372 of Middle East Respiratory Syndrome Coronavirus Infection. Antimicrob Agents
- 373 Chemother 58:4885–4893.
- 374 7. Madrid PB, Panchal RG, Warren TK, Shurtleff AC, Endsley AN, Green CE, Kolokoltsov A,
- 375 Davey R, Manger ID, Gilfillan L, Bavari S, Tanga MJ. 2016. Evaluation of Ebola Virus
- 376 Inhibitors for Drug Repurposing. ACS Infect Dis 1:317–326.
- 8. Xu M, Lee EM, Wen Z, Cheng Y, Huang WK, Qian X, Tcw J, Kouznetsova J, Ogden SC,
- Hammack C, Jacob F, Nguyen HN, Itkin M, Hanna C, Shinn P, Allen C, Michael SG,

379	Simeonov A, Huang W	, Christian KM,	Goate A, Brennand KJ, I	Huang R, Xia M, Ming GL,
	, J	, , ,	, , ,	

- 380 Zheng W, Song H, Tang H. 2016. Identification of small-molecule inhibitors of Zika virus
- infection and induced neural cell death via a drug repurposing screen. Nat Med 22:1101–
 1107.
- Coleman CM, Sisk JM, Mingo RM, Nelson EA, White JM, Frieman MB. 2016. Abl Kinase
 Inhibitors Are Potent Inhibitors of SARS-CoV and MERS-CoV Fusion. J Virol 90:8924–
 8933.
- 386 10. Sisk JM, Frieman MB, Machamer CE. 2018. Coronavirus S protein-induced fusion is
 387 blocked prior to hemifusion by Abl kinase inhibitors. J Gen Virol 1–12.
- 388 11. Roberts A, Deming D, Paddock CD, Cheng A, Yount B, Vogel L, Herman BD, Sheahan T,

Heise M, Genrich GL, Zaki SR, Baric R, Subbarao K. 2007. A mouse-adapted SARS-

- 390 coronavirus causes disease and mortality in BALB/c mice. PLoS Pathog 3:0023–0037.
- 391 12. Coleman CM, Frieman MB. 2015. Growth and Quantification of MERS-CoV Infection.
- 392 Curr Protoc Microbiol 37:15E.2.1-15E.2.9.
- 393 13. Frieman M, Yount B, Agnihothram S, Page C, Donaldson E, Roberts A, Vogel L,
- 394 Woodruff B, Scorpio D, Subbarao K, Baric RS. 2012. Molecular Determinants of Severe
- Acute Respiratory Syndrome Coronavirus Pathogenesis and Virulence in Young and
 Aged Mouse Models of Human Disease. J Virol 86:884–897.
- 14. Dyall J, Johnson JC, Hart BJ, Postnikova E, Cong Y, Zhou H, Gerhardt DM, Michelotti J,
- Honko AN, Kern S, DeWald LE, O'Loughlin KG, Green CE, Mirsalis JC, Bennett RS,
- Olinger GG, Jahrling PB, Hensley LE. 2018. In Vitro and In Vivo Activity of Amiodarone
 Against Ebola Virus. J Infect Dis 218:S592–S596.
- 401 15. Mingo RM, Simmons JA, Shoemaker CJ, Nelson EA, Schornberg KL, D'Souza RS,
- 402 Casanova JE, White JM. 2015. Ebola Virus and Severe Acute Respiratory Syndrome
- 403 Coronavirus Display Late Cell Entry Kinetics: Evidence that Transport to NPC1 +
- 404 Endolysosomes Is a Rate-Defining Step. J Virol 89:2931–2943.

