

1      **Short-term instantaneous prophylaxis and efficient treatment against SARS-**

2      **CoV-2 in hACE2 mice conferred by an intranasal nanobody (Nb22)**

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34 **Abstract** (234 words)

35 Current COVID-19 vaccines need to take at least one month to complete inoculation  
36 and then become effective. Around 51% global population are still not fully vaccinated.  
37 Instantaneous protection is an unmet need among those who are not fully vaccinated.  
38 In addition, breakthrough infections caused by SARS-CoV-2 are widely reported. All  
39 these highlight the unmet need for short-term instantaneous prophylaxis (STIP) in  
40 the communities where SARS-CoV-2 is circulating. Previously, we reported  
41 nanobodies isolated from an alpaca immunized with the spike protein, exhibiting  
42 ultrahigh potency against SARS-CoV-2 and its variants. Herein, we found that Nb22,  
43 among our previously reported nanobodies, exhibited ultrapotent neutralization against  
44 Delta variant with an  $IC_{50}$  value of 0.41 ng/ml (5.13 pM). Furthermore, the crystal  
45 structural analysis revealed that the binding of Nb22 to WH01 and Delta RBDs both  
46 effectively blocked the binding of RBD to hACE2. Additionally, intranasal Nb22  
47 exhibited protection against SARS-CoV-2 Delta variant in the post-exposure  
48 prophylaxis (PEP) and pre-exposure prophylaxis (PrEP). Of note, intranasal Nb22 also  
49 demonstrated high efficacy against SARS-CoV-2 Delta variant in STIP for seven days  
50 administered by single dose and exhibited long-lasting retention in the respiratory  
51 system for at least one month administered by four doses, providing a means of  
52 instantaneous short-term prophylaxis against SARS-CoV-2. Thus, ultrahigh potency,  
53 long-lasting retention in the respiratory system as well as stability at room-temperature  
54 make the intranasal or inhaled Nb22 to be a potential therapeutic or STIP agent against  
55 SARS-CoV-2.

56

57 **Brief summary:** Nb22 exhibits ultrahigh potency against Delta variant in vitro and is  
58 exploited by crystal structural analysis; furthermore, animal study demonstrates high  
59 effectiveness in the treatment and short-term instantaneous prophylaxis in hACE2 mice  
60 via intranasal administration.

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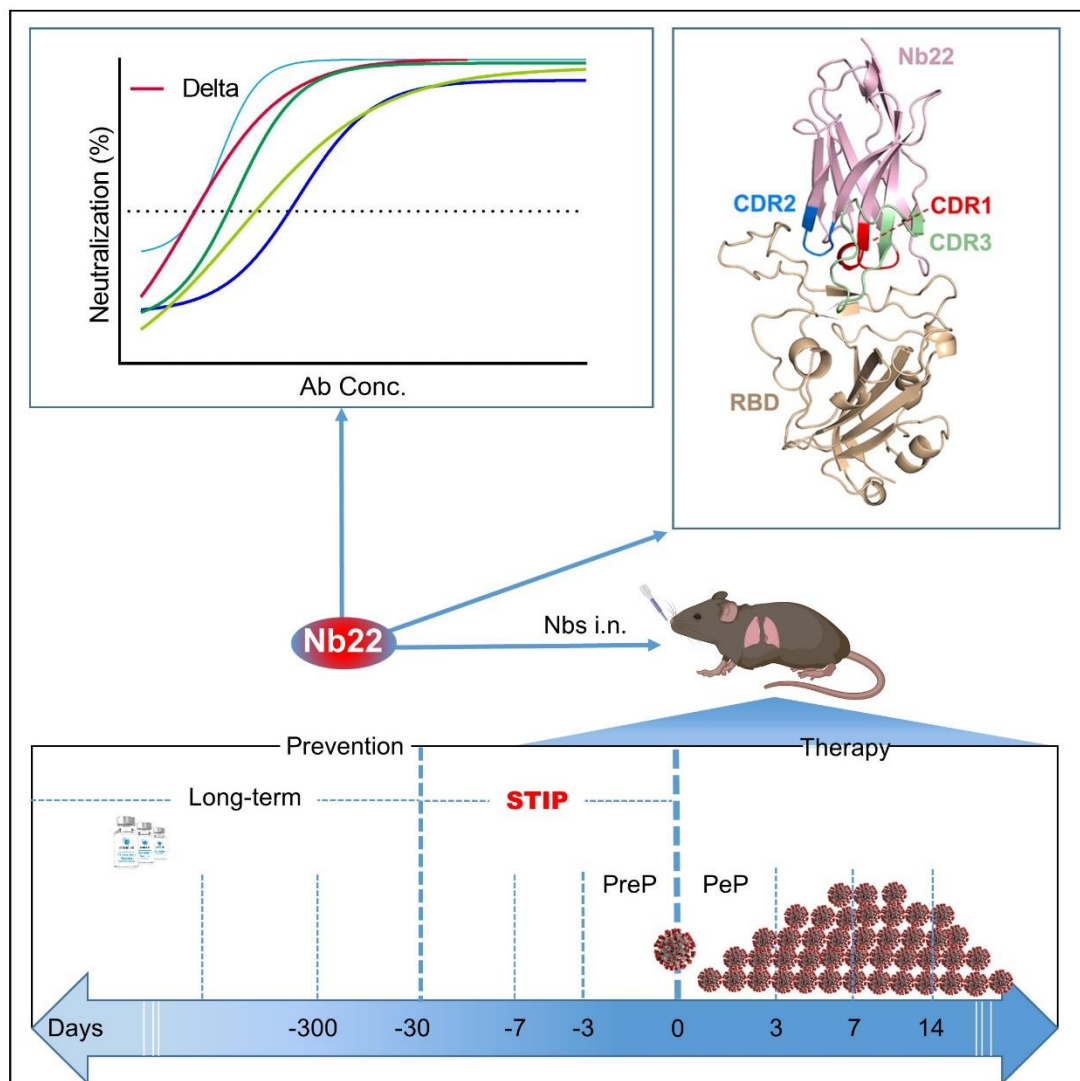
62 **Keywords:** SARS-CoV-2, Delta variant, B.1.617.2, Nanobody, Nb22, STIP, Structure,  
63 Instantaneous prophylaxis, Instantaneous protection

64

65 **Highlights:**

- 66 1) Nb22 exhibits ultrapotent neutralization against Delta variant with an  $IC_{50}$  value of  
67 0.41 ng/ml (5.13 pM).
- 68 2) Structural analysis elucidates the ultrapotent neutralization of Nb22 against Delta  
69 variant.
- 70 3) Nb22 demonstrates complete protection in the treatment of Delta variant infection  
71 in hACE2 transgenic mice.
- 72 4) We complete the proof of concept of STIP against SARS-CoV-2 using intranasal  
73 Nb22 with ultrahigh potency and long-lasting retention in respiratory system.

74 **Graphic Abstract**



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81

82 **Introduction:**

83 SARS-CoV-2 has given rise to the COVID-19 pandemic <sup>1</sup>, resulting in massive  
84 disruption of social and economic activities. Global vaccination has provided protection  
85 against the catastrophic outcome of the pandemic. However, a number of individuals  
86 are either not fully vaccinated or cannot mount adequate responses to the vaccine.  
87 Additionally, current COVID-19 vaccines require multiple doses to achieve full  
88 effectiveness and the immunity wanes within a matter of months, which increases the  
89 risk of infection and demands the use of agents for providing instantaneous protection  
90 at vulnerable times. Several antibodies were approved for emergency use within 7 days  
91 of high-risk exposure in the Post-exposure prophylaxis (PEP) against SARS-CoV-2  
92 infection <sup>2,3</sup>. However, there is no licensed agent in preventing infection before  
93 exposure to SARS-CoV-2 (i.e., as pre-exposure prophylaxis, PrEP). A few PrEP studies  
94 in animal model demonstrated that antibodies exhibited accelerated clearance of SARS-  
95 CoV-2 when administered 1-3 days prior to infection<sup>2,4-6</sup>. The efficacy was not fully  
96 explored when antibodies were administered more than three days prior to SARS-CoV-  
97 2 infection. To the best of our knowledge, there is no effective intervention to prevent  
98 SARS-CoV-2 infection in advance of one week or longer. Therefore, there is a research  
99 gap on short-term instantaneous prophylaxis (STIP) that prevention can take effective  
100 immediately following antibody infusion and last for one week or longer. As such, STIP  
101 is an unmet need for the prevention against SARS-CoV-2.

102

103 The Delta variant, also known as B.1.617.2, was first identified in India in December  
104 2020 and has become predominant in many countries, characterized by the spike protein  
105 mutations T19R, L452R, T478K, D614G, P681R, D950N and a double deletion at 157-  
106 158 <sup>7-11</sup>. It has been designated as a Variant of Concern (VOC) and is believed to be 60%  
107 more transmissible than Alpha variant <sup>12</sup>. Delta variant poses a challenge to the  
108 available COVID-19 vaccines, such as the protective effectiveness of AstraZeneca and  
109 Pfizer vaccines against Delta variant was reduced to 60% and 88%, respectively <sup>11,12</sup>.  
110 More recently, a newly emerged variant, Omicron, has spread rapidly in parts of the  
111 world and drawn attention for its potential impact on the global public health; however,

112 in most of the world including China, Delta variant remains the dominant virus and the  
113 focus of the containment efforts. Recent research indicated that the Delta variant  
114 partially but significantly resisted neutralization by mAbs including Bamlanivimab,  
115 SARS-CoV-2 convalescent sera and vaccine-elicited antibodies <sup>13,14</sup>. While B1-182.1  
116 and A23-58.1, recently isolated from convalescent donors, exhibited ultrapotent  
117 neutralization against Delta variant with IC<sub>50</sub> values of 1.0 and 1.6 ng/ml, respectively  
118 <sup>15</sup>.

