

1 Favipiravir and severe acute respiratory syndrome coronavirus 2 in hamster 2 model

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11 Summary

12 There is a need for safe and effective antiviral molecules with which to combat COVID-19 pandemics.
13 Recently, *in vitro* inhibitory activity of favipiravir against SARS-CoV-2 was reported. Here, we used a
14 Syrian hamster model to explore the pharmacokinetics of this molecule and its *in vivo* efficacy against
15 SARS-CoV-2. Results revealed that high doses (700-1400mg/kg/day) significantly reduced virus
16 replication in the lungs accompanied by clinical alleviation of the disease. However, these high doses
17 were associated with significant toxicity in hamsters. Favipiravir pharmacokinetics displayed non-linear
18 increase in plasma exposure between the doses and good lung penetration. Analysis of viral genomes
19 *in vivo* showed that favipiravir induced a mutagenic effect. Whilst the plasma trough concentrations
20 observed in this study were comparable with those previously found during human clinical trials, this
21 potential toxicity requires further investigation to assess whether a tolerable dosing regimen can be
22 found in humans that effectively reduces virus replication.

23 Keywords

24 COVID-19, SARS-CoV-2, antiviral therapy, favipiravir, animal model, preclinical research

25 Introduction

26 In March 2020, the World Health Organization declared coronavirus disease 2019 (COVID-19) a
27 pandemic (WHO, 2020). The COVID-19 outbreak was originally identified in Wuhan, China, in
28 December 2019 and spread rapidly around the world within a few months. The severe acute
29 respiratory syndrome coronavirus 2 (SARS-CoV-2), the causative agent of COVID-19, belongs to the
30 *Coronaviridae* family and is closely related to the SARS-CoV which emerged in China in 2002 (Zhu et
31 al., 2020). After an incubation period of about 5 days, disease onset usually begins with an influenza-
32 like syndrome associated with high virus replication in respiratory tracts (Huang et al., 2020, He et al.,
33 2020). In some patients, a late acute respiratory distress syndrome, associated with high levels of
34 inflammatory proteins, occurs within one to two weeks (Huang et al., 2020). As of 7 July 2020, more
35 than 11.6 million cases of COVID-19 have resulted in more than 538,000 deaths (Dong et al., 2020). In
36 the face of this ongoing pandemic and its unprecedented repercussions, not only on human health but
37 also on society, ecology and economy, there is an urgent need for effective infection prevention and
38 control measures.

39 Whilst host-directed and immune-based therapies could prove useful for the clinical management of
40 critically ill patients, the availability of safe and effective antiviral molecules would represent an
41 important step towards fighting the current pandemic. As conventional drug development is a slow
42 process, repurposing of drugs already approved for any indication was extensively explored and led to
43 the implementation of many clinical trials for the treatment of COVID-19 (Mercorelli et al., 2018).
44 However, the development of effective antiviral drugs for the treatment of COVID-19, should, as much
45 as possible, rely on robust pre-clinical *in vivo* data, not only on efficacy generated *in vitro*. Accordingly,
46 rapid implementation of rodent and non-human primate animal models should help to assess more
47 finely the potential safety and efficacy of drug candidates and to determine appropriated dose
48 regimens in humans (Chan et al., 2020, Rockx et al., 2020).

49 Favipiravir (6-fluoro-3-hydroxypyrazine-2-carboxamine) is an anti-influenza drug approved in Japan
50 that has shown broad-spectrum antiviral activity against a variety of other RNA viruses (Guedj et al.,
51 2018, Yamada et al., 2019, Segura Guerrero et al., 2018, Tani et al., 2018, Jochmans et al., 2016,
52 Takahashi et al., 2003, Rosenke et al., 2018). Favipiravir is a prodrug that is metabolized intracellularly
53 into its active ribonucleoside 5'-triphosphate form that acts as a nucleotide analogue to selectively
54 inhibit RNA-dependent RNA polymerase and induce lethal mutagenesis (Baranovich et al., 2013,
55 Sangawa et al., 2013). Recently, several studies reported *in vitro* inhibitory activity of favipiravir against
56 SARS-CoV-2 with 50% effective concentrations (EC₅₀) ranging from 62 to >500µM (10 to >78µg/mL)
57 (Wang et al., 2020, Jeon et al., 2020, Shannon et al., 2020). Based on these results, more than 20 clinical
58 trials on the management of COVID-19 by favipiravir are in progress (<https://clinicaltrials.gov/>). In the

59 present study, a Syrian hamster model (*Mesocricetus auratus*) was implemented to explore the *in vivo*
60 safety and efficacy and the pharmacokinetics (PK) of several dosing regimens of favipiravir.

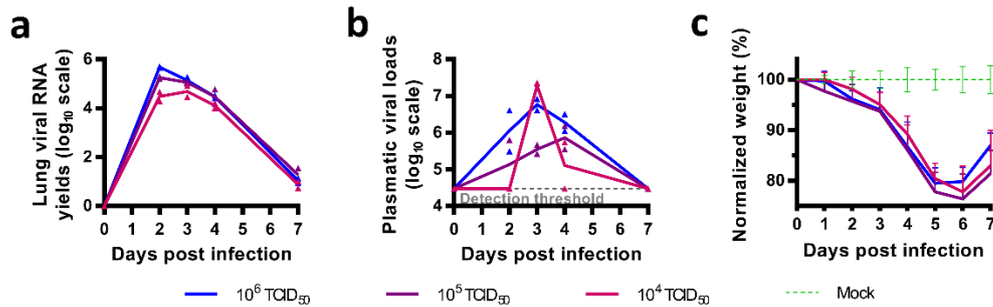
61 Results

62 *In vitro* efficacy of favipiravir

63 Using VeroE6 cells and an antiviral assay based on reduction of cytopathic effect (CPE), we recorded
64 EC_{50} and EC_{90} of 32 and 52.5 $\mu\text{g}/\text{mL}$ using a multiplicity of infection (MOI) of 0.001, 70.0 and $>78.5\mu\text{g}/\text{mL}$
65 with an MOI of 0.01 (Figure S1) in accordance with previous studies (Wang et al., 2020, Jeon et al.,
66 2020, Shannon et al., 2020). Infectious titer reductions (fold change in comparison with untreated cells)
67 were ≥ 2 with 19.6 $\mu\text{g}/\text{mL}$ of favipiravir and ranged between 11 and 342 with 78.5 $\mu\text{g}/\text{mL}$. Using CaCo2
68 cells, which do not exhibit CPE with SARS-CoV-2 BavPat1 strain, infectious titer reductions were around
69 5 with 19.6 $\mu\text{g}/\text{mL}$ of favipiravir and ranged between 144 and 7721 with 78.5 $\mu\text{g}/\text{mL}$ of the drug. 50%
70 cytotoxic concentrations (CC_{50}) in VeroE6 and CaCo2 cells were $>78.5\mu\text{g}/\text{mL}$.

71 Infection of Syrian hamsters with SARS-CoV-2

72 Following Chan *et al.*, we implemented a hamster model to study the efficacy of antiviral compounds
73 (Chan et al., 2020). Firstly, we intranasally infected four-week-old female Syrian hamsters with 10^6
74 $TCID_{50}$ of virus. Groups of two animals were sacrificed 2, 3, 4 and 7 days post-infection (dpi). Viral
75 replication was quantified in sacrificed animals by RT-qPCR in organs (lungs, brain, liver, small/large
76 bowel, kidney, spleen and heart) and plasma. Viral loads in lungs peaked at 2 dpi, remained elevated
77 until 4 dpi and dramatically decreased at 7 dpi (Figure 1a). Viral loads in plasma peaked at 3 dpi and
78 viral replication was detected in the large bowel at 2 dpi (Figure 1b and Table S1). No viral RNA was
79 detected in almost all the other samples tested (Table S1). Subsequently, we infected animals with two
80 lower doses of virus (10^5 and 10^4 $TCID_{50}$). Viral RNA was quantified in lungs, large bowel and plasma
81 from sacrificed animals 2, 3, 4 and 7 dpi (Figure 1a and 1b). Viral loads in lungs peaked at 2 and 3 dpi
82 with doses of 10^5 and 10^4 $TCID_{50}$ respectively. Maximum viral loads in lungs of animals infected with
83 each dose of virus were comparable. Viral RNA yields in plasma and large bowel followed a similar
84 trend but with more variability, with this two lower doses. In addition, clinical monitoring of animals
85 showed no marked symptoms of infection but significant weight losses from 3 dpi when compared to
86 animals intranasally inoculated with sodium chloride 0.9% (Figure 1c).



