

In vitro and in vivo functions of SARS-CoV-2 infection-enhancing and neutralizing antibodies

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# 38 Summary (150 words)

51	
50	antibody-infused macaques.
49	necessarily herald enhanced infection in vivo, increased lung inflammation can rarely occur in SARS-CoV-2
48	bronchoalveolar lavage inflammatory cytokines. Thus, while in vitro antibody-enhanced infection does not
47	higher lung inflammation scores compared to controls. One monkey had alveolar edema and elevated
46	from SARS-CoV-2 replication in monkeys and mice. Three of 46 monkeys infused with enhancing antibodies had
45	FcyR-independent in vitro infection enhancement. However, both types of infection-enhancing antibodies protected
44	(FcγR)-mediated enhancement of virus infection in vitro, while five non-neutralizing NTD antibodies mediated
43	antibodies demonstrated function-specific modes of binding. Select RBD NAbs also demonstrated Fc receptor- $\gamma$
42	convalescent SARS-CoV-2 or a history of SARS-CoV infection. Cryo-electron microscopy of RBD and NTD
41	domain (RBD) and the N-terminal domain (NTD) of SARS-CoV-2 spike from individuals with acute or
40	antibodies is whether they mediate disease enhancement. Here, we isolated NAbs against the receptor-binding
39	SARS-CoV-2 neutralizing antibodies (NAbs) protect against COVID-19. A concern regarding SARS-CoV-2

# 52 Keywords

53 SARS-CoV-2, COVID-19, neutralizing antibody, receptor-binding domain, N-terminal domain, electron

54 micrograph, *in vivo* protection, infection enhancement

### 56 Introduction

57 The severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) has caused a global pandemic with over 58 157 million cases and 3 million deaths (https://coronavirus.jhu.edu). While the ultimate solution to control the 59 COVID-19 pandemic is a safe and effective vaccine, neutralizing Ab (NAb) prophylaxis or treatment of infection 60 may help to control the pandemic (Graham, 2020; Sempowski et al., 2020). Prophylactic or therapeutic use of 61 SARS-CoV-2 NAbs in non-human primates (Baum et al., 2020a; Jones et al., 2020; Zost et al., 2020a) or rodent 62 models (Hassan et al., 2020; Rogers et al., 2020; Wu et al., 2020) have protected against SARS-CoV-2 infection. 63 Potent SARS-CoV-2 NAbs reported to date predominantly target the RBD region (Baum et al., 2020b; Brouwer et 64 al., 2020; Cao et al., 2020; Hansen et al., 2020; Ju et al., 2020; Liu et al., 2020a; Pinto et al., 2020; Robbiani et al., 65 2020; Rogers et al., 2020; Shi et al., 2020; Wrapp et al., 2020a; Wu et al., 2020). In contrast, neutralizing SARS-66 CoV-2 NTD antibodies (Abs) exhibit more modest neutralization potency (Brouwer et al., 2020; Chi et al., 2020; 67 Wec et al., 2020; Zost et al., 2020a; Zost et al., 2020b).

68 A safety concern for clinical use of antibodies is antibody-dependent enhancement (ADE) of infection. ADE 69 in vitro has been reported for respiratory syncytial virus vaccination, dengue virus vaccination, or dengue virus 70 infection (Arvin et al., 2020). ADE is often mediated by Fc receptors for IgG (FcyRs), complement receptors (CRs) 71 or both, and is most commonly observed in monocytes/macrophages and B cells (Iwasaki and Yang, 2020; Ubol 72 and Halstead, 2010). In vitro studies have demonstrated FcyR-mediated ADE of SARS-CoV infection of ACE2-73 negative cells (Jaume et al., 2011; Kam et al., 2007; Wan et al., 2020; Wang et al., 2014; Yilla et al., 2005; Yip et 74 al., 2016; Yip et al., 2014). Additional research has demonstrated FcyR-independent infection enhancement of 75 SARS-CoV in Vero cells, and isolated an Ab that may have enhanced lung viral load and pathology in vivo (Wang 76 et al., 2016). The ability of SARS-CoV-2 S Abs to mediate infection enhancement in vivo is unknown, but is a 77 theoretical concern for COVID-19 vaccine development (Arvin et al., 2020; Bournazos et al., 2020; Haynes et al., 78 2020; Iwasaki and Yang, 2020).

79 Here, we identified potent *in vitro* neutralizing RBD and NTD Abs as well as *in vitro* infection-enhancing

80 RBD and NTD Abs from individuals infected with SARS-CoV or SARS-CoV-2. Negative stain electron

81 microscopy (NSEM) and cryo-electron microscopy (cryo-EM) revealed distinct binding patterns and the precise

82 epitopes of infection-enhancing and neutralizing Abs. In vitro studies demonstrated that select RBD Abs mediated

83 FcyR-dependent infection enhancement, whereas the NTD Abs induced FcyR-independent infection enhancement.

84	However, using monkey and mouse models of SARS-CoV-2 infection, none of the <i>in vitro</i> infection-enhancing
85	Abs enhanced SARS-CoV-2 virus replication or infectious virus in the lung in vivo. Three of 46 monkeys had lung
86	pathology or bronchoalveolar lavage (BAL) cytokine levels greater than controls. However, repeat studies with
87	dose ranges of in vitro enhancing Abs did not increase lung pathology. Thus, in vitro infection-enhancing RBD and
88	NTD Abs controlled virus in vivo and was rarely associated with enhanced lung pathology.
89	
90	RESULTS
91	Isolation of neutralizing and infection-enhancing SARS-CoV-2 Abs
92	SARS-CoV-2-reactive monoclonal Abs from plasmablasts or SARS-CoV-2-reactive memory B cells were
93	isolated (Liao et al., 2009; Liao et al., 2013) from a SARS-CoV-2 infected individual 11, 15 and 36 days post-onset
94	of symptoms. To identify neutralizing Abs against both SARS-CoV and SARS-CoV-2, SARS-CoV-2 S-reactive B
95	cells were isolated from an individual infected with SARS-CoV ~17 years prior to sample collection (Figures 1A-
96	<b>B</b> , S1A-D). From 1,737 total B cells, we isolated 463 Abs that bound to SARS-CoV-2 S or nucleocapsid proteins in
97	high-throughput binding screens (Figure 1C; Table S1). We selected 187 Abs using high binding magnitude,
98	cross-reactivity with human CoVs, high somatic mutation frequency, and long HCDR3 as selection criteria.
99	Downselected Abs were examined for neutralization of SARS-CoV-2 pseudovirus and replication-competent
100	SARS-CoV-2. Forty-four of 81 RBD Abs exhibited neutralization of SARS-CoV-2 pseudovirus or replication-
101	competent virus (Figures S1E-J; Tables S2). Ten of forty-one NTD Abs neutralized SARS-CoV-2 in the
102	293T/ACE2 pseudovirus and plaque reduction assays, at an IC <sub>50</sub> as low as 39 ng/mL ( <i>Figures S1K-M; Tables S2</i> ).
103	In addition, 5 non-neutralizing NTD Abs enhanced SARS-CoV-2 pseudovirus infection in 293T/ACE2 and
104	replication-competent SARS-CoV-2 nano-luciferase virus infection of Vero cells (Figure 1D,E) (Huo et al., 2020).
105	NTD Ab infection enhancement was dependent on ACE2 expression. Both ACE2-expressing 293T cells used for
106	pseudovirus assays and Vero cells lack FcyR expression (Takada et al., 2007). Thus, NTD enhancement of SARS-
107	CoV-2 infection was FcyR-independent.
108	To assess FcyR-dependent infection enhancement, 100 S-reactive IgG1 Abs were tested for their ability to

109 facilitate SARS-CoV-2 infection of TZM-bl cells expressing various FcγRs, but lacking ACE2 and TMPRSS2

110 (*Tables S2*). Three or five Abs enabled SARS-CoV-2 infection of TZM-bl cells expressing either FcyRI or FcyRIb

- 111 respectively (*Figures 1F-J*). The antigen-binding fragments (Fabs) of these Abs did not mediate infection
- 112 enhancement of TZM-bl cells expressing FcyRI or FcyRIIb, demonstrating Fc-dependence for enhancement
- 113 (Figures 1K-L). Thus, RBD Abs can be either neutralizing in ACE2-expressing 293T cells, infection-enhancing in
- 114 the FcγR-expressing TZM-bl cells, or both (*Figure 2A*). NTD Abs can either be neutralizing or infection-
- enhancing in the ACE2<sup>+</sup> 293T cells or VeroE6 cells (*Figure 2A*).
- 116

## 117 Characterization of infection-enhancing Spike Abs

118 We compared the phenotypes and binding modes of RBD Abs that either did not enhance or enhanced

- 119 infection in order to elucidate differences between them. The selected RBD Abs neutralized SARS-CoV-2
- 120 pseudovirus and/or replication-competent virus in ACE2-expressing cells (Figures 2A and S2), despite five of
- 121 these Abs mediating infection enhancement in ACE2-negative, FcyR-positive TZM-bl cells (*Figures 1F-L, 2A*,

122 *and S2*). Both types of selected RBD Abs blocked ACE2 binding to S protein and both types of RBD Abs bound to

- 123 S with high affinities (range = 0.1 to 9 nM)(*Table S3, Figure 2A*). Thus, the infection-enhancing or non-enhancing
- 124 RBD Abs showed similarities in ACE2 blocking, affinity, and neutralization of ACE2-dependent SARS-CoV-2
- 125 infection (*Figure 2A*).

126 For six representative RBD Abs, we obtained NSEM reconstructions of Fabs in complex with stabilized S 127 ectodomain trimer. Infection-enhancing RBD Abs DH1041 and DH1043 bound with a vertical approach (Figure 128 2B), parallel to the central axis of the S trimer, similar to non-infection-enhancing Abs DH1042 and DH1044 129 (Figure 2C). The epitopes of Abs DH1041, DH1042, and DH1043 overlapped with that of the ACE-2 receptor 130 (Wec et al., 2020), consistent with their ability to block ACE-2 binding to S protein (Figures 2A and S3A-B). 131 Their epitopes were similar to those of three previously described Abs, P2B-2F6 (Ju et al., 2020), H11-H4, and 132 H11-D4 (Figure S3C) (Huo et al., 2020; Zhou et al., 2020a). The epitope of another non-infection-enhancing RBD 133 Ab DH1044 was only slightly shifted relative to DH1041, DH1042 and DH1043 (Figure 2C), but resulted in 134 DH1044 not blocking ACE2 binding (Figures 2A and S3A-B). The remaining two RBD Abs, DH1045 and 135 DH1047, cross-reacted with both SARS-CoV and SARS-CoV-2 S (Figures 2A and S2A-B). DH1047 also reacted 136 with bat and pangolin CoV spike proteins (Figures 2A and S2A). Although DH1047 mediated FcyR-dependent 137 infection of TZM-bl cells and DH1045 did not, both Abs bound to RBD-up S conformations with a more horizontal

angle of approach (*Figures 2B-C and S3A*) (Pak et al., 2009). Thus, epitopes and binding angles of RBD Abs
determined by NSEM did not discriminate between Abs that mediated FcγR-dependent infection enhancement and
those that did not.

141 Next we characterized the Fabs of neutralizing NTD Abs DH1050.1 and DH1051 bound to stabilized S 142 ectodomain with affinities of 16 and 19 nM respectively, whereas the infection-enhancing Ab DH1052 bound with 294 nM affinity (Table S3). NSEM reconstructions obtained for nine NTD Abs showed that the FcyR-independent, 143 144 infection-enhancing NTD Abs (DH1053-DH1056) bound to S with their Fab constant domains directed downward 145 toward the virus membrane (*Figure 2D*), whereas the five neutralizing NTD-directed Abs (DH1048-DH1051) 146 bound to S with the constant domain of the Fab directed upward away from the virus membrane (Figure 2E). The 147 five neutralizing Abs bound the same epitope as Ab 4A8 (Chi et al., 2020), with three of the five having the same 148 angle of approach and heavy chain gene segment (V<sub>H</sub>1-24) as 4A8 (Figure S3D-F and Table S2) (Chi et al., 2020). 149 These NTD Abs may constitute a neutralizing Ab class that can be elicited in multiple individuals. Thus, S protein 150 Ab epitopes and binding modes were associated with infection-enhancing activity of NTD Abs.

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# 152 Competition between infection-enhancing and non-infection enhancing Abs

To determine whether infection-enhancing Abs could compete with non-infection-enhancing Abs for binding to S ectodomain, we performed surface plasmon resonance (SPR) competitive binding assays. RBD Abs segregated into two clusters, where Abs within a cluster blocked each other and Abs in different clusters did not block each other (*Figures 3A*). One cluster included Abs DH1041, DH1043 and DH1044, and the other cluster included Abs DH1046 and DH1047. NSEM reconstructions showed combinations of DH1041 and DH1047 Fabs or DH1043 and DH1047 Fabs bound simultaneously to different epitopes of the stabilized S trimer (*Figure 3B*).

159 NTD Abs also segregated into two clusters where one cluster included neutralizing NTD Abs and a second

160 cluster included non-neutralizing NTD Abs (*Figures 3A and 3C*). NSEM reconstructions confirmed that the Fabs

- 161 of neutralizing NTD Ab DH1050.1 and infection-enhancing NTD Ab DH1052 could simultaneously bind to
- 162 distinct epitopes on a single SARS-CoV-2 S trimer (*Figure 3D*). DH1054 was unique as it was able to block both
- 163 infection-enhancing and neutralizing NTD Abs (*Figures 3C*).

164 NTD Abs did not compete with RBD Abs for binding to S trimer (*Figure 3A*), suggesting in a polyclonal
 165 mixture of Abs, the SARS-CoV-2 S trimer could bind both RBD and NTD Abs. NSEM showed that 1 or 2

- different neutralizing RBD Abs (DH1043 and DH1047) could bind to the same S protomer as neutralizing NTD
  Abs DH1050.1 or DH1051 (*Figure 3E*,*F*). Thus, in the presence of a polyclonal Ab response, S trimer could be
  bound by multiple RBD and NTD neutralizing Ab Fabs.
- 169

# 170 FcyR-independent infection-enhancement in the presence of neutralizing Abs

171 Structural determination of Ab binding modes demonstrated that certain infection-enhancing Abs and non-172 infection enhancing Abs bound to distinct epitopes on the same S protomer (Figures 3A-F). Infection-enhancing 173 Ab DH1052 and neutralizing RBD Ab DH1041 were isolated from the same individual. We hypothesized that 174 infection outcome would be dependent on which Ab was present at the highest concentration. When DH1041 175 neutralization was assessed in the presence of 1,325-fold excess of Ab DH1052, infection enhancement was 176 observed when DH1041 concentration was below 10 ng/mL (Figures 3G and S4A-C). A nearly identical result 177 was obtained when we examined neutralization by DH1043 (Figures 3H and S4A-C). In 21 SARS-CoV-2-infected humans, RBD and NTD serum IgG titers were comparable (*Figures S4D-E*). Moreover, the prevalence of DH1052 178 versus DH1041 Abs was assessed using blocking assays and found to be only modestly higher for DH1052 (Figure 179 180 S4F). Thus, a ~1000-fold excess of infection-enhancing NTD Ab was required to out-compete the effect of a 181 potent RBD neutralizing Ab in vitro, but such excess amounts of DH1052 was not observed during natural infection (Figures 3G-H and S4D-F). 182

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# 184 Cryo-EM structural determination of RBD and NTD-directed Ab epitopes

185 To visualize atomic level details of their interactions with the S protein, cryo-EM was used for structural 186 determination of selected representative Abs from the panels of RBD and NTD-directed Abs. For all three RBD-187 directed Abs, the cryo-EM datasets revealed heterogeneous populations of S ectodomain "2P" (S-2P) (Wrapp et al., 188 2020b) with at least one RBD in the "up" position (Figure 4 and Data S1). We did not find any unliganded S or 189 any 3-RBD-down S population, although unliganded S-2P consistently shows a 1:1 ratio of 1-RBD-up and 3-RBD-190 down populations (Henderson et al., 2020; Walls et al., 2020). All S-2P trimers were stoichiometrically bound to 191 three Fabs, with Abs bound to both up and down RBDs in an S-2P trimer. 192 We observed that the primary epitopes of DH1041 and DH1043 were centered on the Receptor Binding Motif

193 (RBM; residues 483-506) of the RBD (*Figures 4A-B and Data S1*), providing structural basis for the ACE-2

194 blocking phenotype of these Abs. While DH1041 utilized its heavy chain complementarity determining regions (CDRs) to contact the RBM, the DH1043 paratope included both its heavy and light chains. In contrast, the epitope 195 196 of Ab DH1047 was focused around the  $\alpha^2$  and  $\alpha^3$  helices and  $\beta^2$  strand that are located outside the N-terminus of 197 the RBM (Figure 4C and Data S1). DH1047 also contacted RBD residues 500-506 outside the RBM, and stacked 198 against the N-terminal end of the  $\alpha$ 3 helix. The DH1047 paratope included heavy chain HCDR2, HCDR3 and light 199 chain LCDR1 and LCDR3. The HCDR3 stacks against and interacts with the residues in the ß2 strand. Interactions 200 with the ß2 strand are also mediated by HCDR2. Similar to DH1041 and DH1043, the DH1047 interacted with an 201 "up" RBD conformation from an adjacent protomer although these interactions were not well-characterized due to 202 disorder in that region.

203 We next determined cryo-EM structures of the NTD-directed neutralizing Abs, DH1050.1 (Figure 4D) and NTD-204 directed infection-enhancing Ab, DH1052 (Figure 4E), at 3.4 Å and 3.0 Å resolutions, respectively. The cryo-EM 205 datasets of DH1050.1- and DH1052-bound complexes showed Fab bound to both 3-RBD-down and 1-RBD-up S-2P spikes (Data S1). Consistent with the NSEM reconstructions, the neutralizing Ab DH1050.1 and the non-206 207 neutralizing, infection-enhancing Ab DH1052 bound opposite faces of the NTD, with the epitope for the 208 neutralizing Ab DH1050.1 facing the host cell membrane and the epitope for the non-neutralizing, infection-209 enhancing Ab DH1052 facing the viral membrane. The dominant contribution to the DH1050.1 epitope came from 210 NTD loop region 140-158 that stacks against the Ab HCDR3 and extends farther into a cleft formed at the interface 211 of the DH1050.1 HCDR1, HCDR2 and HCDR3 loops. The previously described NTD Ab 4A8 interacts with the 212 same epitope in a similar manner as DH1050.1, with its elongated HCDR3 dominating interactions. Although, 213 DH1050.1 and 4A8 (Chi et al., 2020) show a rotation relative to each other about the stacked HCDR3 and NTD 214 140-158 loops. The light chains of DH1050.1 and 4A8 do not contact the S protein, which is consistent with their 215 diverse light chain gene origins (Figure 4E and Data S1). The infection enhancing NTD-directed Ab DH1052 216 bound the NTD at an epitope facing the viral membrane and composed of residues spanning 27-32, 59-62 and 211-217 218, with all the CDR loops of both heavy and light chains involved in contacts with the NTD. We also observed 218 contact of the Ab with the glycan at position 603, as well as the conformationally invariant SD2 region. Thus, we 219 found that the RBD-directed antibodies isolated in this study influenced RBD dynamics and bound only to spike 220 with at least one RBD in the up conformations, and in some cases, also induced the 2-RBD-up and 3-RBD-up spike

conformations. In contrast, the NTD-directed antibodies bound to both the 3-RBD-down and 1-RBD-up spikes that
 are present in the unliganded S-2P.

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### 224 Effect of *in vitro* infection-enhancing and neutralizing NTD Abs in mouse and macaque models

225 Next, we assessed the effect of NTD infection-enhancing Ab DH1052 in a COVID-19 disease mouse model 226 where aged BALB/c mice were challenged with the mouse-adapted SARS-CoV-2 MA10 strain (Leist et al., 2020a). 227 DH1052 lacked neutralization of SARS-CoV-2 MA10 (Figure S4G-H). DH1052 or a control influenza Ab CH65 228 was given 12 hours prior to SARS-CoV-2 MA10 infection (Figure 5A). Throughout the four days of infection, 229 DH1052-infused mice exhibited similar levels of body weight loss and higher survival than mice given CH65 230 (Figures 5B-C). In addition, DH1052-treated mice exhibited lower lung hemorrhagic scores, lower lung viral 231 plaque-forming unit (PFU) titers and lower lung tissue subgenomic RNA (sgRNA) levels compared to control mice 232 (Figures 5D-F). Therefore, DH1052 treatment resulted in less severe disease and reduced viral replication. FcR-233 mediated effector functions may have been the mechanism of suppression since DH1052 bound to mouse FcyRI 234 and FcyRIV (*Table S4*).

235 We next examined the effect of infusion of NTD infection-enhancing Ab DH1052, NTD neutralizing Ab 236 DH1050.1, or control Ab CH65 on SARS-CoV-2 infection in monkeys (Leist et al., 2020b; Rockx et al., 2020). 237 Cynomolgus macaques were infused with 10 mg of Ab per kg body weight and three days later challenged 238 intranasally and intratracheally with 10<sup>5</sup> PFU of SARS-CoV-2 (*Figure 5G*) (54). Human Ab infusion resulted in 239 circulating concentrations ranging from 11 to 238 µg/mL in serum at day 2 post-challenge (Figures 5H-I). Sera 240 with DH1050.1 neutralized SARS-CoV-2 pseudovirus and replication-competent virus, while serum containing 241 DH1052 or CH65 did not neutralize (Figures 5J-K). Four of 5 macaques that received DH1052 had comparable 242 lung inflammation to control CH65-infused macaques four days after infection (Figures 5L and S5A). However, 243 one macaque (BB536A) administered DH1052 showed increased perivascular mononuclear inflammation, 244 perivascular and alveolar edema (*Figure S5B*), and multiple upregulated BAL cytokines (*Table S5*). Immunohistologic analysis demonstrated alveolar and perivascular infiltration of M2-type macrophages in both 245 246 monkey BB536A and a control monkey BB785E (Figures S5C-E). In contrast, macaques administered DH1050.1,

a neutralizing NTD Ab, had lower lung inflammation (*Figures 5L and S5A*) and fewer infiltrating macrophages

- 248 (*Figures S5C-E*). Infusion of either DH1050.1 or DH1052 reduced viral nucleocapsid antigen (*Figures 5M and*
- 249 *S5A*), Envelope (E) gene sgRNA and nucleocapsid (N) gene sgRNA in the BAL (*Figures 5N-O*). In nasal swab
- 250 fluid, DH1050.1 and DH1052 reduced E and N gene sgRNA in macaques with the reduction being significant
- when neutralizing Ab DH1050.1 was infused (*Figures 5P-Q*).
- 252 Since DH1052-mediated *in vitro* infection-enhancement increased as the Ab concentration increased (*Figures*
- 253 *ID-E*), we infused an additional 6 cynomolgus macaques with either 30 mg/kg of DH1052 or CH65 control Ab
- 254 (Figure S6A). DH1052 infusion suppressed BAL viral load (Figures S6B-D), significantly reduced virus
- 255 replication in nasal swab samples (*Figures S6E-G*), and showed no enhanced immunopathology or cytokine
- 256 secretion (*Figures S6H-K, Table S5*). Thus, with high dose (30mg/kg) of DH1052 Ab, there was no infection
- enhancement. These results suggested that the lung pathology seen in monkey BB536A was rare and may not havebeen caused by Ab infusion.
- 259

# 260 FcyR-dependent, *in vitro* infection-enhancing RBD Abs do not enhance SARS-CoV-2 infection in mice

261 Next, we used a SARS-CoV-2 acquisition mouse model to investigate the *in vivo* relevance of RBD

262 neutralizing Abs that also mediated in vitro infection-enhancement (Figures 6A-B). Aged BALB/c mice were 263 injected intraperitoneally with 300 µg of Ab, and challenged with a SARS-CoV-2 mouse-adapted 2AA MA isolate 264 12 hours later (Dinnon et al., 2020). Mice received either in vitro infection-enhancing Ab DH1041, non-infection 265 enhancing Ab DH1050.1, or a combination of both Abs. Administration of DH1041 alone or in combination with 266 DH1050.1 protected all mice from detectable infectious virus in the lungs 48h after challenge (Figure 6A). In the 267 setting of therapeutic treatment, administration of DH1041 alone or in combination with DH1050.1 12 hours after 268 SARS-CoV-2 challenge significantly reduced lung infectious virus titers (*Figure 6B*). Thus, while RBD Ab 269 DH1041 could mediate FcyR-dependent, in vitro infection enhancement, it protected mice from SARS-CoV-2

- 270 infection when administered prophylactically or therapeutically.
- 271 DH1046 and DH1047 are RBD Abs that cross-neutralize SARS-CoV, SARS-CoV-2 and bat WIV1-CoV
- 272 (Figures 2A, S2A-B, S2I-L and 6C-E). Both RBD Abs mediated FcyR-dependent, in vitro SARS-CoV-2 infection
- 273 enhancement (*Figures 1F-L*). We assessed the ability of either DH1046 or DH1047 to enhance or protect against
- bat WIV1-CoV infection in HFH4-ACE2-transgenic mice (*Figures 6F-G*). Mice administered DH1046 or DH1047

before challenge had no detectable infectious virus in the lung, whereas control IgG administered mice had a mean
titer of 84,896 PFU per lung lobe (*Figure 6F*). Administration of DH1047 after challenge eliminated detectable
infectious virus in the lung in 3 of 5 mice (*Figure 6G*). Therapeutic administration of DH1046 reduced infectious
virus titers 10-fold compared to negative control IgG (*Figure 6G*). Thus, DH1046 and DH1047 did not enhance
infection *in vivo*, but rather protected mice from SARS-related bat coronavirus infection.