119  
120 To date, a growing number of nanobodies, single-domain fragments of camelid heavy-  
121 chain antibodies or VH, were reported for the prophylaxis or treatment of SARS-CoV-  
122 2 infection. However, nanobodies with potent neutralization against Delta variant were  
123 rarely reported <sup>6,16-22</sup>. As SARS-CoV-2 is transmitted through and replicates in  
124 respiratory tract and lungs, and does not transverse in blood<sup>1,23</sup>, we believe that,  
125 nanobodies's exceptional resistance to extreme pH and high temperature<sup>24</sup>, makes them  
126 ideal candidates to be administered via intranasal or oral rout, directly to the site of viral  
127 infection. Previously, we identified three ultrapotent nanobodies against the initial  
128 strain of SARS-CoV-2, Wuhan-Hu-01 (WH01); accordingly, one of the intranasally  
129 delivered nanobody was shown to protect hACE2 mice infected by WH01 strain<sup>25</sup>.

130  
131 Here, we compared the neutralizing potency of the aforementioned nanobodies against  
132 various circulating SARS-CoV-2 variants. Nb22-Fc was identified to exhibit increased  
133 neutralization potency against Delta variant compared to WH01 strain, to which the  
134 antibody was originally raised. The binding characterization and crystal structural  
135 analysis were conducted to further elucidate the potential mechanism. Furthermore,  
136 therapeutic studies demonstrate that intranasal Nb22 exhibited complete protection  
137 against SARS-CoV-2 Delta variant in hACE2 mice. Additionally, we comprehensively  
138 evaluated the prophylactic efficacy of Nb22 when intranasally administered at 1, 3, 5,  
139 7 days prior to SARS-CoV-2 infection. Notably, single dose of intranasal Nb22 still  
140 exhibited protective against hACE2 mice even when administered 7 days prior to  
141 infection of Delta variant. Moreover, four doses of intranasal Nb22 could maintain

142 long-lasting retention in respiratory system for more than one month. All these indicate  
143 that intranasal Nb22 could be applied not only in the PrEP and PEP but also in the STIP,  
144 filling the gap between the long-term lagging prevention and PreP.

145

## 146 **Results**

### 147 **Potent neutralization of Delta variant by nanobodies**

148 We previously reported the discovery and characterization of three potent neutralizing  
149 nanobodies against WH01 strain and its variants with  $IC_{50}$  values of ~1 ng/ml. These  
150 three nanobodies (Nb15-Fc, Nb22-Fc and Nb31-Fc) were identified to bind to RBD<sup>25</sup>.  
151 Neutralization experiments were further conducted to measure their activity against the  
152 circulating variants including variants of concern (VOC) comprising Alpha (B.1.1.7  
153 with N501Y), Beta (B.1.351 with E484K and N501Y), Delta (B.1.617.2 with L452R  
154 and T478K) and Gamma (P.1 with K417T, E484K and N501Y), as well as variants of  
155 interest (VOI) comprising Eta (B.1.525 with E484K), Iota (B.1.526 with E484K),  
156 Epsilon (B.1.429 with L452R), and Kappa (B.1.617.1 with L452R and E484Q)<sup>7-10</sup>.  
157 Nb15-Fc exhibited increased potency against Alpha variant, but decreased potency  
158 against Delta variant or Epsilon as compared with WH01, the RBD used to select the  
159 nanobodies. Nb31-Fc exhibited reduced potency against Alpha, Delta and Epsilon  
160 variants relative to WH01 or D614G variant (Fig. 1A-G). Interestingly, Nb22-Fc  
161 exhibited about 2.5-fold increased neutralizing potency against Delta variant with an  
162  $IC_{50}$  value of 0.41 ng/ml (5.13 pM) compared to WH01 with an  $IC_{50}$  of 1.01 ng/ml  
163 (12.63 pM). Notably, Nb22-Fc also exhibited around 8.4-fold increased neutralization  
164 potency against Delta variant relative to variant Alpha with an  $IC_{50}$  of 3.45 ng/ml (43.13  
165 pM) (Fig. 1A-G). Impressively, Nb22-Fc also exhibited outstanding neutralizing curve  
166 against authentic Delta variant compared to Nb15-Fc and Nb31-Fc (Fig. 1F-G). All  
167 three nanobodies failed to neutralize variants containing E484K/Q mutation, suggesting  
168 that E484K/Q mutation in RBD could lead to the resistance to all three nanobodies.  
169 Altogether, Nb15-Fc presented the most potent neutralization against variant Alpha  
170 with an  $IC_{50}$  of 0.18 ng/ml, and Nb15-Fc and Nb31-Fc still retained potent  
171 neutralization of variants containing L452R and T478K mutations in RBD (Fig. 1E-F),

172 though with reduced potency like most other anti-RBD antibodies<sup>7,12</sup>. Of note, Nb22-  
173 Fc exhibited the most potent neutralization against pseudotyped or authentic Delta  
174 variant virus among three nanobodies (Fig. 1A-G).

175

## 176 **Characterization of Nb22-Fc binding to RBD**

177 To explore antibody binding characteristics to the RBD with respect to their  
178 neutralization of Delta variant, the interactions of three nanobodies with variant RBDs  
179 were analyzed using biolayer interferometry (BLI). Nb15-Fc, Nb22-Fc and Nb31-Fc  
180 showed high affinity interactions with RBD of Delta variant at 1.86 nM, 0.31 nM and  
181 0.31 nM, respectively (Fig. 2A-D). However, the ultrahigh affinity of Nb22-Fc and  
182 Nb31-Fc to the RBD of Delta variant did not fully reflect the neutralization potency as  
183 Nb22-Fc neutralized Delta variant with markedly more potency than that of Nb31-Fc,  
184 suggesting that affinity is not the only factor dictating the neutralization activity.  
185 Furthermore, Nb22-Fc exhibited increased affinity with Delta variant RBD relative to  
186 other variant RBDs (Fig. 2A-D), which is in line with the increased potency conferred  
187 by Nb22-Fc against Delta variant as compared with other variants.

188

189 Moreover, immunofluorescence analysis revealed that Nb22-Fc specifically interacted  
190 with spike protein from WH01, D614G, Alpha, Epsilon and Delta variants on the  
191 surface of transfected 293T cells, whereas no binding with the spike protein from other  
192 variants containing E484K/Q mutation (Fig. 2E). These results were substantiated by  
193 flow cytometric results (Fig. 2F). Overall, these specific binding characteristics are  
194 consistent with its specific neutralization properties.

195

## 196 **Structural analysis of RBD-Nb22 complex**

197 Structural analysis of Nb22 interaction with RBD was performed to address the  
198 ultrahigh potency of Nb22 against WH01 strain and Delta variant. Initially, we  
199 determined the crystal structure of WH01 RBD-Nb22 complex at a resolution of 2.7 Å  
200 (Fig. 3A and Table S1). Nb22 adopts a typical β-barreled structure, and contains three  
201 variable complementarity-determining regions (CDR) binding to RBD. The buried

202 surface area (BSA) was 800 Å<sup>2</sup>, mainly constituting of hydrogen bonds and  
203 hydrophobic interactions. 14 residues constituting epitope of three CDRs were  
204 identified using a distance of <4 Å as the cutoff (Fig. 3B). For CDR1, T30 and S33  
205 formed two hydrogen bonds with S494 of RBD, while the hydrophobic interactions  
206 included A32 and F34 of Nb22 and Y449, L452, F490 and Q493 of RBD (Fig. 3C and  
207 Table S2). N57 of CDR2 interacted with G485 by hydrogen bond, and the hydrophobic  
208 interactions were mediated by I56 and Y489 (Fig. 3D). CDR3 is a relatively longer  
209 region with only one hydrogen bond (Y119 and G446). The side chain of P104 inserted  
210 into the hydrophobic cavity formed by F101, R107, Y453, F456 and Y495 (Fig. 3E).  
211 Apart from the five hydrogen bonds in CDR regions, the interface of Nb22 and RBD  
212 was stabilized by two additional hydrogen bonds consisting of G1, S75, N450 and E484  
213 (Fig. 3F). Interactions were also facilitated by the hydrophobic network constituted by  
214 P2, Q3, V4, G28, G29, R73 and D74 of Nb22 (Fig. 3G and Table S2).

215  
216 Superimposition of the structure of WH01 RBD-Nb22 complex and RBD-hACE2  
217 (PDB code: 6MOJ) immediately elucidates the structural basis of neutralization, in  
218 which the binding of Nb22 to RBD effectively blocks the binding of RBD to hACE2  
219 during virus infection. Firstly, the binding site of Nb22 on RBD partially overlaps with  
220 that of hACE2 (Fig. S1A). Secondly, the loop (V102-Y117) of Nb22 clashes with two  
221 α-helices of the N-terminus of hACE2 (Fig. S1B).

222  
223 To elucidate the increased potency of Nb22 in neutralizing Delta variant, we determined  
224 the Delta RBD-Nb22 complex structure at a resolution of 2.9 Å, which revealed that  
225 two distinct mutations, T478K and L452R, had differing impact on the binding K478  
226 locates outside the CDR binding regions, and does not disturb the interaction surface  
227 (Fig. 4A and Table S1). Therefore, T478K mutation does not affect Nb22 neutralization  
228 (Fig. 4A). However, the mutation of hydrophilic leucine to positively charged arginine  
229 (R) at position 452 significantly enhances the interactions of RBD with the CDR3  
230 region of Nb22. Two additionally formed hydrogen bonds, T30-R452 and S33-Q493,  
231 pull the CDR3 loop of Nb22 closer to R452 region of RBD, as revealed by the

232 superimposition of the structures of WH01 and Delta RBD-Nb22 ([Fig. 4B, 4C](#) and  
233 [Table S2](#)), explaining enhanced neutralization activity of Nb22 against the Delta variant.

234

235 **Nb22 exhibits room-temperature stability in vitro and long-lasting retention in**  
236 **vivo**

237 Nanobodies exhibit various advantages including thermostability. We reported that  
238 nanobodies could retain 100% activity even after being incubated at 70 °C for one hour  
239 and retain 83% activity after 80 °C treatment for one hour<sup>25</sup>. Further evaluation showed  
240 that Nb22 could maintain full activity for at least two months at room temperature and  
241 did not lose any activity even undergoing five rounds of freeze-thawing ([Fig. 5A](#) and  
242 [5B](#)), indicating that Nb22-Fc is highly stable and idea for non-cold chain storage and  
243 use at room-temperature.

