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41	Low neutralization of SARS-CoV-2 Omicron BA.2.75.2, BQ.1.1, and XBB.1 by parental
42	mRNA vaccine or a BA.5-bivalent booster
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59	Abstract
60	The newly emerged SARS-CoV-2 Omicron sublineages, including the BA.2-derived
61	BA.2.75.2 and the BA.5-derived BQ.1.1 and XBB.1, have accumulated additional spike mutations
62	that may affect vaccine effectiveness. Here we report neutralizing activities of three human serum
63	panels collected from individuals 23-94 days after dose 4 of a parental mRNA vaccine, 14-32

64 days after a BA.5-bivalent-booster from individuals with 2-4 previous doses of parental mRNA 65 vaccine, or 15-32 days after a BA.5-bivalent-booster from individuals with previous SARS-CoV-2 infection and 2-4 doses of parental mRNA vaccine. The results showed that a BA.5-bivalent-66 booster elicited a high neutralizing titer against BA.4/5 measured at 14- to 32-day post-boost; 67 68 however, the BA.5-bivalent-booster did not produce robust neutralization against the newly emerged BA.2.75.2, BQ.1.1, or XBB.1. Previous infection significantly enhanced the magnitude 69 70 and breadth of BA.5-bivalent-booster-elicited neutralization. Our data support a vaccine update strategy that future boosters should match newly emerged circulating SARS-CoV-2 variants. 71

72

73 Main text

The continuous emergence of new variants of severe acute respiratory syndrome 74 coronavirus 2 (SARS-CoV-2) has caused successive global waves of infection. Since its first 75 report in November 2021 in South Africa, Omicron has become the dominating variant due to its 76 77 high transmissibility and immune evasion^{1,2}, with many Omicron sublineages emerging over time. The initial Omicron BA.1 was displaced by BA.2, which has further evolved to sublineages 78 BA.2.12.1, BA.2.75, BA.2.75.2, BA.4, and BA.5, among which BA.5 is currently dominant in many 79 countries. BA.4 and BA.5 have an identical spike sequence (defined as BA.4/5 hereafter) and 80 their offspring BA.4.6, BF.7, and BQ.1.1 are expanding in prevalence. As of November 19, 2022, 81 82 the BA.2-derived sublineage BA.2.75.2 accounted for 0.8% of the total SARS-CoV-2 infection in the United States; whereas the BA.4/5-derived sublineages BA.4.6, BF.7, BQ.1, and BQ.1.1 83 accounted for 4.4%, 7.8%, 25.5%, and 24.2% of total cases, respectively. In addition, another 84 85 BA.5-derived sublineage XBB, first identified in India in August 2022, is rapidly spreading in 86 Europe and has been detected in the United States. XBB was predominant in Singapore, 87 accounting for 54% of SARS-CoV-2 infections during the week of October 3-9, 2022 (Ministry of 88 Health, Singapore- https://www.moh.gov.sg/).

89 SARS-CoV-2 spike mutations often contribute to immune evasion and/or transmission 90 efficiency³⁻⁹. Previous studies showed that 3 or 4 doses of parental mRNA vaccine did not elicit robust neutralization against BA.4/5, supporting the development of bivalent vaccines that target 91 92 both the ancestral spike and the BA.4/5 spike protein¹⁰⁻¹². Since the newly emerged Omicron 93 sublineages have accumulated additional spike mutations (Fig. 1A), it is important to examine the vaccine-elicited neutralization against these new sublineages. The goal of this study was to 94 compare the neutralizing activities against six newly emerged Omicron sublineages (BA.5, BF.7, 95 BA.4.6, BA.2.75.2, BQ.1.1, and XBB.1) using human sera collected from individuals who received 96 4 doses of parental mRNA vaccine or a BA.5-bivalent-booster after 2-4 doses of parental mRNA 97 vaccine. 98

To facilitate neutralization measurement, we engineered the complete spike gene from 99 Omicron sublineage BA.4/5, BF.7, BA.4.6, BA.2.75.2, BQ.1.1, or XBB.1 into the backbone of 100 101 mNeonGreen (mNG) reporter USA-WA1/2020 SARS-CoV-2 (Fig. 1A). Compared with wild-type USA-WA1/2020 (a strain isolated in January 2020), insertion of mNG gene at open-reading-102 frame-7 of the viral genome attenuated the virus in vivo13. So, the engineered live-attenuated 103 mNG viruses can be used safely in a BSL3 facility with the correct procedures for neutralization 104 and antiviral testing¹⁴. Passage 1 of recombinant BA.4/5-, BF.7-, BA.4.6-, BA.2.75.2-, BQ.1.1-, 105 106 and XBB.1-spike mNG USA-WA1/2020 viruses were sequenced to ensure no undesired 107 mutations. Only the passage 1 virus stocks were used to determine the 50% fluorescent focusreduction neutralization titers (FFRNT₅₀) of vaccinated human sera, to ensure no additional spike 108 109 mutations in the tested recombinant viruses.