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# 281 In vitro infection-enhancing RBD Abs in SARS-CoV-2-challenged nonhuman primates

282 Finally, we assessed RBD Ab infection-enhancement in cynomolgus macaques (Figures 7A). After Ab 283 infusion at 10 mg/kg of body weight, serum human IgG concentrations reached 11-228 µg/mL at day 2 post-284 challenge (Figures 7B-C) and exhibited a wide range of neutralization potencies against SARS-CoV-2 (Figures 285 7D-E). Infusion of RBD Ab DH1041, DH1043, or DH1047 resulted in reduced lung inflammation, undetectable 286 lung viral antigen (Figures 7F-G and S5A), and reduced sgRNA in the upper and lower respiratory tracts (Figures 287 7H-K). RBD Ab DH1046, a weaker neutralizing Ab compared to DH1041, DH1043 or DH1047 (Figure 2A), did 288 not enhance sgRNA E or N in BAL or nasal swab samples (Figures 7H-K), but protected only a subset of infused 289 monkeys. Two DH1046-infused monkeys had increased lung inflammation scores due to increased total areas of 290 inflammation compared to control Ab monkeys (Figures 7F and S5A), but had no evidence of perivascular or 291 alveolar edema nor evidence of abnormal BAL cytokines (*Table S5*). Thus, these two animals had more lung 292 involved with inflammatory macrophage infiltration but did not have pathological evidence of vascular leakage. 293 Comparing the DH1046 group to the control IgG group, viral nucleocapsid antigen in the lung was reduced 294 (*Figures 7G and S5A*). Thus, the weakly neutralizing Ab only partially limited virus replication and lung 295 inflammation.

In vitro infection enhancement by RBD Abs was dependent on Ab concentration, with lower levels of Abshowing the highest magnitude of infection enhancement (*Figure 1G*). Therefore, we performed an additionalpassive infusion study with a series of different concentrations of DH1047 (*Figure S7A*). Cynomolgus macaqueswere infused with 5, 1, or 0.1 mg of DH1047 per kg of body weight resulting in a wide range of DH1047concentrations in serum (*Figures S7B-C*). However, none of the groups of macaques had enhanced virusreplication (although one monkey in the 0.1mg/kg group had higher BAL sgRNA E and N than controls) (*Figures* 

302 *S7D-G*), lung inflammation (*Figures S7H-I*), lung viral antigen (*Figures S7J-K*), or higher BAL inflammatory
303 cytokines (*Table S5*) compared to the control IgG group.

304 Overall, 45 of 46 spike enhancing Ab-infused monkeys did not show enhanced virus replication in vivo, while 305 3 of 46 Ab-treated monkeys exhibited enhancement of lung pathology, with 1 of 46 Ab-treated monkeys had 306 alveolar and perivascular edema and with elevated BAL inflammatory cytokines. In the case of the latter monkey, a 307 follow-up study with 3 times the initial DH1052 Ab dose did not confirm DH1052 result in enhanced lung 308 pathology after SARS-CoV-2 challenge.

309

## 310 **DISCUSSION**

311 Here, we assessed infection-enhancement by SARS-CoV-2 Abs and observed two different types of in vitro 312 infection enhancement. First, RBD Abs mediated classical ADE that required FcyRs and Ab Fc for virus uptake 313 (Lee et al., 2020). Previous studies have demonstrated that uptake of MERS-CoV or SARS-CoV has mostly been 314 mediated by FcyRIIa on the surface of macrophages (Bournazos et al., 2020; Wan et al., 2020; Yip et al., 2016). In 315 contrast, we identified SARS-CoV-2 RBD Abs utilized FcyRIIb or FcyRI. Second, non-neutralizing NTD Abs 316 mediated FcyR-independent infection-enhancement in two different FcyR-negative, ACE2-expressing cell types. 317 The mechanism of FcyR-negative *in vitro* enhancement remains unclear, but one previous study has reported that 318 select NTD Abs can enhance S binding to ACE2 (Liu et al., 2020b). 319 Macrophages and other phagocytes are the target cells that take up MERS-CoV leading to infection-

320 enhancement (Hui et al., 2020; Wan et al., 2020; Zhou et al., 2014). In contrast, neither SARS-CoV nor SARS-

321 CoV-2 productively infect macrophages (Bournazos et al., 2020; Hui et al., 2020; Yip et al., 2016). However, a

322 recent study demonstrated that alveolar macrophages harboring SARS-CoV-2 RNA produce T cell

323 chemoattractants leading to T cell IFN-γ production that in turn, stimulates inflammatory cytokine release from

324 alveolar macrophages (Grant et al., 2021). Why severe lung pathology and inflammatory cytokine production

325 occurred in only 1 of 46 monkeys is unknown, but may relate to host-specific differences regulating inflammatory

326 cytokine production (Bastard et al., 2020; Zhang et al., 2020). It is important to note that the one monkey that

327 developed alveolar and perivascular edema and elevated BAL inflammatory cytokines could have been caused by

328 Ab enhancement of disease, or could have been due to unknown factors that caused more severe disease in animal

329 BB536A that were unrelated to DH1052 administration. That none of 6 animals infused with a higher dose

(30mg/kg) of DH1052 had enhanced pulmonary disease supports the hypothesis that the lung pathology may have
been a severe case of COVID-19 lung disease unrelated to Ab infusion.

332 Previous studies with vaccine-induced Abs against SARS-CoV have also shown in vitro infection-

333 enhancement, but no in vivo infection enhancement in hamsters (Kam et al., 2007). One explanation for this results 334 may be that *in vitro* enhancing Abs may have the ability to suppress SARS-CoV-2 replication *in vivo* through non-335 neutralizing FcR-mediated Ab effector functions (Bournazos et al., 2020; Schafer et al., 2021). A recent study in a 336 SARS-CoV-2 mouse model of acquisition suggested that Fc effector functions contribute to the protective activity 337 of SARS-CoV-2 neutralizing Abs C104, C002, and C110 (Schafer et al., 2021). Thus, Ab effector functions may 338 contribute to the outcome in vivo, but not be accounted for in SARS-CoV-2 enhancement or neutralization assays 339 in vitro. Consistent with previous findings for human IgG (Dekkers et al., 2017), we observed that DH1052 Ab can 340 bind to select murine FcyRs.

341 In vivo, SARS-CoV-2 S trimers circulate in the presence of a polyclonal Ab response. We observed bivalent 342 and trivalent combinations of Fabs from RBD and NTD neutralizing Abs can recognize the same protomer of the S trimer. We speculate given the direction of the C-termini of the Fabs and molecular modeling that three IgGs 343 344 targeting distinct epitopes may be able to interact with the same protomer, if the IgG hinge region is sufficiently 345 flexible and the RBD is in an optimal up conformation for simultaneous engagement. Simultaneous engagement by 346 RBD and NTD Abs could improve synergism of neutralization (Zost et al., 2020a), and avidity of the immune 347 complexes for FcyRs on effector cells (Nagashima et al., 2011; Nagashima et al., 2008; Wang et al., 2017). These 348 results indicate three epitopes that Ab prophylactics could target on RBD and NTD in order to occupy S trimers 349 with multiple IgGs.

## 350 Limitations of the Study

Although rare enhanced immunopathology was observed in monkeys, it is difficult to predict whether this phenomenon will occur in the setting of human infection or vaccination. Furthermore RBD and NTD antibodies were the focus of this study, therefore whether antibodies of other specificities mediate ADE warrants further study. Additionally, the macaque model has a rather short course of infection, thus effects of SARS-CoV-2 antibody on persistent SARS-CoV-2 infection was not examined here.

356	Finally, administration of COVID-19 convalescent sera to over 35,000 COVID-19 patients have demonstrated
357	the treatment to be safe and is not associated with enhanced disease (Joyner et al., 2020). Of greater importance is
358	that both the Pfizer/BioNTech and Moderna mRNA-lipid nanoparticle (LNP) vaccine efficacy trials have
359	completed and showed ~95% vaccine efficacy (Jackson et al., 2020; Polack et al., 2020). That the Moderna
360	mRNA-LNP COVID-19 vaccine efficacy trial had 30 severe cases of COVID-19 occur—all in the placebo group
361	(Baden et al., 2021), demonstrated that if ADE of infection or lung pathology will occur in humans with
362	vaccination, it will be rare. A recent study demonstrated that suboptimal neutralizing Ab level is a significant
363	predictor of severity for SARS-CoV-2 (Garcia-Beltran et al., 2020). Thus, even with the rarity of severe lung
364	pathology associated with presence of anti-spike Ab in animal model studies reported here, it will be important to
365	continue to monitor on-going COVID-19 vaccination for the possibility of vaccine associated enhanced disease
366	when suboptimal neutralizing Ab titers are induced (Haynes et al., 2020).
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- 400 experiments and analysis. W.R. and Y.W. provided statistical analyses. K.W., C.T.D., T.N.D., G.D.S., D.C.M.,
- 401 C.W.W., E.P., M.A.M., I.N.M., R.S., H.A., M.G.L., and R.S.B. oversaw studies. A.G.S., J.F., B.M.H, T.M.C., I.T.,
- 402 T.Z., P.D.K., J.M. and B.G. provided key reagents. K.O.S. and B.F.H. conceived, designed and supervised the
- study, and evaluated all data. D.L., R.J.E., P.A., K.O.S., and B.F.H. wrote the paper. All authors reviewed and 403 approved the manuscript.
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#### 406 **DECLARATION OF INTERESTS**

- 407 B.F.H., G.D.S. K.O.S., R.P., D.L., P.A. and X.L. have applied for patents concerning SARS-CoV-2 Abs that 408 are related to this work. All other authors declare no conflict of interest.
- 409
- 410 **FIGURE LEGENDS**

### 411 Figure 1. SARS-CoV-2 receptor-binding domain (RBD) and N-terminal domain (NTD) Abs mediate

# 412 enhancement of infection.

- 413 (A-B) Timeline of blood sampling, plasmablasts and/or antigen-specific memory B cells (MBC) sorting, and Ab
- 414 isolation from convalescent (A) SARS-CoV-2 and (B) SARS-CoV donors.
- 415 (C) Summary of number and specificity of Abs isolated from each donor.
- 416 (**D-E**) *In vitro* neutralization curves for NTD infection-enhancing Abs against (D) pseudotyped SARS-CoV-2
- 417 D614G in 293T-hACE2 cells, and (E) replication-competent nano-luciferase (nLuc) SARS-CoV-2 in Vero cells.
- 418 (**F-J**) FcγR-dependent pseudotyped SARS-CoV-2 infection-enhancement when RBD Abs or mock medium control
- 419 was added to (F) parental TZM-bl cells, and TZM-bl cells stably expressing human FcyR receptors (G) FcyRI, (H)
- 420 FcγRIIa, (I) FcγRIIb or (J) FcγRIII.
- 421 (K-L) The effect of RBD Ab fragment antigen-binding regions (Fabs) on pseudotyped SARS-CoV-2 D614G
- 422 infection was tested in (K) FcyRI-expressing TZM-bl cells and (L) FcyRIIb-expressing TZM-bl cells. Data are
- 423 represented as mean±SEM. Three or four independent experiments were performed and representative data are
- 424 shown.

# 425 Figure 2. Structural and phenotypic characterization of infection-enhancing and non-infection-enhancing

# 426 **RBD and NTD Abs.**

- 427 (A) Summary of Ab epitope, binding, and neutralizing or infection-enhancing activity in ACE2-positive/FcγR-
- 428 negative cells or ACE2-negative/Fc $\gamma$ R-positive cells. Ab functions are color-coded based on the key shown at the 429 right. MN titer, micro-neutralization titer; ND, not determined.
- 430 (B-E) 3D reconstruction of negative stain electron microscopy images of stabilized SARS-CoV-2 S ectodomain
- 431 trimers (S-2P; gray) bound to the Fabs (various colors) of (B,D) infection-enhancing or (C,E) non-infection-
- 432 enhancing RBD or NTD antibodies.

# 433 Figure 3. Simultaneous binding of infection-enhancing and non-infection enhancing Abs to individual S

- 434 trimers.
- 435 (A) Cross-blocking activity of RBD and NTD neutralizing Abs tested by surface plasmon resonance (SPR). S-2P
- 436 was captured by one Ab (Y-axis) followed by binding by the second Ab (X-axis).
- 437 (B) 3D reconstruction of simultaneous recognition of SARS-CoV-2 S-2P by two RBD Abs DH1041+DH1047, or
- 438 DH1043+DH1047.

- 439 (C) Cross-blocking activity of neutralizing Abs and infection-enhancing NTD Abs tested by SPR and shown as in440 (A).
- 441 (**D-F**) 3D reconstruction of SARS-CoV-2 S-2P simultaneously bound (D) NTD Abs DH1053 and DH1050.1, (E)
- 442 RBD infection-enhancing Ab and a NTD non-infection-enhancing Ab, or (F) triple-Ab combinations of RBD Ab
- 443 DH1043, RBD Ab DH1047, and either NTD Ab DH1051 (left) or DH1050.1 (right).
- 444 (G-H) RBD Ab neutralization of SARS-CoV-2 D614G pseudovirus infection of 293T/ACE2 cells in the presence
- 445 of 1:132 or 1:1,325 ratios of excess infection-enhancing NTD Ab DH1052.
- 446 Figure 4. Cryo-electron microscopy of neutralizing and non-neutralizing Abs in complex with SARS-CoV-2
- 447 Spike ectodomain. Structures of SARS-CoV-2 S protein in complex with RBD Abs (A) DH1041 (red), (B)
- 448 DH1043 (pink), (C) DH1047 (magenta), (D) neutralizing NTD Ab DH1050.1 (blue), and (E) infection-enhancing
- 449 NTD Ab DH1052 (green). Each Ab is bound to S-2P shown in gray with its RBM colored purple blue. (Right)
- 450 Zoomed-in views of the Ab interactions with S-2P trimers. The Ab complementarity determining (CDR) loops are
- 451 colored: HCDR1 yellow, HCDR2 limon, HCDR3 cyan, LCDR1 orange, LCDR2 wheat and LCDR3 light blue. See
- 452 also Supplemental Data 1.
- 453 Figure 5. NTD Ab DH1052 does not always enhance SARS-CoV-2 replication or disease *in vivo*.
- 454 (A-F) DH1052 passive immunization and murine SARS-CoV-2 challenge (A) study design, (B) body weight, (C)
- 455 survival, (**D**) Hemorrhagic scores, (**E**) lung viral titers, and (**F**) SARS-CoV-2 envelope (E) and nucleocapsid (N)
- 456 gene subgenomic RNA (sgRNA).
- 457 (G-Q) Reduction of SARS-CoV-2 replication and disease in cynomolgus macaques by prophylactic administration
- 458 of an NTD neutralizing Ab DH1050.1 or an NTD *in vitro* infection-enhancing Ab DH1052.
- (G) DH1050.1 and DH1052 prophylaxis cynomolgus macaque (n=5 per group) study design. CH65 was used as a
   negative control Ab.
- 461 (H-I) Serum human IgG concentrations at (H) Day -5 and (I) Day 2.
- 462 (J-K) Day 2 serum neutralization titers shown as the reciprocal serum dilution that inhibits 50% ( $ID_{50}$ ) of (J)
- 463 pseudotyped SARS-CoV-2 replication in 293T/ACE2 cells or (K) SARS-CoV-2 replication in Vero cells.
- 464 (L-M) Lung histopathology four days post infection. Lung sections were scored for (L) inflammation by
- 465 hematoxylin and eosin (H&E) staining, and for (M) the presence of SARS-CoV-2 nucleocapsid by
- 466 immunohistochemistry (IHC) staining.

- 467 (N-Q) Viral load quantified as SARS-CoV-2 E gene sgRNA and N gene sgRNA in (N-O) bronchoalveolar lavage
- 468 (BAL) or (P-Q) nasal swab fluid on Day 2 and Day 4 post challenge. LOD, limit of detection. Statistical
- 469 significance in all the panels were determined using Wilcoxon rank sum exact test. Horizontal bars are the group
- 470 mean. Asterisks show the statistical significance between indicated group and CH65 control group: ns, not
- 471 significant, \*P<0.05, \*\*P<0.01, \*\*\*P<0.001.

# 472 Figure 6. RBD Abs that mediate FcyR-dependent infection-enhancement in vitro, protect mice from SARS-

- 473 CoV-2 or bat WIV1-CoV challenge.
- 474 (A-B) Protection of BALB/c mice (n=5 per group) from mouse-adapted SARS-CoV-2 (SARS-CoV-2 2AA MA) by
- 475 (A) prophylactic or (B) therapeutic RBD and/or NTD Ab administration. Ab CH65 served as a negative control.
- 476 Titers of infectious virus in the lung were examined 48h post-infection.
- 477 (C) Maximum likelihood tree of Spike amino acid sequences for SARS-related group 2B and group 2C
- 478 coronaviruses.
- (D) Monoclonal RBD, NTD and S2 Ab ELISA binding titer for soluble S protein ectodomains from human and
   animal coronaviruses. Titers are log area-under-the-curve (AUC).
- 481 (E) SARS-CoV and bat WIV1-CoV cross-neutralization titers for cross-reactive RBD and S2 Abs.
- 482 (F-G) Protection of HFH4-hACE2-transgenic mice (n=5 per group) from SARS-related bat WIV1-CoV challenge
- 483 by (A) prophylactic or (B) therapeutic RBD Ab administration. Lung viral titers were examined at 48 post-infection.
- 484 Statistical significance in all the panels were determined using Wilcoxon rank sum exact test. Horizontal bars are
- 485 the group mean. Asterisks show the statistical significance between indicated group and CH65 control group: ns,
- 486 not significant, \*P<0.05, \*\*P<0.01.

# 487 Figure 7. RBD Abs that mediate FcyR-dependent infection enhancement *in vitro*, protect non-human

- 488 primates from SARS-CoV-2 challenge.
- 489 (A) Cynomolgus macaques (n=5 per group) RBD Ab SARS-CoV-2 challenge study design. DH1041, DH1043,
- 490 DH1046, DH1047 or an irrelevant CH65 were infused into macaques.
- 491 (**B-C**) Serum human IgG concentrations at Day -5 (**B**) and Day 2 (**C**).
- 492 (**D-E**) Day 2 serum neutralization titers shown as the reciprocal serum dilution that inhibits 50% ( $ID_{50}$ ) of (**D**)
- 493 pseudotyped SARS-CoV-2 replication in 293T/ACE2 cells or (E) SARS-CoV-2 replication in Vero cells.

- 494 (F-G) Lung histopathology for (F) inflammation by H&E staining and (G) the presence of SARS-CoV-2
- 495 nucleocapsid by IHC staining 4 days post-challenge.
- 496 (H-K) Viral load quantified as SARS-CoV-2 E gene sgRNA and N gene sgRNA in (H-I) bronchoalveolar lavage
- 497 (BAL) or (J-K) nasal swab fluid on Day 2 and Day 4 post challenge.
- 498 Statistical significance in all the panels were determined using Wilcoxon rank sum exact test. Horizontal bars are
- 499 the group mean. Asterisks show the statistical significance between indicated group and CH65 control group: ns,
- 500 not significant, \*P<0.05, \*\*P<0.01.
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- 504 STAR METHODS

# 505 **KEY RESOURCES TABLE**

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Antibodies		
PE-Cy5 Mouse Anti-Human CD3, Clone# HIT3a	BD Biosciences	Cat#555341; RRID:
		AB_10698936
BV605 Mouse Anti-Human CD14, Clone# M5E2	Biolegend	Cat#301834, RRID:
	-	AB_2563798
BV570 Mouse Anti-Human CD16, Clone# 3G8	Biolegend	Cat# 302035, RRID:
	-	AB_2632790
APC-Cy7 Mouse Anti-Human CD19, Clone# SJ25C1	BD Biosciences	Cat# 557791, RRID:
		AB_396873
FITC Mouse Anti-Human IgD, Clone# IA6-2	BD Biosciences	Cat# 555778, RRID:
		AB_396113
PerCp-Cy5.5 Mouse Anti-Human IgM, Clone# G20-127	BD Biosciences	Cat# 561285,
		RRID:AB_10611998
PE-CF594, Mouse Anti-Human CD10, Clone# HI10A	BD Biosciences	Cat# 562396, RRID:
		AB_11154416
PE-Cy5 Mouse Anti-Human CD235a, Clone# GA-R2	BD Biosciences	Cat# 559944, RRID:
		AB_397387
PE-Cy7 Mouse Anti-Human CD27, Clone# O323	eBioscience	Cat# 25-0279, RRID:
		AB_1724039
APC-AF700 Mouse Anti-Human CD38, Clone# LS198-	Beckman Coulter	Cat# B23489, RRID:
4-2		NA
SARS-CoV/SARS-CoV-2 Spike Ab, Clone# D001	Sino Biological	Cat #40150-D001
Anti-influenza virus hemagglutinin human IgG CH65	(Whittle et al., 2011)	NA
Rabbit polyclonal SARS-CoV-2 nucleocapsid Ab	GeneTex	Cat #GTX135357,
		RRID:AB_2868464
Rat anti-human CD3, Clone# CD3-12	Bio-Rad	Cat #MCA1477,
		RRID:AB_321245
Rabbit anti-human Iba1 polyclonal Ab	Wako	Cat# 019-19741,
The set and hannah tour porjoional the		RRID: AB_839504
	1	

Rabbit anti-human CD68 polyclonal Ab	Sigma-Millipore	Cat# HPA048982,
		RRID: AB_2680587
Rabbit anti-human CD163, Clone# EPR19518	Abcam	Cat# ab182422,
		RRID: AB_2753196
Mouse anti-human HLA-DP/DQ/DR, Clone# CR3/43	Dako	Cat# M0775, RRID:
		AB_2313661
Rabbit anti-human CD11b, Clone# EP1345Y	Abcam	Cat# ab52478, RRID:
		AB_868788
HRP goat anti-human IgG	SouthernBiotech	Cat #2040-05,
		RRID:AB_2795644
HRP goat anti-rabbit IgG	Abcam	Cat #ab97080,
		RRID:AB_10679808
Biotin mouse anti-human IgG Fc, Clone# H2	Southern Biotech	Cat# 9042-08,
		RRID:AB_2796608
Bacterial and Virus Strains	•	
SARS-CoV-2 D614G pseudotyped virus	(Korber et al., 2020)	NA
SARS-CoV-2 virus, Isolate USA-WA1/2020	BEI Resources	Cat #NR-52281
SARS-CoV-2 nanoLuc virus	(Hou et al., 2020)	NA
SARS-CoV nanoLuc virus	(Sheahan et al.,	NA
Shirds Cov handbac viras	(Sheahan et al., 2017)	1111
WIV1-CoV nanoLuc virus	(Menachery et al.,	NA
	2016)	1111
SARS-CoV-2 moues-adapted virus 2AA MA	(Dinnon et al., 2020)	NA
SARS-CoV-2 modes-adapted virus 2747 MAT	(Leist et al., 2020a)	NA
Biological Samples	(Leist et al., 2020a)	
Plasma, PBMCs, nasal swabs and bronchoalveolar	This nonon	NA
	This paper	NA
lavage (BAL) from macaques		
Chemicals, Peptides, and Recombinant Proteins	Thermo Fisher	Cat#L 24072
LIVE/DEAD Fixable Red Dead Cell Stain Kit		Cat#L34972
	Scientific	<u>C</u> , #10000007
SuperScript III Reverse Transcriptase	Invitrogen	Cat #18080085
dNTP Set, PCR Grade	New England	Cat # N0447L
	Biolabs	
UltraPure DNase/RNase-Free Distilled Water	Invitrogen	Cat #10977
GeneLink Random Hexamer Primers	GeneLink	Cat #26-4000-03
AmpliTaq Gold 360 Mastermix	Thermo Fisher	Cat #4398881
	Scientific	
Expi293 media	Invitrogen	Cat #A1435102
Expifectamine	Life Technologies	Cat #A14524
Protein A beads	Pierce	Cat #PI-20334
Mfal IIE	New England	R3589L
MfeI-HF	Biolabs	K3389L
	New England	D21091
MluI-HF	Biolabs	R3198L
SureBlue Reserve tetramethylbenzidine substrate	KPL	Cat #5120-0081
TaqMan Fast Virus 1-Step Master Mix	ThermoFisher	4444434
QIAsymphony DSP Virus/Pathogen Midi Kit	Qiagen	937055
NucleoSpin Gel and PCR Clean-Up	Takara	740609.5
MEGAscript T7 Transcription Kit	ThermoFisher	AM1334
MEGAclear Transcription Clean-Up Kit	ThermoFisher	AM1908
Luciferase Cell Culture Lysis 5x Reagent	Promega	Cat# E1531
Background Reducing Ab Diluent		Cat# E1551 Cat# S3022
	Agilent	
PowerVision Poly-HRP anti-Rabbit IgG IHC Detection	Leica	Cat# PV6121

