

1           **ChAdOx1 nCoV-19 (AZD1222) protects against SARS-CoV-2 B.1.351 and B.1.1.7**

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18

19 **Abstract**

20 We investigated ChAdOx1 nCoV-19 (AZD1222) vaccine efficacy against SARS-CoV-2 variants  
21 of concern (VOCs) B.1.1.7 and B.1.351 in Syrian hamsters. We previously showed protection  
22 against SARS-CoV-2 disease and pneumonia in hamsters vaccinated with a single dose of  
23 ChAdOx1 nCoV-19. Here, we observed a 9.5-fold reduction of virus neutralizing antibody titer  
24 in vaccinated hamster sera against B.1.351 compared to B.1.1.7. Vaccinated hamsters challenged  
25 with B.1.1.7 or B.1.351 did not lose weight compared to control animals. In contrast to control  
26 animals, the lungs of vaccinated animals did not show any gross lesions. Minimal to no viral  
27 subgenomic RNA (sgRNA) and no infectious virus was detected in lungs of vaccinated animals.  
28 Histopathological evaluation showed extensive pulmonary pathology caused by B.1.1.7 or  
29 B.1.351 replication in the control animals, but none in the vaccinated animals. These data  
30 demonstrate the effectiveness of the ChAdOx1 nCoV-19 vaccine against clinical disease caused  
31 by B.1.1.7 or B.1.351 VOCs.

32

33 **Main**

34 The COVID-19 pandemic produced an unprecedented development of SARS-CoV-2 vaccines,  
35 and just over a year after the beginning of the outbreak a total of 12 vaccines have been  
36 authorized or approved globally. As the pandemic progressed, several VOCs have been detected.  
37 These include the B.1.1.7 and B.1.351 VOCs. The B.1.1.7 VOC was first detected in the United  
38 Kingdom and has seven amino acid (AA) substitutions and two deletions in the spike protein<sup>1,2</sup>  
39 compared to the original Wuhan isolate, Wuhan-Hu-1. The B.1.351 VOC was first detected in  
40 South Africa and has eight AA substitutions and one deletion in the spike protein<sup>3</sup> (Table 1). All  
41 currently licensed vaccines are based on the spike protein of Wuhan-Hu-1, thus, concerns have

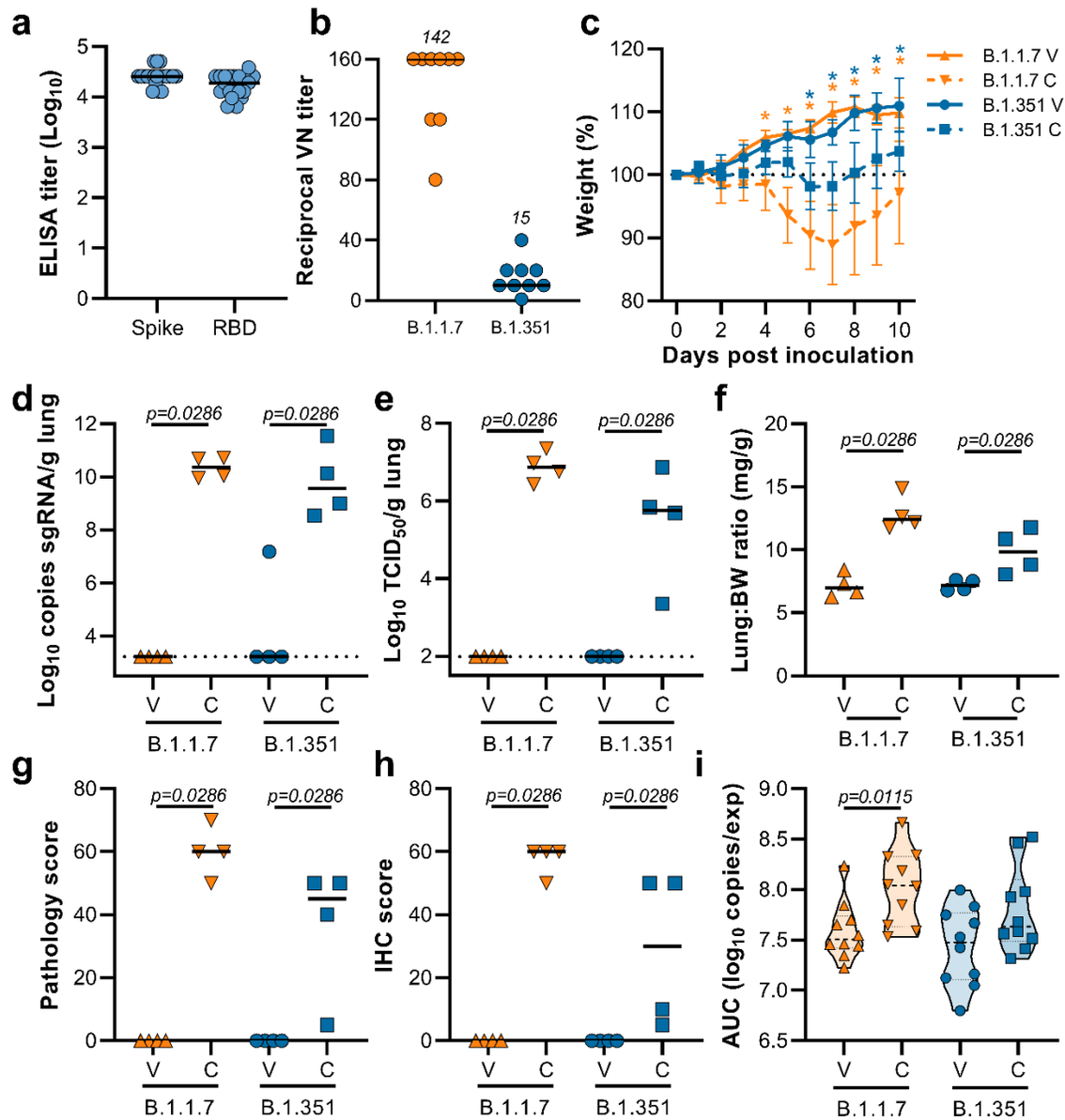
42 been raised that the presence of these changes may affect vaccine efficacy. The goal of this study  
43 was to evaluate ChAdOx1 nCoV-19 (AZD1222) vaccine efficacy in Syrian hamsters, when  
44 challenged using naturally occurring isolates of the VOCs B.1.1.7 and B.1.351.  
45