Three human serum panels with distinct vaccination and/or SARS-CoV-2 infection history
were analyzed. The first panel consisted of 25 sera obtained from individuals 23-94 (median 47)
days post dose 4 of parental monovalent mRNA-1273 or BNT162b2 vaccine (post-dose-4 sera);
these sera were collected from March 16 to June 30, 2022 (Extended Data Table 1). The second

114 panel consisted of 29 sera collected from individuals 14-32 (median 22) days post BA.5-bivalent-115 booster (BA.5-bivalent-booster sera); these specimens were collected from September 30 to October 22, 2022 (Extended Data Table 2). All sera from the first and second panels tested 116 negative against viral nucleocapsid protein (Extended Data Figure 1), suggesting no previous or 117 118 recent SARS-CoV-2 infection. The third panel consisted of 23 sera collected from individuals who were previously infected by SARS-CoV-2 (nucleocapsid antibody positive; Extended Data Figure 119 120 1) and received a BA.5-bivalent-booster 15-32 (median 21) days ago (BA.5-bivalent-boosterinfection sera); the viral infection time and genotype could not be determined because most 121 infections were asymptomatic; these samples were collected from October 4 to 22, 2022 122 (Extended Data Table 3). All participants from the second and third panels had also received 2, 123 3, or 4 doses of parental monovalent mRNA vaccine before receiving the BA.5-bivalent-booster. 124 125 Extended Data Table 1-3 summarize the serum information and neutralization for each serum 126 panel.

Post-dose-4 sera neutralized USA-WA1/2020-, BA.4/5-, BF.7-, BA.4.6-, BA.2.75.2-, 127 BQ.1.1-, and XBB.1-spike mNG SARS-CoV-2 with geometric mean titers (GMTs) of 1533, 95, 69, 128 62, 26, 22, and 15, respectively (Figure 1B and Extended Data Table 1). The neutralizing GMTs 129 against BA.4/5-, BF.7-, BA.4.6-, BA.2.75.2-, BQ.1.1-, and XBB.1-spike viruses were 16.1-, 22.2-, 130 24.7-, 59-, 69.7-, and 102-fold lower than the GMT against the USA-WA1/2020-spike virus, 131 respectively (Figure 1B). Compared with the GMT against the current dominant BA.4/5, the 132 neutralizing GMTs against BF.7-, BA.4.6-, BA.2.75.2-, BQ.1.1-, and XBB.1-spike viruses were 133 134 reduced by 1.4-, 1.5-, 3.7-, 4.3-, and 6.3-fold, respectively. The GMTs against BA.2.75.2 (26) and BQ.1.1 (22) were barely above 20, the detection limit of FFRNT; whereas the GMT against XBB.1 135 (15) was below the FFRNT detection limit. These results indicate that (i) 4 doses of parental 136 mRNA vaccine do not elicit robust neutralization against the newly emerged Omicron sublineages 137

when measured at 23-94 (median 47) days post-dose-4 and (ii) the rank of neutralization evasion is in the order of BA.4/5 < BF.7 \leq BA.4.6 < BA.2.75.2 \leq BQ.1.1 < XBB.1.

BA.5-bivalent-booster sera, collected at 14-32 (median 22) days post-boost, neutralized 140 USA-WA1/2020-, BA.4/5-, BF.7-, BA.4.6-, BA.2.75.2-, BQ.1.1-, and XBB.1-spike SARS-CoV-2s 141 142 with GMTs of 3620, 298, 305, 183, 98, 73, and 35, respectively (Figure 1C and Extended Data Table 2). The neutralizing GMTs against BA.4/5-, BF.7-, BA.4.6-, BA.2.75.2-, BQ.1.1-, and XBB.1-143 spike viruses were 12.1-, 11.9-, 19.8-, 36.9-, 49.6-, and 103-fold lower than the GMT against the 144 USA-WA1/2020, respectively (Figure 1C). The data indicate that although BA.5-bivalent-booster 145 146 elicits high neutralizing titers against BA.4/5 measured at 14-32 days post-boost, the neutralization against BA.2.75.2 (98), BQ.1.1 (73), and XBB.1 (35) remains low after BA.5-147 148 bivalent-booster.