Systems		
Human ACE2 soluble protein	(Edwards et al.,	NA
Fidihari Field solutio protein	(2021)	1111
SARS-CoV-2 Spike S1+S2 ectodomain (ECD)	Sino Biological	Cat #40589-V08B1
SARS-CoV-2 Spike S2 ECD	Sino Biological	Cat #40590-V08B
SARS-CoV-2 Spike RBD from insect cell sf9	Sino Biological	Cat #40592-V08B
SARS-CoV-2 Spike RBD from mammalian cell 293	Sino Biological	Cat #40592-V08H
SARS-CoV Spike Protein DeltaTM	BEI Resources	Cat #NR-722
SARS-CoV WH20 Spike RBD	Sino Biological	Cat #40150-V08B2
SARS-CoV WH20 Spike S1	Sino Biological	Cat #40150-V08B1
MERS-CoV Spike S1+S2	Sino Biological	Cat #40069-V08B
MERS-CoV Spike S1	Sino Biological	Cat #40069-V08B1
MERS-CoV Spike S2	Sino Biological	Cat #40070-V08B
MERS-CoV Spike RBD	Sino Biological	Cat #40071-V08B1
SARS-CoV CL Protease protein	BEI Resources	Cat #30105
SARS-CoV Membrane (M) protein	BEI Resources	Cat #110705
SARS-CoV-2 Spike NTD	(Zhou et al., 2020b)	NA
SARS-CoV Spike RBD	(Hauser et al., 2020)	NA
MERS-CoV Spike RBD	(Hauser et al., 2020) (Hauser et al., 2020)	NA
SARS-CoV-2 Spike-2P	(Edwards et al.,	NA
SAR5-C0V-2 Spike-21	(Ldwards et al., 2021)	
SARS-CoV-2 Spike-HexaPro	(Edwards et al.,	NA
SARS-COV-2 Spike-Hexario	(Ldwards et al., 2021)	147 X
Critical Commercial Assays		
MILLIPLEX MAP Non-Human Primate	Millipore	Cat
Cytokine/Chemokine Panel, 25-analyte multiplex bead	r	#PRCYT2MAG40K
array		
Bright-Glo Luciferase Assay System	Promega	Cat #2650
Britelite Luminescence Reporter Gene Assay System	PerkinElmer Life	Cat #6066761
	Sciences	
Nano-Glo Luciferase Assay System	Promega	Cat #N1150
Deposited Data		
Structure of SARS-CoV-2 S protein in complex with	This paper	PDB 7LAA, EMD-
Receptor Binding Domain Ab DH1041		23246
Structure of SARS-CoV-2 S protein in complex with	This paper	PDB 7LD1, EMD-
Receptor Binding Domain Ab DH1047		23279
Structure of SARS-CoV-2 S protein in complex with N-	This paper	PDB 7LCN, EMD-
terminal domain Ab DH1050.1		23277
Structure of SARS-CoV-2 S protein in complex with N-	This paper	PDB 7LAB, EMD-
terminal domain Ab DH1052		23248
SARS-CoV-2 Spike Protein Trimer bound to DH1043	This paper	PDB 7LJR, EMD-
fab		23400
Negative stain EM structure of Ab DH1041 Fab in	This paper	EMD-22920
complex with SARS-CoV-2 Hexapro spike		
Negative stain EM structure of Ab DH1042 Fab in	This paper	EMD-22921
complex with SARS-CoV-2 2P spike		
Negative stain EM structure of Ab DH1043 Fab in	This paper	EMD-22923
complex with SARS-CoV-2 Hexapro spike		EMD 22020
Negative stain EM structure of Ab DH1044 Fab in	This paper	EMD-22929
complex with SARS-CoV-2 2P spike	This second	EMD 22020
Negative stain EM structure of Ab DH1045 Fab in	This paper	EMD-22930
complex with SARS-CoV-2 Hexapro spike	This paper	EMD 22022
Negative stain EM structure of Ab DH1047 Fab in	This paper	EMD-22933

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complex with SARS-CoV-2 Hexapro spike		
Negative stain EM structure of Ab DH1048 Fab in	This paper	EMD-22936
complex with SARS-CoV-2 Hexapro spike		
Negative stain EM structure of Ab DH1049 Fab in	This paper	EMD-22942
complex with SARS-CoV-2 2P spike		
Negative stain EM structure of Ab DH1050.1 Fab in	This paper	EMD-22944
complex with SARS-CoV-2 Hexapro spike		
Negative stain EM structure of Ab DH1050.2 Fab in	This paper	EMD-22945
complex with SARS-CoV-2 2P spike		
Negative stain EM structure of Ab DH1051 Fab in	This paper	EMD-22946
complex with SARS-CoV-2 Hexapro spike		
Negative stain EM structure of Ab DH1053 Fab in	This paper	EMD-22947
complex with SARS-CoV-2 2P spike in the 1-RBD-up		
state		
Negative stain EM structure of Ab DH1053 Fab in	This paper	EMD-22948
complex with SARS-CoV-2 2P spike in the 3-RBD-	rins pup ti	
down state		
Negative stain EM structure of Ab DH1054 Fab in	This paper	EMD-22951
complex with SARS-CoV-2 2P spike	rins puper	
Negative stain EM structure of Ab DH1055 Fab in	This paper	EMD-22952
complex with SARS-CoV-2 2P spike	This paper	EMID-22932
<u>^</u>	This paper	EMD-22953
Negative stain EM structure of Ab DH1056 Fab in	This paper	EMD-22935
complex with SARS-CoV-2 2P spike	T1.:	EMD 22055
Negative stain EM structure of Ab Fabs DH1043 and	This paper	EMD-22955
DH1051 in complex with SARS-CoV-2 2P spike		
Negative stain EM structure of Ab Fabs DH1041 and	This paper	EMD-22956
DH1051 in complex with SARS-CoV-2 2P spike		EN (D. 22055
Negative stain EM structure of Ab Fabs DH1043 and	This paper	EMD-22957
DH1047 in complex with SARS-CoV-2 2P spike		
Negative stain EM structure of Ab Fabs DH1047 and	This paper	EMD-22958
DH1051 in complex with SARS-CoV-2 2P spike		
Negative stain EM structure of Ab Fabs DH1045 and	This paper	EMD-22969
DH1050.1 in complex with SARS-CoV-2 2P spike		
Negative stain EM structure of Ab Fabs DH1043 and	This paper	EMD-22970
DH1050.1 in complex with SARS-CoV-2 2P spike		
Negative stain EM structure of Ab Fabs DH1041 and	This paper	EMD-22971
DH1047 in complex with SARS-CoV-2 2P spike		
Negative stain EM structure of Ab Fabs DH1050.1 and	This paper	EMD-22984
DH1053 in complex with SARS-CoV-2 2P spike		
Negative stain EM structure of Ab Fabs DH1043,	This paper	EMD-22985
DH1047 and DH1050.1 in complex with SARS-CoV-2		
2P spike		
Negative stain EM structure of Ab Fabs DH1043,	This paper	EMD-22986
DH1047 and DH1051 in complex with SARS-CoV-2 2P		
spike		
Experimental Models: Cell Lines		<b>.</b>
TZM-bl	NIH, ARRRP	Cat #8129
TZM-bl expressing FcyRI	(Perez et al., 2009)	NA
TZM-bl expressing FcyRIIa	(Perez et al., 2009)	NA
	· · · · · ·	
TZM-bl expressing FcγRIIb	(Perez et al., 2009)	NA
TZM-bl expressing FcγRIII	(Perez et al., 2009)	NA
Expi 293i	Invitrogen	Cat #14527
293T/ACE2	(Korber et al., 2020)	NA

Vero E6	ATCC	Cat# CRL-1586
Experimental Models: Organisms/Strains		
BALB/c mouse	Envigo	NA
HFH4-hACE2 transgenic mice	(Menachery et al.,	NA
	2016)	
Cynomolgus macaques	BioQUAL	NA
Oligonucleotides		
VH1 Leader-A 5'- ATGGACTGGACCTGGAGGAT -3'	Thermo Fisher	NA
(PCRa primer)	Scientific	
VH1 Leader-A 5'- ATGGACTGGACCTGGAGCAT -3'	Thermo Fisher	NA
(PCRa primer)	Scientific	
VH1 Leader-A 5'- ATGGACTGGACCTGGAGAAT -3'	Thermo Fisher	NA
(PCRa primer)	Scientific	
VH1 Leader-A 5'- GGTTCCTCTTTGTGGTGGC -3'	Thermo Fisher	NA
(PCRa primer)	Scientific	
VH1 Leader-A 5'- ATGGACTGGACCTGGAGGGT -3'	Thermo Fisher	NA
(PCRa primer)	Scientific	
VH1 Leader-A 5'- ATGGACTGGATTTGGAGGAT -3'	Thermo Fisher	NA
(PCRa primer)	Scientific	
VH1 Leader-A 5'- AGGTTCCTCTTTGTGGTGGCAG	Thermo Fisher	NA
-3' (PCRa primer)	Scientific	
VH1 Leader-A 5'-	Thermo Fisher	NA
ATGGACATACTTTGTTCCACGCTC -3' (PCRa	Scientific	
primer)		
VH1 Leader-A 5'-	Thermo Fisher	NA
ATGGACACACTTTGCTCCACGCT -3' (PCRa	Scientific	
primer)		
VH1 Leader-A 5'-	Thermo Fisher	NA
ATGGACACACTTTGCTACACACTC -3' (PCRa	Scientific	
primer)		
VH1 Leader-A 5'- CCGACGGGGAATTCTCACAG -3'	Thermo Fisher	NA
(PCRa primer)	Scientific	
VH1 Leader-A 5'-	Thermo Fisher	NA
CTGTTATCCTTTGGGTGTCTGCAC -3' (PCRa	Scientific	
VH1 Leader-A 5'- GGTGGCATTGGAGGGAATGTT -	Thermo Fisher Scientific	NA
3' (PCRa primer) VH1 Leader-A 5'- CGAYGACCACGTTCCCATCT -3'	Thermo Fisher	NA
(PCRa primer)	Scientific	INA
VH1 Leader-A 5'- TAGTCCTTGACCAGGCAGC -3'	Thermo Fisher	NA
(PCRa primer)	Scientific	NA (
VH1 Leader-A 5'- TAAAAGGTGTCCAGTGT -3'	Thermo Fisher	NA
(PCRa primer)	Scientific	na –
VH1 Leader-A 5'- TAAGAGGTGTCCAGTGT -3'	Thermo Fisher	NA
(PCRa primer)	Scientific	1111
VH1 Leader-A 5'- TAGAAGGTGTCCAGTGT -3'	Thermo Fisher	NA
(PCRa primer)	Scientific	
VH1 Leader-A 5'- TACAAGGTGTCCAGTGT -3'	Thermo Fisher	NA
(PCRa primer)	Scientific	
VH1 Leader-A 5'- TTAAAGCTGTCCAGTGT -3'	Thermo Fisher	NA
(PCRa primer)	Scientific	
VH1 Leader-A 5'- ATGAAACATCTGTGGTTCTT -3'	Thermo Fisher	NA
(PCRa primer)	Scientific	
	Scientific	1

VH1 Leader-A 5'- TTCTCCAAGGAGTCTGT -3'	Thermo Fisher	NA
(PCRa primer)	Scientific	
VH1 Leader-A 5'-	Thermo Fisher	NA
GCTATTTTTAAAGGTGTCCAGTGT -3' (PCRa	Scientific	
VH1 Leader-A 5'- ATGAAACACCTGTGGTTCTTCC	Thermo Fisher	NA
-3' (PCRa primer)	Scientific	
VH1 Leader-A 5'- ATGAAACACCTGTGGTTCTT -3'	Thermo Fisher	NA
(PCRa primer)	Scientific	
VH1 Leader-A 5'- ATGAAGCACCTGTGGTTCTT -3'	Thermo Fisher	NA
(PCRa primer)	Scientific	
VH1 Leader-A 5'- CCTCCACAGTGAGAGTCTG -3'	Thermo Fisher	NA
(PCRa primer)	Scientific	
VH1 Leader-A 5'- ATGTCTGTCTCCTTCCTCATC -3'	Thermo Fisher	NA
(PCRa primer)	Scientific	
VH1 Leader-A 5'- GGCAGCAGCAACAGGTGCCCA -	Thermo Fisher	NA
3' (PCRa primer)	Scientific	
VH1 Leader-A 5'- GCTCAGCTCCTGGGGCT -3'	Thermo Fisher	NA
(PCRa primer)	Scientific	
VH1 Leader-A 5'- GGAARCCCCAGCDCAGC -3'	Thermo Fisher	NA
(PCRa primer)	Scientific	
VH1 Leader-A 5'- CTSTTSCTYTGGATCTCTG -3'	Thermo Fisher	NA
(PCRa primer)	Scientific	
VH1 Leader-A 5'- CTSCTGCTCTGGGYTCC -3'	Thermo Fisher	NA
(PCRa primer)	Scientific	
VH1 Leader-A 5'- GAGGCAGTTCCAGATTTCAA -3'	Thermo Fisher	NA
(PCRa primer)	Scientific	
VH1 Leader-A 5'- CCTGGGCCCAGTCTGTG -3'	Thermo Fisher	NA
(PCRa primer)	Scientific	1111
VH1 Leader-A 5'- CTCCTCASYCTCCTCACT -3'	Thermo Fisher	NA
(PCRa primer)	Scientific	1474
VH1 Leader-A 5'- GGCCTCCTATGWGCTGAC -3'	Thermo Fisher	NA
(PCRa primer)	Scientific	INA
VH1 Leader-A 5'- GTTCTGTGGTTTCTTCTGAGCTG	Thermo Fisher	NA
-3' (PCRa primer)	Scientific	INA
VH1 Leader-A 5'- ACAGGGTCTCTCTCCCAG -3'	Thermo Fisher	NA
		NA
(PCRa primer)	Scientific	
VH1 Leader-A 5'- ACAGGTCTCTGTGCTCTGC -3'	Thermo Fisher	NA
(PCRa primer)	Scientific	
VH1 Leader-A 5'- CCCTCTCSCAGSCTGTG -3'	Thermo Fisher	NA
(PCRa primer)	Scientific	
VH1 Leader-A 5'- TCTTGGGCCAATTTTATGC -3'	Thermo Fisher	NA
(PCRa primer)	Scientific	
VH1 Leader-A 5'- ATTCYCAGRCTGTGGTGAC -3'	Thermo Fisher	NA
(PCRa primer)	Scientific	
VH1 Leader-A 5'- CAGTGGTCCAGGCAGGG -3'	Thermo Fisher	NA
(PCRa primer)	Scientific	
VH1 Leader-A 5'- AGGCCACTGTCACAGCT -3'	Thermo Fisher	NA
(PCRa primer)	Scientific	
VH1-Int tag 5'-	Thermo Fisher	NA
CTGGGTTCCAGGTTCCACTGGTGACCAGGTGCA	Scientific	
GCTGGTRCAGTCTGGG -3' (PCRb primer)		
VH2-Int tag 5'-	Thermo Fisher	NA
CTGGGTTCCAGGTTCCACTGGTGACCAGRGCAC	Scientific	
<u></u>		

CTTGARGGAGTCTGGTCC -3' (PCRb primer)		
VH3-Int tag 5'-	Thermo Fisher	NA
CTGGGTTCCAGGTTCCACTGGTGACGAGGTKCA	Scientific	
GCTGGTGGAGTCTGGG -3' (PCRb primer)		
VH4-Int tag 5'-	Thermo Fisher	NA
CTGGGTTCCAGGTTCCACTGGTGACCAGGTGCA	Scientific	
GCTGCAGGAGTCGG -3' (PCRb primer)	~	
VH5-Int tag 5'-	Thermo Fisher	NA
CTGGGTTCCAGGTTCCACTGGTGACGARGTGCA	Scientific	
GCTGGTGCAGTCTGGAG -3' (PCRb primer)	Selentific	
VH6-Int tag 5'-	Thermo Fisher	NA
CTGGGTTCCAGGTTCCACTGGTGACCAGGTACA	Scientific	
GCTGCAGCAGTCAGGTCC -3' (PCRb primer)	berentific	
IgG-int 5'-	Thermo Fisher	NA
GGGCCGCTGTGCCCCCAGAGGTGCTCYTGGA -3'	Scientific	11/1
(PCRb primer)	Scientific	
IgM-int 5'-	Thermo Fisher	NA
GGGCCGCTGTGCCCCCAGAGGTGGAATTCTCAC	Scientific	INA
AGGAGACGAGG -3' (PCRb primer)	Scientific	
	Thermo Fisher	NA
IgD-int 5'-		INA
GGGCCGCTGTGCCCCCAGAGGTGTGTCTGCACC	Scientific	
CTGATATGATGG -3' (PCRb primer)		
IgA1-int 5'-	Thermo Fisher	NA
GGGCCGCTGTGCCCCCAGAGGTGCTGGTGCTGC	Scientific	
AGAGGCTCAG -3' (PCRb primer)		
IgA2-int 5'-	Thermo Fisher	NA
GGGCCGCTGTGCCCCCAGAGGTGCTGGTGCTGT	Scientific	
CGAGGCTCAG -3' (PCRb primer)		
VK1-Int tag 5'-	Thermo Fisher	NA
CTGGGTTCCAGGTTCCACTGGTGACGACATCCA	Scientific	
GWTGACCCAGTCTC -3' (PCRb primer)		
VK2-Int tag 5'-	Thermo Fisher	NA
CTGGGTTCCAGGTTCCACTGGTGACGATATTGT	Scientific	
GATGACCCAGWCTCCAC -3' (PCRb primer)		
VK3-Int tag 5'-	Thermo Fisher	NA
CTGGGTTCCAGGTTCCACTGGTGACGAAATTGT	Scientific	
GTTGACRCAGTCTCCA -3' (PCRb primer)		
VK4-Int tag 5'-	Thermo Fisher	NA
CTGGGTTCCAGGTTCCACTGGTGACGACATCGT	Scientific	
GATGACCCAGTCTC -3' (PCRb primer)		
VK5-Int tag 5'-	Thermo Fisher	NA
CTGGGTTCCAGGTTCCACTGGTGACGAAACGAC	Scientific	
ACTCACGCAGTCTC -3' (PCRb primer)		
VK6-Int tag 5'-	Thermo Fisher	NA
CTGGGTTCCAGGTTCCACTGGTGACGAAATTGT	Scientific	
GCTGACWCAGTCTCCA -3' (PCRb primer)		
VK7-Int tag 5'-	Thermo Fisher	NA
CTGGGTTCCAGGTTCCACTGGTGACGACATTGT	Scientific	
GCTGACCCAGTCT -3' (PCRb primer)		
CK-int 5'- GGGAAGATGAAGACAGATGGT -3'	Thermo Fisher	NA
(PCRb primer)	Scientific	
VL1-Int tag 5'-	Thermo Fisher	NA
CTGGGTTCCAGGTTCCACTGGTGACCAGTCTGT	Scientific	
GYTGACKCAGCC -3' (PCRb primer)		
	1	

VI 2 Int too 5!	Thermo Fisher	NA
VL2-Int tag 5'- CTGGGTTCCAGGTTCCACTGGTGACCAGTCTGC	Scientific	INA
	Scientific	
CCTGACTCAGCC -3' (PCRb primer)	The sum of <b>D</b> ' all and	NA
VL3-Int tag 5'- CTGGGTTCCAGGTTCCACTGGTGACTCYTATGA	Thermo Fisher Scientific	NA
	Scientific	
GCTGACWCAGCCAC -3' (PCRb primer)		
VL31-Int tag 5'-	Thermo Fisher	NA
CTGGGTTCCAGGTTCCACTGGTGACTCTTCTGA	Scientific	
GCTGACTCAGGACCC -3' (PCRb primer)		
VL4ab-Int tag 5'- CTGGGTTCCAGGTTCCACTGGTGACCAGCYTGT	Thermo Fisher	NA
	Scientific	
GCTGACTCAATC -3' (PCRb primer)		
VL4c-Int tag 5'-	Thermo Fisher	NA
CTGGGTTCCAGGTTCCACTGGTGACCTGCCTGT	Scientific	P
GCTGACTCAGC -3' (PCRb primer)	<b>T</b> 1 <b>T</b> ' 1	
VL5,9-Int tag 5'-	Thermo Fisher	NA
CTGGGTTCCAGGTTCCACTGGTGACCAGSCTGT	Scientific	
GCTGACTCAGCC -3' (PCRb primer)		
VL6-Int tag 5'-	Thermo Fisher	NA
CTGGGTTCCAGGTTCCACTGGTGACAATTTTAT	Scientific	
GCTGACTCAGCCCCACT -3' (PCRb primer)		
VL7,8-Int tag 5'-	Thermo Fisher	NA
CTGGGTTCCAGGTTCCACTGGTGACCAGRCTGT	Scientific	
GGTGACYCAGGAG -3' (PCRb primer)		
VL10-Int tag 5'-	Thermo Fisher	NA
CTGGGTTCCAGGTTCCACTGGTGACCAGGCAGG	Scientific	
GCWGACTCAG -3' (PCRb primer)		
CL-int 5'- GGGYGGGAACAGAGTGACC -3' (PCRb	Thermo Fisher	NA
primer)	Scientific	
VH_Tag fwd seq 5'-	Thermo Fisher	NA
	Scientific	
CTGGGTTCCAGGTTCCACTGGTGAC -3'	Scientific	
(Sequencing primer)		
(Sequencing primer) CK_int 5'- GGGAAGATGAAGACAGATGGT -3'	Thermo Fisher	NA
(Sequencing primer) CK_int 5'- GGGAAGATGAAGACAGATGGT -3' (Sequencing primer)		
(Sequencing primer)CK_int 5'- GGGAAGATGAAGACAGATGGT -3'(Sequencing primer)CL_int 5'- GGGYGGGAACAGAGTGACC -3'	Thermo Fisher Scientific Thermo Fisher	NA NA
(Sequencing primer) CK_int 5'- GGGAAGATGAAGACAGATGGT -3' (Sequencing primer) CL_int 5'- GGGYGGGAACAGAGTGACC -3' (Sequencing primer)	Thermo Fisher Scientific Thermo Fisher Scientific	NA
(Sequencing primer)CK_int 5'- GGGAAGATGAAGACAGATGGT -3'(Sequencing primer)CL_int 5'- GGGYGGGAACAGAGTGACC -3'(Sequencing primer)HV13221H_R474 5'- GCTGTGCCCCCAGAGGTG -3'	Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher	
(Sequencing primer) CK_int 5'- GGGAAGATGAAGACAGATGGT -3' (Sequencing primer) CL_int 5'- GGGYGGGAACAGAGTGACC -3' (Sequencing primer) HV13221H_R474 5'- GCTGTGCCCCCAGAGGTG -3' (Sequencing primer)	Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific	NA NA
(Sequencing primer) CK_int 5'- GGGAAGATGAAGACAGATGGT -3' (Sequencing primer) CL_int 5'- GGGYGGGAACAGAGTGACC -3' (Sequencing primer) HV13221H_R474 5'- GCTGTGCCCCCAGAGGTG -3' (Sequencing primer) VH1 Leader-A 5'- ATGGACTGGACCTGGAGGAT -3'	Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher	NA
(Sequencing primer) CK_int 5'- GGGAAGATGAAGACAGATGGT -3' (Sequencing primer) CL_int 5'- GGGYGGGAACAGAGTGACC -3' (Sequencing primer) HV13221H_R474 5'- GCTGTGCCCCCAGAGGTG -3' (Sequencing primer)	Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific	NA NA
(Sequencing primer) CK_int 5'- GGGAAGATGAAGACAGATGGT -3' (Sequencing primer) CL_int 5'- GGGYGGGAACAGAGTGACC -3' (Sequencing primer) HV13221H_R474 5'- GCTGTGCCCCCAGAGGTG -3' (Sequencing primer) VH1 Leader-A 5'- ATGGACTGGACCTGGAGGAT -3'	Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher	NA NA
(Sequencing primer)CK_int 5'- GGGAAGATGAAGACAGATGGT -3'(Sequencing primer)CL_int 5'- GGGYGGGAACAGAGTGACC -3'(Sequencing primer)HV13221H_R474 5'- GCTGTGCCCCCAGAGGTG -3'(Sequencing primer)VH1 Leader-A 5'- ATGGACTGGACCTGGAGGAT -3'(PCRa primer)VH1 Leader-A 5'- ATGGACTGGACCTGGAGCAT -3'(PCRa primer)	Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific	NA NA NA
(Sequencing primer) CK_int 5'- GGGAAGATGAAGACAGATGGT -3' (Sequencing primer) CL_int 5'- GGGYGGGAACAGAGTGACC -3' (Sequencing primer) HV13221H_R474 5'- GCTGTGCCCCCAGAGGTG -3' (Sequencing primer) VH1 Leader-A 5'- ATGGACTGGACCTGGAGGAT -3' (PCRa primer) VH1 Leader-A 5'- ATGGACTGGACCTGGAGCAT -3'	Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher	NA NA NA
(Sequencing primer)CK_int 5'- GGGAAGATGAAGACAGATGGT -3'(Sequencing primer)CL_int 5'- GGGYGGGAACAGAGTGACC -3'(Sequencing primer)HV13221H_R474 5'- GCTGTGCCCCCAGAGGTG -3'(Sequencing primer)VH1 Leader-A 5'- ATGGACTGGACCTGGAGGAT -3'(PCRa primer)VH1 Leader-A 5'- ATGGACTGGACCTGGAGCAT -3'(PCRa primer)	Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific	NA NA NA NA
(Sequencing primer)CK_int 5'- GGGAAGATGAAGACAGATGGT -3'(Sequencing primer)CL_int 5'- GGGYGGGAACAGAGTGACC -3'(Sequencing primer)HV13221H_R474 5'- GCTGTGCCCCCAGAGGTG -3'(Sequencing primer)VH1 Leader-A 5'- ATGGACTGGACCTGGAGGAT -3'(PCRa primer)VH1 Leader-A 5'- ATGGACTGGACCTGGAGCAT -3'(PCRa primer)VH1 Leader-A 5'- ATGGACTGGACCTGGAGCAT -3'(PCRa primer)VH1 Leader-A 5'- ATGGACTGGACCTGGAGCAT -3'	Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher	NA NA NA NA
(Sequencing primer) CK_int 5'- GGGAAGATGAAGACAGATGGT -3' (Sequencing primer) CL_int 5'- GGGYGGGAACAGAGTGACC -3' (Sequencing primer) HV13221H_R474 5'- GCTGTGCCCCCAGAGGTG -3' (Sequencing primer) VH1 Leader-A 5'- ATGGACTGGACCTGGAGGAT -3' (PCRa primer) VH1 Leader-A 5'- ATGGACTGGACCTGGAGCAT -3' (PCRa primer) VH1 Leader-A 5'- ATGGACTGGACCTGGAGAAT -3' (PCRa primer)	Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific	NA NA NA NA NA
(Sequencing primer)CK_int 5'- GGGAAGATGAAGACAGATGGT -3'(Sequencing primer)CL_int 5'- GGGYGGGAACAGAGTGACC -3'(Sequencing primer)HV13221H_R474 5'- GCTGTGCCCCCAGAGGTG -3'(Sequencing primer)VH1 Leader-A 5'- ATGGACTGGACCTGGAGGAT -3'(PCRa primer)VH1 Leader-A 5'- ATGGACTGGACCTGGAGCAT -3'(PCRa primer)VH1 Leader-A 5'- ATGGACTGGACCTGGAGCAT -3'(PCRa primer)VH1 Leader-A 5'- ATGGACTGGACCTGGAGAAT -3'(PCRa primer)VH1 Leader-A 5'- ATGGACTGGACCTGGAGCAT -3'(PCRa primer)VH1 Leader-A 5'- ATGGACTGGACCTGGAGCAT -3'(PCRa primer)VH1 Leader-A 5'- ATGGACTGGACCTGGAGAAT -3'(PCRa primer)VH1 Leader-A 5'- GGTTCCTCTTTGTGGTGGC -3'	Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher	NA NA NA NA NA
(Sequencing primer)CK_int 5'- GGGAAGATGAAGACAGATGGT -3'(Sequencing primer)CL_int 5'- GGGYGGGAACAGAGTGACC -3'(Sequencing primer)HV13221H_R474 5'- GCTGTGCCCCCAGAGGTG -3'(Sequencing primer)VH1 Leader-A 5'- ATGGACTGGACCTGGAGGAT -3'(PCRa primer)VH1 Leader-A 5'- ATGGACTGGACCTGGAGCAT -3'(PCRa primer)VH1 Leader-A 5'- ATGGACTGGACCTGGAGCAT -3'(PCRa primer)VH1 Leader-A 5'- ATGGACTGGACCTGGAGAAT -3'(PCRa primer)VH1 Leader-A 5'- ATGGACTGGACCTGGAGCAT -3'(PCRa primer)VH1 Leader-A 5'- ATGGACTGGACCTGGAGCAT -3'(PCRa primer)VH1 Leader-A 5'- ATGGACTGGACCTGGAGAAT -3'(PCRa primer)	Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific	NA NA NA NA NA NA
<ul> <li>(Sequencing primer)</li> <li>CK_int 5'- GGGAAGATGAAGACAGATGGT -3'</li> <li>(Sequencing primer)</li> <li>CL_int 5'- GGGYGGGAACAGAGTGACC -3'</li> <li>(Sequencing primer)</li> <li>HV13221H_R474 5'- GCTGTGCCCCCAGAGGTG -3'</li> <li>(Sequencing primer)</li> <li>VH1 Leader-A 5'- ATGGACTGGACCTGGAGGAT -3'</li> <li>(PCRa primer)</li> <li>VH1 Leader-A 5'- ATGGACTGGACCTGGAGGAAT -3'</li> <li>(PCRa primer)</li> <li>VH1 Leader-A 5'- ATGGACTGGACCTGGAGAAT -3'</li> <li>(PCRa primer)</li> <li>VH1 Leader-A 5'- ATGGACTGGACCTGGAGAAT -3'</li> <li>(PCRa primer)</li> <li>VH1 Leader-A 5'- ATGGACTGGACCTGGAGGAT -3'</li> <li>(PCRa primer)</li> <li>VH1 Leader-A 5'- ATGGACTGGACCTGGAGGAT -3'</li> </ul>	Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher	NA NA NA NA NA NA
<ul> <li>(Sequencing primer)</li> <li>CK_int 5'- GGGAAGATGAAGACAGATGGT -3'</li> <li>(Sequencing primer)</li> <li>CL_int 5'- GGGYGGGAACAGAGTGACC -3'</li> <li>(Sequencing primer)</li> <li>HV13221H_R474 5'- GCTGTGCCCCCAGAGGTG -3'</li> <li>(Sequencing primer)</li> <li>VH1 Leader-A 5'- ATGGACTGGACCTGGAGGAT -3'</li> <li>(PCRa primer)</li> <li>VH1 Leader-A 5'- ATGGACTGGACCTGGAGGAAT -3'</li> <li>(PCRa primer)</li> <li>VH1 Leader-A 5'- ATGGACTGGACCTGGAGAAT -3'</li> <li>(PCRa primer)</li> <li>VH1 Leader-A 5'- ATGGACTGGACCTGGAGAAT -3'</li> <li>(PCRa primer)</li> <li>VH1 Leader-A 5'- ATGGACTGGACCTGGAGGAT -3'</li> <li>(PCRa primer)</li> <li>VH1 Leader-A 5'- ATGGACTGGACCTGGAGGAT -3'</li> <li>(PCRa primer)</li> <li>VH1 Leader-A 5'- ATGGACTGGACCTGGAGGGT -3'</li> <li>(PCRa primer)</li> </ul>	Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific	NA NA NA NA NA NA NA
<ul> <li>(Sequencing primer)</li> <li>CK_int 5'- GGGAAGATGAAGACAGATGGT -3'</li> <li>(Sequencing primer)</li> <li>CL_int 5'- GGGYGGGAACAGAGTGACC -3'</li> <li>(Sequencing primer)</li> <li>HV13221H_R474 5'- GCTGTGCCCCCAGAGGTG -3'</li> <li>(Sequencing primer)</li> <li>VH1 Leader-A 5'- ATGGACTGGACCTGGAGGAT -3'</li> <li>(PCRa primer)</li> <li>VH1 Leader-A 5'- ATGGACTGGACCTGGAGGAAT -3'</li> <li>(PCRa primer)</li> <li>VH1 Leader-A 5'- ATGGACTGGACCTGGAGAAT -3'</li> <li>(PCRa primer)</li> <li>VH1 Leader-A 5'- ATGGACTGGACCTGGAGCAT -3'</li> <li>(PCRa primer)</li> <li>VH1 Leader-A 5'- ATGGACTGGACCTGGAGGAT -3'</li> <li>(PCRa primer)</li> <li>VH1 Leader-A 5'- ATGGACTGGACCTGGAGGGT -3'</li> </ul>	Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher	NA NA NA NA NA NA NA
<ul> <li>(Sequencing primer)</li> <li>CK_int 5'- GGGAAGATGAAGACAGATGGT -3'</li> <li>(Sequencing primer)</li> <li>CL_int 5'- GGGYGGGAACAGAGTGACC -3'</li> <li>(Sequencing primer)</li> <li>HV13221H_R474 5'- GCTGTGCCCCCAGAGGTG -3'</li> <li>(Sequencing primer)</li> <li>VH1 Leader-A 5'- ATGGACTGGACCTGGAGGAT -3'</li> <li>(PCRa primer)</li> <li>VH1 Leader-A 5'- ATGGACTGGACCTGGAGGAT -3'</li> <li>(PCRa primer)</li> <li>VH1 Leader-A 5'- ATGGACTGGACCTGGAGAAT -3'</li> <li>(PCRa primer)</li> <li>VH1 Leader-A 5'- ATGGACTGGACCTGGAGAAT -3'</li> <li>(PCRa primer)</li> <li>VH1 Leader-A 5'- ATGGACTGGACCTGGAGGAT -3'</li> <li>(PCRa primer)</li> <li>VH1 Leader-A 5'- ATGGACTGGACCTGGAGGGT -3'</li> <li>(PCRa primer)</li> </ul>	Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific	NA NA NA NA NA NA NA NA
<ul> <li>(Sequencing primer)</li> <li>CK_int 5'- GGGAAGATGAAGACAGATGGT -3'</li> <li>(Sequencing primer)</li> <li>CL_int 5'- GGGYGGGAACAGAGTGACC -3'</li> <li>(Sequencing primer)</li> <li>HV13221H_R474 5'- GCTGTGCCCCCAGAGGTG -3'</li> <li>(Sequencing primer)</li> <li>VH1 Leader-A 5'- ATGGACTGGACCTGGAGGAT -3'</li> <li>(PCRa primer)</li> <li>VH1 Leader-A 5'- ATGGACTGGACCTGGAGGAAT -3'</li> <li>(PCRa primer)</li> <li>VH1 Leader-A 5'- ATGGACTGGACCTGGAGAAT -3'</li> <li>(PCRa primer)</li> <li>VH1 Leader-A 5'- ATGGACTGGACCTGGAGGAT -3'</li> <li>(PCRa primer)</li> <li>VH1 Leader-A 5'- ATGGACTGGACCTGGAGGAT -3'</li> <li>(PCRa primer)</li> <li>VH1 Leader-A 5'- ATGGACTGGACCTGGAGGGT -3'</li> <li>(PCRa primer)</li> <li>VH1 Leader-A 5'- ATGGACTGGACCTGGAGGAT -3'</li> <li>(PCRa primer)</li> <li>VH1 Leader-A 5'- ATGGACTGGATTGGAGGAT -3'</li> <li>(PCRa primer)</li> <li>VH1 Leader-A 5'- ATGGACTGGATTTGGAGGAT -3'</li> <li>(PCRa primer)</li> </ul>	Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific	NA NA NA NA NA NA NA NA NA
(Sequencing primer) CK_int 5'- GGGAAGATGAAGACAGATGGT -3' (Sequencing primer) CL_int 5'- GGGYGGGAACAGAGTGACC -3' (Sequencing primer) HV13221H_R474 5'- GCTGTGCCCCCAGAGGTG -3' (Sequencing primer) VH1 Leader-A 5'- ATGGACTGGACCTGGAGGAT -3' (PCRa primer) VH1 Leader-A 5'- ATGGACTGGACCTGGAGCAT -3' (PCRa primer) VH1 Leader-A 5'- ATGGACTGGACCTGGAGAAT -3' (PCRa primer) VH1 Leader-A 5'- ATGGACTGGACCTGGAGGAT -3' (PCRa primer) VH1 Leader-A 5'- ATGGACTGGACCTGGAGGGT -3' (PCRa primer) VH1 Leader-A 5'- ATGGACTGGACCTGGAGGGT -3' (PCRa primer) VH1 Leader-A 5'- ATGGACTGGATTTGGAGGAT -3' (PCRa primer)	Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific Thermo Fisher Scientific	NA NA NA NA NA NA NA NA