| <b>Substitution (Wuhan AA numbering)</b> | <b>VOC B.1.1.7<sup>1</sup></b> | <b>VOC B.1.351<sup>3</sup></b> |
|--|--------------------------------|--------------------------------|
| <b>L18F</b>                              | -                              | +                              |
| <b>HV69-70del</b>                        | +                              | -                              |
| <b>D80A</b>                              | -                              | +                              |
| <b>Y144del</b>                           | +                              | -                              |
| <b>D215G</b>                             | -                              | +                              |
| <b>LAL242-244del</b>                     | -                              | +                              |
| <b>K417N</b>                             | -                              | +                              |
| <b>E484K</b>                             | -                              | +                              |
| <b>N501Y</b>                             | +                              | +                              |
| <b>A570D</b>                             | +                              | -                              |
| <b>D614G</b>                             | +                              | +                              |
| <b>P681D</b>                             | +                              | -                              |
| <b>A701V</b>                             | -                              | +                              |
| <b>T716I</b>                             | +                              | -                              |
| <b>S982A</b>                             | +                              | -                              |
| <b>D1118H</b>                            | +                              | -                              |

46 Table 1. AA substitutions detected in the spike protein of VOCs B.1.1.7 (EPI\_ISL\_601443) and  
47 B.1.351 (EPI\_ISL\_678615) compared to Wuhan-Hu-1 (NC\_045512).  
48

49 Syrian hamsters (N=10 per group) were vaccinated intramuscularly with either ChAdOx1 nCoV-  
50 19 or ChAdOx1 green fluorescent protein (GFP,  $2.5 \times 10^8$  IU/hamster) 30 days prior to intranasal  
51 challenge with SARS-CoV-2 VOCs B.1.1.7 or B.1.351. Vaccination with ChAdOx1 nCoV19  
52 resulted in high titers of binding antibodies against the SARS-CoV-2 full-length spike protein  
53 and receptor binding domain (Figure 1a) at 25 days post vaccination. We then investigated  
54 neutralizing antibody titers in serum against infectious virus. Neutralization of B.1.351 was  
55 significantly reduced compared to neutralization of B.1.1.7 (Figure 1b, mean titer of 15 vs 142, p  
56 < 0.0001, Mann-Whitney test).

57 Virus stocks were deep sequenced before inoculation. No mutations were found in B.1.1.7  
58 compared to the isolate's sequence, but two AA substitutions were found in the spike protein of  
59 B.1.351; Q677H (present at 88%) and R682W (present at 89%).  
60 Upon inoculation with virus, weight loss was observed in control hamsters challenged with  
61 B.1.1.7, whereas less pronounced weight loss was observed in control hamsters challenged with  
62 B.1.351 (Figure 1c). In contrast, vaccinated hamsters continued to gain weight throughout the  
63 experiment. A significant difference in weight between vaccinated and control hamsters was  
64 observed starting at 4 days post infection (DPI) for B.1.1.7 and at 6 DPI for B.1.351 (Figure 1c,  
65 Student's t-test corrected for multiple comparisons using the Holm-Šidák method) and continued  
66 throughout the remainder of the experiment. Four out of ten hamsters per group were euthanized  
67 at 5 DPI and lung tissue was harvested. Lung tissue of all control animals contained high levels  
68 of sgRNA (Figure 1d,  $10^8$ - $10^{10}$  copies/gram tissue), and was comparable to sgRNA levels  
69 previously detected in lung tissue of control animals challenged with SARS-CoV-2 D614G  
70 (hCoV-19/USA/MT-RML-7/2020)<sup>4</sup>. Conversely, no sgRNA was detected in lung tissue obtained  
71 from vaccinated hamsters challenged with B.1.1.7, and only one out of four vaccinated hamsters  
72 challenged with B.1.351 had detectable sgRNA (Figure 1d,  $p=0.0286$ , Mann-Whitney test). High  
73 levels of infectious virus were detected in lung tissue of all eight control animals, whereas no  
74 vaccinated animals had detectable infectious virus in lung tissue (Figure 1e,  $p=0.0286$ , Mann-  
75 Whitney test). Lung:body weight ratios on 5 DPI were significantly lower in vaccinated animals  
76 compared to control animals for both VOCs (Figure 1f,  $p=0.0286$ , Mann-Whitney test),  
77 indicating no or reduced pulmonary edema in ChAdOx1 nCoV19-vaccinated animals.  
78