BA.5-bivalent-booster-infection sera, collected at 15-32 (median 21) days post-boost, 149 neutralized USA-WA1/2020-, BA.4/5-, BF.7-, BA.4.6-, BA.2.75.2-, BQ.1.1-, and XBB.1-spike 150 SARS-CoV-2s with GMTs of 5776, 1558, 1223, 744, 367, 267, and 103, respectively (Figure 1D 151 and Extended Data Table 3). The neutralizing GMTs against BA.4/5-, BF.7-, BA.4.6-, BA.2.75.2-152 , BQ.1.1-, and XBB.1-spike viruses were 3.7-, 4.7-, 7.8-, 15.7-, 21.6-, and 56.1-fold lower than the 153 GMT against the USA-WA1/2020-spike SARS-CoV-2, respectively (Figure 1D). Compared with 154 BA.5-bivalent-booster sera without infection history, BA.5-bivalent-booster-infection sera 155 increased the neutralizing GMTs against USA-WA1/2020-, BA.4/5-, BF.7-, BA.4.6-, BA.2.75.2-, 156 BQ.1.1-, and XBB.1-spike viruses by 1.6-, 5.2-, 4.0-, 4.1-, 3.7-, 3.7-, and 2.9-fold, respectively 157 158 (Compare Figures 1C and 1D). The results suggest that (i) previous infection significantly 159 increases the magnitude and breadth of neutralization for BA.5-bivalent-booster and (ii) among the tested Omicron sublineages, XBB.1 exhibits the highest level of immune evasion. 160

161 Collectively, our neutralization results support two conclusions. First, the newly emerged 162 Omicron sublineages continue to increase their immune evasion of vaccine- and/or infection-

elicited neutralization. Among tested Omicron sublineages, BA.2.75.2, BQ.1.1, and XBB.1 exhibit the greatest evasion against vaccine-elicited neutralization, suggesting the potential of these new sublineages to dethrone BA.5 as the dominant lineage in circulation. Second, individuals with SARS-CoV-2 infection history develop higher and broader neutralization against the current circulating Omicron sublineages after the BA.5-bivalent booster.

The study has several limitations. First, we have not examined the antiviral roles of non-168 169 neutralizing antibodies and cell-mediated immunity. These two immune components, together with neutralizing antibodies, protect patients from severe disease and death^{15,16}. Unlike neutralizing 170 171 antibodies, many T cell epitopes after vaccination or natural infection are preserved in Omicron spikes¹⁷. However, robust antibody neutralization is critical to prevent viral infection¹⁸. Second, 172 we have not defined the spike mutations that contribute to the observed immune evasion of the 173 newly emerged Omicron sublineages. Spike mutation F486V was previously shown to drive the 174 175 immune evasion of BA.4/5¹⁰. The new Omicron sublineages BA.2.75.2, BA.4.6, BF.7, BQ.1.1, and XBB.1 share the spike R346T mutation that was reported to confer higher neutralization 176 evasion¹⁹. Third, the current results do not allow a direct comparison of neutralization between 177 parental mRNA vaccine and BA.5-bivalent-booster because of the differences in individuals' 178 179 demographics (e.g., age), numbers of vaccine doses, and serum collection time. Fourth, we don't 180 know (i) how neutralizing titers related to protection against infection, severe disease, or death; (ii) when and which variants infected individuals from the BA.5-bivalent-booster-infection cohort; 181 and (iii) the insight related to the differences in vaccine dose for Moderna's Bivalent (Original and 182 183 Omicron BA. 4/BA. 5) versus Pfizer/BioNTech's BA.4/BA.5-Adapted Bivalent Booster, and (iv) the baseline of the neutralization titers before boost were not determined due to sample unavailability. 184

Our laboratory investigation, along with the recent real-world effectiveness of BA.5bivalent-booster²⁰, supports a vaccine update strategy that future boosters should match new circulating SARS-CoV-2 variants. Given the advantage of mRNA vaccine platform that can rapidly

adapt to new antigen sequences, the key challenge is to determine the future booster sequencebefore new variants become prevalent in circulation.

