

1 **Extensive neutralization against SARS-CoV-2 variants elicited by**  
2 **Omicron-specific subunit vaccine booster**

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4 *Running title:*

5 **Neutralization elicited by Omicron-specific subunit vaccine booster**

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26  
27

28 **Abstract**

29

30 The currently dominant variant of SARS-CoV-2 Omicron, carrying a great number  
31 of mutations, has been verified its strong capacity of immune escape in COVID-19  
32 convalescents and vaccinated individuals. An increased risk of SARS-CoV-2  
33 reinfection or breakthrough infection should be concerned. Here we reported  
34 higher humoral immune response elicited by Delta and Omicron variants after  
35 breaking through previous infection and cross-neutralization against VOCs,  
36 compared to the ancestral wild-type (WT) virus infection. To overcome the  
37 immune escape of Omicron, Omicron-specific vaccine was considered as a novel  
38 and potential strategy. Mouse models were used to verify whether Omicron-specific  
39 RBD subunit boost immune response by immunizing Omicron-RBD recombinant  
40 proteins. Three doses of Omicron-RBD immunization elicit comparable neutralizing  
41 antibody (NAb) titers with three doses of WT-RBD immunization, but the neutralizing  
42 activity was not cross-active. By contrast, two doses of WT-RBD with an  
43 Omicron-RBD booster increased the NAb geometric mean titers against Omicron by  
44 9 folds. Moreover, an additional boost vaccination with Omicron-RBD protein could  
45 increase humoral immune response against both WT and current VOCs. These  
46 results suggest that the Omicron-specific subunit booster shows its advantages in the  
47 immune protection from both WT and current VOCs , and that SARS-CoV-2 vaccines  
48 administration using two or more virus lineages as antigens might improve the NAb  
49 response.

50

51

## 52 **Introduction**

53 Since the coronavirus disease 2019 (COVID-19) pandemic caused by  
54 severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) began in 2019, it  
55 had experienced several waves driven by variants of this virus. At present, five  
56 variants including Alpha, Beta, Gamma, Delta and Omicron were designated as  
57 “variants of concern” (VOCs). As a dominate strain, even though the pandemic of  
58 Delta has lasted over one year, it has been replaced swiftly by Omicron (B.1.1.529)  
59 causing a new round of pandemic within a short time due to its rapid spread with a  
60 great deal of mutations<sup>1</sup>. More than 30 mutations have been accumulated in the  
61 spike (S) protein of Omicron variant, especially 15 of those occurs on  
62 receptor-binding domain (RBD), which is not only the vital binding site to the host  
63 receptor angiotensin-converting enzyme 2 (ACE2) for the entry of SARS-CoV-2,  
64 but also the key target of neutralizing antibodies produced by immune response  
65 and therapeutic antibodies. Spike mutation has been well documented to be  
66 correlated to its infectivity alteration and immune evasion<sup>2-6</sup>. The neutralizing  
67 activity of Omicron of sera has been suggested an extensive reduction from  
68 convalescents or vaccinees who received various types of SARS-CoV-2 vaccines  
69 3,7,8.

70

71 Reduced neutralization elicited by infection or vaccination shows that Omicron  
72 has an increased risk of SARS-CoV-2 reinfection or breakthrough infection. In  
73 31220 Norwegian households, secondary attack rate caused by Omicron was  
74 25.1% (95% CI, 24.4%-25.9%)<sup>9</sup>. A study based on the Qatar national database

75 suggests that the effective protection of previous infection against reinfection with  
76 the omicron variant was approximately 60%, which is lower than alpha, beta and  
77 delta variants (at approximately 90%)<sup>10</sup>. The vaccine effectiveness against  
78 Omicron after two BNT162b2 doses was 65.5% (95% confidence interval [CI],  
79 63.9 to 67.0) at 2 to 4 weeks, and dropping to 8.8% (95% CI, 7.0 to 10.5) at 25 or  
80 more weeks<sup>11</sup>. Currently, an Omicron sub-variant BA.2 shows faster spread and  
81 similar resistance to immunity with high rate of breakthrough infection<sup>12-14</sup>. After  
82 breaking through previous immune protection by vaccine based on the ancestral  
83 wild-type (WT) virus, how the immune response elicited by Omicron breakthrough  
84 infection is needs to be delineated. It will give hints for the development of  
85 protective vaccine and vaccination strategies.