87

88 **Figure 1: Implementation of hamster model**

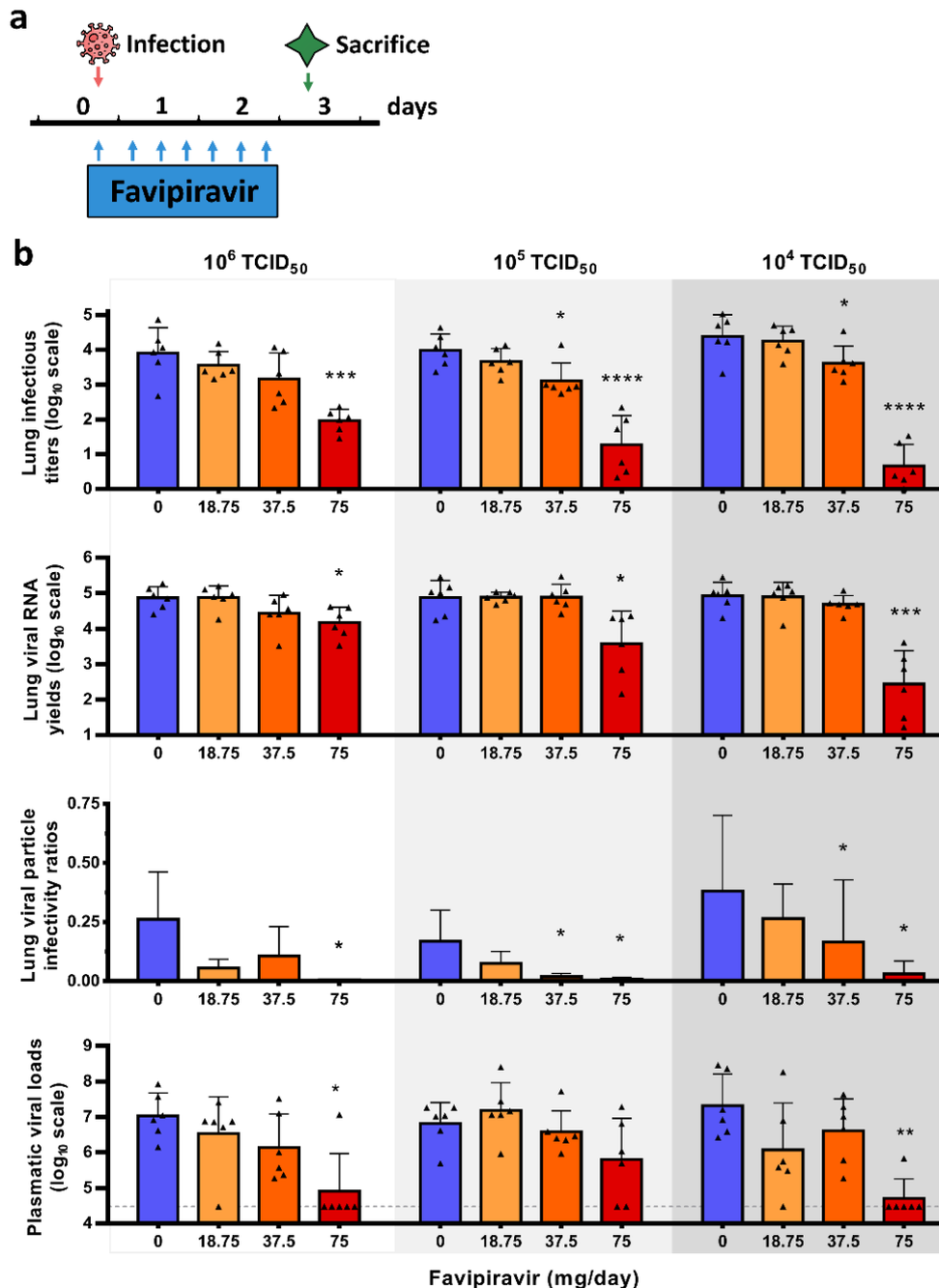
89 Hamsters were intranasally infected with 10⁶, 10⁵ or 10⁴ TCID₅₀ of virus. Viral replication was quantified using
90 an RT-qPCR assay. **a** Lung viral RNA yields. **b** Plasmatic viral loads. **c** Clinical course of the disease. Normalized
91 weight at day *n* was calculated as follows: (% of initial weight of the animal at day *n*)/(mean % of initial weight
92 for mock-infected animals at day *n*). Data represent mean ±SD (details in Table S1).

93 *In vivo* efficacy of favipiravir

94 To assess the efficacy of favipiravir, hamsters received the drug, intraperitoneally, three times a day
95 (TID). We used three doses of favipiravir: 18.75, 37.5 and 75mg/day (corresponding to 340±36, 670±42
96 and 1390±126 mg/kg/day respectively).

97 In a first set of experiments, treatment was initiated at day of infection (preemptive antiviral therapy)
98 and ended at 2 dpi. We infected groups of 6 animals intranasally with three doses of virus (10⁶, 10⁵ and
99 10⁴ TCID₅₀) and viral replication was measured in lungs and plasma at 3 dpi (Figure 2a). When analysis
100 of virus replication in clarified lung homogenates was based on infectious titers (as measured using
101 TCID₅₀ assay), an inverse relationship was observed between infectious titers and the dose of
102 favipiravir (Figure 2b). This trend was even more important when low doses of virus were used to infect
103 animals. At each dose of virus, mean infectious titers for groups of animals treated with 75mg/day TID
104 were significantly lower than those observed with untreated groups ($p \leq 0.0001$): reduction of infectious
105 titers ranged between 1.9 and 3.7 log₁₀. For animals infected with 10⁵ or 10⁴ TCID₅₀, significant
106 infectious titer reductions of around 0.8 log₁₀ were also observed with the dose of 37,5mg/day TID
107 ($p \leq 0.038$). Drug 90% and 99% effective doses (ED₉₀ and ED₉₉) were estimated based on these results
108 and ranged between 31-42mg/day and 53-70mg/day respectively (Table 2). When analysis of virus
109 replication in clarified lung homogenates were assessed on viral RNA yields (as measured using
110 quantitative real time RT-PCR assay), significant differences with groups of untreated animals, ranging
111 between 0.7 and 2.5 log₁₀, were observed only with the higher dose of favipiravir ($p \leq 0.012$). Once
112 again, this difference was more noticeable with lower doses of virus (Figure 2b). Since we found higher
113 reductions of infectious titers than those observed with viral RNA yields, we estimated the relative
114 infectivity of viral particle (*i.e.* the ratio of the number of infectious particles over the number of viral
115 RNA molecules). Decreased infectivity was observed in all treated groups of animals. These differences

116 were always significant with the higher dose of favipiravir ($p \leq 0.031$) and were significant with the dose
117 of 37.5mg/day TID for animals infected with 10^5 or 10^4 TCID₅₀ of virus ($p \leq 0.041$). We then measured
118 plasmatic viral loads using quantitative real time RT-PCR assay and found, with the higher dose of
119 favipiravir and the groups of animals infected with 10^6 or 10^4 TCID₅₀, significant reductions of 2.1 and
120 2.62 log₁₀, respectively ($p \leq 0.022$) (Figure 2b).