primer)		
VH1 Leader-A 5'-	Thermo Fisher	NA
ATGGACACACTTTGCTCCACGCT -3' (PCRa	Scientific	
primer)		
VH1 Leader-A 5'-	Thermo Fisher	NA
ATGGACACACTTTGCTACACACTC -3' (PCRa	Scientific	
primer)		
VH1 Leader-A 5'- CCGACGGGGAATTCTCACAG -3'	Thermo Fisher	NA
(PCRa primer)	Scientific	
VH1 Leader-A 5'-	Thermo Fisher	NA
CTGTTATCCTTTGGGTGTCTGCAC -3' (PCRa	Scientific	
primer)		
VH1 Leader-A 5'- GGTGGCATTGGAGGGAATGTT -	Thermo Fisher	NA
3' (PCRa primer)	Scientific	1 11 1
VH1 Leader-A 5'- CGAYGACCACGTTCCCATCT -3'	Thermo Fisher	NA
(PCRa primer)	Scientific	NA
VH1 Leader-A 5'- TAGTCCTTGACCAGGCAGC -3'	Thermo Fisher	NA
(PCRa primer)	Scientific	INA
VH1 Leader-A 5'- TAAAAGGTGTCCAGTGT -3'	Thermo Fisher	NA
		NA
(PCRa primer) VH1 Leader-A 5'- TAAGAGGTGTCCAGTGT -3'	Scientific Eicher	NT A
	Thermo Fisher	NA
(PCRa primer)	Scientific	
VH1 Leader-A 5'- TAGAAGGTGTCCAGTGT -3'	Thermo Fisher	NA
(PCRa primer)	Scientific	
VH1 Leader-A 5'- TACAAGGTGTCCAGTGT -3'	Thermo Fisher	NA
(PCRa primer)	Scientific	
VH1 Leader-A 5'- TTAAAGCTGTCCAGTGT -3'	Thermo Fisher	NA
(PCRa primer)	Scientific	
VH1 Leader-A 5'- ATGAAACATCTGTGGTTCTT -3'	Thermo Fisher	NA
(PCRa primer)	Scientific	
VH1 Leader-A 5'- TTCTCCAAGGAGTCTGT -3'	Thermo Fisher	NA
(PCRa primer)	Scientific	
VH1 Leader-A 5'-	Thermo Fisher	NA
GCTATTTTTAAAGGTGTCCAGTGT -3' (PCRa	Scientific	
primer)		
VH1 Leader-A 5'- ATGAAACACCTGTGGTTCTTCC	Thermo Fisher	NA
-3' (PCRa primer)	Scientific	
VH1 Leader-A 5'- ATGAAACACCTGTGGTTCTT -3'	Thermo Fisher	NA
(PCRa primer)	Scientific	
VH1 Leader-A 5'- ATGAAGCACCTGTGGTTCTT -3'	Thermo Fisher	NA
(PCRa primer)	Scientific	
VH1 Leader-A 5'- CCTCCACAGTGAGAGTCTG -3'	Thermo Fisher	NA
(PCRa primer)	Scientific	
VH1 Leader-A 5'- ATGTCTGTCTCCTTCCTCATC -3'	Thermo Fisher	NA
(PCRa primer)	Scientific	1 11 1
VH1 Leader-A 5'- GGCAGCAGCAACAGGTGCCCA -	Thermo Fisher	NA
3' (PCRa primer)	Scientific	1 12 E
VH1 Leader-A 5'- GCTCAGCTCCTGGGGCT -3'	Thermo Fisher	NA
(PCRa primer)	Scientific	
VH1 Leader-A 5'- GGAARCCCCAGCDCAGC -3'	Thermo Fisher	NA
(PCRa primer)	Scientific	
VH1 Leader-A 5'- CTSTTSCTYTGGATCTCTG -3'	Thermo Fisher	NA
	Scientific	INA
(PCRa primer)		NA
VH1 Leader-A 5'- CTSCTGCTCTGGGYTCC -3'	Thermo Fisher	NA

ner) Scientific	
er-A 5'- GAGGCAGTTCCAGATTTCAA -3' Thermo Fisher NA	
ner) Scientific er-A 5'- CCTGGGCCCAGTCTGTG -3' Thermo Fisher NA	
ner) Scientific	
er-A 5'- CTCCTCASYCTCCTCACT -3' Thermo Fisher NA	
ner) Scientific	
er-A 5'- GGCCTCCTATGWGCTGAC -3' Thermo Fisher NA	
ner) Scientific	
er-A 5'- GTTCTGTGGGTTTCTTCTGAGCTG Thermo Fisher NA	
primer) Scientific	
er-A 5'- ACAGGGTCTCTCCCAG -3' Thermo Fisher NA	
ner) Scientific	
er-A 5'- ACAGGTCTCTGTGCTCTGC -3' Thermo Fisher NA	
ner) Scientific	
er-A 5'- CCCTCTCSCAGSCTGTG -3' Thermo Fisher NA	
ner) Scientific	
er-A 5'- TCTTGGGCCAATTTTATGC -3' Thermo Fisher NA	
ner) Scientific	
er-A 5'- ATTCYCAGRCTGTGGTGAC -3' Thermo Fisher NA	
ner) Scientific	
er-A 5'- CAGTGGTCCAGGCAGGG -3' Thermo Fisher NA	
ner) Scientific	
er-A 5'- AGGCCACTGTCACAGCT -3' Thermo Fisher NA	
ner) Scientific	
g 5'- Thermo Fisher NA	
TCCAGGTTCCACTGGTGACCAGGTGCA Scientific	
RCAGTCTGGG -3' (PCRb primer)	
g 5'- Thermo Fisher NA	
TCCAGGTTCCACTGGTGACCAGRGCAC Scientific	
GGAGTCTGGTCC -3' (PCRb primer)	
g 5'- Thermo Fisher NA	
TCCAGGTTCCACTGGTGACGAGGTKCA Scientific	
GGAGTCTGGG -3' (PCRb primer)	
g 5'- NA	
TCCAGGTTCCACTGGTGACCAGGTGCA Scientific	
GGAGTCGG -3' (PCRb primer)	
g 5'- Thermo Fisher NA	
TCCAGGTTCCACTGGTGACGARGTGCA Scientific	
GCAGTCTGGAG -3' (PCRb primer)	
g 5'- Thermo Fisher NA	
TCCAGGTTCCACTGGTGACCAGGTACA Scientific	
GCAGTCAGGTCC -3' (PCRb primer)	
Thermo Fisher NA	
CTGTGCCCCAGAGGTGCTCYTGGA -3' Scientific	
ner)	
Thermo Fisher NA	_
CTGTGCCCCCAGAGGTGGAATTCTCAC Scientific	
CGAGG -3' (PCRb primer)	
Thermo Fisher NA	
CTGTGCCCCAGAGGTGTGTCTGCACC Scientific	
CGATGG -3' (PCRb primer)	
- Thermo Fisher NA	
CTGTGCCCCAGAGGTGCTGGTGCTGC Scientific	
CGAGG -3' (PCRb primer)     Thermo Fisher     NA       CTGTGCCCCCAGAGGTGTGTCTGCACC     Scientific     NA       CGATGG -3' (PCRb primer)     Hermo Fisher     NA	
	NA

AGAGGCTCAG -3' (PCRb primer)		
IgA2-int 5'-	Thermo Fisher	NA
GGGCCGCTGTGCCCCCAGAGGTGCTGGTGCTGT	Scientific	
CGAGGCTCAG -3' (PCRb primer)	~	
VK1-Int tag 5'-	Thermo Fisher	NA
CTGGGTTCCAGGTTCCACTGGTGACGACATCCA	Scientific	
GWTGACCCAGTCTC -3' (PCRb primer)	belentine	
VK2-Int tag 5'-	Thermo Fisher	NA
CTGGGTTCCAGGTTCCACTGGTGACGATATTGT	Scientific	na –
GATGACCCAGWCTCCAC -3' (PCRb primer)	Scientific	
VK3-Int tag 5'-	Thermo Fisher	NA
CTGGGTTCCAGGTTCCACTGGTGACGAAATTGT	Scientific	NA
	Scientific	
GTTGACRCAGTCTCCA -3' (PCRb primer)		
VK4-Int tag 5'-	Thermo Fisher	NA
CTGGGTTCCAGGTTCCACTGGTGACGACATCGT	Scientific	K
GATGACCCAGTCTC -3' (PCRb primer)		
VK5-Int tag 5'-	Thermo Fisher	NA
CTGGGTTCCAGGTTCCACTGGTGACGAAACGAC	Scientific	
ACTCACGCAGTCTC -3' (PCRb primer)		
VK6-Int tag 5'-	Thermo Fisher	NA
CTGGGTTCCAGGTTCCACTGGTGACGAAATTGT	Scientific	
GCTGACWCAGTCTCCA -3' (PCRb primer)		
VK7-Int tag 5'-	Thermo Fisher	NA
CTGGGTTCCAGGTTCCACTGGTGACGACATTGT	Scientific	
GCTGACCCAGTCT -3' (PCRb primer)		
CK-int 5'- GGGAAGATGAAGACAGATGGT -3'	Thermo Fisher	NA
(PCRb primer)	Scientific	
VL1-Int tag 5'-	Thermo Fisher	NA
CTGGGTTCCAGGTTCCACTGGTGACCAGTCTGT	Scientific	
GYTGACKCAGCC -3' (PCRb primer)		
VL2-Int tag 5'-	Thermo Fisher	NA
CTGGGTTCCAGGTTCCACTGGTGACCAGTCTGC	Scientific	
CCTGACTCAGCC -3' (PCRb primer)		
VL3-Int tag 5'-	Thermo Fisher	NA
CTGGGTTCCAGGTTCCACTGGTGACTCYTATGA	Scientific	
GCTGACWCAGCCAC -3' (PCRb primer)		
VL3I-Int tag 5'-	Thermo Fisher	NA
CTGGGTTCCAGGTTCCACTGGTGACTCTTCTGA	Scientific	
GCTGACTCAGGACCC -3' (PCRb primer)		
VL4ab-Int tag 5'-	Thermo Fisher	NA
CTGGGTTCCAGGTTCCACTGGTGACCAGCYTGT	Scientific	1 14 1
GCTGACTCAATC -3' (PCRb primer)		
VL4c-Int tag 5'-	Thermo Fisher	NA
CTGGGTTCCAGGTTCCACTGGTGACCTGCCTGT	Scientific	
GCTGACTCAGC -3' (PCRb primer)	Scientific	
VL5,9-Int tag 5'-	Thermo Fisher	NA
CTGGGTTCCAGGTTCCACTGGTGACCAGSCTGT	Scientific	INA
	Scientific	
GCTGACTCAGCC -3' (PCRb primer)	The same of Eliste	NA
VL6-Int tag 5'-	Thermo Fisher	NA
CTGGGTTCCAGGTTCCACTGGTGACAATTTTAT	Scientific	
GCTGACTCAGCCCCACT -3' (PCRb primer)		
VL7,8-Int tag 5'-	Thermo Fisher	NA
CTGGGTTCCAGGTTCCACTGGTGACCAGRCTGT	Scientific	
GGTGACYCAGGAG -3' (PCRb primer)		

		N. 4
VL10-Int tag 5'-	Thermo Fisher	NA
CTGGGTTCCAGGTTCCACTGGTGACCAGGCAGG	Scientific	
GCWGACTCAG -3' (PCRb primer)		
CL-int 5'- GGGYGGGAACAGAGTGACC -3' (PCRb	Thermo Fisher	NA
primer)	Scientific	
VH_Tag fwd seq 5'-	Thermo Fisher	NA
CTGGGTTCCAGGTTCCACTGGTGAC -3'	Scientific	
(Sequencing primer)		
CK_int 5'- GGGAAGATGAAGACAGATGGT -3'	Thermo Fisher	NA
(Sequencing primer)	Scientific	
CL_int 5'- GGGYGGGAACAGAGTGACC -3'	Thermo Fisher	NA
(Sequencing primer)	Scientific	
HV13221H_R474 5'- GCTGTGCCCCCAGAGGTG -3'	Thermo Fisher	NA
(Sequencing primer)	Scientific	
SARS-CoV-2 or WIV1-CoV E gene subgenomic RNA	Taqman	NA
primer/probe:		
forward primer: 5'-CGATCTCTTGTAGATCTGTTCT		
C-3';		
reverse primer: 5'-ATATTGCAGCAGTACGCACACA		
Probe: 5'-FAM-ACACTAGCCATCCTTACT		
GCGCTTCG-BHQ1-3'.	T	
SARS-CoV-2 N gene subgenomic RNA primer/probe:	Taqman	NA
forward primer: 5'-CGATCTCTTGTAGATCTGTTCT		
C-3';		
reverse primer: 5'-GGTGAACCAAGACGCAGTAT-		
3'; Droba: 5' EAM TAACCACAATCCACAACCCACTC		
Probe: 5'-FAM-TAACCAGAATGGAGAACGCAGTG		
GG-BHQ1-3'.	Taaman	NA
WIV1-CoV N gene subgenomic RNA primer/probe:	Taqman	NA
forward primer: 5'-CGATCTCTTGTAGATCTGTTCT C-3';		
c-5, reverse primer: 5'-TGTGAACCAAGACGCAGTATTA		
T-3';		
Probe: 5'-FAM-TAACCAGAATGGAGGACGCAATG		
GG-BHQ1-3';		
SARS-CoV-2 total viral RNA E gene primer/probe:	Tagman	NA
forward primer: 5'-ACAGGTACGTTAATAGTTAATA	Taqman	INA
GCGT-3',		
reverse primer: 5'-ATATTGCAGCAGTACGCACACA		
-3';		
probe: 5'-6FAM/ACACTAGCCATCCTTACTGCGCT		
TCG/IABkFQ-3'.		
Recombinant DNA		
HV1301409_4A (human IgG1_4A heavy chain	Genscript	NA
backbone)	Genseripi	
pH510049_VRC_LS.v2 (human IgG1_LS heavy chain	Genscript	NA
backbone)	Genseripi	11/1
HV1301410 (human kappa chain backbone)	Genscript	NA
HV1301410 (numan kappa chain backbone) HV1301414.v2 (human lambda chain backbone)	<b>^</b>	NA NA
	Genscript	
pcDNA3.1-SARS-CoV-2_SgE (for making subgenomic	Genscript	NA
RNA standard RNA)	Conconint	
pcDNA3.1-SARS-CoV-2_SgN (for making subgenomic	Genscript	NA
RNA standard RNA)		

	re-proof	
pcDNA3.1-WIV1-CoV_SgN (for making subgenomic RNA standard RNA)	Genscript	NA
Software and Algorithms		
Diva	BD Biosciences	http://www.bdbioscien ces.com/us/instrument s/clinical/software/flo w-cytometry- acquisition/bd- facsdiva- software/m/333333/ov erview
FlowJo v9.9.4	FlowJo, LLC	https://www.flowjo.co m
GraphPad Prism v8.3.1	GraphPad Software Inc	https://www.graphpad. com/scientific- software/prism/
SAS v9.4	SAS Institute	NA
Cloanalyst Program	(Kepler et al., 2014)	NA
Biacore S200 Evaluation software	Cytiva	NA
Coot	(Emsley et al., 2010)	Version 0.8.9.2
Relion	(Scheres, 2012; Scheres, 2016)	Version 3.1
Phenix	(Afonine et al., 2018; Liebschner et al., 2019)	Version 1.17
UCSF Chimera	(Pettersen et al., 2004)	http://www.cgl.ucsf.ed u/chimera/
ISOLDE	(Croll, 2018)	Version 1.1
Chimera X	(Goddard et al., 2018)	https://www.rbvi.ucsf. edu/chimerax/
PyMol	The PyMOL Molecular Graphics System (Schrödinger, 2015).	https://www.pymol.or g/2/
Leginon system	(Suloway et al., 2005).	NA
cryoSPARC	(Punjani et al., 2017)	https://cryosparc.com
Bio-Plex Manager Software	Bio-Rad	NA
Adobe Illustrator 2020	Adobe	NA
Adobe Photoshop CC 2019	Adobe	NA
Other		

# **RESOURCE AVAILABILITY**

# 508 Lead contact

	Journal Pre-proof
509	Further information and requests for reagents should be directed and will be fulfilled by the Lead Contact
510	Kevin O. Saunders (kevin.saunders@duke.edu).
511	Materials availability
512	Abs and other reagents generated in this study are available from the Lead Contact with a completed Material
513	Transfer Agreement.
514	Data and code availability
515	The data that support the findings of this study are available from the corresponding authors on request.
516	Additional Supplemental Items are available from Mendeley Data at <u>http://dx.doi.org/10.17632/9y6p7shshy.1</u> .
517	
518	EXPERIMENTAL MODEL AND SUBJECT DETAILS
519	Human Subjects
520	Nasopharyngeal swabs and peripheral blood samples were collected from a convalescent COVID-19 donor
521	(MESSI ID #450905) on designated days after reported onset of COVID symptoms. The SARS-CoV donor PBMO
522	were provided by NIH/VRC. Human subject specimens were collected and used with the informed consent of stud
523	participants and in compliance with the Duke University Medical Center Institutional Review Board (DUHS IRB
524	Pro00100241).
525	
526	Symptom data collections
527	Participant self-reported symptoms were recorded at each time-point for 39 symptom categories (nasal
528	discharge, nasal congestion, sneezing, coughing, shortness of breath, malaise, throat discomfort, fever, headache,
529	shaking chills, loss of smell, loss of taste, excessive sweating, dizziness, pain behind the eyes, itchy/watery eyes,
530	visual blurring, hearing problems, ear pain, confusion, stiff neck, swollen glands, palpitations, chest pain, pain in
531	joints, muscle soreness, fatigue, loss of appetite, abdominal pain, nausea/vomiting, diarrhea, swelling, itchy skin,
532	rash, skin lesions, unusual bleeding, red fingers or toes, red eyes, other: specify). Each symptom was scored on a

- 533 scale of 0–4, with 0 indicating not present, 1 mild, 2 moderate, 3 severe, and 4 very severe symptoms. Daily
- 534 symptom count (number of non-zero symptom categories) and symptom severity (sum of all symptom scores) were
- 535 determined for each survey timepoint. At enrollment, date of symptom onset was determined, and an initial

536 "historical" symptom survey recorded maximum score for each symptom category between symptom onset and537 study enrollment.