86  
87 Due to continuous appearance of SARS-CoV-2 variants, reduced efficiency of  
88 existing vaccines accelerates the need of new vaccine strategies. A booster  
89 following primary vaccination series showed its potential efficiency of promoting  
90 high neutralizing activity<sup>15</sup> and reducing symptomatic SARS-CoV-2 infection<sup>16</sup>, but  
91 booster shots displayed the failure in the breakthrough infection of some  
92 SARS-CoV-2 VOCs<sup>17</sup>. Moreover, simply additional boosters might not improve  
93 immune protection. A fourth-dose booster using the same antigen could not  
94 generate higher antibody titers than the third-dose vaccination and shows the low  
95 prevention against mild or asymptomatic Omicron infections and breakthrough  
96 infection<sup>18</sup>. Boosting with heterologous vaccines as one of the candidates, it has  
97 been proved to be safe and efficient immune response<sup>19-21</sup>. At present, several

98 vaccination programs with heterologous vaccines have been approved in some  
99 countries. In view of the correlation of the immune escape of Omicron variant with  
100 its great number of mutations, Omicron-specific vaccines has been proposed.  
101 Omicron-specific mRNA vaccine booster could induce neutralizing response to  
102 Omicron itself but fail to previous VOCs<sup>22</sup>. As a booster with Omicron-matched  
103 DNA vaccines, increased width of immune response has been observed<sup>23</sup>.  
104 However, in macaque models, vaccination with Omicron-specific boosters do not  
105 increase neutralizing antibody (NAb) titers against Omicron and remain the equivalent  
106 levels of B cell response<sup>24</sup>. Of note, both two types of vaccines were designed  
107 according to full-length spike proteins. In consideration of the key role of  
108 SARS-CoV-2 RBD as the target of neutralizing antibodies, it has an important  
109 significance to verify whether Omicron-specific RBD subunit boost immune response  
110 after previous WT-RBD doses. Here we report immunogenicity and cross-reactivity  
111 of Omicron-specific RBD subunit proteins in mouse models to highlight the need of  
112 next generation of SARS-CoV-2 vaccines with variants-specific antigens.

113  
114

## 115 **Results**

116 In this study, twenty persons who were infected with Omicron or Delta after  
117 vaccination in each cohort were recruited, while 13 individuals previously infected  
118 with WT and unvaccinated were matched as a control cohort according to ages,  
119 sex and the time of sample collection of other two groups( Supplementary Table1 ).  
120 Delta breakthrough infections occurred 2.5-5 months (median 4.1months) after  
121 the last vaccine doses, while Omicron breakthrough infections occurred 3.4-6.6

122 months (median 5.2 months) after the last vaccine doses. Sera samples were  
123 collected at 3-4 time points within 50 days post symptom onset. Their  
124 anti-WT-RBD IgG binding antibody and neutralizing antibodies were determined  
125 by Enzyme-linked immunosorbent assay (ELISA) and the pseudotype-based  
126 neutralizing assay. Within the acute phase of COVID-19 infection, anti-WT-RBD  
127 IgG levels of sera in all three cohorts gradually raised to peak, then the antibody  
128 trends remained steady (Fig. 1A and Extended Data Fig. 1). As expected, WT  
129 infection without additional immune protection, lower IgG titers were observed at  
130 the early stages of infection than Omicron or Delta breakthrough infection with  
131 their complicated histories of vaccination in different individuals. At the late stages  
132 of acute infection, anti-WT-RBD IgG binding antibodies reached to comparable  
133 levels among three cohorts.

134

135 On the other hand, neutralization ability of breakthrough infection has been  
136 observed (Figure 1B, Extended Data Fig. 2). In the WT cohort, NAb titers of sera  
137 against four VOCs (Alpha, Beta, Delta and Omicron) are decreased by 1.3 folds,  
138 13.8 folds, 4.1 folds and 19.2 folds, respectively, compared to the NAb titers  
139 against WT itself. In the Delta cohort, except for high NAb titers against Delta itself,  
140 NAb titers of sera against WT, Alpha, Beta and Omicron are decreased by 1.5  
141 folds, 1.4 folds, 4.9 folds and 6.0 folds, respectively. In the Omicron cohort, except  
142 for high NAb titers against Omicron itself, NAb titers of sera against WT, Alpha,  
143 Beta and Delta are decreased by 1.0 folds, 0.9 folds, 2.1 folds and 1.2 folds,

144 respectively, indicating the moderate cross-neutralization elicited by breakthrough  
145 infections.