244 To determine Nb22-Fc distribution in vivo, YF®750 SE-labeled Nb22 (Nb22-YF750)  
245 was administered via intranasal (i.n.) in a mouse model. The fluorescence could be  
246 readily detected in respiratory system including nasopharynx, trachea and lung 2h post  
247 infusion. The fluorescence was detectable for more than seven days after a single dose  
248 of 200 ug (average of 10 mg/kg) Nb22-YF750 administration, which is in agreement  
249 with our previous reports. As expected, the fluorescence could be detected for more  
250 than one month when four doses of 200 µg (average of 10 mg/kg) Nb22-YF750 were  
251 administered every week ([Fig. 5C-E](#) and [Fig. S2](#)). Nb22-Fc could also be detected in  
252 the blood, indicating that Nb22-Fc could also exert its activity in the blood after  
253 bypassing the respiratory system ([Fig. 5F](#)). All these observations of prolonged  
254 retention of Nb22-Fc in the respiratory system suggest the potential application of the  
255 antibody for STIP against SARS-CoV-2 infection. Taken together, intranasal Nb22-Fc  
256 could be developed as an STIP reagent for its long-lasting retention in the respiratory  
257 system and a portable therapeutics thanks to its room-temperature stability.

258

259 **Intranasal Nb22 is highly efficacious in the STIP, PreP and PeP in hACE2**  
260 **transgenic mice challenged by Delta variant**

261

262 To evaluate the efficacy of Nb22-Fc in vivo, hACE2 transgenic mice were challenged  
263 with  $1 \times 10^5$  PFU SARS-CoV-2 Delta variant (CRST: 1633.06.IVCAS 6.7593) and  
264 conducted as we previously reported<sup>25</sup>. hACE2 mice were divided into nine groups  
265 (n=3-6) as shown in [Figure 6A](#) and 200  $\mu$ g (average of 10 mg/kg) Nb22-Fc was  
266 administered via i.n. before or after Delta variant challenge to evaluate the antibody's  
267 prophylactic or therapeutic efficacy against Delta variant infection. Viral RNA in the  
268 lungs was detected in virus control group named as SARS-CoV-2 group (Group 2).  
269 Animals in Control Nb group Group 3 were challenged with Delta variant and received  
270 control nanobody treatment one hour post infection. As expected, high copy numbers  
271 of viral RNA were also detected in control Nb mice without significant difference  
272 compared to SARS-CoV-2 group ([Fig. 6B](#)).

273 In order to evaluate the prophylactic duration conferred by Nb22, a single dose of Nb22  
274 was administered via i.n. at days 1, 3, 5, and 7, respectively, prior to Delta variant  
275 challenge. Viral RNA copies increased over the course of Nb22-Fc administration ([Fig.](#)  
276 [6B](#)). As expected, viral RNA copies in the aforementioned prophylactic groups were  
277 all significantly lower than that in the control Nb group, indicating that even a single  
278 dose of Nb22-Fc could provide protection against Delta variant infection in hACE2  
279 transgenic mice for 7 days. Nb22 administered in -7d and -5d before challenge  
280 significantly reduced viral load though failed to provide complete protection ([Fig. 6B](#)).  
281 Notably, Nb22 exhibited significant prevention against SARS-CoV-2 infection in the -  
282 3d Nb22 (Group 6) and -1d Nb22 group (Group 7) ([Fig. 6B](#)). All these indicate that  
283 Nb22-Fc provides better protection at the earlier time points upon infection.

284 In the therapeutic group, viral RNA copies in the animals treated with Nb22 at 1 hour  
285 or 1 day postinfection were undetectable in 5/5 mice in the groups 8 and 9, suggesting  
286 that Nb22 had complete protection of hACE2 mice against Delta variant infection ([Fig.](#)  
287 [6B](#)). The viral RNA results were also validated by immunofluorescence (IF) staining  
288 and HE staining in the lungs ([Fig. 6C and 7E](#)). We noted that mice challenged by Delta  
289 variant did not show obvious weight loss even in 4 days post infection ([Fig. S3](#)). In  
290 summary, Nb22 exhibited high efficacy in both the prevention and therapy of hACE2  
291 transgenic mice challenged with Delta variant. Nb22 provided complete protection in

292 PEP (in 1h Nb22 group and 1d Nb22 group) and exhibited high efficacy in PrEP (in -  
293 1d Nb22 and -3d Nb22 group). Impressively, a single dose of Nb22 could maintain  
294 effectiveness in the prevention against Delta variant infection for at least seven days (in  
295 -7d Nb22 group), indicating the potential application for STIP against SARS-CoV-2.

296

297

## 298 **Discussion**

299 To date, a small number of nanobodies with ultrahigh potency against SARS-CoV-2  
300 and its variants have been reported <sup>15,26,27</sup>, whereas nanobodies with potent  
301 neutralization against the currently dominant Delta variant were rarely reported. Our  
302 results revealed that three previously reported nanobodies <sup>25</sup>, retained ultrahigh potency  
303 in neutralization against the Delta variant. Among them, Nb22-Fc with an IC<sub>50</sub> value of  
304 0.41ng/ml (5.13 pM) is outstanding with increased neutralization of Delta variant  
305 relative to Alpha variant. The Nb22 binding to RBD provides mechanistic insight into  
306 the enhanced neutralization against Delta variant, suggesting that the increased binding  
307 affinity enhanced the neutralizing potency against Delta variant relative to Alpha  
308 variant (Fig. 1). Given that most anti-RBD, anti-NTD antibodies or convalescent sera  
309 or vaccine-elicited antibodies showed reduced neutralization of Delta variant relative  
310 to that of Alpha variant <sup>7,12</sup>, the increased neutralization activity of Nb22-Fc against  
311 Delta variant is particularly striking and the structural basis of the phenomenon is of  
312 interest for understanding the neutralization mechanisms.

313

314 The structural analysis further illustrated the characteristics of Nb22 binding to WH01  
315 and Delta RBD and the mechanisms of viral inhibition. Nb22 binding to RBD  
316 effectively blocks the binding of RBD to hACE2 during virus infection. The binding  
317 site of Nb22 on RBD overlaps with that of hACE2 (Fig. S1A), and the loop (V102-  
318 Y117) of Nb22 clashes with two  $\alpha$ -helices of the N-terminus of hACE2 (Fig. S1B). In  
319 addition, crystal structural analysis showed that T478K mutation of Delta variant are  
320 located outside 800  $\text{\AA}^2$  BSA of Nb22 interacting with RBD and do not perturb the  
321 interaction between Nb22 and the RBD of Delta variant. Of note, the guanidine moiety

322 in the L452R mutation forms an additional hydrogen bond with the hydroxyl group of  
323 T30 of Nb22, pulling the CDR3 loop of Nb22 closer, and leading to an extra hydrogen  
324 bond between S33 of Nb22 and Q493 of RBD (Fig. 4). Consequently, the BSA extends  
325 from 800 Å<sup>2</sup> to 835 Å<sup>2</sup>, in comparison with that of Nb22-WH01 RBD. All these  
326 contribute to the enhanced binding and neutralizing potency of Nb22 against Delta  
327 variant.

328 Compared to conventional antibodies for passive immunization, nanobodies are  
329 efficiently produced in prokaryotic expression systems at low cost and possess  
330 favorable biophysical properties including high thermostability<sup>28</sup>. We reported that  
331 nanobodies could remain 100% activity even incubated at 70 °C for one hour and are  
332 amenable to engineering of multimeric nanobody constructs<sup>25</sup>. Such nanobodies exhibit  
333 high effectiveness against virus infection via intranasal administration<sup>25</sup>. The results  
334 reveal that Nb22 could maintain full activity for more than two months at room  
335 temperature and does not lose any activity even after undergoing five rounds of freeze-  
336 thawing.

337 Our results demonstrate that a single dose of intranasal Nb22 could exhibit efficacy in  
338 the STIP, PrEP and PEP against SARS-CoV-2 infection in hACE2 mice. Of note, a  
339 single dose of intranasal Nb22 could maintain efficacy against SARS-CoV-2 infection  
340 for at least one week in hACE2 mice, which would readily serve as STIP. Our antibody  
341 distribution results also revealed that Nb22 could retain in respiratory tracts for at least  
342 one month when weekly administered via i.n.. As such, we anticipate that Nb22 could  
343 provide one month prevention against SARS-CoV-2 infection when administered  
344 intranasally every week.

345 VLong-term lagging prevention against SARS-CoV-2 conferred by approved vaccine  
346 usually takes more than one month to be effective and then lasts for months or years<sup>29</sup>.  
347 Vaccine efficacy has been shown to wan within months after vaccination<sup>7,11</sup>.  
348 Instantaneous prevention of SARS-CoV-2 is also needed for individuals who do not  
349 take vaccines when SARS-CoV-2 is circulating. A few studies in animal model just  
350 demonstrated that antibodies exhibited accelerated clearance of SARS-CoV-2 in PrEP  
351 when administered 1-3 days prior to infection<sup>2,4-6</sup>. Whereas, to the best of our

352 knowledge, few studies have fully investigated the STIP that prevention could be  
353 readily effective immediately following inoculation and last for more than one week to  
354 one month for people at high risk of SARS-CoV-2 infection, which can also serve to  
355 reduce transmission during the asymptomatic stage of the infection. As such, our results  
356 demonstrate that intranasal Nb22 with ultrahigh potency and long-lasting retention in  
357 the lung could satisfy the need of STIP against SARS-CoV-2.

358

359 In summary, structural analysis provides a mechanistic explanation to the enhanced  
360 sensitivity of Delta variant and the increased neutralization potency of this antibody.  
361 The structural analysis may further guide the rational design of pan-coronavirus  
362 vaccines and therapeutics. Nb22 exhibited one of the ultrahigh neutralization potencies  
363 among the reported antibodies or nanobodies against Delta variant infection<sup>7,12,26,30,31</sup>.  
364 We presented proof of concept of STIP against SARS-CoV-2 using our Nb22 and  
365 suggest STIP as a new prophylactic strategy for long-lasting antibodies to prevent virus  
366 infection. Although the newly emerged Omicron variant is spreading globally, the Delta  
367 variant remains the dominant variant and likely the deadliest in most of the world;  
368 therefore, the ultrahigh potent, and thermal stable Nb22 is an excellent candidate for  
369 intranasal or inhalable anti-SARS-CoV-2 agent for both therapy or prophylaxis,  
370 especially including STIP.