538

# 539 Non-human primate model

540 In total, 62 outbred adult male and female cynomolgus macaques (Macaca fascicularis), 2-4 kg body weight, 541 were randomly allocated to groups. The study protocol and all veterinarian procedures were approved by the 542 Bioqual IACUC per a memorandum of understanding with the Duke IACUC, and were performed based on 543 standard operating procedures. Macaques studied were housed and maintained in an Association for Assessment 544 and Accreditation of Laboratory Animal Care-accredited institution in accordance with the principles of the 545 National Institute of Health. All studies were carried out in strict accordance with the recommendations in the 546 Guide for the Care and Use of Laboratory Animals of the National Institutes of Health in BIOQUAL (Rockville, 547 MD). BIOOUAL is fully accredited by AAALAC and through OLAW, Assurance Number A-3086. All physical 548 procedures associated with this work were done under anesthesia to minimize pain and distress in accordance with 549 the recommendations of the Weatherall report, "The use of non-human primates in research." Teklad 5038 Primate 550 Diet was provided once daily by animal size and weight. The diet was supplemented with fresh fruit and vegetables. 551 Fresh water was given ad libitum. All monkeys were maintained in accordance with the Guide for the Care and Use 552 of Laboratory Animals.

553

### 554 Mouse models

555 Eleven to twelve-month old female immunocompetent BALB/c mice purchased from Envigo (BALB/c 556 AnNHsd, stock# 047) were used for SARS-CoV-2 in vivo protection experiments as described previously (Dinnon 557 et al., 2020; Leist et al., 2020a). Ten-week-old HFH4-hACE2 transgenic mice were bred and maintained at the 558 University of North Carolina at Chapel Hill and used for WIV-1 in vivo protection experiments. Mice were housed 559 in groups of five animals per cage and fed standard chow diet. The study was carried out in accordance with the 560 recommendations for care and use of animals by the Office of Laboratory Animal Welfare (OLAW), National 561 Institutes of Health and the Institutional Animal Care. All mouse studies were performed at the University of North 562 Carolina (Animal Welfare Assurance #A3410-01) using protocols (19-168) approved by the UNC Institutional 563 Animal Care and Use Committee (IACUC) and all mouse studies were performed in a BSL3 facility at UNC.

### 564

### 565 METHOD DETAILS

### 566 Expression of Recombinant Viral Proteins

567 The SARS-CoV-2 ectodomain constructs were produce and purified as describe previously (Wrapp, D.et al. 568 2020). Plasmids encoding Spike-2P and HexaPro (Hsieh et al., 2020) were transiently transfected in FreeStyle 293 569 cells (Thermo Fisher) using Turbo293 (SpeedBiosystems). The cultures were collected on Day 6 post transfection. 570 The cells were separated from the medium by centrifugation. Protein were purified from filtered cell supernatants 571 by StrepTactin resin (IBA) and additionally by size exclusive chromatography using Superose 6 10/300 increase 572 column (GE Healthcare) in 2mM Tris pH 8, 200mMnNaCl, 0.02% NaN<sub>3</sub>, SARS-CoV-2 NTD was produced as 573 previously described (Zhou et al., 2020b). SARS-CoV RBD and MERS-CoV Spike RBD were cloned into pVRC 574 vector for mammalian expression (FreeStyle 293F or Expi293F suspension cells). The construct contains an HRV 575 3C-cleavable C-terminal SBP-8xHis tag. Supernatants were harvested 5 days post-transfection and passaged 576 directly over Cobalt-TALON resin (Takara) followed by size exclusion chromatography on Superdex 200 Increase 577 (GE Healthcare) in 1x PBS. Typical yields from FreeStyle 293F cells are approximately 50 mg/liter culture. 578 Affinity tags can be removed using HRV 3C protease (ThermoScientific) and the protein repurified using Cobalt-

- 579 TALON resin to remove the protease, tag and non-cleaved protein.
- 580

## 581 Antigen-Specific Single B Cell Sorting

582 Plasmablasts were sorted by flow cytometry from the SARS-CoV-2 donor on Day 11 and Day 15 post

583 symptom onset. PBMCs were stained with optimal concentrations of the following fluorochrome-Ab conjugates:

584 IgD PE (Clone# IA6-2, BD Biosciences, Catalog# 555779), CD3 PE-Cy5 (Clone# HIT3a, BD Biosciences,

585 Catalog# 555341), CD10 PE-CF594 (Clone# HI10A, BD Biosciences, Catalog# 562396), CD27 PE-Cy7 (Clone#

586 O323, eBioscience, Catalog# 25-0279), CD38 APC-Alexa Fluor (AF) 700 (Clone# LS198-4-2, Beckman Coulter,

- 587 Catalog# B23489), CD19 APC-Cy7 (Clone# LJ25C1, BD Biosciences, Catalog# 561743), CD16 BV570 (Clone#
- 588 3G8, Biolegend, Catalog# 302035), CD14 BV605 (Clone# M5E2, Biolegend, Catalog# 301834), and CD20 BV650
- 589 (Clone# 2H7, BD, Catalog# 563780). The cells were then labeled with Fixable Aqua Live/Dead Cell Stain Kit
- 590 (Invitrogen, Catalog# L34957). On a BD FACSAria II flow cytometer (BD Biosciences), plasmablasts were
- 591 identified as viable CD14<sup>-</sup>/CD16<sup>-</sup>/CD19<sup>+</sup>/CD20<sup>low</sup>/IgD<sup>-</sup>/CD27<sup>high</sup>/CD38<sup>high</sup> cells and sorted as single cells into 96-

592	well plates containing lysis buffer. Sorted plates were frozen at -80°C in the DHVI Flow Facility under BSL3
593	precautions in the Duke Regional Biocontainment Laboratory (Durham, NC) until processing.
594	Antigen-specific memory B cells (MBCs) were isolated by flow cytometric sorting from the SARS-CoV-2
595	donor on Day 36 post symptom onset, and a donor with SARS-CoV history. PBMCs were stained with IgD FITC
596	(Clone# IA6-2, BD Biosciences, Catalog# 555778), IgM PerCp-Cy5.5 (Clone# G20-127, BD Biosciences,
597	Catalog# 561285), CD10 PE-CF594 (Clone# HI10A, BD Biosciences, Catalog# 562396), CD3 PE-Cy5 (Clone#
598	HIT3a, BD Biosciences, Catalog# 555341), CD235a PE-Cy5 (Clone# GA-R2, BD Biosciences, Catalog# 559944),
599	CD27 PE-Cy7 (Clone# O323, eBioscience, Catalog# 25-0279), CD38 APC-AF700 (Clone# LS198-4-2, Beckman
600	Coulter, Catalog# B23489), CD19 APC-Cy7 (Clone# LJ25C1, BD Biosciences, Catalog# 561743), CD14 BV605
601	(Clone# M5E2, Biolegend, Catalog# 301834), CD16 BV570 (Clone# 3G8, Biolegend, Catalog# 302035), and
602	fluorescent-labeled SARS-CoV-2 Spike probes (AF647-conjugated Spike-2P, PE-conjugated Spike-2P, AF647-
603	conjugated NTD, AF647-conjugated RBD, VioBright 515-conjugated RBD). The cells were then labeled with
604	Fixable Aqua Live/Dead Cell Stain Kit (Invitrogen, Catalog# L34957). On a BD FACSAria II flow cytometer (BD
605	Biosciences), antigen-specific MBCs were identified as viable CD3 <sup>-</sup> /CD14 <sup>-</sup> /CD16 <sup>-</sup> /CD235a <sup>-</sup> /CD19 <sup>+</sup> /IgD <sup>-</sup> /probe <sup>+</sup>
606	cells and were sorted as single cells into 96-well plates containing lysis buffer. Collection plates were immediately
607	frozen in a dry ice/ethanol bath, and stored at -80 °C in the DHVI Flow Facility under BSL3 precautions in the
608	Duke Regional Biocontainment Laboratory until processing. Flow cytometric data were analyzed using FlowJo
609	version 10.

610

611 PCR Amplification of Human Ab Genes

Ab genes were amplified by RT-PCR from flow cytometry-sorted single B cells using the methods as described previously (Liao et al., 2009; Wrammert et al., 2008) with modification. The PCR-amplified genes were then purified and sequenced with 10  $\mu$ M forward and reverse primers. Sequences were analyzed by using the human library in Cloanalyst for the VDJ arrangements of the immunoglobulin IGHV, IGKV, and IGLV sequences and mutation frequencies (Kepler et al., 2014). Clonal relatedness of V<sub>H</sub>D<sub>H</sub>J<sub>H</sub> and V<sub>L</sub>J<sub>L</sub> sequences was determined as previously described (Liao et al., 2013).

618

# 619 Expression of Ab Viable Region Genes as Full-Length IgG Recombinant mAbs

620	Transient transfection of recombinant Abs was performed as previously described (Liao et al., 2009). Briefly,
621	purified PCR products were used for overlapping PCR to generate linear human IgG expression cassettes. The
622	expression cassettes were transfected into 293i cells using ExpiFectamine (Thermo Fisher Scientific, Catalog#
623	A14525). The supernatant samples containing recombinant IgGs were used for IgG quantification and preliminary
624	ELISA binding screening.
625	The down-selected human Ab genes were then synthesized and cloned (GenScript) in a human IgG1 backbone
626	(HV1301409_4A) with 4A mutations to enhance Ab-dependent cell-mediated cytotoxicity (ADCC) or a human
627	IgG1 backbone (pH510049_VRC_LS.v2) with a LS mutation to extent Ab half-life (Saunders, 2019). Recombinant
628	IgG Abs were then produced in HEK293i suspension cells by transfection with ExpiFectamine and purified using
629	Protein A resin. The purified IgG Abs were run in SDS-PAGE for Coomassie blue staining and western blot for
630	quality control and then used for the downstream experiments.
631	
632	Ab Binding ELISA
633	For ELISA binding assays of Coronavirus Spike Abs, the antigen panel included SARS-CoV-2 Spike S1+S2
634	ectodomain (ECD) (SINO, Catalog # 40589-V08B1), SARS-CoV-2 Spike-2P (Wrapp et al., 2020b), SARS-CoV-2
635	Spike S2 ECD (SINO, Catalog # 40590-V08B), SARS-CoV-2 Spike RBD from insect cell sf9 (SINO, Catalog #
636	40592-V08B), SARS-CoV-2 Spike RBD from mammalian cell 293 (SINO, Catalog # 40592- V08H), SARS-CoV-
637	2 Spike NTD-Biotin, SARS-CoV Spike Protein DeltaTM (BEI, Catalog # NR-722), SARS-CoV WH20 Spike RBD
638	(SINO, Catalog # 40150-V08B2), SARS-CoV WH20 Spike S1 (SINO, Catalog #40150-V08B1), SARS-CoV RBD
639	MERS-CoV Spike S1+S2 (SINO, Catalog # 40069-V08B), MERS-CoV Spike S1 (SINO, Catalog #40069-V08B1),
640	MERS-CoV Spike S2 (SINO, Catalog #40070-V08B), MERS-CoV Spike RBD (SINO, Catalog #40071-V08B1),
641	MERS-CoV Spike RBD. In preliminary ELISA screening of the transient transfection supernatants, we also
642	screened the Abs against SARS-CoV CL Protease protein (BEI, Catalog # 30105) and SARS-CoV Membrane (M)
643	protein (BEI, Catalog # 110705).
644	For binding ELISA, 384-well ELISA plates were coated with 2 $\mu$ g/mL of antigens in 0.1 M sodium
645	bicarbonate overnight at 4°C. Plates were washed with PBS + 0.05% Tween 20 and blocked with blocked with

646 assay diluent (PBS containing 4% (w/v) whey protein, 15% Normal Goat Serum, 0.5% Tween-20, and 0.05%

647 Sodium Azide) at room temperature for 1 hour. Purified mAb samples in 3-fold serial dilutions in assay diluent

- 648 starting at 100  $\mu$ g/mL, or un-diluted transfection supernatant were added and incubated for 1 hour, followed by
- 649 washing with PBS-0.1% Tween 20. HRP-conjugated goat anti-human IgG secondary Ab (SouthernBiotech,
- 650 catalog# 2040-05) was diluted to 1:10,000 and incubated at room temperature for 1 hour. These plates were washed
- 651 four times and developed with tetramethylbenzidine substrate (SureBlue Reserve- KPL). The reaction was stopped
- with 1 M HCl, and optical density at 450 nm ( $OD_{450}$ ) was determined.
- 653

# 654 Affinity Measurements

655 SPR measurements of SARS-CoV-2 Ab Fab binding to Spike-2P or Spike-HexaPro proteins were performed using a Biacore S200 instrument (Cytiva, formerly GE Healthcare, DHVI BIA Core Facility, Durham, NC) in 656 657 HBS-EP+ 1x running buffer. The Spike proteins were first captured onto a Series S Streptavidin chip to a level of 658 300-400 RU for Spike-2P and 350-450 resonance units (RU) for Spike-HexaPro. The Ab Fabs were injected at 0.5 659 to 500 nM over the captured S proteins using the single cycle kinetics injection mode at a flow rate of 50uL/min. 660 Association phase was maintained with either 120 or 240 second injections of each Fab at increasing concentrations followed by a dissociation of 600 seconds after the final injection. After dissociation, the S proteins 661 662 were regenerated from the streptavidin surface using a 30 second pulse of Glycine pH1.5. Results were analyzed 663 using the Biacore S200 Evaluation software (Cytiva). A blank streptavidin surface along with blank buffer binding were used for double reference subtraction to account for non-specific protein binding and signal drift. Subsequent 664 665 curve fitting analyses were performed using a 1:1 Langmuir model with a local Rmax for the Fabs with the 666 exception of DH1050.1 Fab which was fit using the heterogeneous ligand model with local Rmax. The reported 667 binding curves are representative of two data sets.

668

# 669 Surface Plasmon Resonance Ab Blocking Assay

RBD and NTD Abs binding to S protein was measured by surface plasmon resonance (BIAcore 3000; Cytiva,
formerly GE Healthcare, DHVI BIA Core Facility, Durham, NC) analysis. Ab binding competition and blocking
were measured by SPR following immobilization by amine coupling of monoclonal Abs to CM5 sensor chips
(BIAcore/Cytiva). Ab competition experiments were performed by mixing S protein and mAb (30 minutes
incubation) followed by injection for 5 minutes at 50 µL/min. In separate assays and from analysis of binding to an
identical epitope binding ligand, it was determined that S protein at 20 µm and Ab at 200 µm bind to complete

676 saturation. Ab blocking assays were performed by co-injecting S protein (20  $\mu$ M) over mAb immobilized surfaces 677 for 3 minutes at 30  $\mu$ L/min and a test Ab (200  $\mu$ M) for 3 minutes at 30  $\mu$ L/min. The dissociation of the Ab 678 sandwich complex with the spike protein was monitored for 10 minutes with buffer flow and then a 24 second 679 injection of Glycine pH2.0 for regeneration. Blank buffer binding was used for subtraction to account for signal 680 drift. Data analyses were performed with BIA-evaluation 4.1 software (BIAcore/Cytiva).

681

# 682 ACE2-blocking assay

683 For ACE-2 blocking assays, plates were coated as stated above with 2 ug/mL recombinant ACE-2 protein, then 684 washed and blocked with 3% BSA in 1X PBS. While assay plates blocked, purified Abs were diluted as stated 685 above, only in 1% BSA with 0.05% Tween-20. In a separate dilution plate Spike-2P protein was mixed with the 686 Abs at a final concentration equal to the EC50 at which spike binds to ACE-2 protein. The mixture was allowed to 687 incubate at room temperature for 1 hour. Blocked assay plates were then washed and the Ab-spike mixture was 688 added to the assay plates for a period of 1 hour at room temperature. Plates were washed and a polyclonal rabbit 689 serum against the same Spike-2P protein was added for 1 hour, washed and detected with goat anti rabbit-HRP 690 (Abcam cat# ab97080) followed by TMB substrate. The extent to which Abs were able to block the binding spike 691 protein to ACE-2 was determined by comparing the OD of Ab samples at 450 nm to the OD of samples containing 692 spike protein only with no Ab. The following formula was used to calculate percent blocking: blocking = (100 - 100)693 (OD sample/OD of spike only)\*100).

694

# 695 Negative-stain electron microscopy

696 For each Fab-spike complex, an aliquot of spike protein at ~1-5 mg/ml concentration that had been flash frozen and stored at -80 °C was thawed in an aluminum block at 37 °C for 5 minutes; then 1-4 µl of spike was 697 698 mixed with sufficient Fab to give a 9:1 molar ratio of Fab to spike and incubated for 1 hour at 37 °C. The complex 699 was then cross-linked by diluting to a final spike concentration of 0.1 mg/ml into room-temperature buffer 700 containing 150 mM NaCl, 20 mM HEPES pH 7.4, 5% glycerol, and 7.5 mM glutaraldehyde. After 5 minutes cross-701 linking, excess glutaraldehyde was quenched by adding sufficient 1 M Tris pH 7.4 stock to give a final 702 concentration of 75 mM Tris and incubated for 5 minutes. For negative stain, carbon-coated grids (EMS, CF300-703 cu-UL) were glow-discharged for 20s at 15 mA, after which a 5-µl drop of quenched sample was incubated on the

grid for 10-15 s, blotted, and then stained with 2% uranyl formate. After air drying grids were imaged with a
Philips EM420 electron microscope operated at 120 kV, at 82,000x magnification and images captured with a 2k x
2k CCD camera at a pixel size of 4.02 Å.

707

# 708 Image processing of negative stain images

The RELION 3.0 program was used for all negative stain image processing. Images were imported, CTFcorrected with CTFFIND, and particles were picked using a spike template from previous 2D class averages of spike alone. Extracted particle stacks were subjected to 2-3 rounds of 2D class averaging and selection to discard junk particles and background picks. Cleaned particle stacks were then subjected to 3D classification using a starting model created from a bare spike model, PDB 6vsb, low-pass filtered to 30 Å. Classes that showed clearlydefined Fabs were selected for final refinements followed by automatic filtering and B-factor sharpening with the default Relion post-processing parameters.

716

### 717 Cryo-EM sample preparation, data collection and processing

718 To prepare Ab-bound complexes of the SARS-CoV-2 2P spike, the spike at a final concentration of 1-2 719 mg/mL, in a buffer containing 2 mM Tris pH 8.0, 200 mM NaCl and 0.02% NaN3, was incubated with 5-6 fold 720 molar excess of the Ab Fab fragments for 30–60 min. 0.5% final concentration of glycerol was added to the sample right before freezing. 2.5 µL of protein was deposited on a Quantifoil-1.2/1.3 holey carbon grid that had been glow 721 722 discharged in a PELCO easiGlow<sup>TM</sup> Glow Discharge Cleaning System for 15s at 15 mA. After a 30 s incubation 723 in >95% humidity and 22 °C, excess protein was blotted away for 2.5 seconds using a Whatman #1 filter paper 724 before being plunge frozen into liquid ethane using a Leica EM GP2 plunge freezer (Leica Microsystems). Cryo-725 EM data were collected on a Titan Krios (Thermo Fisher) equipped with a K3 detector (Gatan). Data were acquired 726 using the Leginon system (Suloway et al., 2005). All the datasets were energy filtered through a either a 20eV or 727 30eV slit. The dose was fractionated over 50 raw frames and collected at 50ms framerate. Individual frames were 728 aligned and dose-weighted (Zheng et al., 2017). CTF estimation, particle picking and all downstream data 729 processing steps were carried out in cryoSparc (Punjani et al., 2017). After two rounds of 2D classifications during 730 which junk particles were discarded, heterogeneous refinement was performed using low pass filtered maps of 731 unliganded spike as inputs. The output maps showed densities of the bound Fabs, which were further classified by

754	
735	Cryo-EM structure fitting and analysis
736	Previously published SARS-CoV-2 ectodomain structures of the all 'down' state (PDB ID 6VXX) and single
737	RBD 'up' state (PDB ID 6VYB), and models of 2-RBD-up and 3-RBD-up states derived from these, were used to
738	fit the cryo-EM maps in Chimera (Pettersen et al., 2004). Models of Fabs were generated in SWIS-MODEL and
739	docked into the cryo-EM reconstructions using Chimera. Mutations were made in Coot (Emsley and Cowtan, 2004).
740	Coordinates were fit to the maps using ISOLDE (Croll, 2018) followed by iterative refinement using Phenix
741	(Afonine et al., 2018) real space refinement and subsequent manual coordinate fitting in Coot as needed. Structure
742	and map analysis were performed using PyMol (Schrodinger, 2015), Chimera (Pettersen et al., 2004) and
743	ChimeraX (Goddard et al., 2018).
744	
745	Live SARS-CoV-2 neutralization assays
746	The SARS-CoV-2 virus (Isolate USA-WA1/2020, NR-52281) was deposited by the Centers for Disease
747	Control and Prevention and obtained through BEI Resources, NIAID, NIH. SARS-CoV-2 Micro-neutralization
748	(MN) assays were adapted from a previous study (Berry et al., 2004). In short, sera or purified Abs are diluted two-
749	fold and incubated with 100 TCID50 virus for 1 hour. These dilutions are used as the input material for a TCID50.
750	Each batch of MN includes a known neutralizing control Ab (Clone D001; SINO, CAT# 40150-D001). Data are
751	reported as the last concentration at which a test sample protects Vero E6 cells.
752	SARS-CoV-2 Plaque Reduction Neutralization Test (PRNT) were performed in the Duke Regional
753	Biocontainment Laboratory BSL3 (Durham, NC) as previously described with virus-specific modifications (Berry
754	et al., 2004). Briefly, two-fold dilutions of a test sample (e.g. serum, plasma, purified Ab) were incubated with 50
755	PFU SARS-CoV-2 virus (Isolate USA-WA1/2020, NR-52281) for 1 hour. The Ab/virus mixture is used to
756	inoculate Vero E6 cells in a standard plaque assay (Coleman and Frieman, 2015; Kint et al., 2015). Briefly,
757	infected cultures are incubated at 37°C, 5% CO2 for 1 hour. At the end of the incubation, 1 mL of a viscous
758	overlay (1:1 2X DMEM and 1.2% methylcellulose) is added to each well. Plates are incubated for 4 days. After
759	fixation, staining and washing, plates are dried and plaques from each dilution of each sample are counted. Data are

heterogeneous refinement, followed by non-uniform refinement to obtain the final reconstructions that were used

733 for model fitting.

734

732

/00	reported as the concentration at which 50% of input virus is neutralized. A known neutralizing control Ab is
761	included in each batch run (Clone D001; SINO, CAT# 40150-D001). GraphPad Prism was used to determine
762	IC/EC <sub>50</sub> values.
763	SARS-CoV-2 nano-luciferase (nanoLuc), SARS-CoV nanoLuc and WIV1-CoV nanoLuc replication-
764	competent virus neutralization assay were described previously (Hou et al., 2020; Menachery et al., 2016; Sheahan
765	et al., 2017).
766	
767	Pseudo-typed SARS-CoV-2 neutralization assay and infection-enhancing assays
768	Neutralization of SARS-CoV-2 Spike-pseudotyped virus was performed by adopting an infection assay
769	described previously (Korber et al., 2020) with lentiviral vectors and infection in either 293T/ACE2.MF (the cell
770	line was kindly provided by Drs. Mike Farzan and Huihui Mu at Scripps). Cells were maintained in DMEM
771	containing 10% FBS and 50 µg/ml gentamicin. An expression plasmid encoding codon-optimized full-length spike
772	of the Wuhan-1 strain (VRC7480), was provided by Drs. Barney Graham and Kizzmekia Corbett at the Vaccine
773	Research Center, National Institutes of Health (USA). The D614G mutation was introduced into VRC7480 by site-
774	directed mutagenesis using the QuikChange Lightning Site-Directed Mutagenesis Kit from Agilent Technologies
775	(Catalog # 210518). The mutation was confirmed by full-length spike gene sequencing. Pseudovirions were
776	produced in HEK 293T/17 cells (ATCC cat. no. CRL-11268) by transfection using Fugene 6 (Promega, Catalog
777	#E2692). Pseudovirions for 293T/ACE2 infection were produced by co-transfection with a lentiviral backbone
778	(pCMV ΔR8.2) and firefly luciferase reporter gene (pHR' CMV Luc) (Naldini et al., 1996). Culture supernatants
779	from transfections were clarified of cells by low-speed centrifugation and filtration (0.45 $\mu$ m filter) and stored in 1
780	ml aliquots at -80 °C.
781	For 293T/ACE2 neutralization assays, a pre-titrated dose of virus was incubated with 8 serial 3-fold or 5-fold

dilutions of mAbs in duplicate in a total volume of 150  $\mu$ l for 1 hr at 37 °C in 96-well flat-bottom poly-L-lysinecoated culture plates (Corning Biocoat). Cells were suspended using TrypLE express enzyme solution (Thermo Fisher Scientific) and immediately added to all wells (10,000 cells in 100  $\mu$ L of growth medium per well). One set of 8 control wells received cells + virus (virus control) and another set of 8 wells received cells only (background control). After 66-72 hr of incubation, medium was removed by gentle aspiration and 30  $\mu$ L of Promega 1x lysis

buffer was added to all wells. After a 10-minute incubation at room temperature, 100 µl of Bright-Glo luciferase reagent was added to all wells. After 1-2 minutes, 110 µl of the cell lysate was transferred to a black/white plate (Perkin-Elmer). Luminescence was measured using a PerkinElmer Life Sciences, Model Victor2 luminometer. Neutralization titers are the mAb concentration (IC50/IC80) at which relative luminescence units (RLU) were reduced by 50% and 80% compared to virus control wells after subtraction of background RLUs. Negative neutralization values are indicative of infection-enhancement. Maximum percent inhibition (MPI) is the reduction in RLU at the highest mAb concentration tested.