121

122 **Figure 2: Virological results with preemptive favipiravir therapy**

123 **a** Experimental timeline. **b** Viral replication in lungs and plasma. Hamsters were intranasally infected with 10^6 ,
124 10^5 or 10^4 TCID₅₀ of virus. Lung infectious titers (measured using a TCID₅₀ assay) and viral RNA yields were
125 (measured using an RT-qPCR assay) expressed in TCID₅₀/copy of γ -actine gene and viral genome copies/copy of

126 γ -actine gene respectively. Relative lung viral particle infectivities were calculated as follows: ratio of lung
127 infectious titer over viral RNA yields. Plasmatic viral loads (measured using an RT-qPCR assay) are expressed in
128 viral genome copies/mL of plasma (the dotted line indicates the detection threshold of the assay). Data represent
129 mean \pm SD. ****, ***, ** and * symbols indicate that the average value for the group is significantly lower than
130 that of the untreated group with a p-value <0.0001 , ranging between 0.0001-0.001, 0.001-0.01 and 0.01-0.05
131 respectively (details in Table S2 and S3).

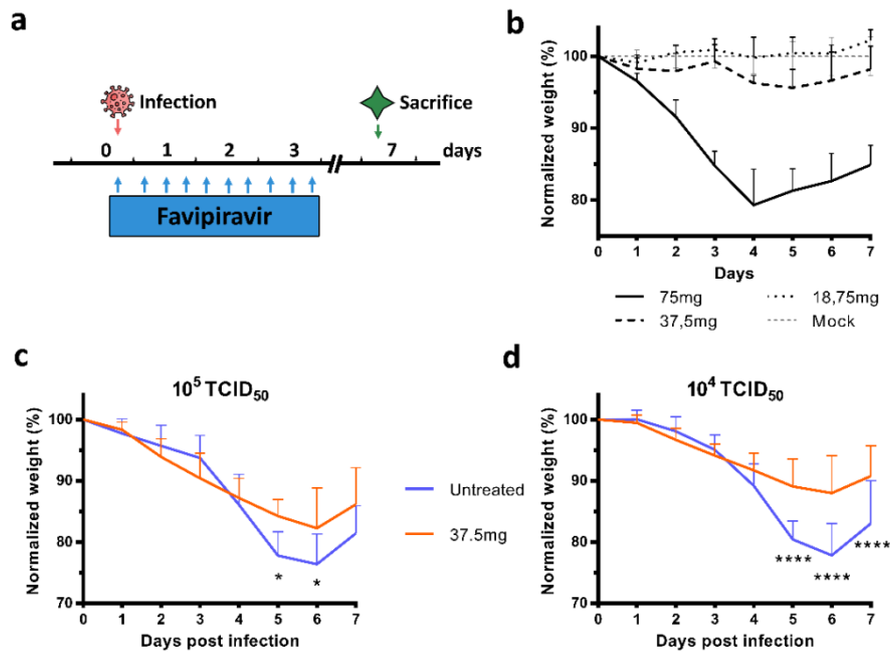
132 **Table 2: 50%, 90% and 99% drug effective doses**

	ED ₅₀ mg/day (95%CI ¹)	ED ₉₀ mg/day (95%CI ¹)	ED ₉₉ mg/day (95%CI ¹)
Preemptive therapy			
10 ⁴ TCID ₅₀	34 (30-37)	42 (38-46)	53 (48-58)
10 ⁵ TCID ₅₀	26 (21-30)	37 (31-44)	56 (46-65)
10 ⁶ TCID ₅₀	15 (10-20)	31 (21-41)	70 (48-93)
Preventive therapy			
10 ⁴ TCID ₅₀	27 (25-29)	35 (32-38)	47 (44-51)

¹: 95% confidence interval

Dose-response curves are presented in Figure S2.

133 In a second set of experiments, we assessed, over a period of 7 days, the impact of treatment on the
134 clinical course of the disease using weight loss as the primary criterion (Figure 3a). Beforehand, we
135 evaluated the toxicity of the three doses of favipiravir with groups of four non-infected animals treated
136 from day 0 to day 3 (Figure 3b). High toxicity was observed with the dose of 75mg/day TID with
137 significant weight loss noticed from the first day of treatment (Table S4). We also found a constant,
138 but moderate, toxicity with the dose of 37.5mg/day TID that was significant at day 4 and 5 only. No
139 toxicity was detected with the lower dose of favipiravir. To assess if the toxicity observed with the
140 highest dose of favipiravir was exacerbated by the infection, we compared weight losses of infected
141 and non-infected animals treated with the dose of 75mg/day TID. Regardless of the dose of virus, no
142 significant difference was observed at 1, 2 and 3 dpi (Figure S3). After this evaluation of favipiravir
143 toxicity, we intranasally infected groups of 10 animals with two doses of virus (10⁵ or 10⁴ TCID₅₀).
144 Treatment with a dose of 37.5mg/day TID was initiated on the day of infection (preemptive antiviral
145 therapy) and ended at 3 dpi (Figure 3a). With both doses of virus, treatment was associated with
146 clinical alleviation of the disease (Figure 3c-d). With the dose of 10⁵ TCID₅₀, mean weights of treated
147 animals were significantly higher than those of untreated animals at 5 and 6 dpi ($p \leq 0.031$). Similar
148 observations were made with the dose of 10⁴ TCID₅₀ at 5, 6 and 7 dpi ($p < 0.0001$).



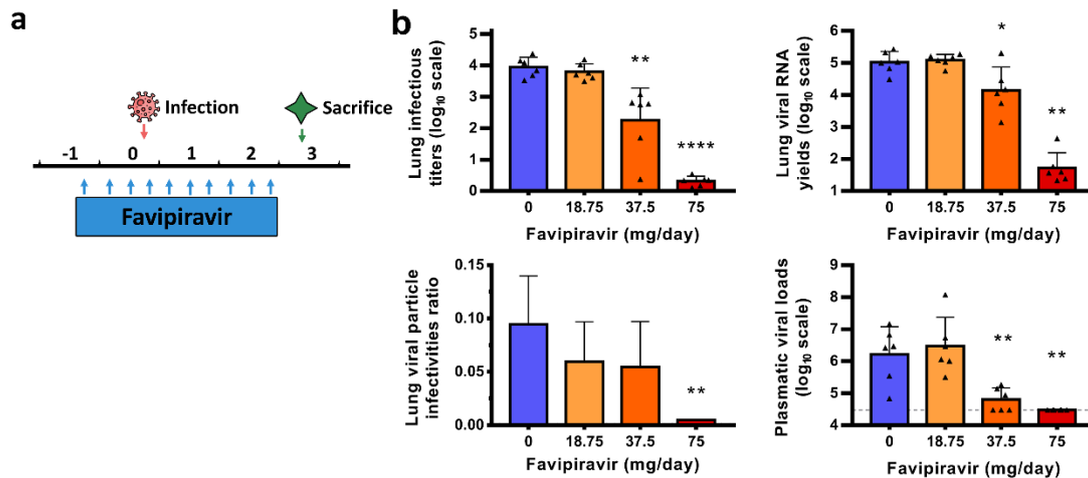
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150 **Figure 3: Clinical follow-up with preemptive favipiravir therapy**

151 **a** Experimental timeline. **b** Evaluation of the toxicity of the three doses of favipiravir (mg/day TID) with uninfected
 152 animals following an identical experimental timeline without infection. **c-d** Clinical follow-up with animals
 153 infected respectively with 10^5 and 10^4 TCID₅₀ of virus and treated with a dose of favipiravir of 37.5mg/day TID.
 154 Normalized weight at day n was calculated as follows: (% of initial weight of the animal at day n)/(mean % of
 155 initial weight for mock-infected animals at day n). Data represent mean \pm SD. **** and * symbols indicate a
 156 significant difference between treated and untreated animals with a p-value <0.0001 and ranging between 0.01-
 157 0.05 respectively (details in Table S2 and S4).