146

147 During the follow-up visit, NAb titers among these three cohorts presents  
148 consistent dynamic changes with that of IgG antibodies (Figure 1C-E, Extended  
149 Data Fig. 3). WT cohort showed various degrees of neutralizing resistance to  
150 SARS-CoV-2 VOCs, especially to Beta and Omicron. In the Delta and Omicron  
151 cohorts, high neutralizing activity against themselves has been seen due to  
152 specific breakthrough infection. In the two breakthrough infection cohorts, even  
153 though NAb titers against WT induced by breakthrough infection did not exceed  
154 NAb levels in the WT cohort, sera displayed decreased neutralizing resistance  
155 against some variants. Taken together, compared with WT infection, breakthrough  
156 infection especially by Omicron could induce wide ranges and high levels of the  
157 humoral immune response to Omicron and other VOCs.

158

159 Widespread neutralizing activity of Omicron or Delta breakthrough infection  
160 against WT and other variants has been seen on the basis of the exposure of two  
161 antigens. Then, mouse models were used to evaluate whether previously  
162 ancestral vaccination or Omicron-specific immune response has the major  
163 contribution to the increased protection. BALB/c mice were distributed into 4  
164 groups and immunized by two doses of SARS-CoV-2 RBD recombinant proteins  
165 subunit with 2-week interval plus one booster one week after the second dose  
166 (Figure 2A): Group1 immunized with 3 doses of WT-based RBD recombinant

167 proteins (WT-RBD, Cat: K1516, Okaybio, China), Group2 immunized with 3 doses  
168 of Omicron- based RBD recombinant proteins  
169 (Omicron-RBD,Cat: 40592-V08H121, Sinobiological, China) , Group3 immunized  
170 with 2 doses of WT-RBD plus one dose of Omicron-RBD and Group4 immunized  
171 by the adjuvant (Cat: KX0210042, Biodragon, China) as the control.

172

173 Mice sera were collected and performed ELISA and the pseudotype-based  
174 neutralizing assay to determine their binding IgG levels and neutralizing effect on  
175 different VOCs. Among all these groups, two-week interval immunization induced  
176 the increased levels of anti-WT-RBD IgG binding antibodies (Fig. 2B). At Day 21  
177 following the first dose of immunization, the third doses of RBD recombinant  
178 subunit proteins boost IgG antibody levels. Of note, higher anti-WT-RBD IgG  
179 levels have been shown in the sera elicited by three doses of WT-RBD proteins or  
180 two doses of WT-RBD proteins plus one dose of Omicron protein in Group1 and  
181 Group3, compared to Omicron-specific antibody response in Group2. It suggests  
182 partial cross-recognition of antibody response elicited by Omicron. On the other  
183 hand, booster by either WT-RBD or Omicron-RBD at the third dose of  
184 immunization produced the comparable levels of anti-WT-RBD IgG. The results  
185 are consistent with that of clinical samples we tested above.

186 Two-week interval with boosting by the third-dose immunization led to 26-fold  
187 increase (GMT from 138 to 3688) of NAbs against WT in Group1, while 30-fold  
188 increase (GMT from 183 to 5525) of NAb titers against Omicron variant in Group2

189 (Figure 2B-C). Omicron-specific NAb titers against Omicron in Group3  
190 (GMT=5525) is comparable with WT-induced immune response (GMT=3688) to  
191 WT itself in Group1. It suggests that the immunogenicity of Omicron does not  
192 change. In contrast, sera from only WT-immunized mice in Group1 or only  
193 Omicron-immunized mice in Group2 do not neutralize other VOCs, except for  
194 itself (Figure 2D). However, it is noteworthy that Omicron-specific boosting after  
195 two-dose WT-RBD immunization can raise NABs against Omicron by 9 folds.  
196 Subsequently, we tested cross-neutralization of these mice among three groups  
197 (Figure 2E). In Group1 only WT-RBD immunization showed high neutralizing  
198 capacity against WT itself and Delta, but weak neutralizing capacity against  
199 Omicron and Beta variants (3.7- and 2.4-fold decline, respectively). Likewise, in  
200 Group2, mice administrated with 3-dose of Omicron-RBD generated extremely  
201 high NABs against Omicron variant itself but fail to neutralize other tested VOCs.  
202 Interestingly, in Group3, the Omicron-specific shot boost NAb titers not only  
203 against Omicron, but also against WT, Beta and Delta variants with GMT of 921,  
204 3140, 962 and 1712, respectively. As NAb titers are found to be correlates with  
205 effective protection<sup>25</sup>, the data presented here suggest that Omicron-specific  
206 boosting will help to improve the immune protection from Omicron and that  
207 immunization by heterologous antigens will be beneficial to obtain wider  
208 protection against different SARS-CoV-2 variants.