371

372

### 373 **Materials and Methods**

#### 374 **Expression of nanobodies**

375 The Fc1 gene (CH2-CH3) of the human monoclonal antibody was fused with the VHH  
376 gene of nanobodies (named as Nb-Fc or Nbs) to assist the purification and prolong the  
377 half-life of the Nb antibody, following our previously published protocol<sup>32</sup>. The Nbs  
378 were finally cloned into the pcDNA3.4 eukaryotic expression vector (Invitrogen),  
379 which were transfected into 293F cells (cat.# R79007, Thermo Scientific) to produce  
380 Nb-Fcs. Nb fused with Fc was purified using Protein G (cat.# 20399, Thermo

381 Scientific).

382

383 **Enzyme linked immunosorbent assay (ELISA) analysis**

384 Antibody quantification and antibody characterization were tested by ELISA as our  
385 previously reported method<sup>33</sup>, with modifications. In brief, the protein was coated to  
386 ELISA plates (Corning) at a concentration of 0.5 µg/ml. After washing 2-4 times, 5%  
387 non-fat milk in PBS was added and incubated for blocking at 37 °C for 1 h. After  
388 washing, 100 µl serially diluted sera or purified antibody was added and incubated at  
389 37 °C for 1 h. Following washing, secondary antibody of goat anti-llama IgG (H+L)  
390 with HRP (Novus, cat.# NB7242, 1:10000 dilution) or goat anti-human IgG with HRP  
391 was added and incubated at 37 °C for 1 h. Accordingly, 3,3',5,5'-Tetramethylbenzidine  
392 (TMB, Sigma) substrate was added at 37 °C for 10 minutes (min); and 10 µl 0.2 M  
393 H<sub>2</sub>SO<sub>4</sub> was added to stop the reaction. The optical densities at 450 nm (OD450) were  
394 measured using the Infinite 200 (Tecan, Ramsey, MN, USA). Antibody quantification  
395 in the sera was calculated according to the standard curve generated by purified  
396 antibody.

397

398 **Neutralization activity of nanobodies against pseudovirus**

399 Pseudovirus neutralization assay was carried out following our previously published  
400 protocol<sup>25</sup>, with the follow modifications. Briefly, pseudovirus of SARS-CoV-2  
401 variants was produced by co-transfection of pNL4-3.Luc.R-E-, an HIV-1 NL4-3  
402 luciferase reporter vector that comprises defective Nef, Env and Vpr (HIV AIDS  
403 Reagent Program), and pVAX1 (Invitrogen) expression vectors encoding the spike  
404 proteins of respective variants into 293T cells (ATCC). Supernatants containing  
405 pseudovirus were collected after 48 hours (h), and viral titers were determined by  
406 luciferase assay in relative light units (Bright-Glo Luciferase Assay Vector System,  
407 Promega Biosciences). Human codon optimized S genes of SARS-CoV-2 variants were  
408 synthesized, and the corresponding pseudoviruses were produced following the above  
409 protocol. For neutralization assay, SNB02, an Nb-Fc specific against SFTSV<sup>32</sup>, served  
410 as a negative control. Neutralization assays were conducted by incubating pseudovirus

411 with serial dilutions of purified nanobodies at 37 °C for 1 h. HEK293T-ACE2 cells  
412 (cat.# 41107ES03, Yeasen Biotech Co., Ltd. China) (approximately  $2.5 \times 10^4$  per well)  
413 were added in duplicate to the virus-antibody mixture. Half-maximal inhibitory  
414 concentrations (IC<sub>50</sub>) of the evaluated nanobodies were determined by luciferase  
415 activity 48 h following exposure to virus-antibody mixture, and analyzed by GraphPad  
416 Prism 8.01 (GraphPad Software Inc.).

417 **Immunofluorescence and flow cytometric analysis**

418 Immunofluorescence and flow cytometric analysis were conducted following our  
419 previously published protocol <sup>34</sup>, with minor modifications. Briefly, S gene sequences  
420 for SARS-CoV-2 spike protein of various SARS-CoV-2 variants were obtained from  
421 the GISAID website (<https://gisaid.org>). S genes were synthesized and constructed as  
422 expression plasmids by GenScript. The plasmids were transfected into 293T cells  
423 (ATCC) cultured in 12-well plates. Next, 48 hours post transfection, the cells were  
424 washed by PBS and fixed with 4% paraformaldehyde for 20 minutes at room  
425 temperature. The purified Nb-Fc was used to stain the 293T cells, followed by Alexa  
426 Fluor 488 AffiniPure goat Anti-human IgG (H+L) (1:500 dilution) (109-545-003,  
427 Jackson ImmunoResearch). For immunofluorescence analysis, the cells on the plate  
428 were examined and the images were acquired using an OLYMPUS IX73. For flow  
429 cytometric analysis, the cells were resuspended in 500 µl PBSF buffer (PBS+2% FBS)  
430 and analyzed using ACEA NovoCyte TM (Agilent Biosciences); non-transfected 293T  
431 cells served as a negative control.

432

433 **Affinity determination by Bio-Layer Interferometry (BLI)**

434 We measured antibody affinity using a ForteBio OctetRED 96 BLI (Molecular Devices  
435 ForteBio LLC, Fremont, CA) with shaking at 1,000 rpm at 25 °C <sup>25</sup>. To determine the  
436 affinity of Nbs with human Fc tag, Nb-Fcs were loaded to anti-human Fc (AHC)  
437 biosensors (cat.# 18-5060, Fortebio) in a kinetic buffer (PBS, 0.02% (v/v) Tween-20,  
438 pH 7.0) for 200 sec prior to baseline equilibration for 180 sec in a kinetic buffer.  
439 Association of SARS-CoV-2 RBD in a three-fold dilution series from 33.3 nM to 1.2  
440 nM was performed prior to dissociation for 180 sec. After each cycle, the biosensors

441 were regenerated through 3 brief pulses of 5 sec each with 100 mM pH 2.7 glycine-  
442 HCL followed by a running buffer. The data were baseline subtracted before fitting  
443 using a 1:1 binding model and the ForteBio data analysis software.  $K_D$ ,  $K_a$  and  $K_d$   
444 values were determined by applying a global fit to all data.

445 **Expression and purification of WH01 and Delta RBD protein for crystal  
446 structural analysis**

447 The WH01 and Delta RBD were expressed using the Bac-to-Bac baculovirus system.  
448 The two pAcgp67-RBD (residues 333–530) plasmid with a C-terminal 8×His tag were  
449 transfected into Sf9 cells using Cellfectin II Reagent (Invitrogen) to produce the  
450 recombinant baculoviruses. After 3 rounds of amplification, Hi5 cells were infected  
451 with baculoviruses at an MOI of 4 at a density of  $2 \times 10^6$  cells/ml. The supernatants of  
452 cell culture containing the secreted RBD were harvested at 60 h after infection. The  
453 RBD was purified by Ni-NTA resin (GE Healthcare). Nonspecific contaminants were  
454 removed by washing the resin with 20 mM Tris-HCl, 150 mM NaCl, pH 7.5, and the  
455 target proteins were eluted with elution buffer containing 20 mM Tris-HCl, 150 mM  
456 NaCl, 500 mM imidazole, pH 7.5. The eluted proteins were further purified by  
457 Superdex 75 (GE Healthcare, USA) and stored in 20 mM Tris-HCl, 150 mM NaCl, pH  
458 7.5.  
459

460 **Expression and purification of Nb22 for crystal structural analysis**

461 The VHH gene for Nb22 was amplified by PCR and cloned into a pET21a vector with  
462 *Bam*H I and *Xho* I restriction sites. The recombinant plasmids were transformed into  
463 *Escherichia coli*. BL21 (DE3). The cells were cultured in LB medium and grown to  
464 OD<sub>600</sub> = 0.8 at 37°C. Isopropyl-D-1-thiogalactopyranoside (IPTG) was added to a final  
465 concentration of 1.0 mM to induce the protein expression, and the cultures were grown  
466 at 16 °C overnight. Cells were harvested by centrifugation at 4,500 rpm for 15 min, re-  
467 suspended and homogenized in the lysis buffer containing 20 mM Tris-HCl, 150 mM

468 NaCl, pH 7.5 using ultrasonic. Cell debris was removed by centrifugation at 18,000  
469 rpm for 30 min. The supernatants were added to Ni- NTA resin (GE Healthcare, USA).  
470 The nonspecific contaminants were eluted by washing the resin with the lysis buffer  
471 containing 10 mM imidazole. The target protein with 6 x His tag, named as Nb22, was  
472 subsequently eluted with the lysis buffer containing 500 mM imidazole. Nb22 was  
473 eluted and purified by Superdex 75 (GE Healthcare, USA).

474

#### 475 **Crystallization, structural determination and data acquisition**

476 The complexes were prepared by mixing WH01 or Delta RBD and Nb22 at a 1:1.2  
477 molar ratio and incubating at 4 °C overnight. The complexes were further purified by  
478 Superdex 75 (GE Healthcare, USA) to remove the excess nanobody. The crystals were  
479 screened by vapor-diffusion sitting-drop method at 16°C. The crystals appeared and  
480 reached their final size within 3 days in a well solution comprising 0.1 M HEPES (pH  
481 7.0), 5% v/v (+/-)-2-Methyl-2,4-pentanediol (MPD), 10% polyethylene glycol (PEG)  
482 10000 (WH01 RBD-Nb22) and 0.1 M Tris (pH 7.0), 37.5% Jeffamine (Delta RBD-  
483 Nb22), respectively.