158

159 In a third set of experiments, treatment was started one day before infection (preventive antiviral
 160 therapy) and ended at 2 dpi. We intranasally infected groups of 6 animals with 10^4 TCID₅₀ of virus and
 161 viral replication was measured in lungs and plasma at 3 dpi (Figure 4a). Once again, an inverse
 162 relationship was observed between lung infectious titers and the dose of favipiravir (Figure 4b). Mean
 163 infectious titers for groups of animals treated with 37.5 and 75mg/day TID were significantly lower
 164 than those observed with untreated groups ($p \leq 0.002$). Of note, undetectable infectious titers were
 165 found for all animals treated with the higher dose. Estimated ED₉₀ and ED₉₉ were 35 and 47mg/day
 166 respectively (Table 2). Significant reductions of viral RNA yields of 0.9 and 3.3 log₁₀, were observed with
 167 animals treated with 37.5 and 75mg/day TID respectively ($p \leq 0.023$). Resulting infectivity of viral
 168 particle was decreased, with a significant reduction only for the higher dose of favipiravir ($p = 0.005$).
 169 Finally, we found significantly reduced plasmatic viral loads with animals treated with 37.5 and
 170 75mg/day TID ($p \leq 0.005$).



170

171 **Figure 4: Virological results with preventive favipiravir therapy**

172 **a** Experimental timeline. **b** Viral replication in lungs and plasma. Hamsters were intranasally infected with 10^4
173 $TCID_{50}$ of virus. Lung infectious titers (measured using a $TCID_{50}$ assay) and viral RNA yields were (measured
174 using an RT-qPCR assay). They are expressed in $TCID_{50}/copy$ of γ -actine gene and viral genome copies/copy of
175 γ -actine gene respectively. Relative lung virus infectivities were calculated as follows: ratio of lung infectious titer
176 over viral RNA yields. Plasmatic viral loads (measured using an RT-qPCR assay) are expressed in viral genome
177 copies/mL of plasma (the dotted line indicates the detection threshold of the assay). Data represent mean \pm SD.
178 ****, ** and * symbols indicate that the average value for the group is significantly different from that of the
179 untreated group with a p-value <0.0001 , ranging between 0.001-0.01 and 0.01-0.05 respectively (details in Table
180 S2 and S3).

181 **Favipiravir pharmacokinetics (PK) in a hamster model**

182 We first assessed the PK and lung distribution of favipiravir in a subgroup of uninfected animals. Groups
183 of animals were treated respectively with a single dose of favipiravir administered intraperitoneally:
184 6.25mg, 12.5 mg and 25 mg. In each dose group, we sacrificed 3 animals at specific time points post-
185 treatment (0.5, 1, 5 or 8 hours) for determination of favipiravir in plasma. Drug concentration in lung
186 tissue was determined at 0.5 and 5 hours post-treatment. Subsequently, we assessed the favipiravir
187 concentration after multiple dose in animals intranasally infected with 10^5 $TCID_{50}$ of virus. Groups of 9
188 animals received the three doses evaluated for 3 days (Figure 2a): 18.75mg/day, 37.5mg/day and
189 75mg/day TID and were sacrificed at 12-hours after the last treatment dose. Favipiravir was quantified
190 in plasma (n=9) and lung tissue (n=3).

191 Results are presented in Table 3 and Figure S4. The single dose PK analysis showed that the maximum
192 concentration of favipiravir was observed at 0.5 hour at all doses, then plasma drug concentrations
193 decreased exponentially to reach concentrations below $10 \mu\text{g}/\text{ml}$ at 12 hours. Favipiravir PK exhibited
194 a non-linear increase in concentration between the doses. After multiple doses, trough concentrations
195 (12 hours) of favipiravir also exhibited a non-linear increase between doses. The extrapolated 12 hours
196 post-treatment concentrations after a single dose were calculated in order to determine the

197 accumulation ratio. Accumulation ratios were respectively 6, 16 and 21 at the 3 doses, confirming the
 198 non-proportional increase between doses. The average concentration after single dose administration
 199 over 0 to 12-hour intervals was calculated and the respective values obtained were 10.1 µg/mL, 38.7
 200 µg/mL and 100.5 µg/mL for the 3 favipiravir doses.
 201 Favipiravir lung concentrations were 1.6 to 2.7-fold lower than in plasma for both administration of
 202 single and multiple doses. After a single dose, the mean lung to plasma ratio ranged from 0.37 to 0.62
 203 according to the time post-treatment and was similar between the 3 doses of favipiravir at 0.5 hours.
 204 A high ratio 5 hours post-treatment was observed at the highest dose (25 mg) with an increase by a
 205 factor 1.6 to 1.8 compared with the lower doses. After multiple doses, the lung penetration of
 206 favipiravir was confirmed with a mean lung to plasma ratio ranging from 0.35 to 0.44. Favipiravir was
 207 not detected in the lungs at the lowest dose (18.75 mg/day).

208 **Table 3: Plasma and lung concentrations of favipiravir after administration of a single dose or multiple**
 209 **dose of favipiravir**

	Single Dose			Multiple Dose ¹ (Day 3)		
	Plasma (µg/mL)	Lung (µg/g)	L/p ratio	Plasma (µg/mL)	Lung (µg/g)	L/p ratio
Dose: 25 mg						
0.5 hr	372 ± 47.5	216 ± 39	0.58 ± 0,04			
1 hr	279 ± 49.9					
5 hr	135 ± 49.0	81,3 ± 24	0.62 ± 0,10			
8 hr	5.77 ± 1.34					
12 hr	1.43 ²			29.9 ± 9.83	16.0 ± 4.87	0.44 ± 0,07
Dose: 12.5 mg						
0.5 hr	166 ± 52.0	90.7 ± 12.7	0.58 ± 0.14			
1 hr	155 ± 20.6					
5 hr	10.7 ± 5.16	3.84 ± 1.49	0.37 ± 0.052			
8 hr	1.94 ± 0.06					
12 hr	0.16 ²			2.57 ± 1.22	1.36 ± 0.14	0.35 ± 0,03
Dose: 6.25 mg						
0.5 hr	86.3 ± 4.11	50.2 ± 16.4	0.58 ± 0.17			
1 hr	35.2 ± 27.8					
5 hr	2.90 ± 0.25	1.09 ± 0.05	0.38 ± 0.05			
8 hr	0.56 ± 0.16					
12 hr	0.05 ²			0.31 ± 0.14	not detected	<i>n.a.</i>

210 Data represent mean ±SD; Three animals for each condition except at multiple dose (n=9 for plasma; n=3 for
 211 lung); details in Table S5

212 ¹: PK realized after 3 days of favipiravir administered three times a day (18.75, 37.5 or 75mg/day TID)