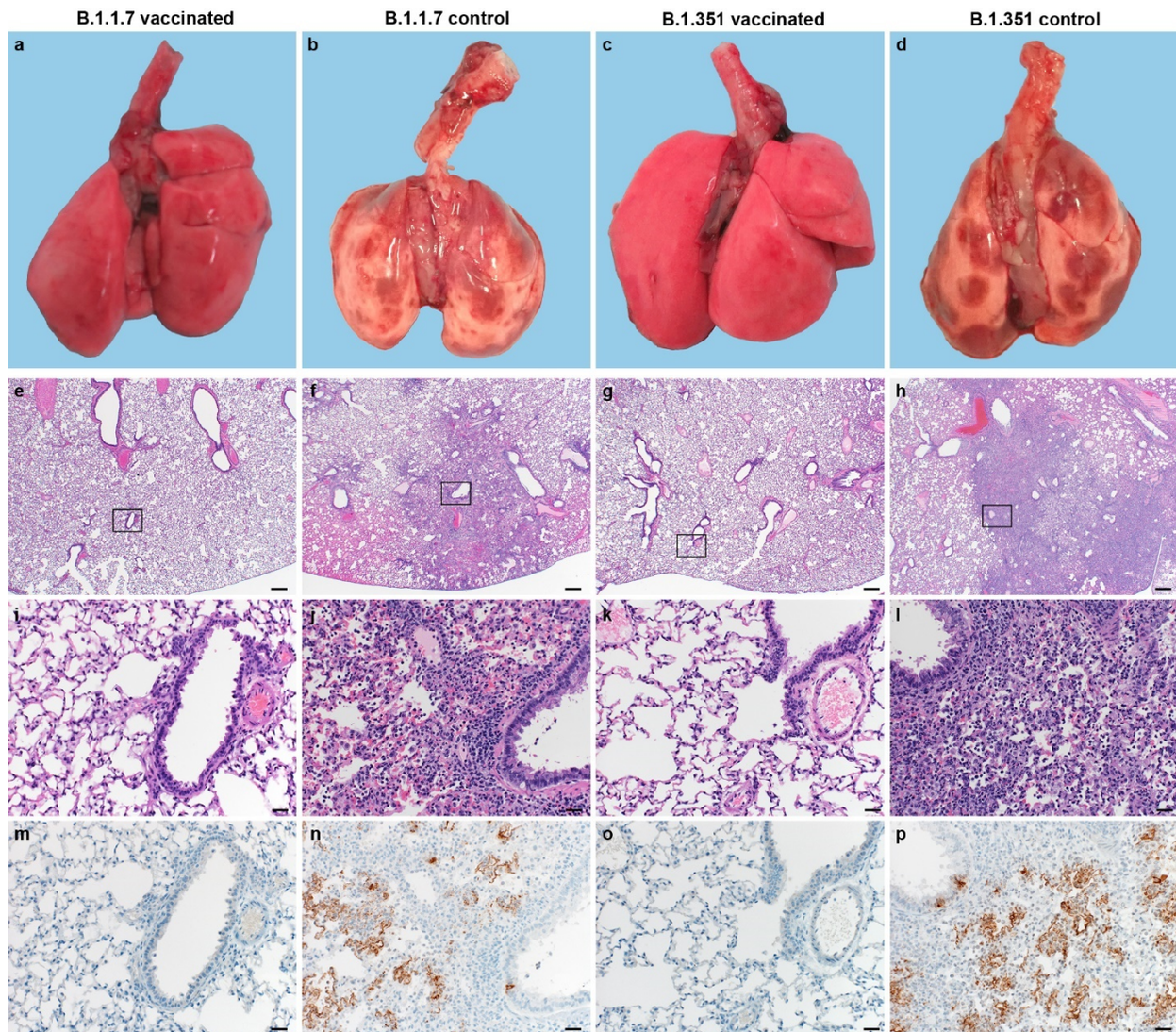
79

80 **Figure 1. Vaccination of Syrian hamsters with ChAdOx1 nCoV-19 prevents lower**  
 81 **respiratory tract infection with SARS-CoV-2 VOCs B.1.1.7 and B.1.351.** a. Binding  
 82 antibodies against spike protein or RBD of SARS-CoV-2 (clade A) in serum obtained 25 days  
 83 post vaccination with ChAdOx1 nCoV-19. Line = median; numbers = mean. b. Virus  
 84 neutralizing antibody titers against B.1.1.7 or B.1.351 VOCs in serum obtained 25 days post  
 85 vaccination with ChAdOx1 nCoV-19. Line = median. c. Relative weight upon intranasal  
 86 challenge with  $10^4$  TCID<sub>50</sub> of B.1.1.7 or B.1.351. Shown is geometric mean with 95%  
 87 confidence interval (CI). \* = p-value < 0.005, corrected for multiple comparisons using the Holm-  
 88 Šidák correction. d. sgRNA viral load in lung tissue obtained at 5 DPI. Line = median. Dotted

89 line = limit of detection. e. Infectious SARS-CoV-2 titer in lung tissue obtained at 5 DPI. Line =  
90 median. Dotted line = limit of detection. f. Lung:body weight (BW) ratio (mg:g) of hamsters  
91 euthanized at 5 DPI. Line = median. g. Percentage affected lung tissue per animal as determined  
92 via histology. Line = median. h. Percentage of lung tissue positive for SARS-CoV-2 antigen per  
93 animal. Line = median. i. Truncated violin-plot of area under the curve (AUC) analysis of  
94 shedding as measured by sgRNA analysis in swabs. Dashed line = median. Dotted line =  
95 quartiles. Statistical significance determined via Kruskal-Wallis test (b), mixed-effect analyses  
96 (c), or Mann-Whitney test (d-i). V = ChAdOx1 nCoV-19 vaccinated; C = ChAdOx1 GFP  
97 vaccinated; Orange upward triangle = Hamsters vaccinated with ChAdOx1 nCoV-19, challenged  
98 with B.1.1.7; Orange downward triangle = Hamsters vaccinated with ChAdOx1 GFP, challenged  
99 with B.1.1.7; Blue circle = Hamsters vaccinated with ChAdOx1 nCoV-19, challenged with  
100 B.1.351; Blue square = Hamsters vaccinated with ChAdOx1 GFP, challenged with B.1.351.  
101