794 For the TZM-bl neutralization assays, a pre-titrated dose of virus was incubated with serial 3-fold dilutions of 795 test sample in duplicate in a total volume of 150 ul for 1 hr at 37 °C in 96-well flat-bottom culture plates. Freshly 796 trypsinized cells (10,000 cells in 100 µl of growth medium containing 75 µg/ml DEAE dextran) were added to each well. One set of control wells received cells + virus (virus control) and another set received cells only 797 798 (background control). After 68-72 hours of incubation, 150 µl of cultured medium was removed from each well, 799 and 100ul of Britelite Luminescence Reporter Gene Assay System (PerkinElmer Life Sciences) were added and 800 plates incubated for 2 min at room temperature. After this period 150 µl of the lysate was transferred to black solid 801 plates (Costar) for measurements of luminescence in a Perking Elmer instrument. Neutralization titers are the 802 serum dilution at which relative luminescence units (RLU) were reduced by 50% and 80% compared to virus 803 control wells after subtraction of background RLUs. MPI is the reduction in RLU at the highest mAb concentration 804 tested. Infection-enhancing assays were performed with the same format but using TZM-bl cell lines stably 805 expressing each of the four human FcyR receptors (Perez et al., 2009). In this assay an increase in RLUs over the 806 virus control signal represents FcR-mediated entry.

807

# 808 Non-human primate protection study

809 Groups of five cynomolgus macaques (2-4 kg) were given intravenous infusion with Abs at 10 mg/kg body

- 810 weight on Day -3, relative to infectious virus challenge. For each animal, 10<sup>5</sup> PFU (~10<sup>6</sup> TCID50) SARS-CoV-2
- 811 virus (Isolate USA-WA1/2020) were diluted in 4 mL, and were given by 1 mL intranasally and 3 mL
- 812 intratracheally on Day 0. Plasma and serum samples were collected on Day -5, 0, 2, and 4. Nasal swabs, nasal
- 813 washes, and bronchoalveolar lavage (BAL) were collected on Day -5, 2, and 4.

# 814

# 815 Histopathology and Immunohistochemistry (IHC)

Lung specimen from nonhuman primates were fixed in 10% neutral buffered formalin, processed, and blocked
in paraffin for histological analysis. All samples were sectioned at 5 µm and stained with hematoxylin-eosin (H&E)
for routine histopathology. Sections were examined under light microscopy using an Olympus BX51 microscope
and photographs were taken using an Olympus DP73 camera.

820

821 Staining for SARS-CoV-2 antigen was achieved on the Bond RX automated system with the Polymer Define 822 Detection System (Leica) used per manufacturer's protocol. Tissue sections were dewaxed with Bond Dewaxing 823 Solution (Leica) at 72 for 30 min then subsequently rehydrated with graded alcohol washes and 1x Immuno 824 Wash (StatLab). Heat-induced epitope retrieval (HIER) was performed using Epitope Retrieval Solution 1 (Leica), 825 heated to 100oC for 20 minutes. A peroxide block (Leica) was applied for 5 min to quench endogenous peroxidase 826 activity prior to applying the SARS-CoV-2 Ab (1:2000, GeneTex, GTX135357). Abs were diluted in Background 827 Reducing Ab Diluent (Agilent). The tissue was subsequently incubated with an anti-rabbit HRP polymer (Leica) 828 and colorized with 3,3'-Diaminobenzidine (DAB) chromogen for 10 min. Slides were counterstained with 829 hematoxylin. For macrophage staining, Abs for the following markers were used: CD3 (T cell marker; Bio Rad, 830 Catalog # MCA1477; 1:600 dilution), Iba1 (macrophage marker; Wako, Catalog # 019-19741; 1:800 dilution), 831 CD68 (M1 macrophage marker, Sigma-Millipore, Catalog # HPA048982; 1:1000 dilution), CD163 (M2 832 macrophage marker; Abcam, Catalog # ab182422; 1:500 dilution), HLA-DP/DO/DR (Catalog # M1 macrophage 833 marker; Dako, Catalog # M0775; 1:100 dilution), CD11b (monocyte/granulocyte marker; Abcam, Catalog # 834 ab52478; 1:1000 dilution). 835 Samples were evaluated by a board-certified veterinary pathologist in a blinded manner. Sections of the left

836 caudal (Lc), right middle (Rm), and right caudal (Rc) lung were evaluated and scored for the presence of

837 inflammation by H&E staining, and for the presence of SARS-CoV-2 nucleocapsid by IHC staining. The sums of

838 Lc, Rm, and Rc scores in each animal shown in figures.

839

840 Luminex assay

841 For cytokine profiling, 7-fold concentrated cynomolgus macaques BAL samples were measured using a 25-842 analyte multiplex bead array (Millipore, catalog # PRCYT2MAG40K) including sCD137, Eotaxin, sFasL, FGF-2, 843 Fractalkine, Granzyme A, Granzyme B, IL-1α, IL-2, IL-4, IL-6, IL-16, IL-17A, IL-17E/IL-25, IL-21, IL-22, IL-23, 844 IL-28A, IL-31, IL-33, IP-10, MIP-3α, Perforin, RANTES, TNFβ. Samples were prepared according to the 845 manufacturer's recommended protocol and read using a Flexmap 3D suspension array reader (Luminex Corp.). 846 Data were analyzed using Bio-Plex manager software v6.2 (Bio-Rad). 847 For human Ab quantification, SARS-CoV-2 Spike-2P protein, A/Solomon Islands/3/2006 hemagglutinin (HA) 848 protein or bovine serum albumen (Sigma) was carbodiimide coupled to MagPlex-C beads (Luminex Corp) 849 according to the bead manufacturer's protocol. Briefly, beads were washed in H<sub>2</sub>O then activated by incubation 850 with 5 mg/mL sulfo-N-hydroxysulfosuccinimide and 5 mg/mL 1-ethyl-3-(3-dimethylaminopropyl) carbodiimide 851 hydrochloride (ThermoFisher) for 20 minutes. Activated beads were washed twice in PBS (ThermoFisher) and then vortexed at 1,500 RPM for two hours at room temperature with 25 µg protein per 5.0 x 10<sup>6</sup> beads. Labelled beads 852 were washed in PBS (ThermoFisher), 1% BSA, 0.02% Tween-20, 0.05% Sodium Azide (all Sigma), counted using 853 854 a hemacytometer and stored at -80°C. NHP sera were diluted 1:200 in assay buffer (PBS, 1% BSA, pH 7.4, Gibco), 855 then 50 µL of diluted sera or monoclonal Ab 3-fold serially diluted in assay buffer (1000-0.45ng/mL) was added to a 96-well plate and mixed with 50 µL assay buffer containing 2500 BSA-conjugated beads (negative control) plus 856 2500 HA or Spike-conjugated beads. The plate was shaken at 800RPM for 60 minutes at room temperature, 857 858 washed twice in assay buffer and then 100  $\mu$ L 4  $\mu$ g/mL biotin-conjugated mouse anti-human IgG Fc clone H2 859 (Southern Biotech) in assay buffer was added to every well. The plate was shaken at 800 RPM for 30 minutes at 860 room temperature, washed two times in assay buffer and then 50  $\mu$ L 4  $\mu$ g/mL streptavidin-r-phycoerythrin 861 (Invitrogen) in assay buffer was added to every well. The plate was shaken at 800 RPM for 30 minutes at room 862 temperature and washed twice in assay buffer. Beads were resuspended in 150 µL/well assay buffer, shaken at 800 863 RPM for 15 minutes at room temperature and then analyzed on a BioPlex 200 bead reader (Bio-Rad). Sera antigen-864 specific Ab concentrations were calculated using Bio-Plex Manager software (Bio-Rad) by extrapolating from the 865 results of the serially-diluted monoclonal Ab. Sera with Abs above the upper limit of quantitation were re-assayed 866 at 1:1000 or 1:5000. The limit of detection (LOD) for this assay is 0.278 µg/mL. 867

# 868 Mouse protection study

869 Eleven to twelve-month old female immunocompetent BALB/c mice purchased from Envigo (BALB/c 870 AnNHsd, stock# 047) were used for SARS-CoV-2 in vivo protection experiments as described previously (Dinnon 871 et al., 2020). Ten-week-old HFH4-hACE2 transgenic mice were bred and maintained at the University of North 872 Carolina at Chapel Hill and used for WIV-1 in vivo protection experiments. Mice were housed in groups of five 873 animals per cage and fed standard chow diet. Virus inoculations were performed under anesthesia (Ketamine and 874 Xylazine) and effort was taken to minimize animal suffering. For evaluating the prophylactic efficacy of mAbs, 875 mice were intraperitoneally treated with 300 µg of each mAb or 150 µg of each mAb in combination 12 hours prior 876 to infection. Mice were infected intranasally with 1X10<sup>5</sup> PFU of mouse-adapted SARS-CoV-2 2AA MA (Dinnon 877 et al., 2020) or WIV-1. For evaluating the therapeutic efficacy of mAbs, mice were intraperitoneally treated with 878 300 µg of each mAb or 150 µg of each mAb in combination 12 hours following infection. Forty-eight hours post 879 infection, mice were sacrificed, and lungs were harvested for viral titer as measured by plaque assays and RNA 880 analysis. In another study, fifty-two weeks old female BALB/c mice were i.p. injected with DH1052 (200ug/mice, n=10) or CH65 control Ab (200ug/mice, n=9). After 12 hours, mice were challenged with 1X10^4 PFU of mouse-881 882 adapted SARS-CoV-2 MA10 virus (Leist et al., 2020a). Mice were sacrificed at day 4 post infection, and lungs 883 were harvested for viral titer as measured by plaque assays and RNA analysis. The study was carried out in 884 accordance with the recommendations for care and use of animals by the Office of Laboratory Animal Welfare 885 (OLAW), National Institutes of Health and the Institutional Animal Care. All mouse studies were performed at the 886 University of North Carolina (Animal Welfare Assurance #A3410-01) using protocols (19-168) approved by the 887 UNC Institutional Animal Care and Use Committee (IACUC) and all mouse studies were performed in a BSL3 888 facility at UNC.

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# 890 Viral RNA Extraction and Quantification

The assay for SARS-CoV-2 quantitative Polymerase Chain Reaction (qPCR) detects total RNA using the
WHO primer/probe set E\_Sarbeco (Charité/Berlin). A QIAsymphony SP (Qiagen, Hilden, Germany) automated
sample preparation platform along with a virus/pathogen DSP midi kit and the *complex800* protocol were used to
extract viral RNA from 800 µL of pooled samples. A reverse primer specific to the envelope gene of SARS-CoV-2
(5'-ATA TTG CAG CAG TAC GCA CAC A-3') was annealed to the extracted RNA and then reverse transcribed

into cDNA using SuperScript<sup>TM</sup> III Reverse Transcriptase (Thermo Fisher Scientific, Waltham, MA) along with 896 RNAse Out (Thermo Fisher Scientific, Waltham, MA). The resulting cDNA was treated with RNase H (Thermo 897 Fisher Scientific, Waltham, MA) and then added to a custom 4x TaqMan<sup>TM</sup> Gene Expression Master Mix (Thermo 898 899 Fisher Scientific, Waltham, MA) containing primers and a fluorescently labeled hydrolysis probe specific for the 900 envelope gene of SARS-CoV-2 (forward primer 5'-ACA GGT ACG TTA ATA GTT AAT AGC GT-3', reverse 901 primer 5'-ATA TTG CAG CAG TAC GCA CAC A-3', probe 5'-6FAM/AC ACT AGC C/ZENA TCC TTA CTG 902 CGC TTC G/IABkFQ-3'). The qPCR was carried out on a QuantStudio 3 Real-Time PCR System (Thermo Fisher 903 Scientific, Waltham, MA) using the following thermal cycler parameters: heat to 50°C, hold for 2 min, heat to 904 95°C, hold for 10 min, then the following parameters are repeated for 50 cycles: heat to 95°C, hold for 15 seconds, 905 cool to 60°C and hold for 1 minute. SARS-CoV-2 RNA copies per reaction were interpolated using quantification 906 cycle data and a serial dilution of a highly characterized custom DNA plasmid containing the SARS-CoV-2 907 envelope gene sequence. Mean RNA copies per milliliter were then calculated by applying the assay dilution factor 908 (DF=11.7). The limit of detection (LOD) for this assay is approximately 62 RNA copies per milliliter of sample.

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# 910 Subgenomic mRNA assay

911 SARS-CoV-2 E gene and N gene subgenomic mRNA (sgRNA) was measured by a one-step RT-qPCR 912 adapted from previously described methods (Wolfel et al., 2020; Yu et al., 2020). To generate standard curves, a 913 SARS-CoV-2 E gene sgRNA sequence, including the 5'UTR leader sequence, transcriptional regulatory sequence 914 (TRS), and the first 228 bp of E gene, was cloned into a pcDNA3.1 plasmid. For generating SARS-CoV-2 N gene 915 sgRNA, the E gene was replaced with the first 227 bp of N gene. The respectively pcDNA3.1 plasmids were 916 linearized, transcribed using MEGAscript T7 Transcription Kit (ThermoFisher, Catalog # AM1334), and purified 917 with MEGAclear Transcription Clean-Up Kit (ThermoFisher, Catalog # AM1908). The purified RNA products

918 were quantified on Nanodrop, serial diluted, and aliquoted as E sgRNA or N sgRNA standards.

919 RNA extracted from animal samples or standards were then measured in Taqman custom gene expression

920 assays (ThermoFisher Scientific). For these assays we used TaqMan Fast Virus 1-Step Master Mix (ThermoFisher,

921 catalog # 4444432) and custom primers/probes targeting the E gene sgRNA (forward primer: 5'

922 CGATCTCTTGTAGATCTGTTCTCE 3'; reverse primer: 5' ATATTGCAGCAGT ACGCACACA 3'; probe: 5'

923 FAM-ACACTAGCCATCCTTACTGCGCTTCG-BHQ1 3') or the N gene sgRNA (forward primer: 5'

924	CGATCTCTTGTAGATCTGTTCTC 3'; reverse primer: 5' GGTGAA CCAAGACGCAGTAT 3'; probe: 5' FAM-								
925	TAACCAGAATGGAGAACGCAGTG GG-BHQ1 3'). RT-qPCR reactions were carried out on a QuantStudio 3								
926	Real-Time PCR System (Applied Biosystems) or a StepOnePlus Real-Time PCR System (Applied Biosystems)								
927	using a program below: reverse transcription at 50°C for 5 minutes, initial denaturation at 95°C for 20 seconds,								
928	then 40 cycles of denaturation-annealing-extension at 95°C for 15 seconds and 60°C for 30 seconds. Standard								
929	curves were used to calculate E or N sgRNA in copies per ml; the limit of detections (LOD) for both E and N								
930	sgRNA assays were 12.5 copies per reaction or 150 copies per mL of BAL/nasal swab/nasal wash.								
931									
932	QUANTIFICATION AND STATISTICAL ANALYSIS								
933	Data were plotted using Prism GraphPad 8.0. Wilcoxon rank sum exact test was performed to compare								

differences between groups with p-value < 0.05 considered significant using SAS 9.4 (SAS Institute, Cary, NC).

935 No adjustments were made to the p-values for multiple comparisons.

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- 1208
- 1209

1210 Supplementary Figure Legends

1211 Figure S1. Isolation of SARS-CoV-2-reactive Abs from single cell-sorted plasmablasts and memory B cells of

- 1212 SARS-CoV-2 and SARS-CoV-1 convalescent donors. Related to Figure 1.
- 1213 (A) Symptom severity scores of the COVID-19 convalescent donor. The method to determine severity score is in
- supplementary online material. Red arrows indicate the blood sampling time points that we used to isolate Abs.
- 1215 (B) Viral load from nasopharyngeal (NP) swabs.
- 1216 (C) Serum micro-neutralization titer. Micro-Neutralization titers were defined as the highest serum dilution that
- 1217 neutralize all the virus, or 99% inhibitory concentration (IC<sub>99</sub>).
- 1218 (D) Flow cytometry gating strategy for unbiased plasmablasts sorting or antigen specific-memory B cells sorting.
- 1219 At day 11 and day 15 post onset of COVID-19 symptom, plasmablasts (CD14<sup>-</sup>/CD16<sup>-</sup>/CD3<sup>-</sup>/CD235a<sup>-</sup>
- 1220 /CD19<sup>+</sup>/CD20<sup>low</sup>/IgD<sup>-</sup>/CD27<sup>high</sup>/CD38<sup>high</sup>) from a SARS-CoV-2 donor. Antigen specific B cells from SARS-CoV-1
- 1221 and SARS-CoV-2 donors were sorted with different combinations of the SARS-CoV-2 S-2P, RBD, NTD probes.
- 1222 Representative data for sorting Spike double positive, Spike<sup>+</sup> or NTD<sup>+</sup>, as well as  $RBD^+$  or NTD<sup>+</sup> subsets were
- 1223 shown.
- 1224 (E-H) Neutralization activity of RBD Abs. (E) Proportion of SARS-CoV-2 RBD Abs (n=81) that exhibited
- 1225 detectable neutralization in the microneutralization assay. (F) Neutralization IC<sub>50</sub> and IC<sub>80</sub> of RBD neutralizing Abs
- 1226 (NAbs) against pseudotyped SARS-CoV-2. (G) Microneutralization titer, plaque reduction neutralization test
- 1227 (PRNT) IC<sub>50</sub> and IC<sub>80</sub> of RBD NAbs against replication-competent SARS-CoV-2. Microneutralization titer was
- defined as the lowest Ab concentration that neutralize all the virus, or 99% inhibitory concentration ( $IC_{99}$ ). Abs
- 1229 with undetectable microneutralization titers are shown as gray symbols and nAbs are represented by blue symbols.
- 1230 (H) RBD NAbs blocking of ACE2 binding to SARS-CoV-2 Spike (S) protein. Blocking titer is shown as IC<sub>50</sub>.
- 1231 (I-J) Correlation analysis of RBD Abs between neutralization and ACE2 blocking activities. Spearman correlation
- 1232 analysis were performed for (I) ACE2 blocking IC50 vs. PV neutralization IC50, as well as (J) for ACE2 blocking
- 1233 IC50 vs. SARS-CoV-2 neutralization titers (indicated by the lowest concentration that shows no CPE). Purified
- 1234 RBD Abs in Table S1 and S2 that have pseudovirus neutralization data (n=59) or SARS-CoV-2 micro-
- 1235 neutralization assay data (n=80) were used in this analysis. P-value and r were indicated for each figures.
- 1236 (K-M) Neutralization activity of NTD Abs. (K) Proportion of SARS-CoV-2 NTD Abs (n=41) that exhibited
- 1237 detectable neutralization in the microneutralization assay. (L) Neutralization IC<sub>50</sub> and IC<sub>80</sub> of NTD neutralizing Abs

1238	against pseudotyped SARS-CoV-2. (M) Microneutralization titer, PRNT IC <sub>50</sub> and IC <sub>80</sub> of NTD neutralizing Abs
1239	against replication-competent SARS-CoV-2. Abs with undetectable microneutralization titers are shown as gray
1240	symbols and neutralizing Abs are represented by orange symbols. Horizontal bars represent the geometric means
1241	for each group of Abs.

1242

# 1243 Figure S2. Binding and neutralization activities of down-selected SARS-CoV-2 Abs. Related to Figure 2.

- 1244 (A-D) ELISA binding curves of down-selected Abs. Different SARS-CoV-2 or other CoV viral antigens were
- 1245 coated on plates and detected with serial diluted (A) RBD infection-enhancing Abs, (B) RBD non-infection-
- 1246 enhancing Abs, (C) NTD infection-enhancing Abs, and (D) NTD non-infection-enhancing Abs.
- 1247 (E-F) Neutralization curves for RBD Abs against pseudotyped (E) and replication-competent (F) SARS-CoV-2.
- 1248 (G-H) Neutralization curves for NTD Abs against pseudotyped (G) and replication-competent (H) SARS-CoV-2.
- 1249 (I-L) Neutralization curves for cross-neutralizing Abs against pseudotyped (I) and replication-competent (J) SARS-
- 1250 CoV-2, SARS-CoV-1 nanoluciferase (nLuc) virus (L), and Bat WIV1-CoV nLuc virus (L).
- 1251

# 1252 Figure S3. Comparison of RBD and NTD epitopes from NSEM. Related to Figure 2.

- 1253 (A) A spike model (PDB 6ZGE) and corresponding Fab homology models were manually docked and rigidly fit
- 1254 into each negative stain density map.
- (B) The RBD of each model is enlarged and shown as a white surface, with the putative epitope of each Ab colored.Black outline indicates the ACE2 binding footprint.
- (C) Comparison to ACE2 footprint and epitopes of three published Abs with similar epitopes. See main text forreferences.
- 1259 (D) A spike model (PDB 6ZGE) and corresponding Fab homology models were manually docked and rigidly fit
- 1260 into each negative stain density map.
- 1261 (E) The NTD of each model is enlarged and shown as a white surface, with the epitope of each Ab colored. Orange
- 1262 outline indicates the epitope of Ab 4A8, shown at bottom right. Outlines illustrate that the neutralizing Abs
- 1263 DH1048-51 share the same epitope, whereas the infection-enhancing Abs DH1053-56 bind a distinct epitope.

1264 (F) The model of spike complex with Fab 4A8 (orange ribbons, PDG 7C2L) is rigidly fit into each of the NSEM

1265 maps (transparent surfaces). The close fit of 4A8 into DH1049, DH1050.1 and DH1050.2 indicate theses have the

- same approach angle as 4A8, whereas DH1048 and DH1051 have slightly different approaches.
- 1267

# 1268 Figure S4. In vitro analysis of human Abs and SARS-CoV-2 infected serum samples. Related to Figure 3,

# 1269 **Figure 5 and Figure 7.**

- 1270 (A-C) Effect of combining infection-enhancing RBD and NTD Abs on SARS-CoV-2 pseudovirus infection in
- 1271 ACE2-expressing cells. The infection-enhancing NTD Ab DH1052 was tested alone (A) or mixed with infection-
- 1272 enhancing RBD Abs DH1041 (B) or DH1043 (C) in 1:13 ratio or 1:13250 ratio, respectively. The NTD:RBD Ab
- 1273 mixtures (orange), as well as RBD Ab alone (blue), were five-fold serially diluted and tested for neutralization
- against SARS-CoV-2 D614G pseudovirus in 293T/ACE2 cells.
- 1275 (D-F) Comparison of RBD and NTD directed serum Ab responses in SARS-CoV-2 infected humans.
- 1276 (D) Serum IgG binding titers to RBD (blue) and NTD (salmon) as measured by ELISA as log area-under-curve
- 1277 (AUC). Each symbol represents an individual study participant, with the mean binding titer for the visit day shown
- 1278 as a black horizontal bar.
- 1279 (E) Percent decrease in binding to NTD relative to RBD binding titer. Each symbol represents the change in
- 1280 binding titer for an individual study subject. Mean decrease is shown as a black horizontal bar.
- 1281 (F) Serum blocking of RBD neutralizing Ab DH1041 (blue) or NTD neutralizing Ab DH1050.1 (salmon), or non-
- 1282 neutralizing Ab DH1052 (burgundy) binding to SARS-CoV-2 spike. Black symbols show individual study
- 1283 participants. Mean blocking percentage for the visit day is shown as a filled bar.
- 1284 (G-H) Neutralization activities of neutralizing and enhancing Abs against mouse adapted SARS-CoV-2.
- 1285 (G) NTD neutralizing Abs DH1050.1, RBD neutralizing and enhancing Abs DH1041 were tested for neutralization
- 1286 activities against wild-type (WT) virus, mouse adapted 2AA MA virus, and mouse adapted MA10 virus in live
- 1287 virus neutralization assay. CH65 Ab was used as a control. Mean value of neutralization (%) from duplicate wells
- 1288 were shown.
- 1289 (H) NTD enhancing Ab DH1052 and control Ab CH65 were tested for neutralization activities against wild-type
- 1290 (WT) virus, mouse adapted 2AA MA virus, and mouse adapted MA10 virus in live virus neutralization assay.
- 1291 Mean value of neutralization (%) from duplicate wells were shown.

1292

# Figure S5. Lung histopathology of Ab-treated and SARS-CoV-2 challenged cynomolgus macaques. Related to Figure 5 and Figure 7.

- 1295 (A) Representative images of hematoxylin and eosin (H&E) staining and SARS-CoV-2 antigen
- 1296 immunohistochemistry (IHC) staining from each group. All images were taken at 10x magnification. The images in
- 1297 this presentation are representative of the average severity of pathologic processes observed and recorded during
- 1298 microscopic evaluation. Red arrows indicate SARS-CoV-2 infection foci.
- 1299 (B) Following microscopic evaluation of DH1052, 1 animal (BB536A) out of 5 animals in this group exhibited
- 1300 histologic features that was substantially more severe than the rest of the cohort and may suggest some degree of
- 1301 Ab-mediated disease enhancement. The features were characterized by prominent perivascular mononuclear
- 1302 inflammation (\*) and a substantial amount of perivascular and alveolar edema (fluid; X). These findings suggest a
- 1303 vaso-centric process with some degree of altered vascular permeability. The remaining 4 animals in DH1052 group
- 1304 had inflammatory changes that ranged from minimal to moderate severity and more infiltrates were mixed and
- 1305 predominantly polymorphonuclear with lesser mononuclear cell involvement and present in the alveolar spaces.
- 1306 (C-E) Expression of macrophage activation markers in macaque lung tissues. An animal from the CH65 control
- 1307 group (C), the DH1052-treated animal (BB536A) that exhibited substantially more severe lung inflammation (D),
- 1308 and an animal from the NTD NAb DH1050.1 group (E) were selected for Immunohistochemistry (IHC) staining.
- 1309 Immunohistochemical staining was performed using MHCII, CD68, IBA1 and CD163 to detect classically
- 1310 activated macrophages (M1) and/or alternatively activated macrophages (M2). CD11b is a macrophage/monocyte
- 1311 marker and CD3 is a T cell marker. All images are 10x magnification; scale bars=  $100\mu$ m.
- 1312

# Figure S6. High dose NTD enhancing Ab DH1052 does not enhance SARS-CoV-2 replication or disease *in vivo*. Related to Figure 5.