209

## 210 **Discussion**

211 Widespread immune escape in COVID-19 convalescents and vaccinees has  
212 been reported extensively. Furthermore, the fast transmission of Omicron and a

213 surge of Omicron infected cases indicated the high risk of reinfection and  
214 breakthrough infection<sup>17,26</sup>. In a study about the influenza vaccine, authors  
215 suggest that prior infection enhances antibody responses to inactivated vaccine  
216 and is important to attain protective antibody titers<sup>27</sup>. In the Delta breakthrough  
217 infection after fully vaccination, 31-fold higher neutralizing antibody titers against  
218 the SARS-CoV-2 delta variant than vaccinees without infection was observed<sup>28</sup>.  
219 To evaluate the effect of prior vaccination on breakthrough infection, we analyzed  
220 the characteristics of humoral immune response elicited by Omicron variant after  
221 breakthrough infection. Compared to the previous infection with the ancestral  
222 strain WT, NAb titers against several variants from individuals infected with Delta  
223 or Omicron after breaking through the early immune protection generated by  
224 vaccines is significantly wider. Recent studies reported the consistent result that  
225 vaccination followed by breakthrough Omicron infection improved  
226 cross-neutralization of VOCs<sup>29,30</sup>, while neutralizing capacity of the unvaccinated  
227 individuals, which is triggered by Omicron, do not cross-neutralize other variants.  
228 These results suggest that prior immunity induced by vaccines will be beneficial to  
229 overcome the high neutralization resistance of Omicron<sup>31</sup>.  
230  
231 In light of extensive neutralization observed in Omicron breakthrough infection,  
232 we sought to understand the respective contribution of prior vaccination and  
233 Omicron-specific immunogenicity to this to establish more efficient immune  
234 protection. Therefore, we used immunized animal with WT-RBD and  
235 Omicron-RBD recombinant proteins to exhibit immune response elicited by  
236 Omicron-specific booster and heterologous antigens. Compared to NAb titers

237 against WT pseudovirus produced by 3-dose of WT-RBD, Omicron-RBD alone  
238 can induce comparable NAbs against Omicron pseudovirus itself. It indicates that  
239 in mouse models, Omicron-RBD has the similar immunogenicity with WT-RBD.  
240 Furthermore, Omicron-RBD booster following two-dose of WT-RBD can induce  
241 9-fold higher levels of NAbs against Omicron pseudovirus than the WT-RBD  
242 booster. We showed that Omicron-RBD boost following primary series could  
243 produce wider protection against the SARS-CoV-2 WT strain and circulating  
244 variants, which is consistent with what we observed above in Omicron  
245 breakthrough infection.

246

247 However, Omicron-mRNA boost in vaccinated macaques has not displayed  
248 significantly different NAb titers and B cells response<sup>24</sup>. This could be due to  
249 different immunization intervals and antigen epitopes. The time interval between  
250 vaccination and infection has been shown significant correlation with the potency  
251 of Omicron-neutralizing antibodies<sup>32</sup>. In our animal models, there is only 7-day  
252 interval between primary series (two doses) and booster. By contrast,  
253 immunization by Omicron-RBD subunit proteins shows the advantage that  
254 produce more NAbs to specially target against Omicron variant, while mRNA  
255 vaccine targeting full-length spike protein may produce more irrelevant antibodies,  
256 instead of targeting Omicron RBD<sup>33</sup>.

257

258 Our results indicate that heterologous antigens with various epitopes, which is  
259 different from single antigen as we have been vaccinated, may help to improve

260 the height and width of NAb activity<sup>34,35</sup>. Except the booster vaccination strategies,  
261 A “bivalent” lipid nanoparticle (LNP) mRNA vaccine containing both Omicron and  
262 Delta RBD-LNP in half dose has been observed cross-neutralization against WT  
263 and three SARS-Cov-2 variants<sup>36</sup>. Multivalent vaccines could be alternative  
264 choice for the future development of SARS-CoV-2 vaccines and vaccination  
265 programs.