102 Upon necropsy, lungs from control animals showed gross lesions previously observed in  
103 hamsters inoculated with SARS-CoV-2 WA1 or a D614G isolate, with focal areas of hilar  
104 consolidation and hyperemia<sup>4,5</sup>. No gross lesions were observed in lung tissue obtained from any  
105 of the vaccinated animals (Figure 2a-d). Microscopically, pulmonary lesions of control animals  
106 consisted of a moderate to marked broncho-interstitial pneumonia extending into the adjacent  
107 alveoli. Bronchi and bronchioles had multifocal necrotic epithelial cells and moderate numbers  
108 of infiltrating neutrophils and macrophages. Alveolar septa were expanded by edema fluid and  
109 leucocytes. In contrast, vaccinated animals did not show any evidence of SARS-CoV-2  
110 pathology (Figure 2e-l). Immunohistochemistry using a monoclonal antibody against SARS-  
111 CoV-2 demonstrated viral antigen in bronchial and bronchiolar epithelium, type I and II  
112 pneumocytes as well as pulmonary macrophages within the control animals, but not in  
113 vaccinated animals (Figure 2m-p). For the sections of lung evaluated, the percentage of lung  
114 tissue showing pathology and the percentage of lung tissue positive for SARS-CoV-2 antigen  
115 was determined by a veterinary pathologist blinded to the study group allocations. No pathology  
116 nor SARS-CoV-2 antigen was found in lungs of vaccinated animals, but both were abundantly  
117 present in lungs of control animals (Figure 1g,h,  $p=0.0286$ , Mann-Whitney test). Finally,

118 oropharyngeal swabs were collected on 1 to 5 DPI, evaluated for sgRNA, and an area under the  
119 curve was calculated per animal to determine the cumulative amount of virus shed. We observed  
120 a decrease in the total amount of respiratory shedding for both vaccinated groups compared to  
121 control animals, although this was statistically significant for B.1.1.7 only (Figure 1i,  $p=0.0115$ ,  
122 Mann-Whitney test).  
123



124

125 **Figure 2. Pulmonary effects of direct intranasal challenge with SARS-CoV-2 variants**  
126 **B.1.1.7 and B.1.351 in Syrian hamsters at 5 DPI.** a-d. Gross pathology of hamster lungs; a/c.  
127 Normal lungs. b/d. Multifocal and focally extensive areas of consolidation. e-h. H&E staining,

128 20x; e/g. No pathology. f/h. Focally extensive areas of bronchointerstitial pneumonia. i-l. H&E  
129 staining, 200x; i/k. No pathology. j/l. Bronchointerstitial pneumonia with alveolar histiocytosis,  
130 fibrin and edema. m-p. IHC staining against N protein SARS-CoV-2 (brown). m/o. No staining.  
131 n/p. Staining of bronchiolar epithelial cells, type I&II pneumocytes and rare macrophages.  
132

133 We investigated the presence of the AA substitutions Q677H and R682W observed in 88-89% of  
134 the spike protein of our B.1.351 stock in swabs and lung tissue obtained from control animals  
135 challenged with B.1.351. Whereas we did find these substitutions in swabs obtained 1 DPI, they  
136 were not present in swabs obtained at 5 DPI. Likewise, the substitutions were only found in lung  
137 tissue of one out of four control hamsters (Table 2).

138

| AA substitutions | Presence in swabs<br>(1 DPI, N=5) | Presence in swabs<br>(5 DPI, N=3) | Presence in lungs<br>(N=4) |
|------------------|-----------------------------------|-----------------------------------|----------------------------|
| Q677H            | 44.1-65.7%                        | 0%                                | 0 (N=3), 81% (N=1)         |
| R682W            | 44.9-65.8%                        | 0%                                | 0 (N=3), 81% (N=1)         |

139 Table 2. Presence of substitutions Q677H and R682W in swabs and lung tissue of hamsters  
140 directly inoculated with B.1.351.

141

142 This study demonstrates efficacy of the ChAdOx1 nCoV-19 vaccine against circulating variants  
143 of concern in the SARS-CoV-2 Syrian hamster model. The Syrian hamster SARS-CoV-2  
144 infection model is characterized by natural susceptibility to SARS-CoV-2 and development of a  
145 robust upper and lower respiratory tract infection<sup>6</sup>. The hamster model has been successfully  
146 used for the preclinical development of several vaccines including the Ad26 and mRNA-1273  
147 vaccines by Janssen<sup>7</sup> and Moderna<sup>8</sup>, respectively. Several groups have reported the effect of  
148 spike protein substitutions observed in B.1.1.7 and B.1.351 VOCs on the virus neutralizing  
149 capacity of serum obtained from vaccinated or convalescent individuals. In general, these studies  
150 conclude that the substitutions found in the B.1.1.7 spike protein have limited to no effect on  
151 virus neutralization titres<sup>9-15</sup>. Data from a UK phase III trial taken from a time when B.1.1.7