- 1315 (A) Diagram of the macaque study design showing cynomolgus macaques (n=5 per group) were infused with high
- 1316 dose (30 mg/kg body weight) DH1052 or an irrelevant control CH65 Ab 3 days before 10<sup>5</sup> PFU of SARS-CoV-2
- 1317 challenge via intranasal and intratracheal routes. Viral load including viral RNA and subgenomic RNA (sgRNA)
- 1318 were measured at the indicated pre-challenge and post-challenge timepoints. Lungs were harvested on Day 4 post-
- 1319 challenge for histopathology analysis.

- 1320 (B-D) SARS-CoV-2 (B) E gene sgRNA, (C) N gene sgRNA and (D) E gene total viral RNA in bronchoalveolar
- 1321 lavage (BAL) on Day 2 and Day 4 post challenge.
- (E-G) SARS-CoV-2 (E) E gene sgRNA, (F) N gene sgRNA and (G) E gene total viral RNA in nasal swab on Day 2
  and Day 4 post challenge.
- 1324 (H-I) Lung inflammation. Sections of the left caudal (Lc), right middle (Rm), and right caudal (Rc) lung were
- evaluated and scored for the presence of inflammation by hematoxylin and eosin (H&E) staining. (H) Summary of
- 1326 inflammation scores. Symbols indicate the sums of Lc, Rm, and Rc scores in each animal. (I) Representative
- 1327 images of lung H&E staining.
- 1328 (J-K) Immunohistochemistry (IHC) staining for the presence of SARS-CoV-2 nucleocapsid in lungs. (J) Summary
- 1329 of IHC scores. Symbols indicate the sums of Lc, Rm, and Rc scores in each animal. (K) Representative images of
- 1330 lung IHC staining. Red arrows indicate SARS-CoV-2 infection foci.
- 1331 LOD, limit of detection. Statistical significance in all the panels were determined using Wilcoxon rank sum exact
- 1332 test. Asterisks show the statistical significance between indicated group and CH65 control group: ns, not significant,
- 1333 \*P<0.05, \*\*P<0.01, \*\*\*P<0.001.
- 1334

# 1335 Figure S7. Different doses of a cross-neutralizing Ab DH1047 treatments do not enhance SARS-CoV-2

- 1336 replication *in vivo*. Related to Figure 7.
- 1337 (A) Diagram of the macaque study design. Cynomolgus macaques (n=5 per group) were infused with DH1047 at
- the dose of 10 mg/kg, 5 mg/kg, 1 mg/kg, 0.1 mg/kg weight. Macaques treated with 10 mg/kg weight of DH65 Ab
- 1339 were set as the control group. Three days post-infusion,  $10^5$  PFU of SARS-CoV-2 challenge via intranasal and
- 1340 intratracheal routes. Viral load including viral RNA and subgenomic RNA (sgRNA) were measured at the
- 1341 indicated pre-challenge and post-challenge timepoints. Lungs were harvested on Day 4 post-challenge for
- 1342 histopathology analysis.
- 1343 (B) Serum human IgG concentrations at Day 2.
- 1344 (C) Day 2 serum neutralization titers shown as the reciprocal serum dilution that inhibits 50% (ID<sub>50</sub>) of SARS-
- 1345 CoV-2 replication in Vero cells.
- 1346 (D-E) SARS-CoV-2 (D) E gene sgRNA and (E) N gene sgRNA in bronchoalveolar lavage (BAL) on Day 2 and
- 1347 Day 4 post challenge.

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348	(F-G) SARS-CoV-2 (F) E gene sgRNA and (G) N gene sgRNA in nasal swab on Day 2 and Day 4 post challenge.								
349	(H-I) Lung inflammation. Sections of the left caudal (Lc), right middle (Rm), and right caudal (Rc) lung were								
350	evaluated and scored for the presence of inflammation by hematoxylin and eosin (H&E) staining. (H) Summary of								
351	inflammation scores. Symbols indicate the sums of Lc, Rm, and Rc scores in each animal. (I) Representative								
352	images of lung H&E staining.								
353	(J-K) Immunohistochemistry (IHC) staining for the presence of SARS-CoV-2 nucleocapsid in lungs. (J) Summary								
354	of IHC scores. Symbols indicate the sums of Lc, Rm, and Rc scores in each animal. (K) Representative images of								
355	lung IHC staining. Red arrows indicate SARS-CoV-2 infection foci.								
356	LOD, limit of detection. Statistical significance in all the panels were determined using Wilcoxon rank sum exact								
357	test. Asterisks show the statistical significance between indicated group and CH65 control group: ns, not significan								
358	*P<0.05, **P<0.01, ***P<0.001.								
359									
360									
361	Supplementary Items (available online as excel sheet or PDF format)								
362									
363	Table S1. High-throughput ELISA binding screen of Abs recovered from SARS-CoV-2 and SARS-CoV-1								
364	donors. Related to Figure 1.								
365									
366	Supplementary Table S2. Immunogenetic analysis of select neutralizing and non-neutralizing SARS-CoV-2								
367	Abs. Related to Figure 2.								
368									
369	Table S3. RBD and NTD Fabs affinity for Spike proteins. Related to Figure 2.								
370									
371	Table S4. RBD and NTD Ab affinity for mouse CD16/FcγR3, CD16-2/FcγR4, CD32B/FcγR2b and								
372	CD64/FcγR1. Related to Figure 5 and Figure 6.								
373									
374	Table S5. Luminex cytokine profiling of BAL samples from cynomolgus macaques. Data set related to								
375	Figure 5 and 7, Figure S6 and Figure S7 were shown in different tabs. BAL samples collected on Day -5 (pre-								

- 1376 challenge), Day 2 and Day 4 post-challenge were concentrated (x10) and measured using a 25-analyte multiplex
- 1377 bead array by Luminex assay. The animal (BB536A) in DH1052 group that exhibited substantially more severe
- 1378 disease and cytokine expression was marked in red.
- 1379
- 1380 Supplementary Data 1. Cryo-EM information. Related to Figure 4.
- 1381

Journal Proproof

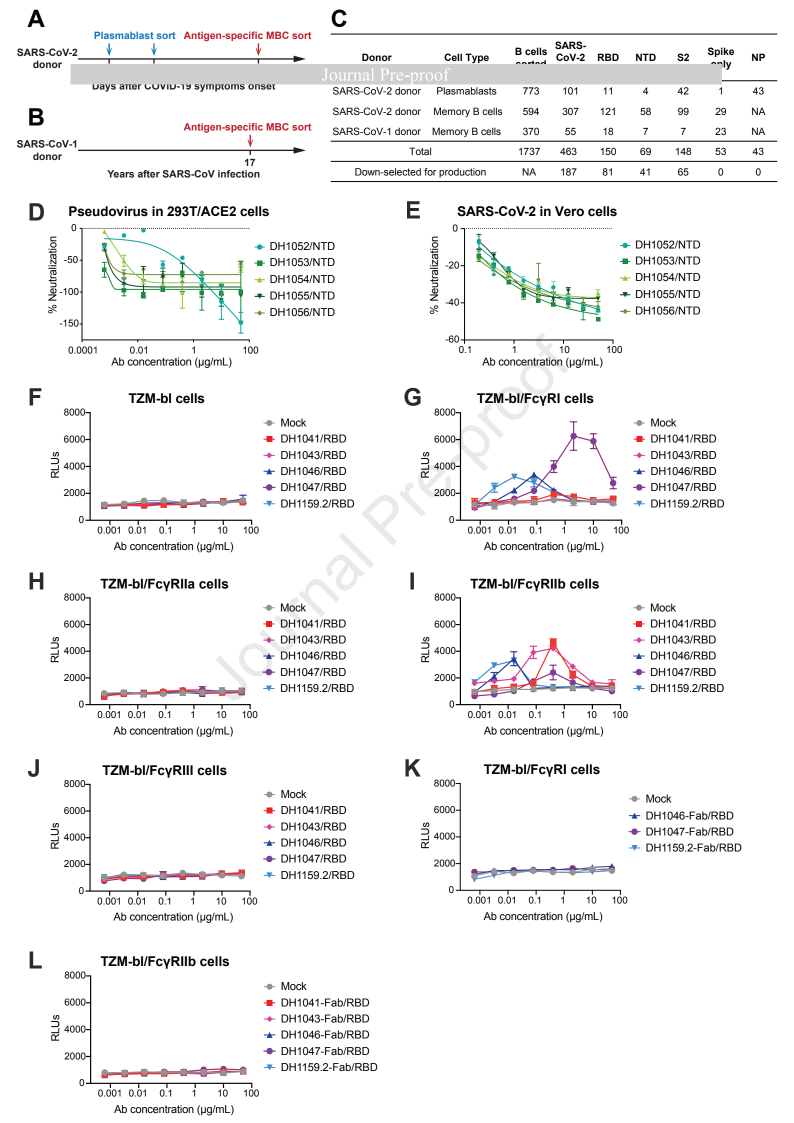
# **Highlights:**

- RBD or NTD antibodies exhibited infection enhancement in vitro but not in vivo
- Neutralizing or infection-enhancing NTD antibodies bound distinct epitopes
- In vitro infection-enhancing antibodies protected from SARS-CoV-2 in vivo
- Cross-reactive RBD neutralizing antibodies were protective--most potent, DH1047

# eTOC Blurb:

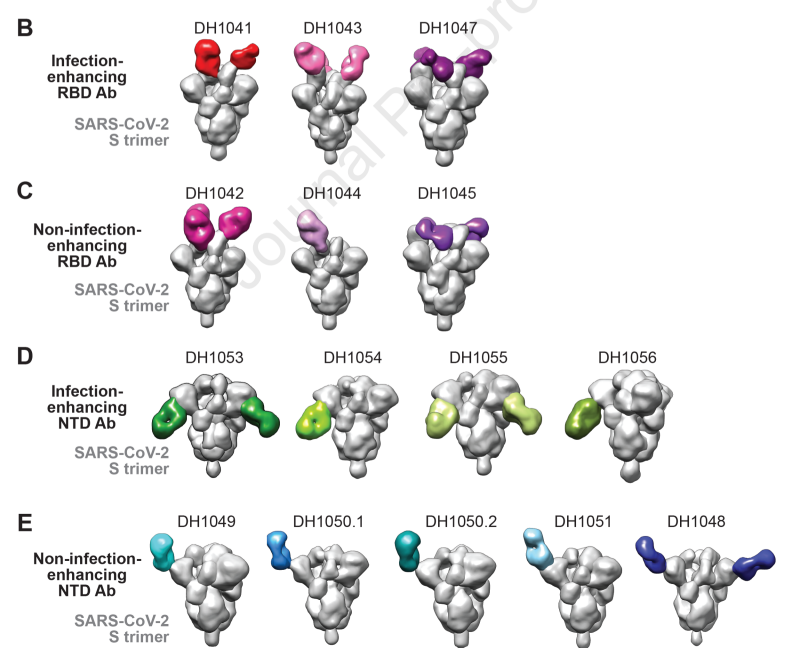
Convalescent human-derived SARS-CoV-2 RBD and NTD antibodies mediated neutralization as well as infection-enhancement *in vitro*, yet infusion of these antibodies in mice or cynomolgus macaques resulted in protection from viral replication.

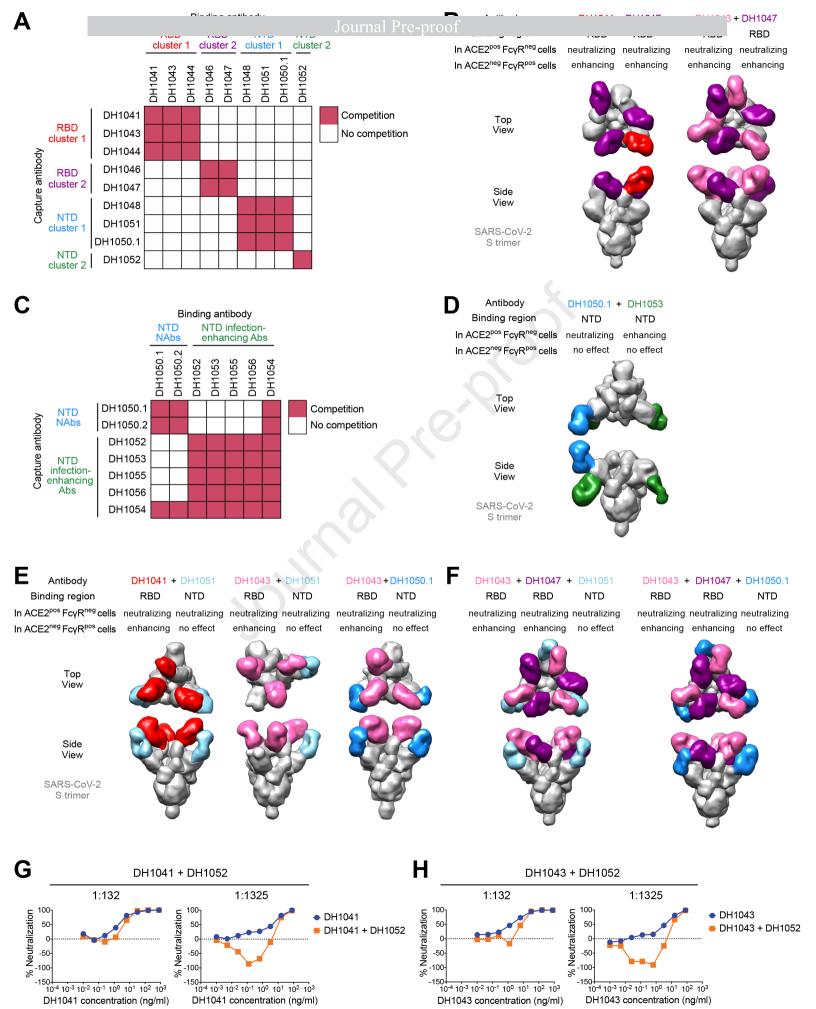
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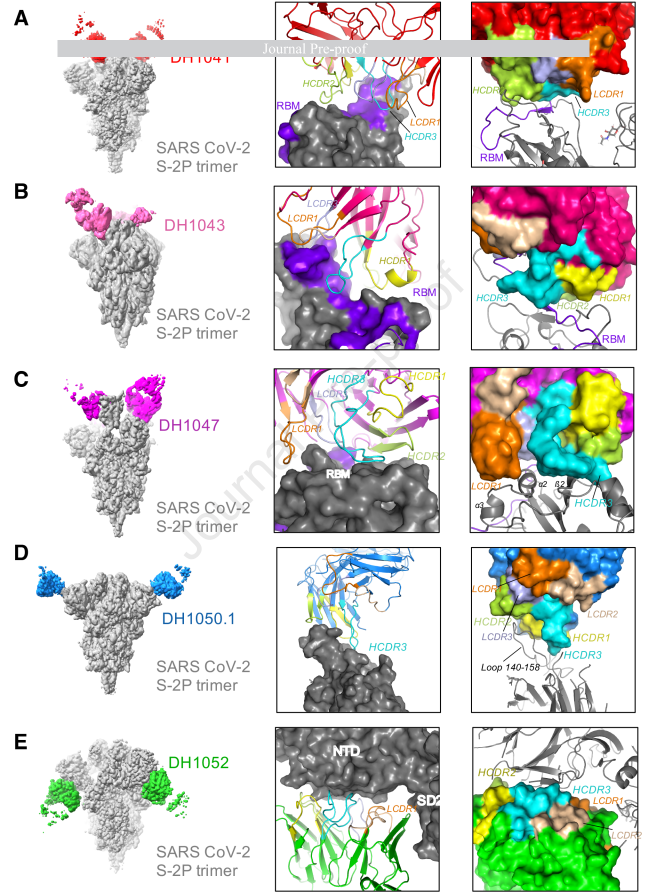


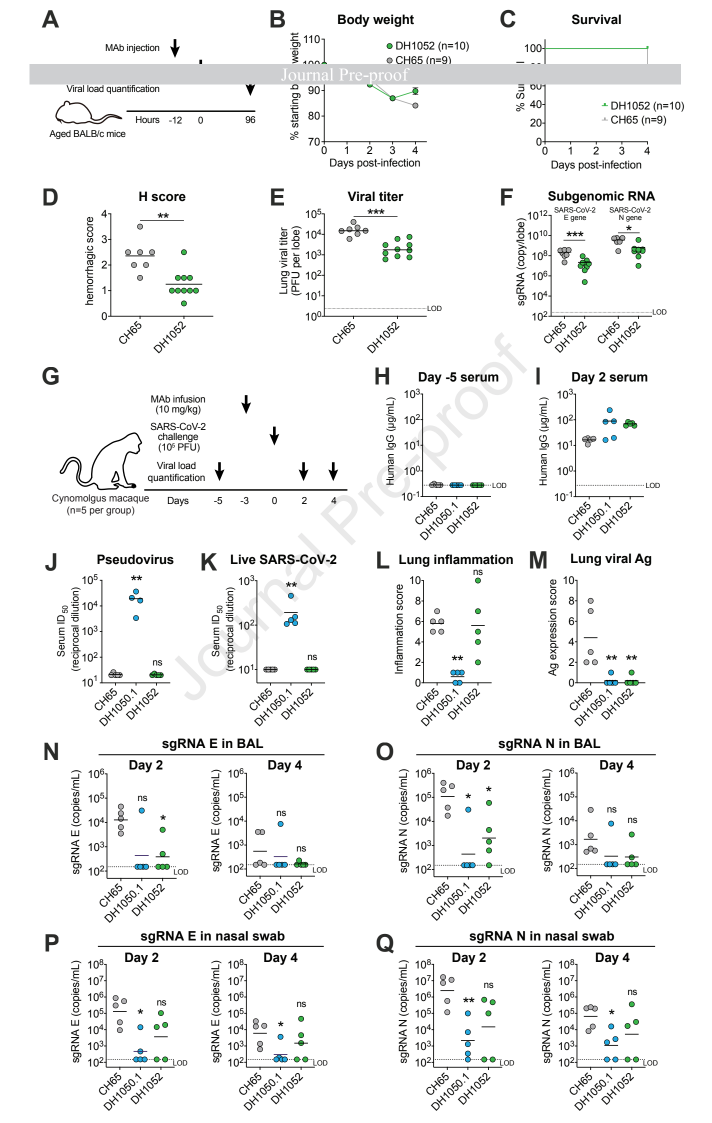
Α

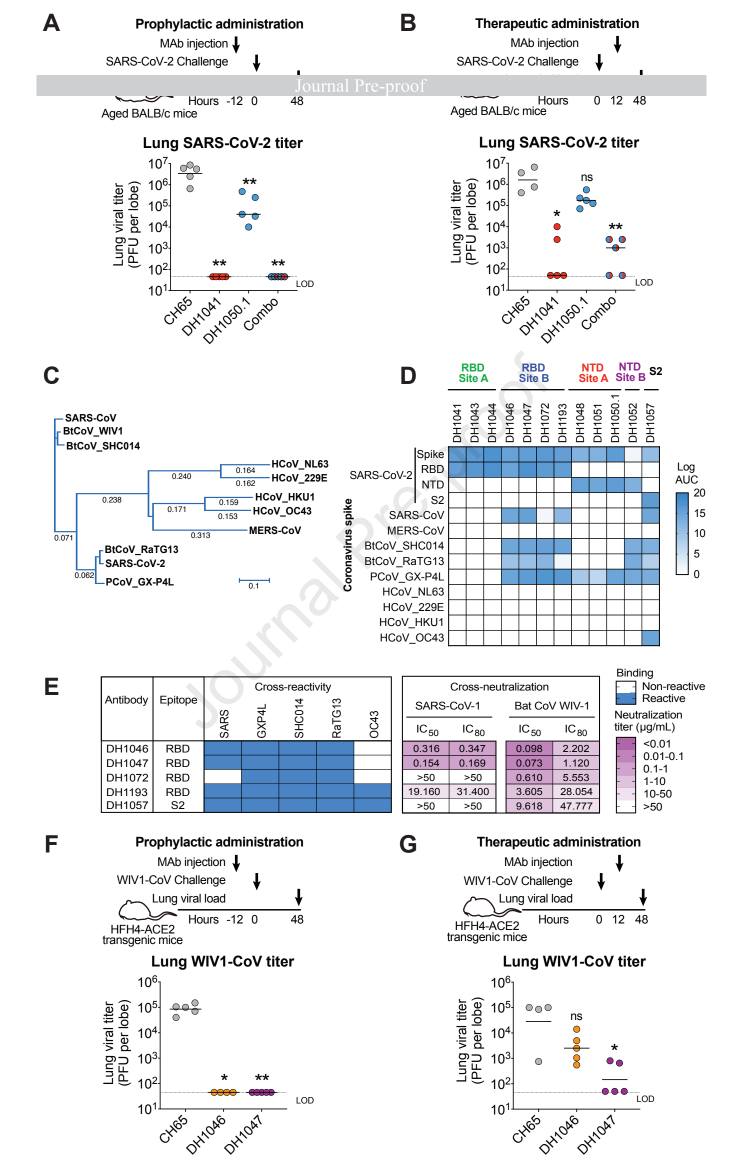
		Effect on SARS- CoV-2 infection	Cross- reactivity with	ACE2 blocking		SARS-CoV-2 pseudovirus			SARS-CoV-2 replication competent virus			Effect on SARS- CoV-2 infection	
Antibody	Epitope	FcyR <sup>neg</sup>	ACE2 <sup>neg</sup> FcγR <sup>pos</sup> cells	SARS -CoV-1	IC 50	IC <sub>80</sub>	IC <sub>50</sub>	IC <sub>80</sub>	Maximal% inhibition	MN titer	IC 50	IC <sub>80</sub>	No effect Infection-enhancing
DH1041	RBD				0.036	0.497	0.017	0.049	>100	0.098	0.016	0.063	Neutralizing
DH1043	RBD				0.043	0.280	0.0015	0.020	>100	0.098	0.034	0.099	
DH1159.2	RBD				>50	>50	11.900	>50	77.0	>100	ND	ND	Cross-reactivity
DH1046	RBD				0.201	6.512	0.396	2.030	>100	12.500	1.086	9.383	Non-reactive
DH1047	RBD				0.078	0.567	0.090	0.360	>100	1.100	0.124	0.666	Reactive
DH1042	RBD				0.059	0.371	0.011	0.053	>100	0.280	0.071	0.269	Reactive
DH1044	RBD				>50	>50	0.021	0.080	98.0	0.550	0.076	0.273	
DH1045	RBD				0.226	24.353	0.380	2.260	>100	6.250	1.437	4.827	Neutralization titer
DH1052	NTD				>50	>50	>50	>50	-148.0	>100	>100	>100	or blocking titer
DH1053	NTD				>50	>50	>50	>50	-99.0	>100	>100	>100	(µg/mL)
DH1054	NTD				>50	>50	>50	>50	-63.0	>100	>100	>100	
DH1055	NTD				>50	>50	>50	>50	-106.0	>100	>100	>100	<0.01
DH1056	NTD				>50	>50	>50	>50	-56.0	>100	>100	>100	0.01-0.1
DH1048	NTD				>50	>50	0.520	>50	72.0	0.390	0.608	2.232	0.1-1
DH1049	NTD				>50	>50	>50	>50	49.0	0.390	0.385	3.539	1-10
DH1050.2	NTD				>50	>50	0.280	>50	67.0	0.780	0.087	1.187	10-50
DH1051	NTD				>50	>50	0.049	>50	68.0	0.780	0.134	0.737	>50
DH1050.1	NTD				>50	>50	0.039	>50	62.0	0.780	0.161	0.614	