- 405 16. Pastick KA, Okafor EC, Wang F, Lofgren SM, Skipper CP, Nicol MR, Pullen MF,
- 406 Rajasingham R, Mcdonald EG, Lee TC, Schwartz IS, Kelly LE, Lother SA, Mitjà O,
- 407 Letang E, Abassi M, Boulware DR. 2020. Review: Hydroxychloroquine and Chloroquine
- 408 for Treatment of SARS-CoV-2 (COVID-19). Open Forum Infect Dis.
- 409 17. Wang LH, Rothberg KG, Anderson RGW. 1993. Mis-assembly of clathrin lattices on
- 410 endosomes reveals a regulatory switch for coated pit formation. J Cell Biol 123:1107–
- 411 1117.
- 412 18. Baird JK. 2005. Effectiveness of antimalarial drugs. N Engl J Med 352.
- 413 19. Li W, Moore MJ, Vasllieva N, Sui J, Wong SK, Berne MA, Somasundaran M, Sullivan JL,
- Luzuriaga K, Greeneugh TC, Choe H, Farzan M. 2003. Angiotensin-converting enzyme 2
- 415 is a functional receptor for the SARS coronavirus. Nature 426:450–454.
- 416 20. Zhou P, Yang X Lou, Wang XG, Hu B, Zhang L, Zhang W, Si HR, Zhu Y, Li B, Huang CL,
- 417 Chen HD, Chen J, Luo Y, Guo H, Jiang R Di, Liu MQ, Chen Y, Shen XR, Wang X, Zheng
- 418 XS, Zhao K, Chen QJ, Deng F, Liu LL, Yan B, Zhan FX, Wang YY, Xiao GF, Shi ZL.
- 419 2020. A pneumonia outbreak associated with a new coronavirus of probable bat origin.
- 420 Nature 579:270–273.
- 421 21. Wan Y, Shang J, Graham R, Baric RS, Li F. 2020. Receptor Recognition by the Novel
- 422 Coronavirus from Wuhan: an Analysis Based on Decade-Long Structural Studies of 423 SARS Coronavirus. J Virol 94.
- 424 22. Hoffmann M, Kleine-Weber H, Schroeder S, Krüger N, Herrler T, Erichsen S, Schiergens
- 425 TS, Herrler G, Wu NH, Nitsche A, Müller MA, Drosten C, Pöhlmann S. 2020. SARS-CoV-
- 426 2 Cell Entry Depends on ACE2 and TMPRSS2 and Is Blocked by a Clinically Proven
 427 Protease Inhibitor. Cell 181:271-280.e8.
- 428 23. Ziegler HK, Unanue ER. 1982. Decrease in macrophage antigen catabolism caused by
- 429 ammonia and chloroquine is associated with inhibition of antigen presentation to T cells.
- 430 Proc Natl Acad Sci U S A 79:175–178.

	431	24.	Al-Bari MAA. 20	015. Chloroc	uine analogue	es in drug	discovery: r	new directions of use	es,
--	-----	-----	-----------------	--------------	---------------	------------	--------------	-----------------------	-----

- 432 mechanisms of actions and toxic manifestations from malaria to multifarious diseases. J
 433 Antimicrob Chemother 70:1608–21.
- 434 25. Rainsford KD, Parke AL, Clifford-Rashotte M, Kean WF. 2015. Therapy and
- 435 pharmacological properties of hydroxychloroquine and chloroquine in treatment of
- 436 systemic lupus erythematosus, rheumatoid arthritis and related diseases.
- 437 Inflammopharmacology 23:231–269.
- 438 26. Rockx B, Baas T, Zornetzer GA, Haagmans B, Sheahan T, Frieman M, Dyer MD, Teal
- 439 TH, Proll S, van den Brand J, Baric R, Katze MG. 2009. Early Upregulation of Acute
- 440 Respiratory Distress Syndrome-Associated Cytokines Promotes Lethal Disease in an
- 441 Aged-Mouse Model of Severe Acute Respiratory Syndrome Coronavirus Infection. J Virol
 442 83:7062–7074.
- 443 27. Smits SL, De Lang A, Van Den Brand JMA, Leijten LM, Van Ijcken WF, Eijkemans MJC,
- 444 Van Amerongen G, Kuiken T, Andeweg AC, Osterhaus ADME, Haagmans BL. 2010.
- 445 Exacerbated innate host response to SARS-CoV in aged non-human primates. PLoS
- 446 Pathog 6.
- 28. Channappanavar R, Perlman S. 2017. Pathogenic human coronavirus infections: causes
 and consequences of cytokine storm and immunopathology. Semin Immunopathol
 39:529–539.
- 450 29. Wang M, Cao R, Zhang L, Yang X, Liu J, Xu M, Shi Z, Hu Z, Zhong W, Xiao G. 2020.
- 451 Remdesivir and chloroquine effectively inhibit the recently emerged novel coronavirus
 452 (2019-nCoV) in vitro. Cell Res 0.
- 453 30. Brown AJ, Won JJ, Graham RL, Dinnon KH, Sims AC, Feng JY, Cihlar T, Denison MR,
- 454 Baric RS, Sheahan TP. 2019. Broad spectrum antiviral remdesivir inhibits human
- 455 endemic and zoonotic deltacoronaviruses with a highly divergent RNA dependent RNA
- 456 polymerase. Antiviral Res 169.