152 predominated, showed minimal impact on ChAdOx1 nCoV-19 vaccine efficacy<sup>15</sup>. Likewise, in  
153 an observational study of vaccine effectiveness in adults aged over 70 years in the UK, a single  
154 dose of either ChAdOx1 nCoV-19 or the Pfizer/BioNTech vaccine BNT162b2 reduced  
155 hospitalization in elderly adults with co-morbidities by 80%<sup>16</sup>. In contrast, the substitutions  
156 found in the B.1.351 spike protein (Table 1) result in a significant reduction of virus neutralizing  
157 capacity with pseudotype or infectious virus neutralization assays<sup>9-15,17,18</sup>. In a phase II study of  
158 ChAdOx1 nCoV-19 in South Africa in 2000 adults with a median age of 31 years, vaccine  
159 efficacy against mild to moderate disease was reduced when the virus recovered after infection  
160 was B.1.351 (19 cases in the vaccinated group and 20 in the placebo group)<sup>19</sup>. Vaccine efficacy  
161 against severe disease could not be determined as no severe cases occurred in this young cohort.  
162 The South African arm of the ENSEMBLE study which tested vaccine efficacy after a single  
163 dose of Janssen's COVID-19 vaccine candidate enrolled 6,576 participants in South Africa, out  
164 of a total of 43,783 in multiple countries, with 34% of participants across the study aged over 60  
165 years. Vaccine efficacy against moderate to severe disease was 64% (CI 41.2%, 78.7%) in South  
166 Africa compared to 72% (CI 58.2%, 81.7%) in the USA at 28 days post vaccination<sup>20</sup>. Efficacy  
167 of the vaccine against severe to critical disease was 81.7% in South Africa, which was similar to  
168 the reported 85.9% and 87.6% in the USA and Brazil, respectively<sup>20</sup>. Vaccine efficacy against  
169 mild disease was not reported. These clinical trial results are consistent with the findings of the  
170 preclinical study reported here; ChAdOx1 nCov-19 may be less effective at reducing upper  
171 respiratory tract infection caused by B.1.351 than by B.1.1.7, consistent with reduced efficacy  
172 against mild disease. However, complete protection against lower respiratory tract disease was  
173 observed in this challenge study, consistent with protection against severe disease. Based on our

174 data, we hypothesize that the currently available vaccines will likely still protect against severe  
175 disease and hospitalization caused by VOC B.1.351.

176 Limited data on the immunological determinants of protection are available, however recent data  
177 from rhesus macaques indicate that relatively low neutralizing antibody titers are sufficient for  
178 protection against SARS-CoV-2, and that cellular immune responses may contribute to  
179 protection if antibody responses are suboptimal<sup>21</sup>. Induction of binding and neutralizing  
180 antibodies as well as SARS-CoV-2 cellular spike protein-specific T cell responses after  
181 vaccination have been reported<sup>22,23</sup> and most SARS-CoV-2 specific T cell epitopes in both  
182 convalescent and vaccinated individuals are not affected by the AA substitutions found in the  
183 spike protein of the B.1.1.7 and B.1.351 variants<sup>24</sup>. Protection against severe COVID-19 disease  
184 might be mediated by T cells and would therefore not be different between the current variants.  
185 However, as T cell-mediated protection can only act after the initial infection has occurred, mild,  
186 polymerase chain reaction (PCR)-positive disease may still occur in the upper respiratory tract.

187 It should be noted that the B.1.351 virus stock used to challenge hamsters contained two  
188 additional non-fixed AA substitutions; Q677H and R682W at 88% and 89%, respectively.

189 Interestingly, the relative presence of these two AA substitutions was markedly reduced at 1 DPI  
190 and absent on 5 DPI. In addition, they were only present in lung tissue of one control hamster at  
191 5 DPI, suggesting that they are rapidly selected against in the SARS-CoV-2 hamster model over  
192 the course of infection. Nonetheless, since the substitutions thought to be important in immune  
193 evasion, such as E484K<sup>18</sup>, are still present in the virus stock, efficient replication and lung  
194 pathology was observed in infected hamsters, we do not believe the presence of these additional  
195 AA substitutions present as a quasispecies affect the data interpretation.

196 Based on the current studies and healthcare priorities in real-world settings, we believe it is  
197 essential to focus on prevention of moderate to severe disease requiring hospitalization. We show  
198 that ChAdOx1 nCoV-19 vaccination resulted in complete protection against disease in hamsters.  
199 As implied by the data presented by Janssen<sup>20</sup>, viral vectored vaccines may provide substantial  
200 protection against lower respiratory tract infection caused by the B.1.351 variant and subsequent  
201 hospitalization and death. With the ongoing evolution of SARS-CoV-2, the readily available and  
202 cost-effective hamster model allows rapid evaluation of the protective efficacy of novel VOCs.  
203 In addition, it will allow rapid preclinical benchmarking of existing vaccines against preclinical  
204 vaccines with updated antigen designs.

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## 206 **References**

- 207 1. Chand, M. *et al.* Investigation of novel SARS-COV-2 variant Variant of Concern 202012/01.
- 208 2. Davies, N. G. *et al.* Estimated transmissibility and impact of SARS-CoV-2 lineage B.1.1.7 in  
209 England. *Science* eabg3055 (2021) doi:10.1126/science.abg3055.
- 210 3. Tegally, H. *et al.* *Emergence and rapid spread of a new severe acute respiratory syndrome-*  
211 *related coronavirus 2 (SARS-CoV-2) lineage with multiple spike mutations in South Africa.*  
212 <http://medrxiv.org/lookup/doi/10.1101/2020.12.21.20248640> (2020)  
213 doi:10.1101/2020.12.21.20248640.
- 214 4. van Doremalen, N. *et al.* *Intranasal ChAdOx1 nCoV-19/AZD1222 vaccination reduces*  
215 *shedding of SARS-CoV-2 D614G in rhesus macaques.*  
216 <http://biorxiv.org/lookup/doi/10.1101/2021.01.09.426058> (2021)  
217 doi:10.1101/2021.01.09.426058.