484 To collect data, a single crystal was mounted on a nylon loop and was flash-cooled with  
485 a nitrogen gas stream at 100 K. Diffraction data of WH01 RBD-Nb22 was collected on  
486 BL18U1 at Shanghai Synchrotron Radiation Facility (SSRF) at a wavelength of  
487 0.97915 Å. While, the Delta RBD-Nb22 was collected on BL02U1 at a wavelength of  
488 0.97918 Å. Data were processed and scaled using the HKL3000 package and  
489 autoPROC<sup>35</sup>. The structures were elucidated using the molecular replacement (MR)  
490 method in PHASER program<sup>36</sup> with the structure of SARS-CoV-2 RBD (PDB code:  
491 7CJF)<sup>37</sup> as the initial searching model. The model was built into the modified  
492 experimental electron density using COOT<sup>38</sup> and further refined in PHENIX<sup>39</sup>. The  
493 final refinement statistics are summarized in Table S1. Structural figures were prepared  
494 by PyMOL. Epitope and paratope residues, as well as their interactions, were identified  
495 by PISA ([http://www.ebi.ac.uk/pdbe/prot\\_int/pistart.html](http://www.ebi.ac.uk/pdbe/prot_int/pistart.html)) at the European  
496 Bioinformatics Institute.

497

498 **Pharmacokinetics (PK) of Nb22-Fc *in vivo*.**

499 Purified Nb22-Fc were injected via intranasal (*i.n.*) into BALB/c mice (Qing Long Shan  
500 Animal Center, Nanjing, China) at a dose of 10 mg/kg. The concentration of Nb22-Fc  
501 in serum was measured by ELISA. The  $T_{1/2}$  of Nb22-Fc was caculated as  $\ln(2)/k$ , where  
502 k is a rate constant expressed reciprocally of the x axis time units by the plateau  
503 followed one phase decay or one phase decay equation in the GraphPad software.

504

505 **Spatial distribution of Nb22-Fc *in vivo***

506 Spatial distribution of Nb22-Fc were conducted following our previously published  
507 protocol<sup>25</sup>, with minor modifications. Nb22-Fc labeled with far infrared dye YF®750  
508 SE (US EVERBRIGHT INC, YS0056) were named as Nb22-YF750. 10 mg/kg Nbs-  
509 YF750 were administered via intranasal into nude mice (18-22g, Qing Long Shan  
510 Animal Center, Nanjing, China). Images were recorded at Ex:740 nm/Em:780 nm by  
511 NightOWL LB 983 (Berthold, Germany) at the indicated time point. Images were  
512 analyzed using Indigo imaging software Ver. A 01.19.01.

513

514 **Evaluating the efficacy of Nb22-Fc in SARS-CoV-2 infected hACE2 mice.**

515 The efficacy of Nb22-Fc against SARS-CoV-2 were evaluated according to our  
516 previously published protocol<sup>25</sup>, with minor modifications. In brief, A total of 43 8-  
517 week-old male transgenic hACE2 mice (C57BL/6J) (cat.# T037630, GemPharmatech  
518 Co., Ltd., Nanjing, China) were challenged with  $1 \times 10^5$  PFU SARS-CoV-2 Delta  
519 variant (CRST: 1633.06.IVCAS 6.7593) per mouse. The mice were randomly divided  
520 into nine groups (n=3-6) for either prophylactic or therapeutic evaluation, as described  
521 in [Figure 5A](#). Mice without any treatment and challenge were taken as blank control  
522 (No SARS-CoV-2, n=3). Mice challenged with SARS-CoV-2 were taken as infection  
523 control (SARS-CoV-2, n=5). 250  $\mu$ g/mouse (average of 10 mg/kg) SNB02 (Y-Clone,  
524 China), an anti-SFTSV antibody constructed by Nb fused with human Fc1 (Nb-Fc)<sup>32</sup>,  
525 was intranasally injected 1 hour (h) after infection and was taken as an isotype antibody  
526 treated control (Control Nb, n=3). For the prophylactic group, mice were intranasally  
527 injected with Nb22-Fc at a dose of 250  $\mu$ g/mouse (average of 10 mg/kg) at 7 days (d),

528 5d, 3d, 1d before infection (named as -7d Nb22, -5d Nb22, -3d Nb22, -1d Nb22,  
529 respectively, n=5-6). For the therapeutic group, mice were intranasally injected with  
530 Nb22-Fc at a dose of 250 µg/mouse (average of 10 mg/kg) 1 h or 24 h after infection  
531 (named as 1h Nb22 and 1d Nb22, n=5, respectively). Body weight of mouse was  
532 recorded daily. Given that hACE2 transgenic mice typically clear virus within 5-7 days  
533 after SARS-CoV-2 infection<sup>40</sup>, the mice were sacrificed at 4 days post infection (dpi).  
534 Subsequently, lung tissues were harvested for viral load determination and tissue  
535 sections for immunofluorescence (IF) and hematoxylin and eosin (H&E) staining. All  
536 experiments were conducted in a Biosafety Level 3 (BSL-3) facility.

537

538

539 **Viral load measurement by quantitative RT-PCR.**

540 Viral load was measured by quantitative real-time PCR (qRT-PCR) on RNA extracted  
541 from the supernatant of lung homogenates as reported previously<sup>41</sup>. Briefly, lung  
542 homogenates were prepared by homogenizing perfused lung using an electric  
543 homogenizer. The inactivated samples were transferred from the BSL-3 to BSL-2  
544 laboratory and total RNA was extracted from the collected supernatant. Each RNA  
545 sample was reverse transcribed to 50 µl cDNA with HiScript II Q RT SuperMix for  
546 qPCR (+gDNA wiper) (R223-01). 5 µl cDNA was added into a 25 µl qRT-PCR reaction  
547 containing the ChamQ SYBR qPCR Master Mix (High ROX Premixed) (Q341-02,  
548 Vazyme Biotech, China) and primers designed to target the nucleocapsid protein of  
549 SARS-CoV-2 (5' - GGGAACTTCTCCTGCTAGAAT -3' and 5' -  
550 CAGACATTTGCTCTCAAGCTG -3' ). The samples were run in triplicate on an  
551 ABI 7900 Real-Time System (Applied Biosystems, Thermo Fisher Scientific). The  
552 following cycling conditions were performed: 1 cycle of 50 °C for 2 min, 1 cycle of  
553 95 °C for 10 min, and 40 cycles of 95 °C for 15 sec and 58 °C for 1 min.

554

555 **Immunofluorescence staining of SARS-CoV-2-infected cells and H&E staining in  
556 tissues.**

557 The lung tissues were immersed in 10% neutral buffered formalin (cat.# Z2902, Sigma)

558 for 24 h. After the formalin fixation, the tissues were placed in 70% ethanol (Merck)  
559 and subsequently embedded with paraffin. Tissue sections (5- $\mu$ m thick) were prepared  
560 for H&E staining and immunofluorescence staining for SARS-CoV-2 detection using  
561 the Coronavirus nucleocapsid protein (NP) antibody (cat. 40143-MM05, Sino  
562 Biological). Images were collected under a Pannoramic MIDI system (3DHISTECH,  
563 Thermo) using Pannoramic scanner software and analyzed by ImageJ (NIH).

564

### 565 **Quantification and statistical analysis**

566 All statistical analyses were carried out using GraphPad Prism 8.01 software (GraphPad)  
567 or OriginPro 8.5 software (OriginLab). ANOVA or Mann-Whitney test was performed  
568 for group comparisons.  $P < 0.05$  was considered as statistically significant with mean  
569  $\pm$ SEM or mean  $\pm$ SD.

570

### 571 **Study approval**

572 The study and the protocol for this research were approved by the Center for Public  
573 Health Research, Medical School, Nanjing University. All animal experimental  
574 procedures without infection were approved by the Committee on the Use of Live  
575 Animals by the Ethics Committee of Nanjing University. All animals infected by  
576 SARS-CoV-2 were performed in Biosafety Level 3 animal facilities in accordance with  
577 the recommendations for care and use of the Institutional Review Board of Wuhan  
578 Institute of Virology of the Chinese Academy of Sciences (Ethics Number:  
579 WIVA11202111). All the authors declare their compliance with publishing ethics.

580

581

### 582 **Author contributions**

583 XW conducted most experiments, analyzed the data and wrote the draft manuscript. LC  
584 conducted all the neutralization experiments. LZ, BH, MJ, SX, HS, DZ, LL, WN  
585 provided technical assistance. YW, SM and SY conducted the structural analysis. FN,  
586 YL, HH, QH and YL, evaluated the efficacy of Nb22 in SARS-CoV-2 infected  
587 transgenic hACE2 mice. ZW designed the study, directed and revised the manuscript.

588 All authors critically reviewed the draft manuscript and approved the final version.

589

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702  
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705 **Figure Legends**

706 **Figure 1 Characterizing nanobodies neutralizing circulating variants of SARS-  
707 CoV-2.** The neutralization curve of Nb15-Fc (A), Nb22-Fc (B), Nb31-Fc(C) and  
708 SNB02 (D) inhibiting SARS-CoV-2 pseudovirus of circulating variants. Nb-Fcs and  
709 SNB02 were all constructed as the format of VHH fused with human Fc1. SNB02 was  
710 taken as an antibody control specific for SFTS virus. (E) The summary curve of IC<sub>50</sub>  
711 of Nb-Fcs exhibiting potent neutralization against SARS-CoV-2 variants. (F) The  
712 neutralization potency of Nb-Fcs was evaluated based on authentic SARS-CoV-2 Delta  
713 variant plaque reduction neutralization test. (G) The summary table of IC<sub>50</sub> and IC<sub>80</sub> of  
714 Nb-Fcs in A-C and F, displaying potent neutralization. Data are represented as mean ±  
715 SD. All experiments were repeated at least twice.

716  
717

718 **Figure 2. Characterizing the binding of Nbs.** Kinetic binding curve of Nb15-Fc (A),  
719 Nb22-Fc (B) and Nb31-Fc (C) at the concentration 33.3 nM, 11.1nM, 3.7nM and 1.2  
720 nM with RBD of Delta variant, respectively, detected by BLI. Binding curves are  
721 colored black, and fit of the data to a 1:1 binding model is colored red. (D)  
722 Representative binding curve of various RBD as indicated to Nb22-Fc tested by BLI.  
723 Nb22-Fc binding with RBD from representative SARS-CoV-2 variants detected by  
724 immunofluorescence assay (E) and flow cytometric analysis (F), respectively. Mock  
725 served as a cell control without plasmid transfection. Images were visualized under the  
726 ×10 objective. All experiments were repeated at least twice.