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199 Author contributions

200 Conceptualization, P.R., X.X., P.-Y.S.; Methodology, C.K., J.Z., H.X., M.L., H.C.C., P.R., X.X., P.-

201 Y.S.; Investigation, C.K., J.Z., H.X., M.L., H.C.C., P.R., X.X., P.-Y.S.; Resources, P.R., X.X., P.-

202 Y.S.; Data Curation, C.K., J.Z., P.R., X.X.; Writing-Original Draft, P.R., X.X., P.-Y.S.; Writing-

203 Review & Editing, C.K., J.Z., H.X., M.L., H.C.C., P.R., X.X., P.-Y.S.; Supervision, P.R., X.X., P.-

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- 205

206 Competing interests

X.X. and P.-Y.S. have filed a patent on the reverse genetic system. X.X., J.Z., and P.-Y.S.
 received compensation from Pfizer for COVID-19 vaccine development. Other authors declare
 no competing interests.

211 Figure Legends

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213 Figure 1. Neutralization against Omicron sublineages. (A) Construction of Omicron sublineagespike mNG SARS-CoV-2. Amino acid mutations, deletions (Δ), and insertions (Ins) are indicated 214 in reference to the USA-WA1/2020 spike. L: leader sequence; ORF: open reading frame; NTD: 215 N-terminal domain of S1; RBD: receptor binding domain; S: spike glycoprotein; S1: N-terminal 216 217 furin cleavage fragment of S; S2: C-terminal furin cleavage fragment of S; E: envelope protein; M: membrane protein; N: nucleocapsid; UTR: untranslated region. (B) FFRNT₅₀s of human sera 218 after dose 4 parental mRNA vaccine. The p values (two-sided) for group comparison of GMTs are 219 220 the following. USA-WA1/2020 versus others: <0.0001; BA.4/5-spike versus BF.7-, BA.4.6-, BA.2.75.2-, BQ.1.1-, XBB.1-spike: 0.029, 0.001, <0.0001, <0.0001, <0.0001; BF.7-spike versus 221 BA.4.6-, BA.2.75.2-, BQ.1.1-, XBB.1-spike: 0.103, <0.0001, <0.0001, <0.0001; BA.4.6-spike 222 versus BA.2.75.2-, BQ.1.1-, XBB.1-spike: 0.0001, <0.0001, and <0.0001; BA.2.75.2-spike versus 223 BQ.1.1-, XBB.1-spike: 0.24, <0.0001; BQ.1.1-spike versus XBB.1-spike: 0.0028. The FFRNT₅₀ 224 225 values against BA.4/5-spike were reported previously¹¹. (C) FFRNT₅₀ of 29 sera collected after BA. 5-bivalent booster from individuals without infection history. The p values (two-sided) for 226 group comparison of GMTs are the following. USA-WA1/2020 versus others: <0.0001; BA.4/5-227 spike versus BF.7-, BA.4.6-, BA.2.75.2-, BQ.1.1-, XBB.1-spike: 0.844, <0.0001, <0.0001, 228 <0.0001, <0.0001; BF.7-spike versus BA.4.6-, BA.2.75.2-, BQ.1.1-, XBB.1-spike: all <0.0001; 229 BA.4.6-spike versus BA.2.75.2-, BQ.1.1-, XBB.1-spike: all <0.0001; BA.2.75.2-spike versus 230 231 BQ.1.1-, XBB.1-spike: 0.69, <0.0001; BQ.1.1-spike versus XBB.1-spike: <0.0001. (D) FFRNT₅₀ of 23 sera collected after BA.5-bivalent-booster from individuals with infection history. The p 232 values (two-sided) for group comparison of GMTs are the following. USA-WA1/2020 versus 233 others: <0.0001; BA.4/5-spike versus BF.7-, BA.4.6-, BA.2.75.2-, BQ.1.1-, XBB.1-spike: 0.0049, 234 <0.0001, <0.0001, <0.0001, <0.0001; BF.7-spike versus BA.4.6-, BA.2.75.2-, BQ.1.1-, XBB.1-235 spike: all <0.0001; BA.4.6-spike versus BA.2.75.2-, BQ.1.1-, XBB.1-spike: 0.0005, <0.0001, 236 <0.0001; BA.2.75.2-spike versus BQ.1.1-, XBB.1-spike: 0.114, <0.0001; BQ.1.1-spike versus 237 XBB.1-spike: <0.0001. For figure panels b-d, bar heights and the numbers above indicate GMTs. 238 Error bars indicate 95% CI. The fold of GMT reduction against each Omicron sublineage, 239 compared with the GMT against USA-WA1/2020, is shown in italic font. The dotted line indicates 240 the limit of detection of FFRNT₅₀. Statistic analyses were performed using the Wilcoxon matched-241 pairs signed-rank test for group comparison of GMTs. 242