- 218 5. Port, J. R. *et al.* SARS-CoV-2 disease severity and transmission efficiency is increased for  
219 airborne but not fomite exposure in Syrian hamsters.  
220 <http://biorxiv.org/lookup/doi/10.1101/2020.12.28.424565> (2020)  
221 doi:10.1101/2020.12.28.424565.
- 222 6. Muñoz-Fontela, C. *et al.* Animal models for COVID-19. *Nature* **586**, 509–515 (2020).
- 223 7. Tostanoski, L. H. *et al.* Ad26 vaccine protects against SARS-CoV-2 severe clinical disease  
224 in hamsters. *Nat Med* **26**, 1694–1700 (2020).
- 225 8. Meyer, M. *et al.* mRNA-1273 efficacy in a severe COVID-19 model: attenuated activation of  
226 pulmonary immune cells after challenge.  
227 <http://biorxiv.org/lookup/doi/10.1101/2021.01.25.428136> (2021)  
228 doi:10.1101/2021.01.25.428136.
- 229 9. Liu, Y. *et al.* Neutralizing Activity of BNT162b2-Elicited Serum — Preliminary Report. *N*  
230 *Engl J Med* NEJMc2102017 (2021) doi:10.1056/NEJMc2102017.
- 231 10. Zhou, D. *et al.* Evidence of escape of SARS-CoV-2 variant B.1.351 from natural and vaccine  
232 induced sera. *Cell* S0092867421002269 (2021) doi:10.1016/j.cell.2021.02.037.
- 233 11. Planas, D. *et al.* Sensitivity of infectious SARS-CoV-2 B.1.1.7 and B.1.351 variants to  
234 neutralizing antibodies. <http://biorxiv.org/lookup/doi/10.1101/2021.02.12.430472> (2021)  
235 doi:10.1101/2021.02.12.430472.
- 236 12. Wang, P. *et al.* Antibody Resistance of SARS-CoV-2 Variants B.1.351 and B.1.1.7.  
237 <http://biorxiv.org/lookup/doi/10.1101/2021.01.25.428137> (2021)  
238 doi:10.1101/2021.01.25.428137.

- 239 13. Wu, K. *et al.* *mRNA-1273 vaccine induces neutralizing antibodies against spike mutants*  
240 *from global SARS-CoV-2 variants*. <http://biorxiv.org/lookup/doi/10.1101/2021.01.25.427948>  
241 (2021) doi:10.1101/2021.01.25.427948.
- 242 14. Xie, X. *et al.* Neutralization of SARS-CoV-2 spike 69/70 deletion, E484K and N501Y  
243 variants by BNT162b2 vaccine-elicited sera. *Nat Med* (2021) doi:10.1038/s41591-021-  
244 01270-4.
- 245 15. Emary, K. R. W. *et al.* Efficacy of ChAdOx1 nCoV-19/AZD1222 Vaccine Against SARS-  
246 CoV-2 VOC (B.1.1.7). *SSRN Journal* (2021) doi:10.2139/ssrn.3779160.
- 247 16. Hyams, C. *et al.* Assessing the Effectiveness of BNT162b2 and ChAdOx1nCoV-19 COVID-  
248 19 Vaccination in Prevention of Hospitalisations in Elderly and Frail Adults: A Single  
249 Centre Test Negative Case-Control Study. *Lancet*.
- 250 17. Cele, S. *et al.* *Escape of SARS-CoV-2 501Y.V2 from neutralization by convalescent plasma*.  
251 <http://medrxiv.org/lookup/doi/10.1101/2021.01.26.21250224> (2021)  
252 doi:10.1101/2021.01.26.21250224.
- 253 18. Greaney, A. J. *et al.* *Comprehensive mapping of mutations to the SARS-CoV-2 receptor-*  
254 *binding domain that affect recognition by polyclonal human serum antibodies*.  
255 <http://biorxiv.org/lookup/doi/10.1101/2020.12.31.425021> (2021)  
256 doi:10.1101/2020.12.31.425021.
- 257 19. Madhi, S. A. *et al.* *Safety and efficacy of the ChAdOx1 nCoV-19 (AZD1222) Covid-19*  
258 *vaccine against the B.1.351 variant in South Africa*.  
259 <http://medrxiv.org/lookup/doi/10.1101/2021.02.10.21251247> (2021)  
260 doi:10.1101/2021.02.10.21251247.

- 261 20. Janssen. Emergency Use Authorization (EUA) for an Unapproved Product Review  
262 Memorandum.
- 263 21. McMahan, K. *et al.* Correlates of protection against SARS-CoV-2 in rhesus macaques.  
264 *Nature* (2020) doi:10.1038/s41586-020-03041-6.
- 265 22. the Oxford COVID Vaccine Trial Group *et al.* T cell and antibody responses induced by a  
266 single dose of ChAdOx1 nCoV-19 (AZD1222) vaccine in a phase 1/2 clinical trial. *Nat Med*  
267 **27**, 270–278 (2021).
- 268 23. Sahin, U. *et al.* COVID-19 vaccine BNT162b1 elicits human antibody and TH1 T cell  
269 responses. *Nature* **586**, 594–599 (2020).
- 270 24. Tarke, A. *et al.* Negligible impact of SARS-CoV-2 variants on CD4<sup>+</sup> and CD8<sup>+</sup> T cell  
271 reactivity in COVID-19 exposed donors and vaccinees.  
272 <http://biorxiv.org/lookup/doi/10.1101/2021.02.27.433180> (2021)  
273 doi:10.1101/2021.02.27.433180.

274

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290 N.v.D., D.R.A., C.K.Y, J.R.P., M.G.H., J.E.S., S.L.A., C.M., G.S, and V.J.M. analyzed results,  
291 R.J.F., N.v.D and D.R.A. wrote the manuscript, all co-authors reviewed the manuscript.;

292 **Competing interests:** S.C.G. is a board member of Vaccitech and named as an inventor on a  
293 patent covering the use of ChAdOx1-vector-based vaccines and a patent application covering a  
294 SARS-CoV-2 (nCoV-19) vaccine (UK patent application no. 2003670.3). T.L. is named as an  
295 inventor on a patent application covering a SARS-CoV-2 (nCoV-19) vaccine (UK patent  
296 application no. 2003670.3). The University of Oxford and Vaccitech, having joint rights in the  
297 vaccine, entered into a partnership with AstraZeneca in April 2020 for further development,  
298 large-scale manufacture and global supply of the vaccine. Equitable access to the vaccine is a  
299 key component of the partnership. Neither Oxford University nor Vaccitech will receive any  
300 royalties during the pandemic period or from any sales of the vaccine in developing countries.  
301 All other authors declare no competing interests.