727

728 **Figure 3. Structural analysis of Nb22 and WH01 RBD complex.** (A) The overall  
729 complex structure of Nb22 and WH01 RBD. The CDR1 (red), CDR2 (blue), CDR3  
730 (green) of Nb22 (pink) and WH01 RBD (orange) are displayed in cartoon  
731 representation. (B) The epitope of Nb22 shown in surface representation. The CDR  
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785 lung tissue sections were also analyzed by H&E staining. Immunofluorescence and HE  
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787

788 **Supplemental Materials**

789 **Supplemental Figure 1. Nb22 blocks the binding of hACE2 to WH01 RBD. (A)**

790 Overlap of Nb22 and hACE2 binding sites on WH01 RBD. hACE2 binding site on  
791 WH01 RBD is shown in cyan line. Nb22 binding site is shown in pink line. The overlap  
792 region is represented by ellipses with dashed lines. **(B)** The loop (V102-Y117) of Nb22  
793 is clashed with the two helices on N-terminal of hACE2. The loop is colored in red and  
794 helixes are colored in green.

795

796 **Supplemental Figure 2. Spational distribution of Nb22 labeled with dye YF®750**

797 **SE.** Mice were dissected and detected by NightOwl LB 983 after 200  $\mu$ g Nb22-  
798 YF®750 SE infusion into mice as indicated in figure 5C. The fluorescence intensity  
799 was measured at 2 hours (0d-Nb22), 7d (7d-Nb22), 14d (14d-Nb22), 21d (21d-Nb22)  
800 and 28d (8d-Nb22) after infusion of Nb22 via i.n., respectively. Blank, blank mice  
801 without infusion of any antibody, was taken as blank control. The fluorescence intensity  
802 of various organs including trachea (Tr), lung (Lu), heart (H), stomach(St), intestine  
803 (In), liver(Li), spleen (Sp), kidney (Ki), bladder (B), were analyzed

804 **Supplemental Figure 3. Body weight of mice.** Body weight of mice in figure 6 was  
805 recorded at the indicated time point.

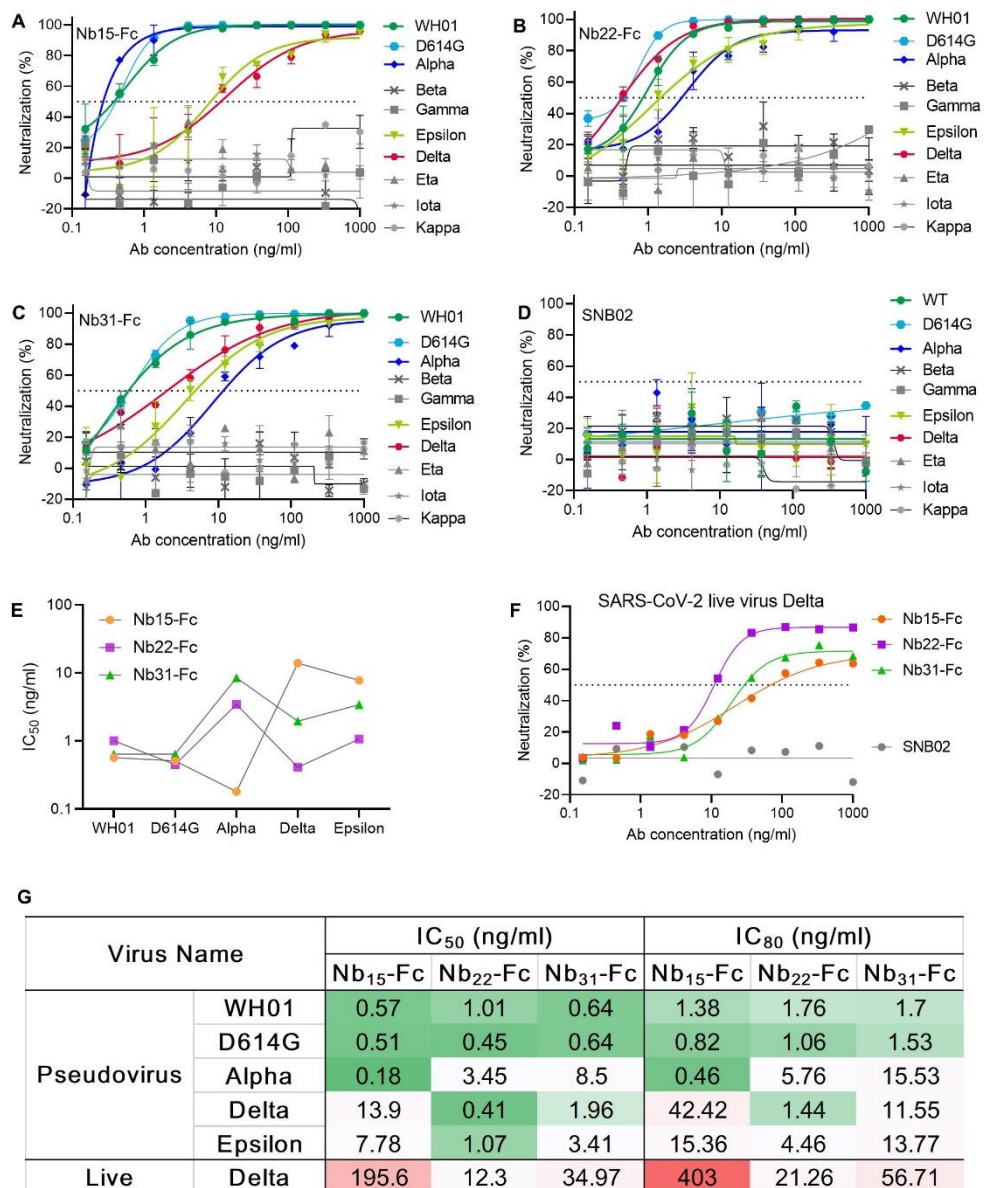
806

807 **Supplemental Table 1.** Data collection and refinement statistics

808 **Supplemental Table 2.** Residues contributed to interaction between Nb22 and RBD  
809 were identified by PISA at the European Bioinformatics Institute.

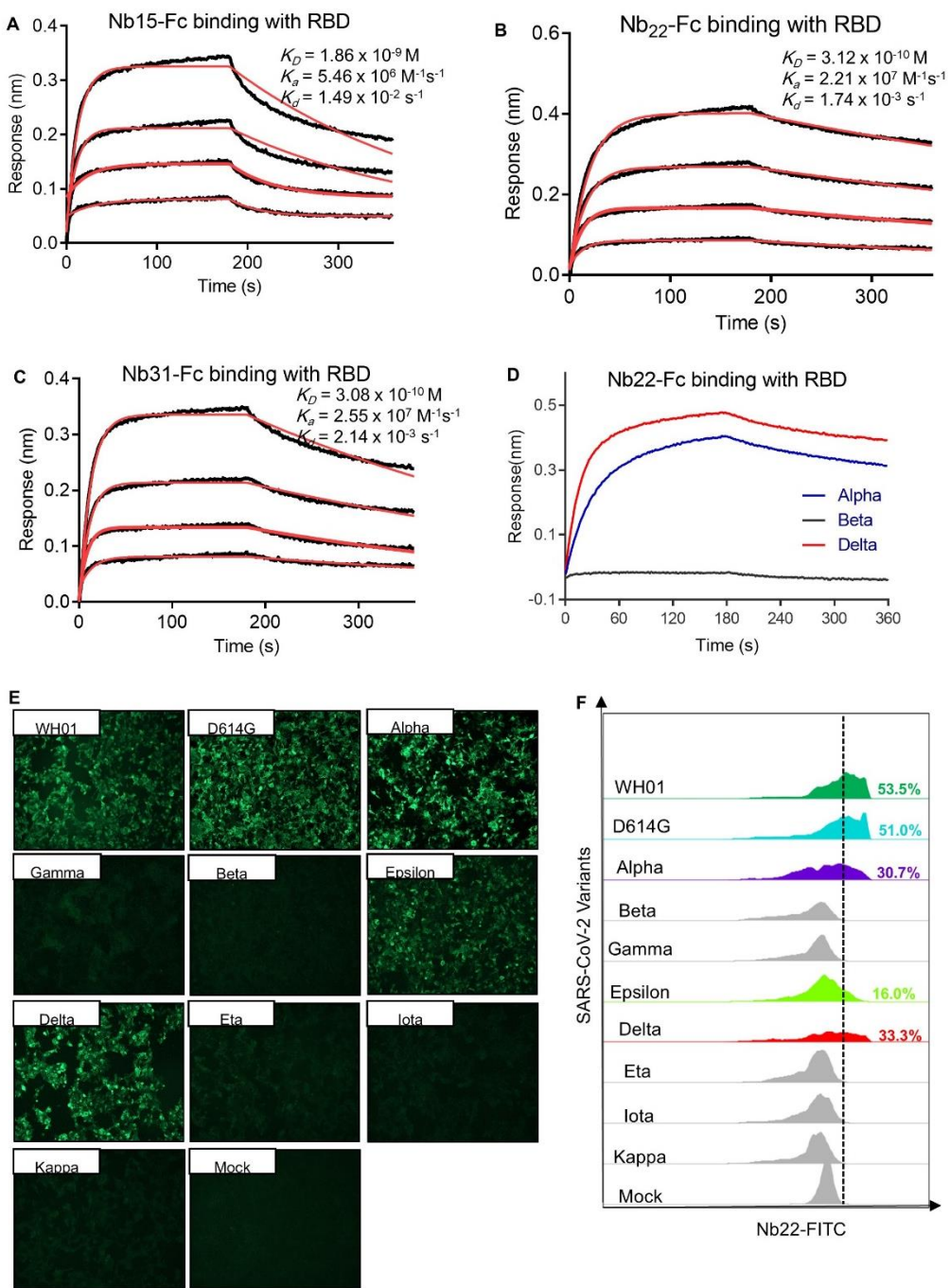
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811 **Figures**