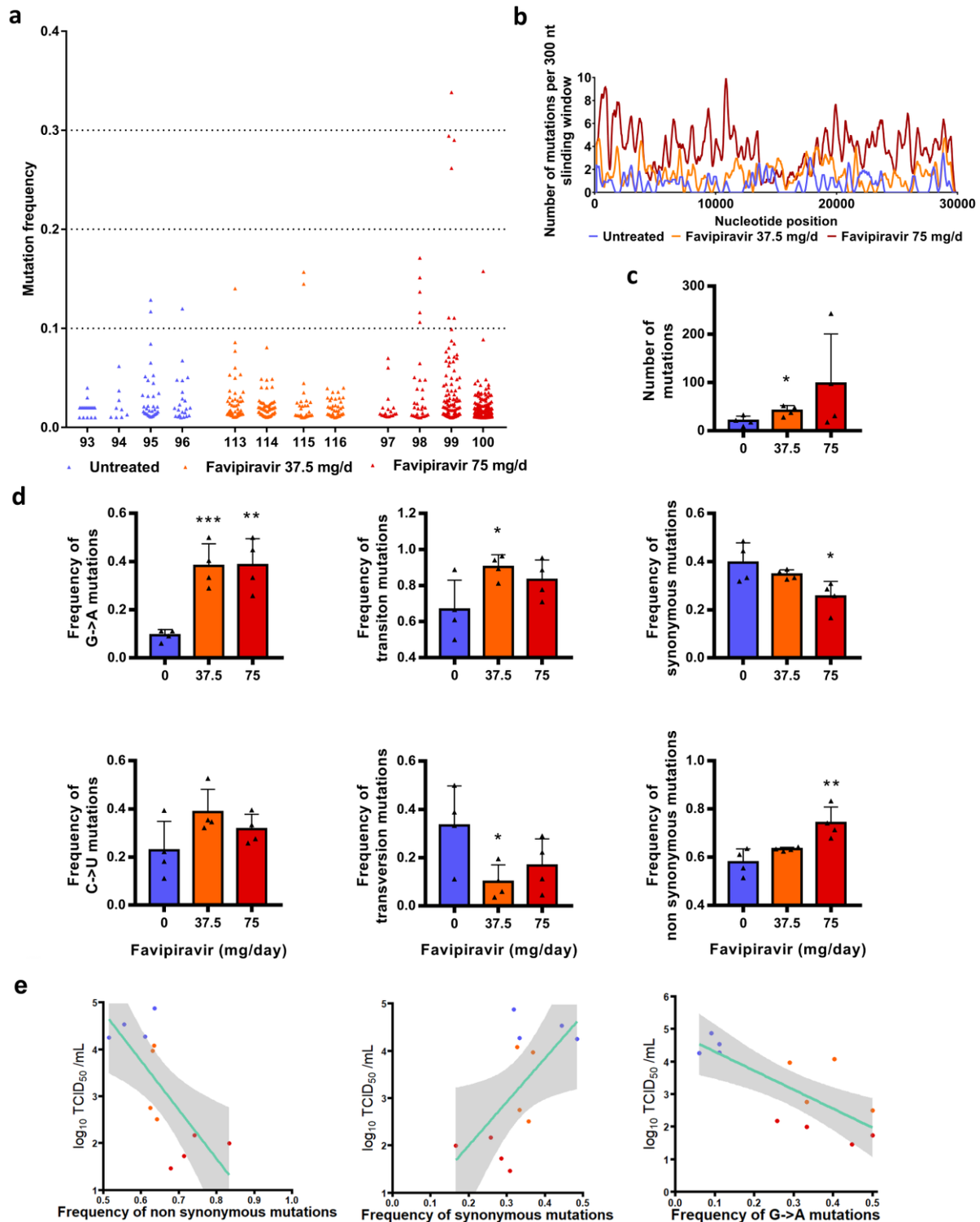
213 ²: extrapolated C_{12h}. *na*: not applicable

214 Mutagenic effect of favipiravir

215 To understand which genomic modifications accompanied favipiravir treatment, direct complete
216 genome sequencing of clarified lung homogenates from animals intranasally infected with 10^6 TCID₅₀
217 of virus and treated with the two highest doses of drug (preemptive antiviral therapy; Figure 2) was
218 performed. Data were generated by next generation sequencing from lung samples of four animals
219 per group (untreated, 37.5mg/day TID and 75mg/day TID). The mean sequencing coverage for each
220 sample ranged from 10,991 to 37,991 reads per genomic position and we subjected substitutions with
221 a frequency $\geq 1\%$ to further analysis. The genetic variability in virus stock was also analyzed: 14
222 nucleotide polymorphisms were detected of which 5 recorded a mutation frequency higher than 10%
223 (Table S6).

224 In order to study the mutagenic effect of favipiravir, we used the consensus sequence from virus stock
225 as reference and all the mutations simultaneously detected in a lung sample and in virus stock were
226 not considered in the further analysis (1 to 4 mutations per sample, see Table S6). Overall, no majority
227 mutations were detected (mutation frequency $>50\%$), mutations were distributed throughout the
228 whole genome and almost all of them exhibited a frequency lower than 10% (Figure 5a and 5b).

229 Results revealed a relationship between the number of mutations detected per sample and the dose
230 of favipiravir (Figure 5c): the mean number of mutations increased by a factor 2 and 4.8 with groups
231 of animals treated with 37.5 and 75mg/day TID, respectively. The difference is significant only with a
232 dose of 37.5mg/day TID ($p=0.029$). This increase of the number of mutations is mainly the consequence
233 of the occurrence of a large number of G→A substitutions and, to a lesser extent, C→U substitutions.
234 Consequently, regardless of the dose of favipiravir, mean frequency of G→A substitutions was
235 significantly increased by a factor of 4.2 ($p\leq 0.009$). This rise of these transition mutations led to
236 increased frequency of all transition mutations (significant only at dose of 37.5mg/day TID; $p=0.037$)
237 and increased frequency of non-synonymous mutations (significant only at dose of 75mg/day TID;
238 $p=0.009$) (Figure 5d). We investigated whether or not effectiveness in treated animals was linked with
239 the characteristics of the mutations detected on viral populations and found that frequency of non-
240 synonymous, synonymous and G→A mutations were associated with infectious titers in lungs ($p<0.03$;
241 Figure 5e). Finally, our experiments revealed some parallel evolution events; 32 substitutions in viral
242 sub-populations were detected in two independent animals. Notably, 18 of these shared mutations
243 were detected only with treated animals, 14 of them being non-synonymous (Table S8). These
244 mutations are located in nsp2, 3, 4, 5, 6, 14, N protein, Matrix, ORF 3a and 8. At this stage, one cannot
245 conclude if these substitutions reflect the adaptation to the hamster model or are the result of the
246 antiviral selection.



247

248 **Figure 5: Mutagenic effect of favipiravir**

249 **a** Viral genetic diversity in clarified lung homogenates. For each condition, four samples were analyzed. Each
 250 triangle represents a mutation (only substitutions with a frequency $\geq 1\%$ were considered). **b** Patterns of mutation
 251 distribution on complete viral genome. Each variable nucleotide position was counted only once when found.
 252 The variability was represented using 75 nt sliding windows. For each condition, variable nucleotide positions
 253 were determined and represented using a 300 nt sliding window. **c** Mean number of mutations. Data represent

254 mean \pm SD. **d** Mutation characteristics. For each sample, the frequency of a given mutation was calculated as
255 follows: number of this kind of mutation detected in the sample divided by the total number of mutations
256 detected in this sample. Data represent mean \pm SD. ** and * symbols indicate that the average value for the group
257 is significantly different from that of the untreated group with a p-value ranging between 0.001-0.01 and 0.01-
258 0.05 respectively (details in details in Table S6 and S7). **e** Association between lung infectious titers (measured
259 using a TCID₅₀ assay) and frequency of non synonymous, synonymous and G→A mutations. Each dot represent
260 data from a given animal.

261 Discussion

262 In the current study, we used a hamster model to assess efficacy of the favipiravir against SARS-CoV-
263 2. Following infection, viral RNA was mainly detected in lungs, blood, and, to a lesser extent, in the
264 large bowel. Peak of viral replication was observed at 2-3 dpi followed by observation of significant
265 weight losses, in line with recently reported investigations that involved 6-10 weeks old hamsters
266 (Kaptein et al., 2020, Chan et al., 2020). Clinically, the main symptom was weight loss, observed from
267 the first day of infection and followed by recovery at 6dpi. This confirmed that the *in vivo* model, with
268 younger animals (4 weeks-old), is suitable for preclinical evaluation of antiviral compounds against
269 SARS-CoV-2.