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304

305 **Materials and Methods**

306 *Ethics Statement*

307 All animal experiments were conducted in an AAALAC International-accredited facility and  
308 were approved by the Rocky Mountain Laboratories Institutional Care and Use Committee  
309 following the guidelines put forth in the Guide for the Care and Use of Laboratory Animals 8<sup>th</sup>  
310 edition, the Animal Welfare Act, United States Department of Agriculture and the United States  
311 Public Health Service Policy on the Humane Care and Use of Laboratory Animals.  
312 The Institutional Biosafety Committee (IBC) approved work with infectious SARS-CoV-2 virus  
313 strains under BSL3 conditions. Virus inactivation of all samples was performed according to  
314 IBC-approved standard operating procedures for the removal of specimens from high  
315 containment areas.

316 *Cells and virus*

317 SARS-CoV-2 variant B.1.351 (hCoV-19/South African/KRISP-K005325/2020,  
318 EPI\_ISL\_678615) was obtained from Dr. Tulio de Oliveira and Dr. Alex Sigal at the Nelson R  
319 Mandela School of Medicine, UKZN. SARS-CoV-2 variant B.1.1.7 (hCoV-  
320 19/England/204820464/2020, EPI\_ISL\_683466) was obtained from Public Health England via  
321 BEI. Virus propagation was performed in VeroE6 cells in DMEM supplemented with 2% fetal  
322 bovine serum, 1 mM L-glutamine, 50 U/ml penicillin and 50 µg/ml streptomycin (DMEM2).  
323 VeroE6 cells were maintained in DMEM supplemented with 10% fetal bovine serum, 1 mM L-  
324 glutamine, 50 U/ml penicillin and 50 µg/ml streptomycin. Mycoplasma testing is performed at  
325 regular intervals and no mycoplasma was detected.

326 *Animal Experiments*

327 ChAdOx1 nCoV-19 was formulated as previously described<sup>25</sup>. Four groups of 10, 4-6-week-old  
328 female Syrian hamsters (Envigo Indianapolis, IN) were vaccinated with  $2.5 \times 10^8$  infectious units

329 of ChAdOx1 nCoV-19 vaccine or ChAdOx1-GFP delivered intramuscularly in two 50  $\mu$ L doses  
330 into the posterior thighs 30 days prior to challenge. Five days prior to challenge a blood sample  
331 was collected via the retro-orbital plexus under isoflurane anesthesia and spun at 2000 g for 10  
332 min to obtain serum. Two groups (10 ChAdOx1 nCoV-19 vaccinated and 10 ChAdOx1 GFP  
333 vaccinated hamsters) were challenged with  $10^4$  TCID<sub>50</sub>/mL B.1.1.7 diluted in sterile Dulbecco's  
334 Modified Eagle's media (DMEM), in a 40  $\mu$ L bolus delivered intranasally, one-half into each  
335 nostril. Two other groups (10 ChAdOx1 nCoV-19 vaccinated and 10 ChAdOx1 GFP vaccinated  
336 hamsters) were similarly challenged with B.1.351 also diluted in sterile DMEM. Weights were  
337 recorded daily until 14 DPI. Oropharyngeal swabs were collected daily in 1 mL of DMEM2 up  
338 until 5 DPI. On 5 DPI 4 animals from each group were euthanized. The lungs were excised,  
339 weighed, and photographed, and samples taken for qRT-PCR analysis, virus titrations and  
340 histopathology. The remaining six animals in each group were monitored daily for signs of  
341 disease and weighed until 14 DPI.

#### 342 *Virus titration*

343 Lung sections were weighed and homogenized in 1 mL of DMEM. Virus titrations were  
344 performed by end-point titration of 10-fold dilutions of virus swab media or tissue homogenates  
345 on VeroE6 cells in 96-well plates. When titrating tissue homogenate, the top 2 rows of cells were  
346 washed 2 times with PBS prior to the addition of a final 100  $\mu$ l of DMEM2. Cells were incubated  
347 at 37°C and 5% CO<sub>2</sub>. Cytopathic effect was read 6 days later.

#### 348 *Virus neutralization*

349 Sera were heat-inactivated (30 min, 56 °C). After an initial 1:10 dilution of the sera, two-fold  
350 serial dilutions were prepared in DMEM2. 100 TCID<sub>50</sub> of SARS-CoV-2 variant B.1.1.7 or  
351 B.1.351 was added to the diluted sera. After a 1hr incubation at 37°C and 5% CO<sub>2</sub>, the virus-

352 serum mixture was added to VeroE6 cells. The cells were incubated for 6 days at 37°C and 5%  
353 CO<sub>2</sub> at which time they were evaluated for CPE. The virus neutralization titer was expressed as  
354 the reciprocal value of the highest dilution of the serum that still inhibited virus replication.

#### 355 *RNA extraction and quantitative reverse-transcription polymerase chain reaction*

356 RNA was extracted from oropharyngeal swabs using the QiaAmp Viral RNA kit (Qiagen)  
357 according to the manufacturer's instructions and following high containment laboratory  
358 protocols. Lung samples were homogenized and extracted using the RNeasy kit (Qiagen)  
359 according to the manufacturer's instructions and following high containment laboratory  
360 protocols. A viral sgRNA<sup>26</sup> specific assay was used for the detection of viral RNA. Five µL of  
361 extracted RNA was tested with the Quantstudio 3 system (Thermofisher) according to  
362 instructions from the manufacturer. A standard curve was generated during each run using  
363 SARS-CoV-2 standards containing a known number of genome copies.