457	31.	Sheahan TP, Sims AC, Graham RL, Menachery VD, Gralinski LE, Case JB, Leist SR,
458		Pyrc K, Feng JY, Trantcheva I, Bannister R, Park Y, Babusis D, Clarke MO, MacKman
459		RL, Spahn JE, Palmiotti CA, Siegel D, Ray AS, Cihlar T, Jordan R, Denison MR, Baric
460		RS. 2017. Broad-spectrum antiviral GS-5734 inhibits both epidemic and zoonotic
461		coronaviruses. Sci Transl Med 9.
462		
463		
464		
465	<u>Figur</u>	e legend
466	Figur	e 1 – Percentage inhibition and percentage cytotoxicity graphs from drug screens
467	starti	ng at 50 µM using an 8-point, 1:2 dilution series. Results from one representative drug
468	scree	n of three showing percentage inhibition and cytotoxicity for each of the tested drugs.
469	Triplic	cate wells of cells were pre-treated with the indicated drug for 2 hours prior to infection with
470	SARS	G-CoV-2 at MOI 0.01. Cells were incubated for 72 hours prior to performing CellTiter-Glo
471	assay	s to assess cytopathic effect. Data are scored as percentage inhibition of relative cell
472	viabili	ty for drug treated versus vehicle control. Data are the mean percentages with error bars
473	displa	ying standard deviation between the triplicate wells.
474		
475	Figur	e 2 – Hydroxychloroquine and chloroquine inhibit production of SARS-CoV-2 N and
476	RdRp	mRNA.
477	Vero	cells were pre-treated with hydroxychloroquine sulfate (A, C and E) or chloroquine
478	phosp	whate (B, D and F) at the indicated concentration (or 0.1% water as vehicle control) for 2 h
479	prior t	o infection with SARS-CoV-2 (WA-1 strain) at MOI 0.1. 24 h post-infection cells were
480	collec	ted in TRIzol. RNA was extracted from TRIzol sample and qRT-PCR was performed for
481	viral F	RdRp (A and B) or N (C and D) mRNA using WHO primers. RNA levels were normalized
482	with 1	8S RNA and fold change for drug treated to vehicle control was calculated (dotted line to

483 denote a fold change of 1 which is no change over control). Data are from 3 independent 484 infections performed on triplicate wells, the fold change was calculated in each independent 485 experiment and the mean fold change is plotted with error bars displaying standard deviation. 486 Along with TRIzol samples for RNA supernatant was collected from cells and used for TCID₅₀ assays to determine infectious virus production following treatment with HCQ (E) or CQ (F) Data 487 488 are from 3 independent infections performed on triplicate wells with the TCID₅₀/ml being 489 averaged across all wells. Error bars are the standard deviation. G) Cells were treated with 50 490 µM HCQ or 0.1% water as control. Drug was either added 2 h prior to infection, at the time of 491 infection or 2 h after infection with MOI 0.1 SARS-CoV-2. After 24 h infection, supernatant was 492 collected and used for TCID₅₀ assays to determine infectious virus production. Data are from 3 493 independent infections performed on triplicate wells with the TCID₅₀/ml being averaged across 494 all wells. Error bars are the standard deviation. H) SARS-CoV spike psuedoviruses (PV) were 495 used for infection of BSC1 cells. The cells were treated with 10 µM of CQ or CPZ for 1 h prior to 496 infection with PV for 3 h. The PV carry BlaM and cells were loaded with CCF2 to monitor 497 cleavage and shift in fluorescence output for evidence of S-mediated entry into cells. Data are 498 normalised to PV alone and are from 3 independent experiments with error bars representing 499 standard deviation.

500

501 Figure 3 – Antiviral activity of additional FDA approved compounds against SARS-CoV-2. 502 Other drugs that showed antiviral activity in our initial CellTiter-Glo screening were tested for 503 inhibition of productive virus infection. Cells were treated with the indicated concentrations of A) 504 amodiaguine dihydrochloride dihydrate, B) amodiaguine hydrochloride, C) chlorpromazine, D) 505 mefloquine and E) imatinib for 2 h prior to infection with SARS-CoV-2 at MOI 0.1 for 24 h. 506 Supernatant was collected and used for TCID₅₀ assay to quantify infectious virus production. 507 Data are from a representative experiment of four performed on triplicate wells. Data are the 508 mean TCID₅₀/ml with error bars being standard deviation.