244 References

245 1 Frederik Plesner Lyngse et al. Transmission of SARS-CoV-2 Omicron VOC subvariants BA.1 and 246 BA.2: Evidence from Danish Households. BioRxiv, doi: https://doi.org/10.1101/2022.1101.1128.22270044 (2022). 247 Xia, H. et al. Neutralization and durability of 2 or 3 doses of the BNT162b2 vaccine against 248 2 249 Omicron SARS-CoV-2. Cell Host Microbe 30, 485-488.e483 (2022). 250 https://doi.org:10.1016/j.chom.2022.02.015 251 3 Zou, J. et al. Neutralization against Omicron SARS-CoV-2 from previous non-Omicron infection. 252 Nat Commun 13, 852 (2022). https://doi.org:10.1038/s41467-022-28544-w 253 Liu, Y. et al. BNT162b2-Elicited Neutralization against New SARS-CoV-2 Spike Variants. N Engl J 4 254 Med 385, 472-474 (2021). https://doi.org:10.1056/NEJMc2106083 255 5 Liu, Y. et al. Neutralizing Activity of BNT162b2-Elicited Serum. N Engl J Med 384, 1466-1468 256 (2021). https://doi.org:10.1056/NEJMc2102017 257 6 Plante, J. A. et al. Spike mutation D614G alters SARS-CoV-2 fitness. Nature 592, 116-121 (2021). 258 https://doi.org:10.1038/s41586-020-2895-3 259 7 Liu, Y. et al. Delta spike P681R mutation enhances SARS-CoV-2 fitness over Alpha variant. Cell 260 Rep 39, 110829 (2022). https://doi.org:10.1016/j.celrep.2022.110829 261 Liu, J. et al. BNT162b2-elicited neutralization of B.1.617 and other SARS-CoV-2 variants. Nature 8 262 **596**, :273-275 (2021). <u>https://doi.org:10.1038/s41586-021-03693-y</u> 263 9 Liu, Y. et al. The N501Y spike substitution enhances SARS-CoV-2 transmission. Nature 602, 294-264 299 (2021). https://doi.org:10.1101/2021.03.08.434499 Kurhade, C. et al. Neutralization of Omicron sublineages and Deltacron SARS-CoV-2 by 3 doses of 265 10 266 BNT162b2 vaccine or BA.1 infection. Emerg Microbes Infect, 1-18 (2022). 267 https://doi.org:10.1080/22221751.2022.2099305 Xie, X. et al. Neutralization of SARS-CoV-2 Omicron sublineages by 4 doses of the original mRNA 268 11 vaccine. Cell Rep, 111729 (2022). https://doi.org:10.1016/j.celrep.2022.111729 269 270 Kurhade, C. et al. Neutralization of Omicron BA.1, BA.2, and BA.3 SARS-CoV-2 by 3 doses of 12 271 BNT162b2 vaccine. Nat Commun 13, 3602 (2022). https://doi.org:10.1038/s41467-022-30681-1 272 13 Johnson, B. A. et al. Nucleocapsid mutations in SARS-CoV-2 augment replication and pathogenesis. PLoS Pathog 18, e1010627 (2022). https://doi.org:10.1371/journal.ppat.1010627 273 274 14 Muruato, A. E. et al. A high-throughput neutralizing antibody assay for COVID-19 diagnosis and 275 vaccine evaluation. Nat Commun 11, 4059 (2020). https://doi.org:10.1038/s41467-020-17892-0 276 15 Bartsch, Y. C. et al. Omicron variant Spike-specific antibody binding and Fc activity are preserved in recipients of mRNA or inactivated COVID-19 vaccines. Sci Transl Med 14, eabn9243 (2022). 277 278 https://doi.org:10.1126/scitranslmed.abn9243 279 16 Grifoni, A. et al. SARS-CoV-2 human T cell epitopes: Adaptive immune response against COVID-280 19. Cell Host Microbe 29, 1076-1092 (2021). https://doi.org:10.1016/j.chom.2021.05.010 281 17 Redd, A. D. et al. Minimal Crossover between Mutations Associated with Omicron Variant of 282 SARS-CoV-2 and CD8(+) T-Cell Epitopes Identified in COVID-19 Convalescent Individuals. mBio, 283 e0361721 (2022). https://doi.org:10.1128/mbio.03617-21 284 Barouch, D. H. Covid-19 Vaccines - Immunity, Variants, Boosters. N Engl J Med (2022). 18 285 https://doi.org:10.1056/NEJMra2206573 286 19 Jian, F. et al. Further humoral immunity evasion of emerging SARS-CoV-2 BA.4 and BA.5 287 subvariants. Lancet Infect Dis 22, 1535-1537 (2022). https://doi.org:10.1016/S1473-288 3099(22)00642-9