#### 364 *Viral RNA sequencing*

365 For sequencing from viral stocks, sequencing libraries were prepared using Stranded Total RNA  
366 Prep Ligation with Ribo-Zero Plus kit per manufacturer's protocol (Illumina) and sequenced on  
367 an Illumina MiSeq at 2 x 150 base pair reads. For sequencing from swab and lung tissue, total  
368 RNA was depleted of ribosomal RNA using the Ribo-Zero Gold rRNA Removal kit (Illumina).  
369 Sequencing libraries were constructed using the KAPA RNA HyperPrep kit following  
370 manufacturer's protocol (Roche Sequencing Solutions). To enrich for SARS-CoV-2 sequence,  
371 libraries were hybridized to myBaits Expert Virus biotinylated oligonucleotide baits following  
372 the manufacturer's manual, version 4.01 (Arbor Biosciences, Ann Arbor, MI). Enriched libraries  
373 were sequenced on the Illumina MiSeq instrument as paired-end 2 X 151 base pair reads. Raw  
374 fastq reads were trimmed of Illumina adapter sequences using cutadapt version 1.12<sup>27</sup> and then

375 trimmed and filtered for quality using the FASTX-Toolkit (Hannon Lab, CSHL). Remaining  
376 reads were mapped to the SARS-CoV-2 2019-nCoV/USA-WA1/2020 genome (MN985325.1) or  
377 hCoV-19/England/204820464/2020 (EPI\_ISL\_683466) or hCoV-19/SouthAfrica/KRISP-  
378 K005325/2020 (EPI\_ISL\_678615) using Bowtie2 version 2.2.9<sup>28</sup> with parameters --local --no-  
379 mixed -X 1500. PCR duplicates were removed using picard MarkDuplicates (Broad Institute)  
380 and variants were called using GATK HaplotypeCaller version 4.1.2.0<sup>29</sup> with parameter -ploidy  
381 2. Variants were filtered for QUAL > 500 and DP > 20 using bcftools.

### 382 *Expression and purification of SARS-CoV-2 S and receptor binding domain*

383 Protein production was performed as described previously<sup>30,31</sup>. Expression plasmids encoding the  
384 codon optimized SARS-CoV-2 full length S and RBD were obtained from Kizzmekia Corbett  
385 and Barney Graham (Vaccine Research Center, Bethesda, USA)<sup>32</sup> and Florian Krammer (Icahn  
386 School of Medicine at Mt. Sinai, New York, USA)<sup>33</sup>. Expression was performed in Freestyle  
387 293-F cells (Thermofisher), maintained in Freestyle 293 Expression Medium (Gibco) at 37°C  
388 and 8% CO<sub>2</sub> shaking at 130 rpm. Cultures totaling 500 mL were transfected with PEI at a  
389 density of one million cells per mL. Supernatant was harvested 7 days post transfection, clarified  
390 by centrifugation and filtered through a 0.22 µm membrane. The protein was purified using Ni-  
391 NTA immobilized metal-affinity chromatography (IMAC) using Ni Sepharose 6 Fast Flow Resin  
392 (GE Lifesciences) or NiNTA Agarose (QIAGEN) and gravity flow. After elution the protein was  
393 buffer exchanged into 10 mM Tris pH8, 150 mM NaCl buffer (S) or PBS (RBD) and stored at –  
394 80°C.

### 395 *ELISA*

396 ELISA was performed as described previously<sup>25</sup>. Briefly, maxisorp plates (Nunc) were coated  
397 overnight at 4°C with 50 ng/well S or RBD protein in PBS. Plates were blocked with 100 µl of

398 casein in PBS (Thermo Fisher) for 1hr at RT. Serum diluted 1:6,400 was further 2-fold serially  
399 diluted in casein in PBS was incubated at RT for 1hr. Antibodies were detected using affinity-  
400 purified polyclonal antibody peroxidase-labeled goat-anti-monkey IgG (Seracare, 074-11-021) in  
401 casein followed by TMB 2-component peroxidase substrate (Seracare, 5120-0047). The reaction  
402 was stopped using stop solution (Seracare, 5150-0021) and read at 450 nm. All wells were  
403 washed 4x with PBST 0.1% tween in between steps. Threshold for positivity was set at 2x OD  
404 value of negative control (serum obtained from unvaccinated hamsters prior to start of the  
405 experiment).

#### 406 *Data availability statement*

407 Data have been deposited in Figshare

408

#### 409 **References Materials and Methods**

410 25. van Doremalen, N. *et al.* ChAdOx1 nCoV-19 vaccine prevents SARS-CoV-2 pneumonia in  
411 rhesus macaques. *Nature* **586**, 578–582 (2020).

412 26. Corman, V. M. *et al.* Detection of 2019 novel coronavirus (2019-nCoV) by real-time RT-  
413 PCR. *Eurosurveillance* **25**, (2020).

414 27. Rothe, C. *et al.* Transmission of 2019-nCoV Infection from an Asymptomatic Contact in  
415 Germany. *The New England journal of medicine* (2020) doi:10/ggivr8.

416 28. Avanzato, V. A. *et al.* Case Study: Prolonged Infectious SARS-CoV-2 Shedding from an  
417 Asymptomatic Immunocompromised Individual with Cancer. *Cell* S0092867420314562  
418 (2020) doi:10.1016/j.cell.2020.10.049.

- 419 29. Stadlbauer, D. *et al.* SARS-CoV-2 Seroconversion in Humans: A Detailed Protocol for a  
420 Serological Assay, Antigen Production, and Test Setup. *Current Protocols in Microbiology*  
421 **57**, (2020).
- 422 30. Wrapp, D. *et al.* Cryo-EM structure of the 2019-nCoV spike in the prefusion conformation.  
423 *Science* **367**, 1260–1263 (2020).
- 424 31. Amanat, F. *et al.* A serological assay to detect SARS-CoV-2 seroconversion in humans. *Nat*  
425 *Med* **26**, 1033–1036 (2020).