510	Figure 4 – CQ and CPZ are protective against SARS-CoV (MA15) infection <i>in vivo</i>
511	Mice were treated with CQ or CPZ 1 day prior to infection with SARS-CoV (MA15) and dosed
512	with each drug across the 4 day infection time course. Water was used as the vehicle control for
513	both drugs and PBS was used as a control for uninfected mice. A) Weight loss of mice treated
514	with CQ at two different dose levels (0.8 mg and 1.6 mg) over the 4 day infection. Data are
515	presented as relative weight loss compared to the mouse weight on day 0. In each treatment
516	group there were 5 mice and the data are mean average and standard deviation. B) At day 4,
517	mice were euthanized and lung sections were used for H&E staining. C) In addition to collecting
518	lungs for section staining, there was also collection to determine titer of virus by plaque assay.
519	D) Weight loss of mice treated with CPZ at three different doses (20 μ g, 100 μ g, and 200 μ g)
520	with the same experimental set up as in A. E and F) As B and C but for CPZ treated mice.
521	
522	Table 1 - IC50 and CC50 values for 20 FDA approved drugs against SARS-CoV-2.
523	Abbreviations: MOI (multiplicity of infection), IC50 (half maximal inhibitory concentration), CC50
524	(half maximal cytotoxic concentration), avg. (average), ND (not determined).
525	A – Run totals listed as IC50,CC50
526	B – At least one CC50 could be extrapolated from the curve fit suggesting toxicity and SI are
527	slightly higher than listed
528	C – No CC50 could be extrapolated from the curve fit suggesting toxicity and SI are much
529	higher than listed.
530	
531	
532	
533	
534	

- 541 Figure 1



554 Figure 2







- 574 Figure 4



Drug	MOI	Plate Replicates	IC50 (avg.)	CC50 (avg.)	SI (avg.)
Amodiaquine Dihydrochloride Dihydrate	0.004	2,3 ^a	2.59	34.42	13.31
	0.01	3	4.94	34.42	6.97
Amodiaquine Hydrochloride	0.004	3	2.36	>38.63 ^b	>16.37 ^b
	0.01	3	5.64	>38.63 ^b	>6.84 ^b
Anisomycin	0.004	3	ND	<0.39	ND
	0.01	3	ND	<0.39	ND
Benztropine Mesylate	0.004	3	13.8	>>50 [°]	>>3.62 ^c
	0.01	2,3 ^a	17.79	>>50 [°]	>>2.81 [°]
Chloroquine Phosphate	0.004	3	42.03	>50 ^b	>1.19 ^b
	0.01	3	46.8	>50 ^b	>1.07 ^b
Chlorpromazine Hydrochloride	0.004	2,3 ^a	3.14	11.88	3.78
	0.01	2,3 ^a	4.03	11.88	2.94
Clomipramine Hydrochloride	0.004	2,3 ^a	5.63	>29.68 ^b	>5.27 ^b
	0.01	3	7.59	>29.68 ^b	>3.91 ^b
Emetine Dihydrochloride Hydrate	0.004	3	ND	<0.39	ND
	0.01	2,3 ^a	ND	<0.39	ND
Fluphenazine Dihydrochloride	0.004	3,2 ^a	6.36	20.02	3.15
	0.01	2	8.98	20.02	2.23
Fluspirilene	0.004	3	3.16	30.33	9.61
	0.01	3	5.32	30.33	5.71
Gemcitabine Hydrochloride	0.004	3	ND	23.22	ND
	0.01	3	ND	23.22	ND
Hydroxychloroquine Sulfate	0.004	3	9.21	>>50 [°]	>>5.43 ^c
	0.01	3	11.17	>>50 ^c	>>4.48 ^c
Imatinib Mesylate	0.004	3	3.24	>30.86 ^b	>9.52 ^b
	0.01	3	5.32	>30.86 ^b	>5.80 ^b
Mefloquine Hydrochloride	0.004	3	7.11	18.53	2.61
	0.01	3	8.06	18.53	2.3
Promethazine Hydrochloride	0.004	3	9.21	>42.59	>4.62 ^b
	0.01	3	10.44	>42.59°	>4.08°
Tamoxifen Citrate	0.004	2	34.12	37.96	1.11
	0.01	1,2 ^a	8.98	37.96	4.23
Terconazole Vetranal	0.004	3	11.92	41.46	3.48
	0.01	2,3 ^a	16.14	41.46	2.57
Thiethylperazine Maleate	0.004	3	7.09	18.37	2.59
	0.01	3	8.02	18.37	2.29
Toremifene Citrate	0.004	2,3 ^a	4.77	20.51	4.3
	0.01	3	11.3	20.51	1.81
Triparanol	0.004	2,3 ^a	4.68	21.21	4.53
	0.01	2,3 ^a	6.41	21.21	3.31