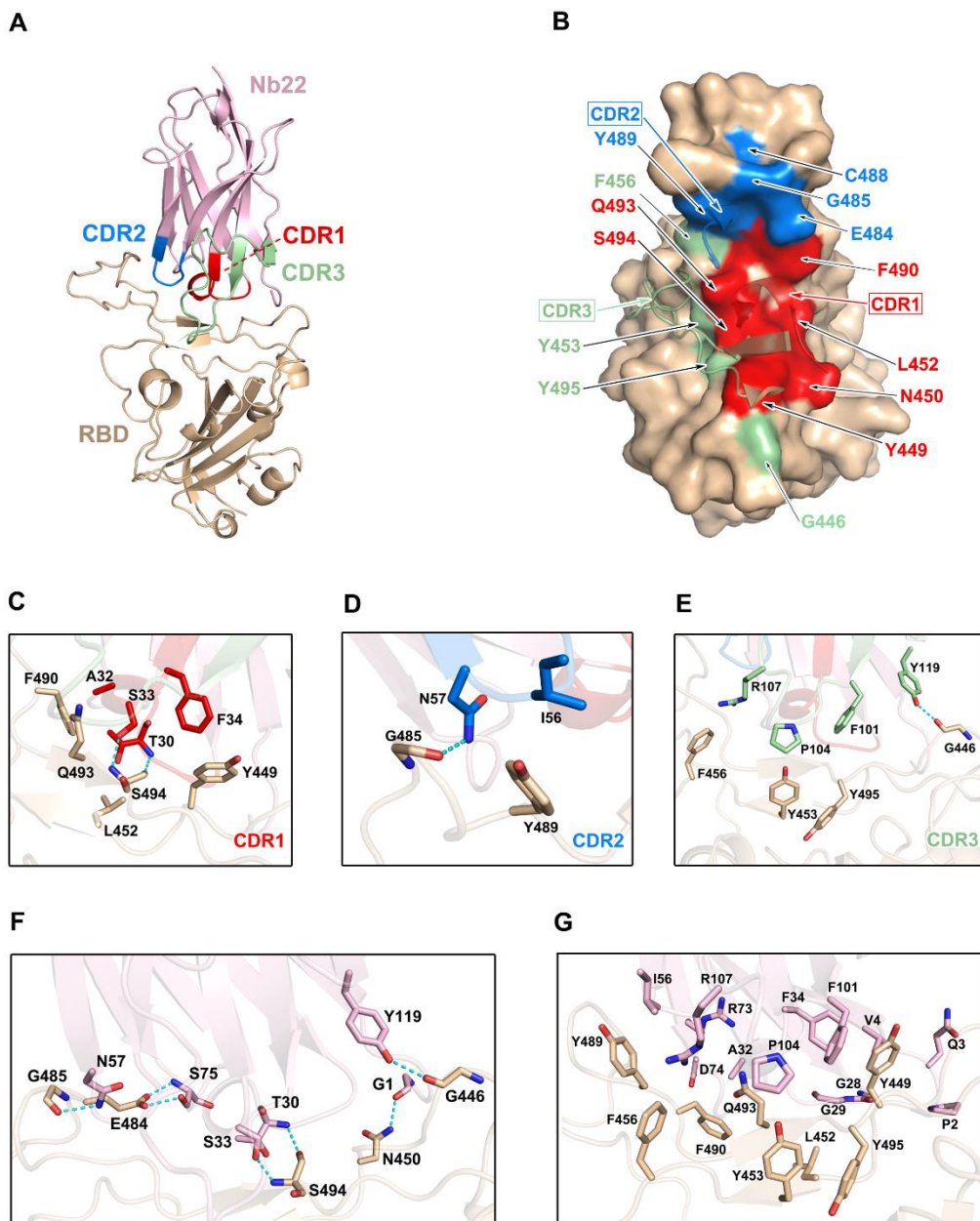
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**Figure 1 Characterizing nanobodies neutralizing circulating variants of SARS-CoV-2.** The neutralization curve of Nb15-Fc (A), Nb22-Fc (B), Nb31-Fc(C) and SNB02 (D) inhibiting SARS-CoV-2 pseudovirus of circulating variants. Nb-Fcs and SNB02 were all constructed as the format of VHH fused with human Fc1. SNB02 was taken as an antibody control specific for SFTS virus. (E) The summary curve of IC<sub>50</sub> of Nb-Fcs exhibiting potent neutralization against SARS-CoV-2 variants. (F) The neutralization potency of Nb-Fcs was evaluated based on authentic SARS-CoV-2 Delta variant plaque reduction neutralization test. (G) The summary table of IC<sub>50</sub> and IC<sub>80</sub> of Nb-Fcs in A-C and F, displaying potent neutralization. Data are represented as mean  $\pm$  SD. All experiments were repeated at least twice.



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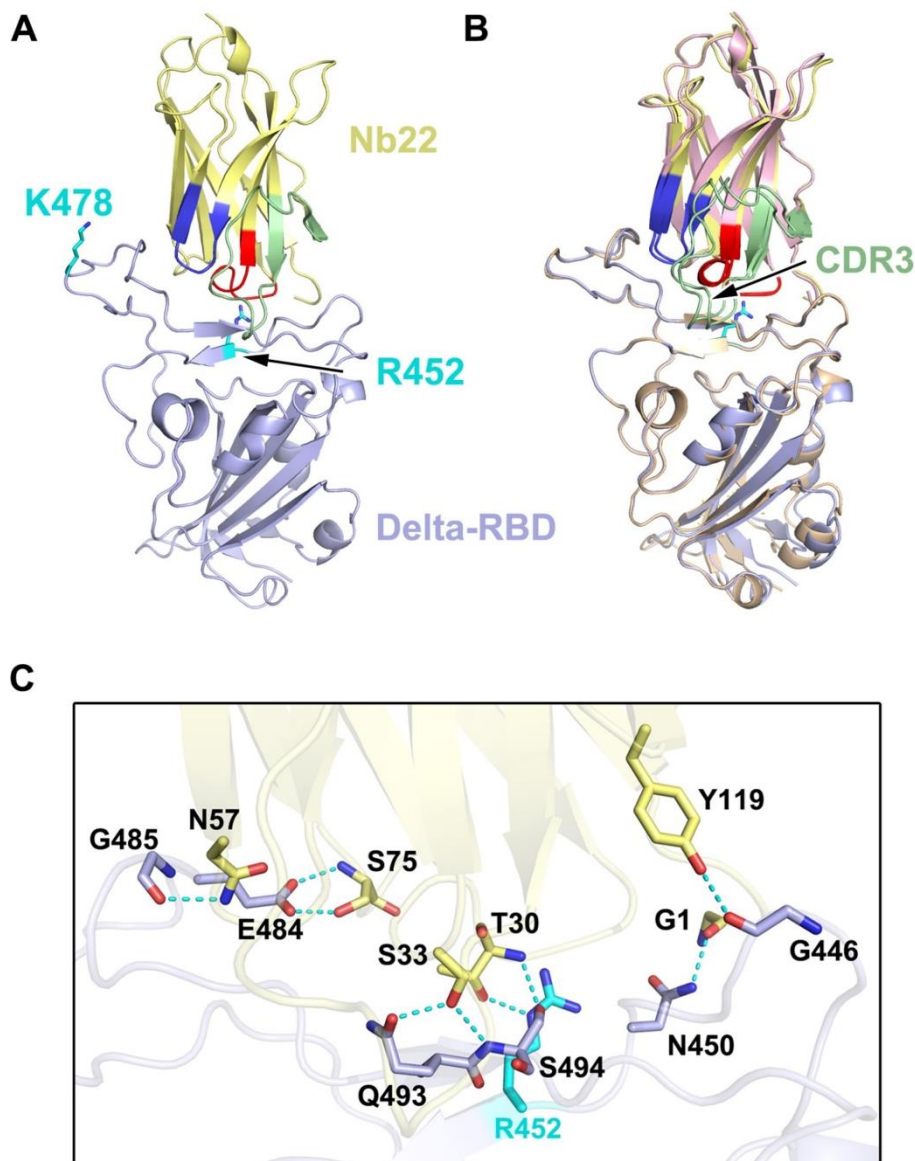
825 **Figure 2. Characterizing the binding of Nbs.** Kinetic binding curve of Nb15-Fc (A),  
 826 Nb22-Fc (B) and Nb31-Fc (C) at the concentration 33.3 nM, 11.1 nM, 3.7 nM and 1.2  
 827 nM with RBD of Delta variant, respectively, detected by BLI. Binding curves are  
 828 colored black, and fit of the data to a 1:1 binding model is colored red. (D)  
 829 Representative binding curve of various RBD as indicated to Nb22-Fc tested by BLI.  
 830 Nb22-Fc binding with RBD from representative SARS-CoV-2 variants detected by  
 831 immunofluorescence assay (E) and flow cytometric analysis (F), respectively. Mock  
 832 served as a cell control without plasmid transfection. Images were visualized under the  
 833  $\times 10$  objective. All experiments were repeated at least twice.