266  
267 There are several limitations in our study. Due to limited participants with Omicron  
268 or Delta breakthrough infection were included in our study, the correlation of  
269 clinical characteristics with antibody response cannot be analyzed. Unvaccinated  
270 individuals who were infected with Omicron had not been recruited, but  
271 Omicron-specific immune response was observed in mouse models.  
272 Intramuscular injection in mouse models is not be completely equivalent to natural  
273 infection, hamster models could be used for virus challenge for further study.

274  
275 Collectively, our data provides hints that the current booster vaccinations using  
276 WT-RBD protein or WT-S mRNA vaccine may be less efficient in preventing  
277 infections with the Omicron variant. Our results support the hypothesis that an  
278 additional boost vaccination with Omicron-RBD protein could increase humoral  
279 immune response against both WT and current VOCs.

280

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292

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294 conceptual ideas and designed the study. P.P., J.H., H.D. and C.H. performed the  
295 experiments and statistical analysis. C.F. provided the essential assistance  
296 through experiments. Q.F., G.T. and M.J. were responsible for sample collection.  
297 All authors provided scientific expertise and the interpretation of data for the work.  
298 P.P. drafted the manuscript. All authors contributed to critical revision of the  
299 manuscript for important intellectual content. All authors reviewed and approved  
300 the final version of the report.

301

302 **Conflict of Interest:** The authors declare no conflicts of interest.

## FIGURE LEGEND

**Fig.1 | The characteristics of immune response elicited by the Omicron breakthrough infection.** **A** dynamic change of anti-WT-RBD IgG binding antibodies in three cohorts: Omicron breakthrough infection or Delta breakthrough infection (n=20 for each cohort, sampled at 4 time points within 46 days after symptom onset), and WT infection without prior vaccination (sampled at 3 time points within 50 days after symptom onset). **B** Pseudovirus-based neutralizing assays were performed to test neutralizing activity of the sera against WT, Alpha, Beta, Delta and Omicron variants from three cohorts. **C-E** longitudinal observation of neutralizing antibody (NAb) titers of individuals experienced WT infection(C), Delta (D) and Omicron (E) breakthrough infection against WT, Alpha, Beta, Delta and Omicron pseudovirus. Statistical data analysis was performed using GraphPad Prism software. The half-maximal inhibitory dose ( $ID_{50}$ ) was calculated as NAb titers. Values above points indicate the geometric mean titers (GMTs). The threshold of  $ID_{50}$  detection was 1:40.

**Fig. 2 | Humoral immune response elicited by Omicron-specific booster in mouse models.** **A** Schematic of BALB/c mice immunization programs. 6 mice for each group were intramuscularly injected with the indicated recombinant proteins or the adjuvant as control. **B** dynamic change of anti-WT-RBD IgG binding antibodies in three groups within 35 days after the first dose of immunization. 1:16000 diluted sera were performed by ELISA. **C-D** Sera were collected and performed the pseudotype-based neutralizing assay to determine their neutralizing capacity to WT or Omicron pseudoviruses at 14 days (squares) and 28 days (triangles) after the first dose of immunization by WT-RBD (C) or Omicron-RBD (D) recombinant subunit proteins. **E** NAbs titers of sera against WT or Omicron from 3 groups collected at 28 days after the first dose of immunization. **F** Cross-neutralization of mice sera of 3 groups against WT, Beta, Delta and Omicron variants at 28 days after the first dose of immunization. Statistical data analysis was performed using GraphPad Prism version 8.0 software. Data on dot-bar plots is shown as GMT  $\pm$  SEM with individual data points in plots. Values above points indicate the GMTs. The threshold of ID<sub>50</sub> detection was 1:40.

## **Materials and Methods**

### **Patients and samples**

We enrolled 53 patients who had been identified to be previously infected with SARS-CoV-2 at the Eighth People's Hospital of Guangzhou from January 2020 to January 2022. Thirteen patients infected with the SARS-COV-2 wild-type virus strain, 20 individuals infected with the SARS-COV-2 Delta virus after vaccination, and 20 patients infected with the SARS-COV-2 omicron virus after vaccination were included in our study. All infections were confirmed by q-PCR and sequenced to identify the genotype. The collection of all samples obtained the consent from subjects according to the protocols approved by the Ethics Review Board of the Eighth People's Hospital of Guangzhou Institutional Review Board. Plasma was isolated from blood samples within 2h after collection according to the following steps: (1) Patient sera were heat incubated for inactivation at 56°C in water bath for 30 min; (2) centrifugation at 3000 rpm for 15 min, followed by transferring to new tubes; (3) Store at -80°C for further use.