289 20 290 291 292 293 294	Link-Gelles, R. <i>et al.</i> Effectiveness of Bivalent mRNA Vaccines in Preventing Symptomatic SARS- CoV-2 Infection — Increasing Community Access to Testing Program, United States, September– November 2022. <i>MMWR Morb Mortal Wkly Rep</i> ePub: 22 November 2022. (2022). <u>https://doi.org</u> : <u>http://dx.doi.org/10.15585/mmwr.mm7148e1</u>
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296 Methods

297 Ethical statement

All virus work was performed in a biosafety level 3 (BSL-3) laboratory with redundant fans in the 298 299 biosafety cabinets at The University of Texas Medical Branch at Galveston. All personnel wore powered air-purifying respirators (Breathe Easy, 3M) with Tyvek suits, aprons, booties, and 300 double gloves. The research protocol regarding the use of human serum specimens was reviewed 301 and approved by the University of Texas Medical Branch (UTMB) Institutional Review Board (IRB 302 number 20-0070). No informed consent was required since these deidentified sera were leftover 303 304 specimens from the routine standard of care and diagnostics before being discarded. No 305 diagnosis or treatment was involved either. The use of human serum specimens in this study was reviewed and approved by the University of Texas Medical Branch (UTMB) Institutional Review 306 Board (IRB number 20-0070). 307

308 Cells

Vero E6 (ATCC® CRL-1586) purchased from the American Type Culture Collection (ATCC, Bethesda, MD) and Vero E6 cells expressing TMPRSS2 (JCRB1819) purchased from SEKISUI XenoTech, LLC were maintained in a high-glucose Dulbecco's modified Eagle's medium (DMEM) containing 10% fetal bovine serum (FBS; HyClone Laboratories, South Logan, UT) and 1% penicillin/streptomycin at 37°C with 5% CO₂. Culture media and antibiotics were purchased from Thermo Fisher Scientific (Waltham, MA). Both cell lines were tested negative for *Mycoplasma*.

315 Human Serum

Three panels of human sera collected at UTMB were used in the study. Samples were collected based on availability. Varied ages with both genders are included. The population contains varied races or ethnicity, including white, Hispanic, black, and Asian. Subjects have received at least

319 two doses of the COVID-19 vaccine with or without evidence of SARS-CoV-2 infection. The first 320 panel consisted of 25 sera collected from individuals 23-94 (median 47) days after receiving dose 4 of parental vaccine mRNA-1273 or BNT162b2. This panel had been tested negative for SARS-321 CoV-2 nucleocapsid protein expression using Bio-Plex Pro Human IgG SARS-CoV-2 322 323 N/RBD/S1/S2 4-Plex Panel (Bio-rad). The second panel consisted of 29 sera collected from individuals 14-32 (median 22) days after BA.5-bivalent-booster from Pfizer (BA.4/BA.5-Adapted 324 325 Bivalent Booster) or Moderna (Bivalent Booster). All sera from this panel were tested negative for antibodies against SARS-CoV-2 nucleocapsid protein. The third panel consisted of 23 sera from 326 individuals who were previously infected by SARS-CoV-2 (as determined by SARS-CoV-2 327 nucleocapsid ELISA), vaccinated with 2-4 doses of parental mRNA vaccine, and received a BA.5-328 bivalent-booster 15-32 (median 21) days before serum collection. The genotypes of the infecting 329 330 SARS-CoV-2 variants could not be determined for the third serum panel. Patient information was 331 completely deidentified from all specimens. No informed consent was required because these deidentified sera were leftover specimens from standard care and diagnostics before being 332 discarded. The use of human sera for this study was reviewed and approved by the UTMB IRB 333 334 (number 20-0070). The de-identified human sera were heat-inactivated at 56°C for 30 min before the neutralization test. The serum information is presented in Extended Data Table 1-3. 335