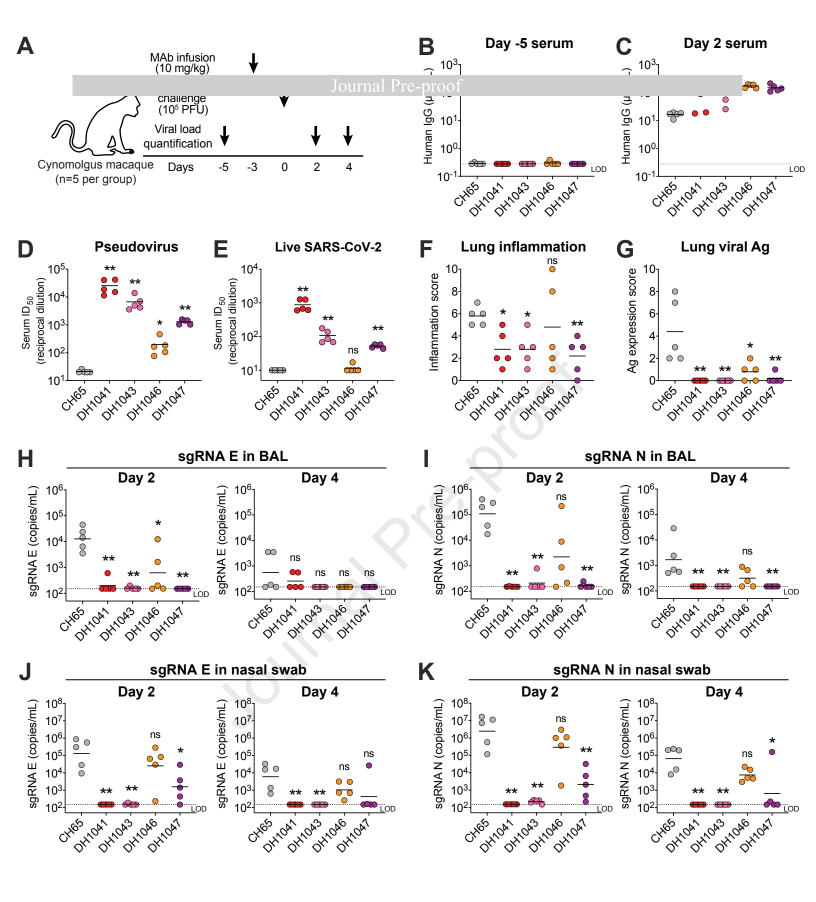


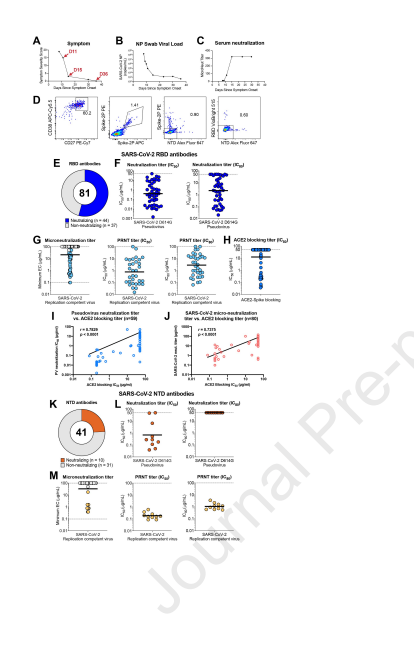


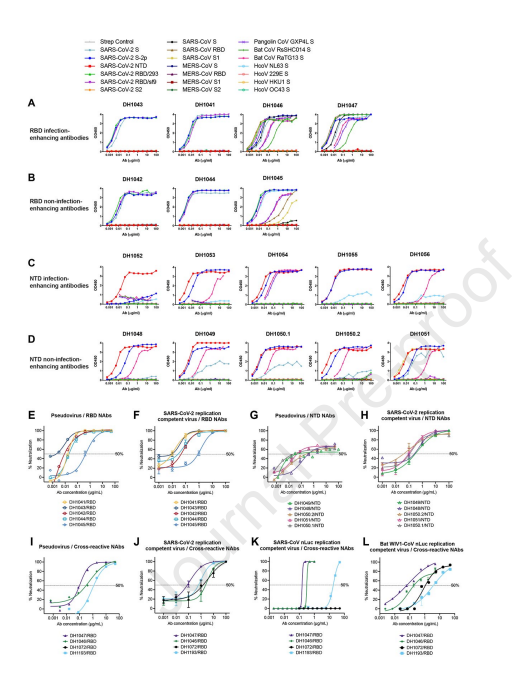








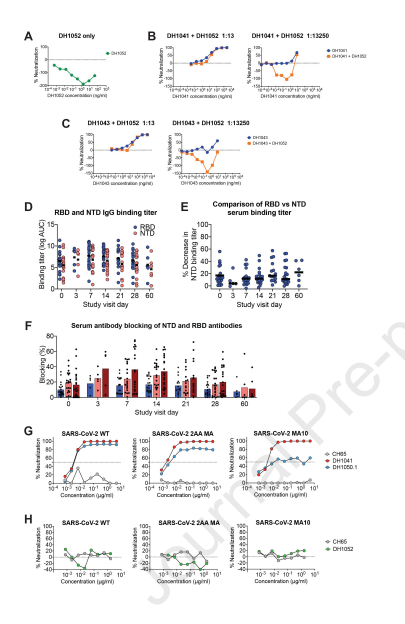




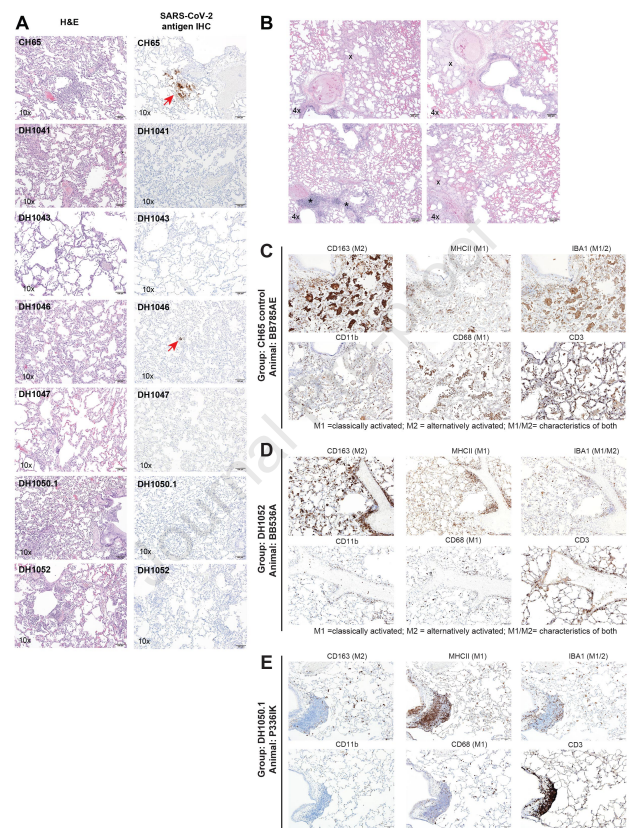
Journal Pre-proof



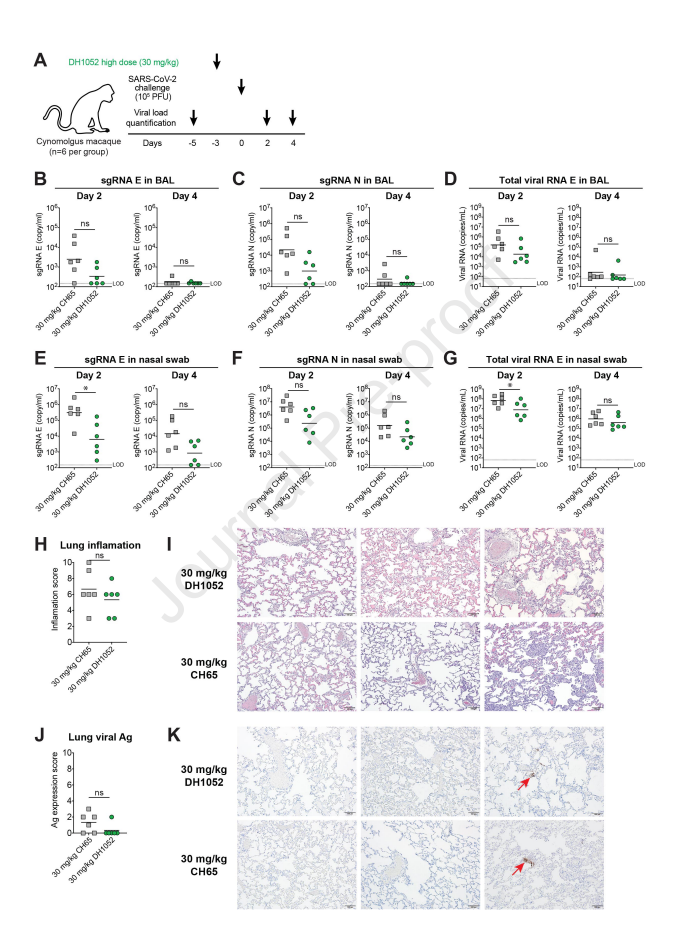
### Iournal Pre-proof



### Journal Pre-proof

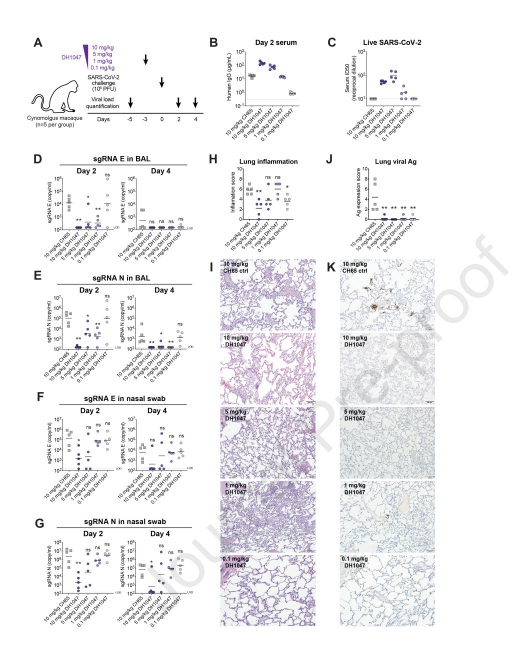


M1 =classically activated; M2 = alternatively activated; M1/M2= characteristics of both



Journal Prevention

## Journal Pre-proof

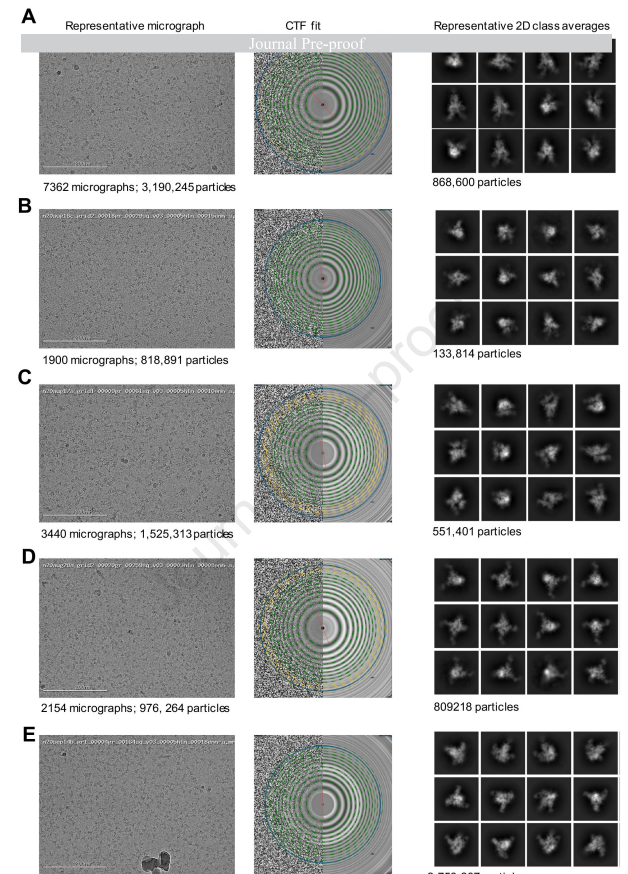


# 1. Cryo-EM data collection and refinement statistics.

	nal Pre-proc	of			_
	24MFK	30MFK	35MFK	113KJ	26MFK
	DH1041	DH1052	DH1047	DH1043	DH1050.1
PDB ID	7LAA	7LAB	7LD1	7LJR	7LCN
EMDB ID	EMD- 23246	EMD- 23248	EMD- 23279	EMD- 23400	EMD- 23277
Data Collection	1				
Microscope	FEI Titan Krios				
Voltage (kV)	300	300	300	300	300
Electron dose $(e^{-}/A^2)$	65.94	66.71	66.77	66.77	65.09
Detector	Gatan K3				
Pixel Size (Å)	1.058	1.058	1.058	1.058	1.058
Defocus Range (µm)	~0.75-2.50	~0.75-2.50	~0.75-2.50	~0.75-2.50	~0.75-2.5
Magnification	81000	81000	81000	81000	81000
Micrographs collected	7362	9375	3440	1900	2154
Reconstruction		0			
Software	cryoSparc	cryoSparc	cryoSparc	cryoSparc	cryoSpar
Particles	151,384	143,4115	127,401	133,814	426,025
Symmetry	C1	C1	C1	C1	C1
Box size (pix)	350	350	350	350	350
Resolution (Å) (FSC 0.143)*	3.42	2.97	3.4	3.66	3.35
Refinement (Phenix)					
Protein residues	2994	4194	4251	3378	4296
Chimera CC	0.72	0.77	0.79	0.68	0.73
R.m.s. deviations					
Bond lengths (Å)	0.014	0.014	0.012	0.012	0.013
Bond angles (°)	2.235	1.958	1.920	1.899	2.006
Validation					
Validation Molprobity score	1.99	1.25	1.41	1.17	1.53
	1.99 2.19	1.25 0.40	1.41 0.15	1.17 0.23	1.53 0.42
Molprobity score					
Molprobity score Clash score	2.19	0.40	0.15	0.23	0.42
Molprobity score Clash score Favored rotamers (%)	2.19	0.40	0.15	0.23	0.42
Molprobity score Clash score Favored rotamers (%) Ramachandran	2.19 96.62	0.40 98.41	0.15 97.65	0.23 98.88	0.42 97.48
Molprobity score Clash score Favored rotamers (%) Ramachandran Favored regions (%)	2.19 96.62 88.15	0.40 98.41 92.97	0.15 97.65 89.16	0.23 98.88 90.03	0.42 97.48 89.05

\*Resolutions are reported according to the FSC 0.143 gold-standard criterion

## 2. Cryo-EM data processing details.

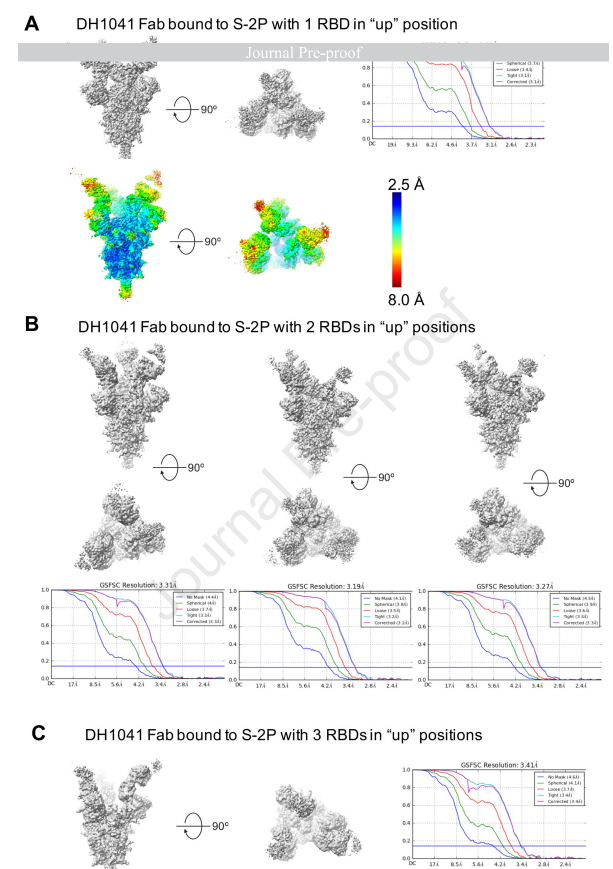


9375 micrographs; 4,412, 857 particles

2,758,267 particles

**2. Cryo-EM data processing details**. (left) Representative micrograph, (middle) CTF fit and (right) Representative 2D class averages for (A) DH1041-Spike-2P (S2P) complex, (B) DH1043-S2P complex, (C) DH1047-S2P complex, (D) DH1050.1-S2P complex, (E) DH1052-S2P complex.

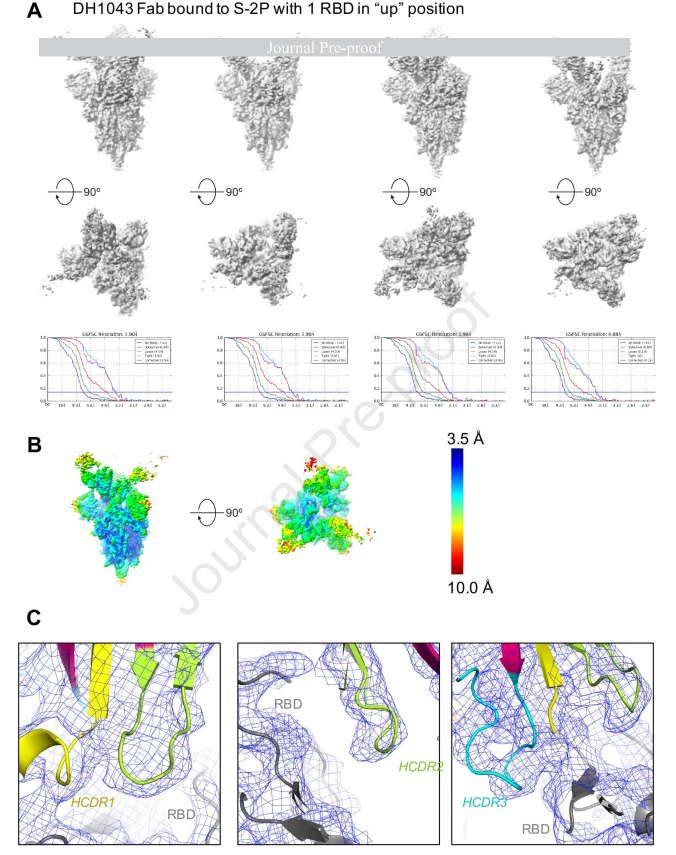
## 3. Global and Local map resolutions for DH1041/S-2P complex.



3. Global and Local map resolutions for DH1041/S-2P complex. (A) Cryo-EM reconstruction of DH1041 bound to 1-RBD-up 2P spike. Top row show refined map and FSC curves. Bottom row shows refined colored by local resolution. Zoomed-in view of the S2 region is shown on the right with cryo-EM map shown as blue mesh and underlying fitted model as sticks and colored by element (B) Cryo-EM reconstruction of DH1041 bound to 2-RBD-up 2P spike. (C) Cryo-EM reconstruction of DH1041 bound to 3-RBD-up 2P spike.

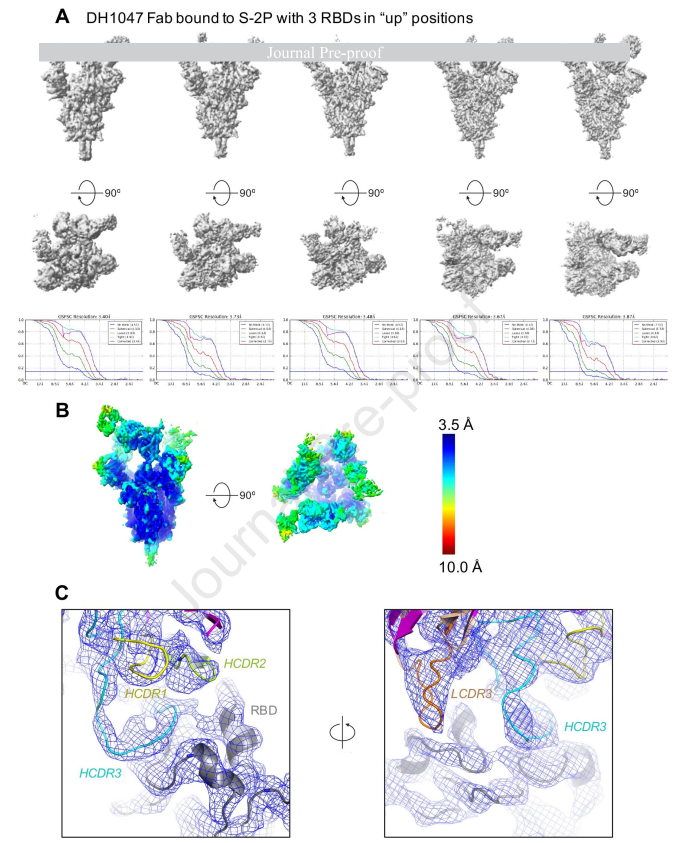
17Å 8.5Å 5.64 4.2Å 3.4Å 2.8Å 2.4Å

4. Global and Local map resolutions for DH1043/S-2P complex.



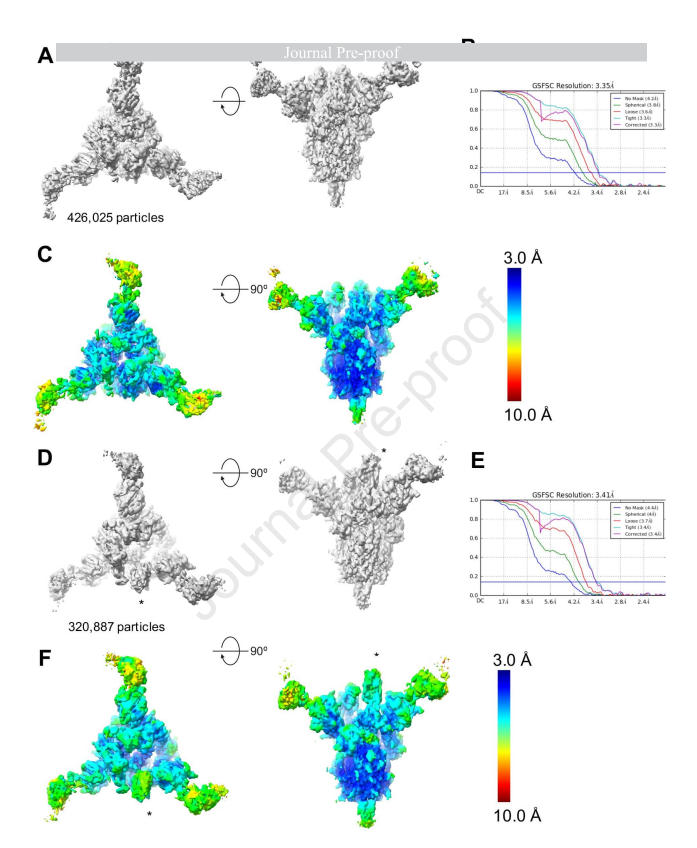
**4. Global and Local map resolutions for DH1043/S-2P complex.** (A) Cryo-EM reconstructions of DH1043 bound to 1-RBD-up 2P spike Top two rows show refined maps, bottom row shows the FSC curve for each corresponding map. (B) Left. Refined cryo-EM map that was used for model building colored by local resolution. Right. Zoomed-in view of the S2 region with cryo-EM map shown as blue mesh and underlying fitted model as sticks and colored by element. (C) Zoomed-in view of the DH1043 interface with RBD. The cryo-EM map is shown as a blue mesh with underlying fitted model shown in cartoon representation, with the DH1047 HCDR1 loop colored yellow, HCDR2 colored limon, HCDR3 cyan, and LCDR3 light blue.

# 5. Global and Local map resolutions for DH1047/S-2P complex.



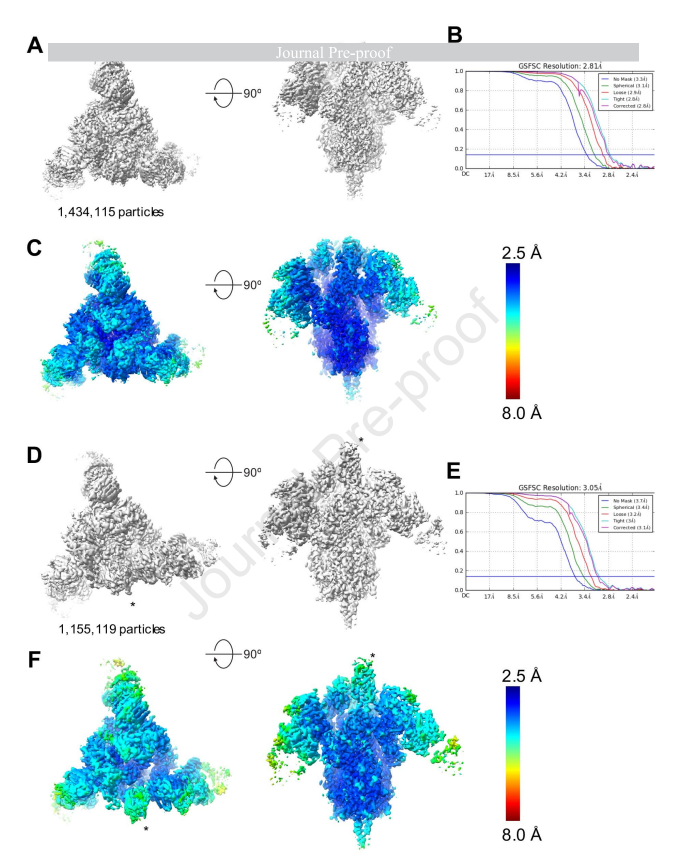
**5. Global and Local map resolutions for DH1047/S-2P complex.** (A) Cryo-EM reconstructions of DH1047 bound to 3-RBD-up 2P spike. Top two rows show refined maps, bottom row shows the FSC curve for each corresponding map. (B) Left. Refined cryo-EM map that was used for model building colored by local resolution. Right. Zoomed-in view of the S2 region with cryo-EM map shown as blue mesh and underlying fitted model as sticks and colored by element. (C) Zoomed-in view of the DH1047 interface with RBD. The cryo-EM map is shown as a blue mesh with underlying fitted model shown in cartoon representation, with the DH1047 HCDR1 loop colored yellow,HCDR2 colored limon, HCDR3 cyan and LCDR1 orange.

6. Global and Local map resolutions for DH1050.1/S-2P complex.



**6. Global and Local map resolutions for DH1050.1/S-2P complex.** (A) Cryo-EM reconstruction of DH1050.1 bound to 3-RBD-down 2P spike. (B) Fourier shell correlation curves. (C) Left. Refined cryo-EM map colored by local resolution for the DH1050.1 bound to 3-RBD-down 2P spike. Right. Zoomed-in view of the S2 region with cryo-EM map shown as blue mesh and underlying fitted model as sticks and colored by element. (D) Cryo-EM reconstruction of DH1050.1 bound to 1-RBD-up 2P spike. (E) Fourier shell correlation curves. (F) Refined cryo-EM map colored by local resolution for the DH1050.1 bound to 1-RBD-up 2P spike.

7. Global and Local map resolutions for DH1052/S-2P complex.



**7. Global and Local map resolutions for DH1052/S-2P complex.** (A) Cryo-EM reconstruction of DH1052 bound to 3-RBD-down stabilized Spike "2P" (S-2P). (B) Fourier shell correlation curves. (C) Left. Refined cryo-EM map colored by local resolution for the DH1052 bound to 3-RBD-down S-2P. Right. Zoomed-in view of the S2 region with cryo-EM map shown as blue mesh and underlying fitted model as sticks and colored by element. (D) Cryo-EM reconstruction of DH1052 bound to 1-RBD-up S-2P. (E) Fourier shell correlation curves. (F) Refined cryo-EM map colored by local resolution for the DH1052 bound to 1-RBD-up S-2P.