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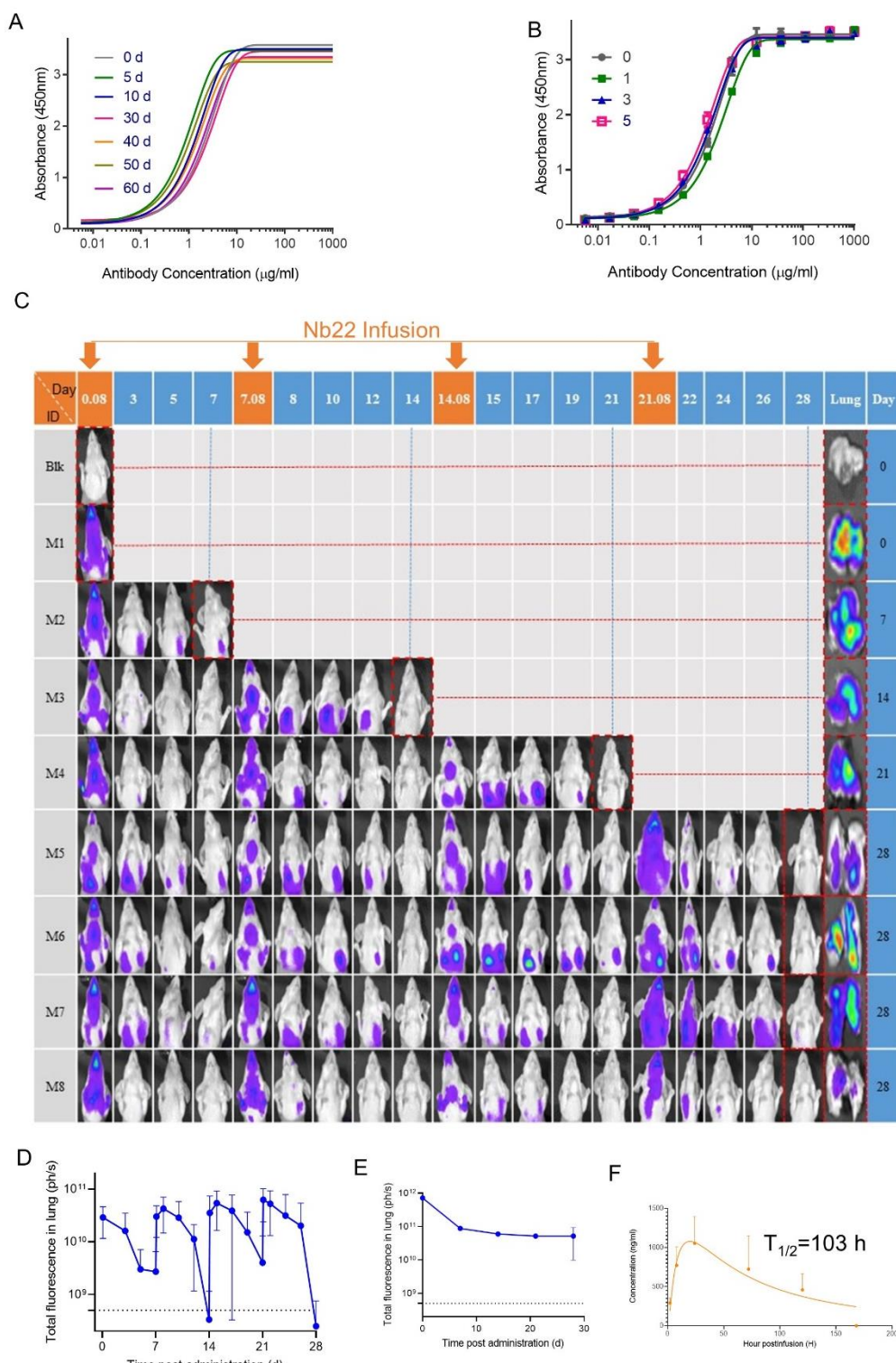
835

836 **Figure 3. Structural analysis of Nb22 and WH01 RBD complex.** (A) The overall  
837 complex structure of Nb22 and WH01 RBD. The CDR1 (red), CDR2 (blue), CDR3  
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845

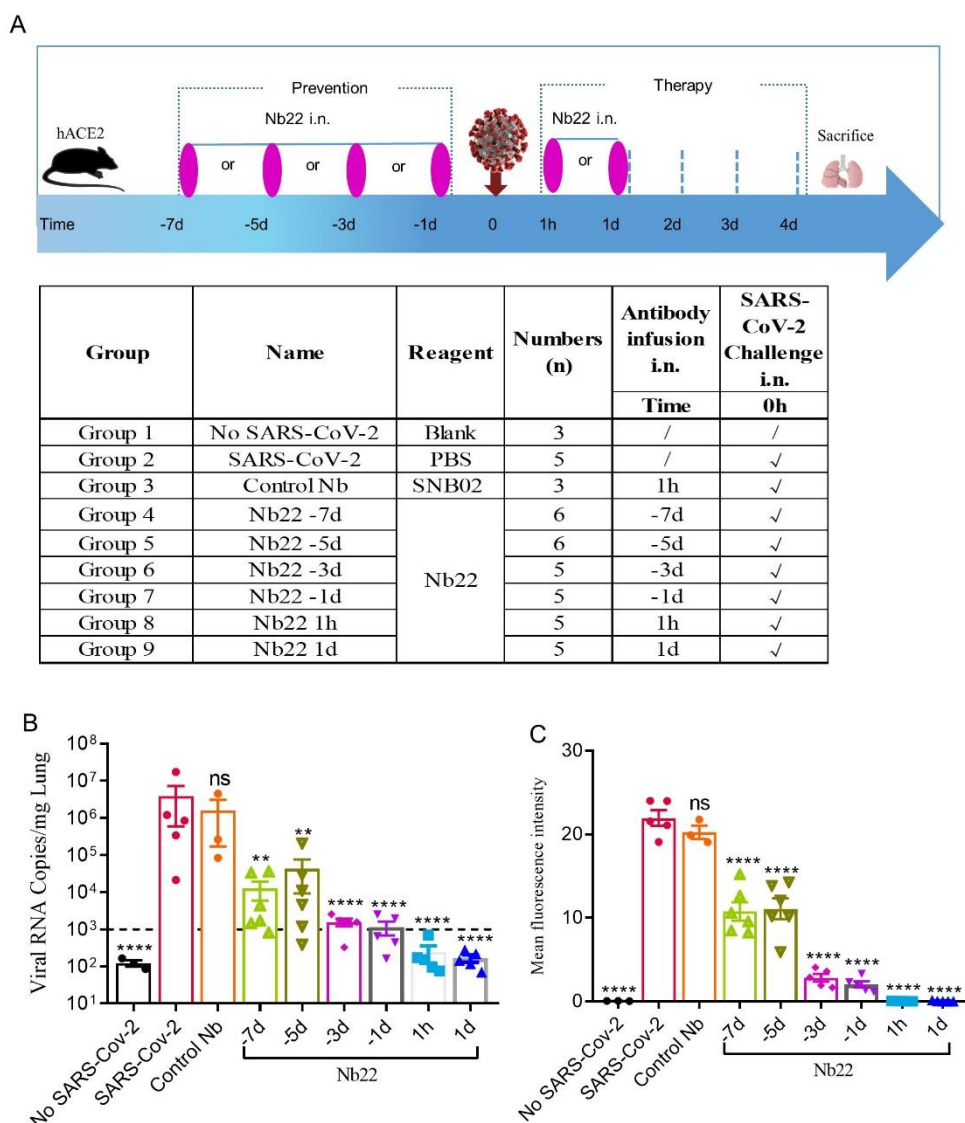
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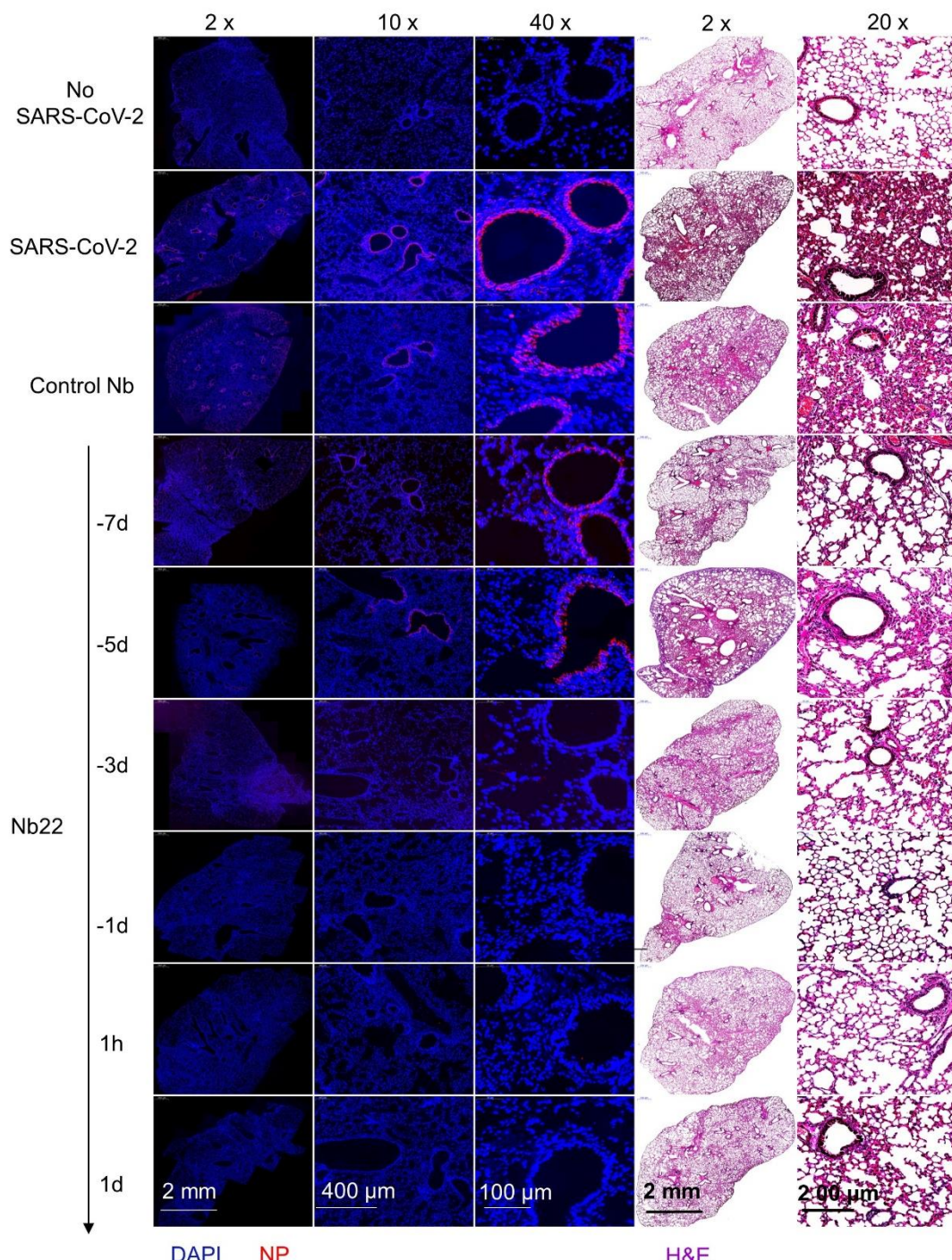
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