270 Using a preemptive strategy, we demonstrated that doses of favipiravir of around 700-1400mg/kg/day
271 TID reduced viral replication in lungs of infected animals and allowed clinical alleviation of the disease.
272 Reduction of viral replication was greater when estimated on the basis of infectious titers than on total
273 viral RNA as previously observed in non-human primates treated with Remdesivir (Williamson et al.,
274 2020). However, the effective doses of favipiravir were higher than those usually used in rodent models
275 (\approx 100-400mg/kg/day) (Sidwell et al., 2007, Smither et al., 2014, Julander et al., 2009, Tani et al., 2018,
276 Oestereich et al., 2016, Yamada et al., 2019). This can be correlated with the high favipiravir EC₅₀ found
277 *in vitro* for SARS-CoV-2. Moreover, effective doses were associated with significant toxicity in our
278 hamster model. This observed toxicity reflected only the adverse effects of favipiravir and was not
279 exacerbated during SARS-CoV-2 infection. Indeed, similar weight losses were measured among
280 infected and non-infected animals treated with the highest dose of favipiravir at 1, 2 and 3dpi.

281 In the present study, reduction of viral replication was correlated with the dose of favipiravir
282 administrated and inversely correlated with the dose of virus inoculated. In a recent study, favipiravir
283 administrated *per os* twice daily (loading dose of 600mg/kg/day followed by 300mg/kg/day) revealed
284 a mild reduction of lung viral RNA yields using a similar hamster model with high doses of virus (2×10^6
285 TCID₅₀) (Kaptein et al., 2020). These results are in accordance with ours at the lower dose of favipiravir
286 (around 340mg/kg/day TID).

287 By characterizing the dose response curve, we estimated that the dose required to reduce by 90%
288 (ED₉₀) the level of infectious titers in lungs is in the range of 570-780mg/kg/day. In the most favourable
289 situation, where high doses were used as a preemptive therapy, favipiravir led to undetectable viral
290 replication in lung and plasma. These results showed that the use of high doses of favipiravir could
291 expand its *in vivo* spectrum against RNA viruses.

292 With influenza viruses, favipiravir acts as a nucleotide analogue. It is metabolized intracellularly to its
293 active form and incorporated into nascent viral RNA strands. This inhibits RNA strand extension and

294 induces abnormal levels of mutation accumulation into the viral genome (Baranovich et al., 2013,
295 Sangawa et al., 2013). Recently, it was shown *in vitro* that favipiravir has a similar mechanism of action
296 with SARS-CoV-2 through a combination of chain termination, reduced RNA synthesis and lethal
297 mutagenesis (Shannon et al., 2020). Our genomic analysis confirmed the mutagenic effect of favipiravir
298 *in vivo*. Indeed, we found that favipiravir treatment induced appearance of a large number of G→A
299 and C→U mutations into viral genomes. This was associated to a decrease of viral infectivity probably
300 because alteration of the genomic RNA disturb the replication capacity. Similar findings were described
301 *in vitro* and *in vivo* with other RNA viruses (Baranovich et al., 2013, Guedj et al., 2018, Escribano-
302 Romero et al., 2017, Arias et al., 2014). Of note, we also observed a strong inverse association between
303 infectious titers in lungs and the proportion of non-synonymous mutations detected in viral
304 populations. Because random non-synonymous mutations are more deleterious than synonymous
305 mutations (Cuevas et al., 2012), this suggests that they were randomly distributed over the three
306 positions of the codons and that no compensatory mechanism was triggered by the virus to eliminate
307 them (*i.e.* negative selection). Finally, the inverse correlation between lung infections titers and the
308 frequency of G→A substitutions showed that an increased proportion of these mutations beyond an
309 error threshold might be expected to cause lethal mutagenesis.

310 Genomic analyses revealed that 18 mutations detected in viral sub-populations were shared only with
311 treated animals. Two of them were located in the nsp14 coding region involved in the proof-reading
312 activity of the viral RNA polymerisation (Eckerle et al., 2007, Ferron et al., 2018). However, they were
313 located in the N7 MTase domain involved in viral RNA capping (Chen et al., 2013, Ma et al., 2015). By
314 comparison, resistance mutations selected against Remdesivir in β -coronavirus murine hepatitis virus
315 model were obtained in the RdRP (nsp12) coding sequence (Agostini et al., 2018). Further
316 investigations are needed to assess the impact of these mutations on the antiviral effect of favipiravir.
317 Favipiravir PK in our hamster model displayed a non-linear increase in plasma exposure between the
318 doses as already reported in nonhuman primates (Madelain et al., 2017). The observed favipiravir
319 concentration versus time profiles were in agreement with previous results of a PK study performed
320 in 7-8 week-old hamsters orally treated with a single dose of 100mg/kg of favipiravir (Gowen et al.,
321 2015). The maximum plasma drug concentration occurred at 0.5 h after oral administration, earlier
322 than in humans, and then decreased rapidly in agreement with its short half-life (Madelain et al., 2016).
323 After repeated doses, plasma exposure confirmed non-linear PK over the entire range of doses, further
324 emphasized by accumulation ratios. The important accumulation observed at the highest dose could
325 explain in part the toxicity observed in hamsters at this dose. Favipiravir undergoes an important
326 hepatic metabolism mainly by aldehyde oxidase producing an inactive M1 metabolite and inhibits
327 aldehyde oxidase activity in a concentration- and time-dependent manner. These properties explain

328 the self-inhibition of its own metabolism as observed in our study in which the highest dose of
329 favipiravir led to a greater increase in favipiravir concentrations (Madelain et al., 2020).

330 A good penetration of favipiravir in lungs was observed with lung/plasma ratios ranging from 35 to
331 44% after repeated doses, consistent with its physicochemical properties. Lung exposure was also in
332 accordance with previous studies (Gowen et al., 2015).

333 How clinically realistic are these results? To address this question we compared the drug
334 concentrations obtained in the hamster model with those obtained in patients. In 2016, a clinical trial
335 evaluated the use of favipiravir in Ebola infected patients (Sissoko et al., 2016). The dose used in Ebola
336 infected patients was 6000mg on day 0 followed by 1200mg BID for 9 days. The median trough
337 concentrations of favipiravir at Day 2 and Day 4 were 46.1 and 25.9µg/mL, respectively. This is within
338 the range observed here in hamsters treated with the highest dose (around 1400mg/kg/day), with a
339 mean trough concentration of 29.9µg/mL. However, additional investigations are required to
340 determine whether or not similar favipiravir plasma exposure in SARS-COV-2 infected patients are
341 associated with antiviral activity. The major differences in PK between hamster and humans, and the
342 toxicity observed at the highest doses in our animal model limits the extrapolation of our results.
343 Therefore, whether safe dosing regimens in humans may achieve similar plasma exposure and
344 recapitulate the profound effect on viral replication is unknown. Further, the intracellular
345 concentration of the active metabolite was not determined and which parameter of the drug
346 pharmacokinetics best drives the antiviral effect remains to be established.

347 In summary, this study establishes that high doses of favipiravir are associated with antiviral activity
348 against SARS-CoV-2 infection in a hamster model. The better antiviral efficacy was observed using a
349 preventive strategy, suggesting that favipiravir could be more appropriate for a prophylactic use. Our
350 results should be interpreted with caution because high doses of favipiravir were associated with signs
351 of toxicity in our model. It is required to determine if a tolerable dosing regimen could generate similar
352 exposure in non-human primates, associated with significant antiviral activity, before testing a high
353 dose regimen in COVID-19 patients. Furthermore, subsequent studies should determine if an increased
354 antiviral efficacy can be reached using favipiravir in association with other effective antiviral drugs,
355 since this strategy may enable to reduce the dosing regimen of favipiravir. Finally, this work reinforces
356 the need for rapid development of animal models to confirm *in vivo* efficacy of antiviral compounds
357 and accordingly, to determine appropriate dose regimens in humans before starting clinical trials.