### **Mouse models and study design**

8-week-old female BALB/c mice (6 mice per group) were provided by the Laboratory Animal Center of Chongqing Medical University (SCXK (YU) 2018-0003). Recombinant WT-RBD (Cat: K1516, Okaybio, China) and Omicron-RBD protein (Cat: 40592-V08H121, Sinobiological, China) as antigens for immunization were diluted with PBS, then mixed with an equal volume of QuickAntibody™-Mouse 3W adjuvant (Cat# KX0210042, Biodragon,

China) and completely emulsified by a syringe. Each mouse was intramuscularly injected with 100  $\mu$ l the antigen/adjuvant mixture. Serum samples were collected from tail tips before each vaccination and at 28 days after the first injection, then measure the antibody titers by ELISA and pseudovirus neutralization assay.

### **Enzyme-linked immunosorbent assay (ELISA)**

The recombinant RBD proteins derived from SARS-CoV-2 Wild-type (WT-RBD, Cat : K1516, Okaybio, China ) and Omicron strains (Omicron-RBD, Cat: 40592-V08H121, Sinobiological) were coated on 96-well microtiter plate (100ng/well) at 4°C overnight. After blocking with 5% skim milk powder and 2% BSA in PBS for 2 hours at room temperature, sera of enrolled patients were diluted and added into the plates and incubated at 37°C for 1 hour. After washing, wells were incubated with goat anti-mouse (Cat: ab6789, abcam, UK)/human (Cat: ab97225, abcam, UK) IgG-Horseradish peroxidase (HRP) antibody (1:10000 dilution) for 1 hour at 37°C. TMB substrate was added and incubated for 15 minutes at 37°C for color development. Reactions were stopped with stop solution, and the absorbance was determined at 450 nm using a microplate reader (Biotek, USA).

### **Pseudovirus neutralization assay**

For the neutralization assay, 50 $\mu$ L pseudoviruses of SARS-CoV-2 ( Alpha, Beta, Delta, Omicron, D614G), equivalent to  $3.8 \times 10^4$  vector genomes, were incubated with serial dilutions of sera samples (dilutions of 1:40, 160, 640, 2560) from patients or mice for 1h at 37°C, then added the mixture into the 96-well plates seeded with 293T-ACE2 cells( $1.6 \times 10^4$  cells/well). The cells were refreshed with DMEM medium 8 h post-infection. Cells were lysed by 30  $\mu$ l lysis buffer (Promega, Madison, WI, USA) at 72 h post-infection to measure RLU with luciferase assay reagent (Promega, Madison, WI, USA) according to the product instruction. Neutralization inhibition rate was calculated using GraphPad Prism 8.0 software (GraphPad Software, San Diego, CA, USA). The titers of neutralizing antibodies were calculated as 50% inhibitory dose (ID<sub>50</sub>).

### **Ethics statement**

Animal studies were approved by and conducted in compliance with the Committee on the Ethics of Animal Experiments of the Institutional Animal Care and Use Committee at the Laboratory Animal Center of Chongqing Medical University.