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369 Author Contributions

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372 Resources, F.T., B.C., J.G., X.d.L., C.S. and A.N. ; Writing – Original Draft, J.S.D., M.C., J.G., C.S. and A.N.
373 ; Writing – Review & Editing, J.G., X.d.L., C.S. and A.N. ; Visualization, J.S.D., M.C., G.L., F.T., P.R.P. and
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375 Declaration of Interests

376 J.G has consulted for F. Hoffman-La Roche. C.S has consulted for ViiV Healthcare, MSD and Gilead.
377 The remaining authors declare no competing interests.

378 Methods

379 Cells

380 VeroE6 cells (ATCC CRL-1586) and Caco-2 cells (ATCC HTB-37) were grown at 37°C with 5% CO₂ in
381 minimal essential medium (MEM) supplemented with 7.5% heat-inactivated fetal bovine serum (FBS),
382 1% Penicillin/Streptomycin and 1% non-essential amino acids (all from ThermoFisher Scientific).

383 Virus

384 All experiments with infectious virus were conducted in biosafety level (BSL) 3 laboratory. SARS-CoV-2
385 strain BavPat1, supplied through European Virus Archive GLOBAL (<https://www.european-virus->
386 [archive.com/](https://www.european-virus-archive.com/)), was provided by Christian Drosten (Berlin, Germany). Virus stocks were prepared by
387 inoculating at MOI of 0.001 a 25cm² culture flask of confluent VeroE6 cells with MEM medium
388 supplemented with 2.5% FBS. The cell supernatant medium was replaced each 24h hours and
389 harvested at the peak of infection, supplemented with 25mM HEPES (Sigma), aliquoted and stored at
390 -80°C.

391 *In vitro* determination of EC₅₀, EC₉₀, CC₅₀ and infectious titer reductions

392 One day prior to infection, 5×10⁴ VeroE6 cells were seeded in 96-well culture plates (5×10⁴ cells/well
393 in 100µL of 2.5% FBS medium (assay medium). The next day, seven 2-fold serial dilutions of favipiravir
394 (Courtesy of Toyama-Chemical; 0.61µg/mL to 78.5µg/mL, in triplicate) were added (25µL/well, in assay
395 medium). Eight virus control wells were supplemented with 25µL of assay medium and eight cell
396 controls were supplemented with 50µL of assay medium. After 15 min, 25µL of virus suspension,
397 diluted in assay medium, was added to the wells at an MOI of 0.01 or 0.001 (except for cell controls).
398 Three days after infection, cell supernatant media were collected to perform TCID₅₀ assay (at
399 concentration of 78.5, 39.3, 19.6µg/mL), as described below, in order to calculate infectious titer
400 reductions and cell viability was assessed using CellTiter-Blue reagent (Promega) following
401 manufacturer's instructions. Fluorescence (560/590nm) was recorded with a Tecan Infinite 200Pro
402 machine (Tecan). The 50% and 90% effective concentrations (EC₅₀, EC₉₀) were determined using
403 logarithmic interpolation (% of inhibition were calculated as follows: $(OD_{\text{sample}} - OD_{\text{virus control}}) / (OD_{\text{cell control}} -$
404 $OD_{\text{virus control}})$). For the evaluation of CC₅₀ (the concentration that induced 50% cytotoxicity), the same
405 culture conditions were set as for the determination of the EC₅₀, without addition of the virus, then
406 cell viability was measured using CellTiter Blue (Promega). CC₅₀ was determined using logarithmic
407 interpolation.

408 *In vivo* experiments

409 Approval and authorization

410 *In vivo* experiments were approved by the local ethical committee (C2EA—14) and the French
411 ‘Ministère de l’Enseignement Supérieur, de la Recherche et de l’Innovation’ (APAFIS#23975) and
412 performed in accordance with the French national guidelines and the European legislation covering
413 the use of animals for scientific purposes. All experiments were conducted in BSL 3 laboratory.

414 Animal handling

415 Three-week-old female Syrian hamsters were provided by Janvier Labs. Animals were maintained in
416 ISOcage P - Bioexclusion System (Techniplast) with unlimited access to water/food and 14h/10h
417 light/dark cycle. Animals were weighed and monitored daily for the duration of the study to detect the
418 appearance of any clinical signs of illness/suffering. Virus inoculation was performed under general
419 anesthesia (isoflurane). Organs and blood were collected after euthanasia (cervical dislocation) which
420 was also realized under general anesthesia (isoflurane).

421 Hamster Infection

422 Anesthetized animals (four-week-old) were intranasally infected with 50µL containing 10⁶, 10⁵ or
423 10⁴ TCID₅₀ of virus in 0.9% sodium chloride solution). The mock group was intranasally inoculated with
424 50µL of 0.9% sodium chloride solution.

425 Favipiravir administration

426 Hamster were intra-peritoneally inoculated with different doses of favipiravir. Control group were
427 intra-peritoneally inoculated with a 0.9% sodium chloride solution.

428 Organ collection

429 Organs were first washed in 10mL of 0.9% sodium chloride solution and then transferred to a 2mL or
430 50mL tube containing respectively 1mL (small/large bowel pieces, kidney, spleen and heart) or 10mL
431 (lungs, brain and liver) of 0.9% sodium chloride solution and 3mm glass beads. They were crushed
432 using a the Tissue Lyser machine (Retsch MM400) for 5min at 30 cycles/s and then centrifuged 5min à
433 1200g. Supernatant media were transferred to a 2mL tube, centrifuged 10 min at 16,200g and stored
434 at -80°C. One milliliter of blood was harvested in a 2mL tube containing 100µL of 0.5M EDTA
435 (ThermoFischer Scientific). Blood was centrifuged for 10 min at 16,200g and stored at -80°C.

436 **Quantitative real-time RT-PCR (RT-qPCR) assays**

437 To avoid contamination, all experiments were conducted in a molecular biology laboratory that is
438 specifically designed for clinical diagnosis using molecular techniques, and which includes separate
439 laboratories dedicated to perform each step of the procedure. Prior to PCR amplification, RNA
440 extraction was performed using the QIAamp 96 DNA kit and the Qiacube HT kit and the Qiacube HT

441 (both from Qiagen) following the manufacturer's instructions. Shortly, 100 μ l of organ clarified
442 homogenates, spiked with 10 μ l of internal control (bacteriophage MS2) (Ninove et al., 2011), were
443 transferred into an S-block containing the recommended volumes of VXL, proteinase K and RNA carrier.
444 RT-qPCR (SARS-CoV-2 and MS2 viral genome detection) were performed with the Express one step RT-
445 qPCR Universal kit (ThermoFisher Scientific) using 3.5 μ l of RNA and 6.5 μ l of RT-qPCR mix that contains
446 250nmol of each primer and 75nmol of probe. Amplification was performed with the QuantStudio 12K
447 Flex Real-Time PCR System (ThermoFisher Scientific) using the following conditions: 50°C for 10min,
448 95°C for 20s, followed by 40 cycles of 95°C for 3s, 60°C for 30s. qPCR (γ -actine gene detection) was
449 performed under the same condition as RT-qPCR with the following modifications: we used the Express
450 one step qPCR Universal kit (ThermoFisher Scientific) and the 50°C step of the amplification cycle was
451 removed. Primers and probes sequences used to detect SARS-CoV-2, MS2 and γ -actine are described
452 in Table S9.

453 Tissue-culture infectious dose 50 (TCID₅₀) assay

454 To determine infectious titers, 96-well culture plates containing confluent VeroE6 cells were
455 inoculated with 150 μ l per well of serial dilutions of each sample (four-fold or ten-fold dilutions when
456 analyzing lung clarified homogenates or cell supernatant media respectively). Each dilution was
457 performed in sextuplicate. Plates were incubated for 4 days and then read for the absence or presence
458 of cytopathic effect in each well. Infectious titers were estimated using the method described by Reed
459 & Muench (REED and MUENCH, 1938).

460 Favipiravir pharmacokinetics

461 Animal handling, hamster infections and favipiravir administrations were performed as described
462 above. A piece of left lung was first washed in 10mL of sodium chloride 0.9% solution, blotted with
463 filter paper, weighed and then transferred to a 2mL tube containing 1mL of 0.9% sodium chloride
464 solution and 3mm glass beads. It was crushed using the Tissue Lyser machine (Retsch MM400) during
465 10min at 30 cycles/s and then centrifuged 5min à 1200g. Supernatant media were transferred to 2mL
466 tubes, centrifuged 10 min at 16,200g and stored at -80°C. One milliliter of blood was harvested in a
467 2mL tube containing 100 μ l of 0.5M EDTA (ThermoFischer Scientific). Blood was centrifuged for 10 min
468 at 16,200g and stored at -80°C.

469 Quantification of favipiravir in plasma and lung tissues was performed by a validated sensitive and
470 selective validated high-performance liquid chromatography coupled with tandem mass spectrometry
471 method (UPLC-TQD, Waters, USA) with a lower limit of quantification of 0.1 μ g/mL. Precision and
472 accuracy of the 3 quality control samples (QCs) were within 15% over the calibration range (0.5 μ g/mL
473 to 100 μ g/mL) (Bekegnran *et al.*, submitted). Favipiravir was extracted by a simple protein precipitation
474 method, using acetonitrile for plasma and ice-cold acetonitrile for clarified lung homogenates. Briefly,

475 50 μ L of samples matrix was added to 500 μ L of acetonitrile solution containing the internal standard
476 (favipiravir- $^{13}\text{C},^{15}\text{N}$, Alsachim), then vortexed for 2min followed by centrifugation for 10min at 4°C.
477 The supernatant medium was evaporated and the dry residues were then transferred to 96-well plates
478 and 50 μ L was injected. To assess the selectivity and specificity of the method and matrix effect, blank
479 plasma and tissues homogenates from 2 control animals (uninfected and untreated) were processed
480 at each run. Moreover, the same control samples spiked with favipiravir concentration equivalent to
481 the QCs (0.75, 50 and 80 $\mu\text{g}/\text{mL}$) were also processed and compared to the QCs samples.
482 Noncompartmental analysis conducted using software Pkanalix2019R2 (www.lixoft.com). Areas
483 under the plasma concentration time curve were computed using medians of favipiravir
484 concentrations at 0.5, 1, 5 and 8 hours, and extrapolated until T=12h. C_{trough} were extrapolated at
485 T=12h using lambda-z loglinear regression on the decreasing slope of concentrations.

486 Sequence analysis of the full-length genome

487 200 μ L of lung clarified homogenate or infectious cell supernatant (virus stock) was inactivated with an
488 equal volume of VXL lysis buffer (Qiagen) and viral RNA was extracted using an EZ1 Advanced XL robot
489 with the EZ1 mini virus 2.0 kit (both from Qiagen) and linear acrylamide (ThermoFisher Scientific) in
490 place of carrier RNA. cDNA was generated in a final volume of 40 μ L using 14 μ L of nucleic acid extract,
491 random hexamer and the Protoscript II First Strand cDNA Synthesis Kit (New England Biolabs). A
492 specific set of primers (Table S10) was used to generate thirteen amplicons covering the entire genome
493 with the Q5 High-Fidelity DNA polymerase (New England Biolabs). PCR mixes (final volume 25 μ L)
494 contained 2.5 μ L of cDNA, 2 μ L of each primer (10 μM) and 12.5 μ L of Q5 High-Fidelity 2X Master Mix.
495 Amplification was performed with the following conditions: 30 sec at 98°C, then 45 cycles of 15 sec at
496 98°C and 5 min à 65°C. Size of PCR products was verified by gel electrophoresis. For each sample, an
497 equimolar pool of all amplicons was prepared and purified using Monarch PCR & DNA Cleanup Kit (New
498 England Biolabs). After DNA quantification using Qubit dsDNA HS Assay Kit and Qubit 2.0 fluorometer
499 (ThermoFisher Scientific), amplicons were fragmented by sonication into fragments of around 200bp
500 long. Libraries were built by adding barcodes, for sample identification, and primers using AB Library
501 Builder System (ThermoFisher Scientific). To pool equimolarly the barcoded samples a quantification
502 step by real time PCR using Ion Library TaqMan Quantitation Kit (ThermoFisher Scientific) was
503 performed. Then, emulsion PCR from pools and loading on 530 chip was performed using the
504 automated Ion Chef instrument (ThermoFisher Scientific). Sequencing was performed using the S5 Ion
505 torrent technology v5.12 (ThermoFisher Scientific) following manufacturer's instructions. Consensus
506 sequence was obtained after trimming of reads (reads with quality score <0.99, and length <100pb
507 were removed and the 30 first and 30 last nucleotides were removed from the reads). Mapping of the
508 reads on a reference (determine following blast of De Novo contigs) was done using CLC genomics

509 workbench software v.20 (Qiagen). A *de novo* contig was also produced to ensure that the consensus
510 sequence was not affected by the reference sequence. Mutation frequency for each position was
511 calculated as the number of reads with a mutation compared to the reference divided by the total
512 number of reads at that site. Only substitutions with a frequency of at least 1% were taken into account
513 for the analysis (Table S6).

514 **ED₅₀, ED₉₀ and ED₉₉ determination**

515 We conducted a nonlinear regression of infectious viral load against dose, using an E_{max} model, giving

516
$$VL = VL_0 \times \left(1 - \left(\frac{D^\gamma}{D^\gamma + D_{50}^\gamma} \right) \right)$$
 with VL_0 being infectious viral load of untreated animals. We estimated

517 D_{50} the dose required to decrease viral load by 50%, using a coefficient γ to account for the high
518 sigmoidicity of the relation between dose and titers. γ coefficient was chosen as the one maximizing

519 likelihood of the model. We extrapolated the D_{90} and D_{99} using $D_{90} = \sqrt[\gamma]{9 \times D_{50}^\gamma}$ and $D_{99} =$

520 $\sqrt[\gamma]{99 \times D_{50}^\gamma}$, as well as their 95% confidence interval using the delta method.

521 **Statistical analysis**

522 Graphical representations and statistical analyses were performed with Graphpad Prism 7 (Graphpad
523 software) except linear/nonlinear regressions and their corresponding graphical representations that
524 were performed using R statistical software (<http://www.R-project.org>). Statistical details for each
525 experiments are described in the figure legends and in corresponding supplemental tables. P-values
526 lower than 0.05 were considered statistically significant.

527 Supplemental Data

528 Supplemental figure 1: In vitro efficacy of favipiravir

529 Supplemental figure 2: Dose-response curves

530 Supplemental figure 3: Evaluation of the toxicity for animals infected and treated with high doses of
531 favipiravir

532 Supplemental figure 4: Plasma concentrations of favipiravir after administration of a single dose of
533 favipiravir

534 Supplemental table 1: Implementation of hamster model

535 Supplemental table 2: Individual data from in vivo experiments

536 Supplemental table 3: Statistical analysis of in vivo experiments

537 Supplemental table 4: Statistical analysis of clinical monitoring

538 Supplemental table 5: Individual data of favipiravir pharmacokinetics

539 Supplemental table 6: Individual data for analysis of mutagenic effect of favipiravir

540 Supplemental table 7: Statistical analysis of mutagenic effect of favipiravir

541 Supplemental table 8: Shared mutations detected in lung clarified homogenates

542 Supplemental table 9: (RT)-qPCR systems

543 Supplemental table 10: Primer sequences used to produce overlapping amplicons for next generation
544 sequencing

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