Fig. 1

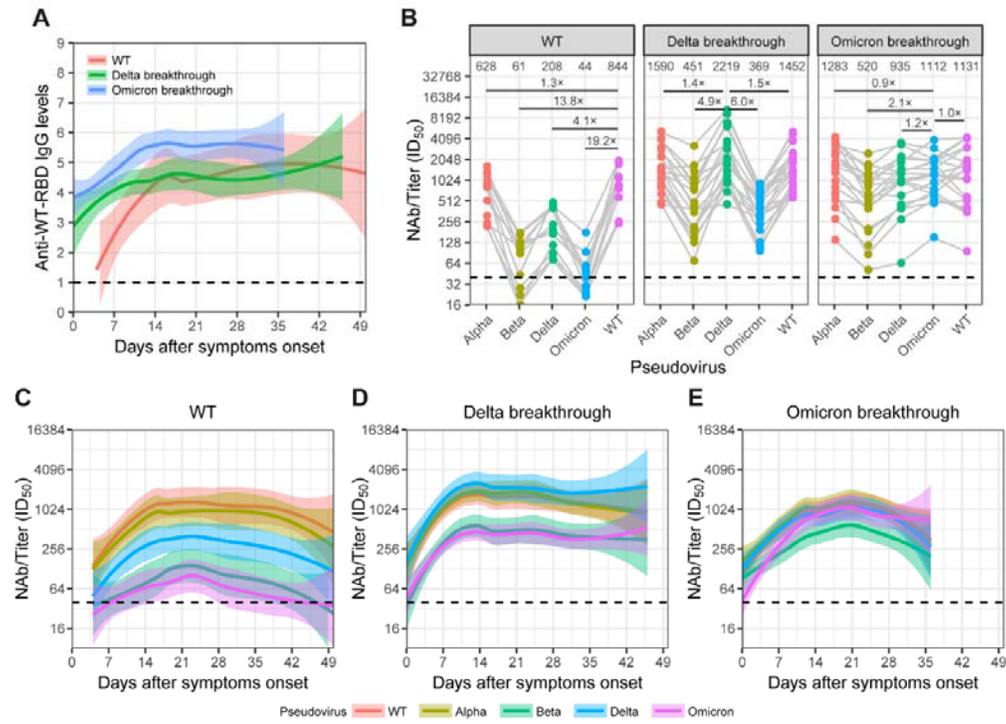
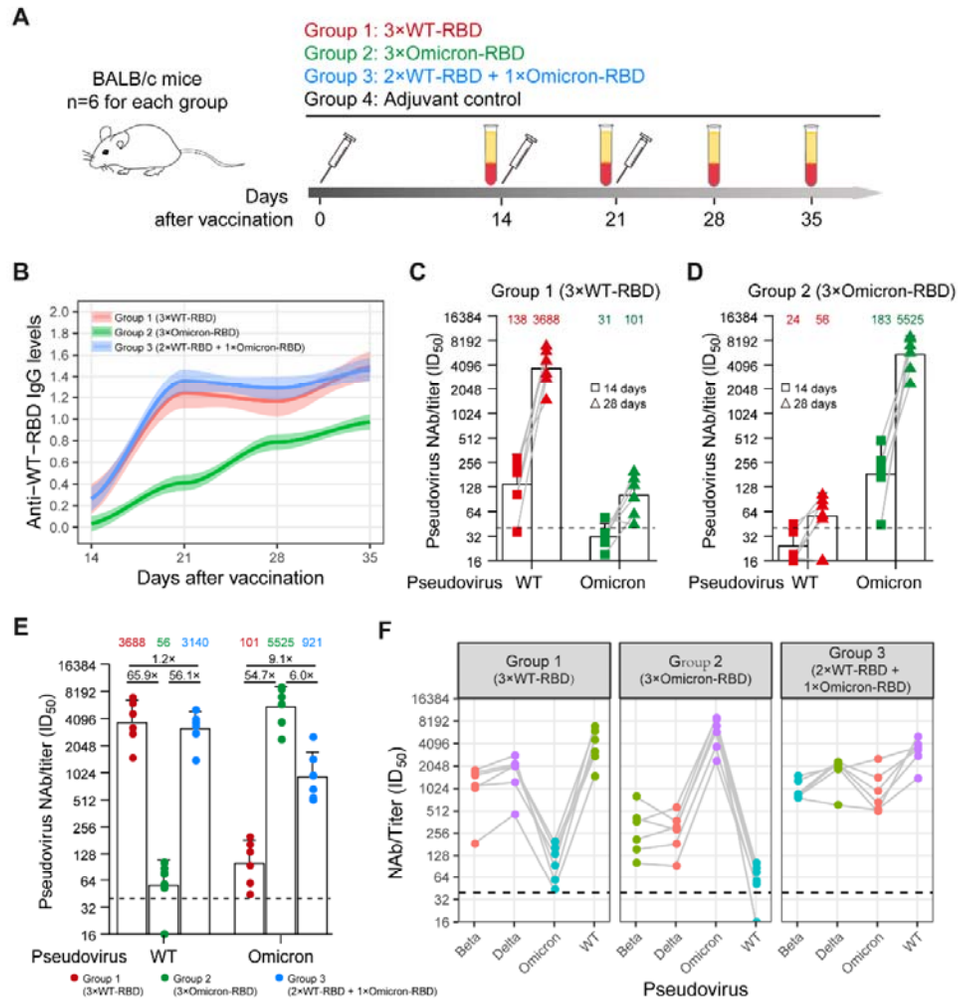


Fig. 2



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