336 Generation of recombinant Omicron sublineages-mNG SARS CoV-2

Recombinant Omicron sublineage BA.4/5-, BF.7-, BA.4.6-, BA.2.75.2-, BQ.1.1-, and XBB.1-spike 337 mNG SARS-CoV-2s was constructed by engineering the complete spike gene from the indicated 338 339 variants into an infectious cDNA clone of mNG USA-WA1/2020 as reported previously^{21,22}. Spike 340 sequences were based on BA.4/5 (BA.4: GISAID EPI_ISL_11 542270; BA.5: GISAID 341 EPI ISL 11542604; BA.4 and BA.5 have the identical spike sequence), BA.4.6 (GISAID EPI_ISL_15380489), 342 BA.2.75.2 (GISAID EPI_ISL_14458978), BF.7 (GISAID EPI_ISL_14425795), EPI_ISL_15542649) 343 BQ.1.1 (GISAID XBB.1 (GISAID and

344 EPI_ISL_15232105). The full-length infectious cDNA clone of SARS-CoV-2 was assembled by in 345 vitro ligation followed by in vitro transcription to synthesize the viral genomic RNA. The full-length RNA transcripts were electroporated in Vero E6-TMPRSS2 cells to recover the viruses. Viruses 346 347 were rescued post 2-3 days after electroporation and served as P0 stock. P0 stock was further 348 passaged once on Vero E6 cells to produce P1 stock. The reason for using Vero E6 cells (rather than using Vero E6-TMPRSS2) to prepare the P1 virus is that the infectivity of the P1 virus can 349 350 be affected by the cell types; since our established FFRNT assay uses Vero E6 cells, we chose 351 to prepare the P1 viruses using the same Vero E6 cells. The spike gene was sequenced from all P1 stock viruses to ensure no undesired mutation. The infectious titer of the P1 virus was 352 quantified by fluorescent focus assay on Vero E6 cells. The P1 virus was used for the 353 neutralization test. The protocols for the mutagenesis of mNG SARS-CoV-2 and virus production 354 355 were reported previously²³. All virus preparation and neutralization assays were carried out at the biosafety level 3 (BSL-3) facility at the University of Texas Medical Branch at Galveston. 356

357 Fluorescent focus reduction neutralization test (FFRNT)

Neutralization titers of human sera were measured by FFRNT using the USA-WA1/2020-, BA.4/5-358 , BF.7-, BA.4.6-, BA.2.75.2-, BQ.1.1- and XBB.1-spike mNG SARS-CoV-2s at BSL-3. The details 359 of the FFRNT protocol were reported previously²³. Briefly, 2.5×10^4 Vero E6 cells per well were 360 seeded in 96-well plates (Greiner Bio-one[™]). The cells were incubated overnight. On the next 361 day, each serum was 2-fold serially diluted in the culture medium with the first dilution of 1:20 362 (final dilution range of 1:20 to 1:20,480). The diluted serum was incubated with 100-150 FFUs of 363 364 mNG SARS-CoV-2 at 37 °C for 1 h, after which the serum virus mixtures were loaded onto the 365 pre-seeded Vero E6 cell monolayer in 96-well plates. After 1 h infection, the inoculum was 366 removed and 100 µl of overlay medium (supplemented with 0.8% methylcellulose) was added to each well. After incubating the plates at 37 °C for 16 h, raw images of mNG foci were acquired 367 368 using CytationTM 7 (BioTek) armed with 2.5× FL Zeiss objective with a wide-field of view and

369 processed using the Gen 5 software settings (GFP [469,525] threshold 4000, object selection size 370 50-1000 µm). The foci in each well were counted using the Gen5 software and normalized to the non-serum-treated controls to calculate the relative infectivities. The FFRNT₅₀ value was defined 371 as the minimal serum dilution that suppressed >50% of fluorescent foci. The neutralization titer of 372 each serum was determined in duplicate assays, and the geometric mean was taken. Tables S1-373 3 summarize the FFRNT₅₀ results. Data were initially plotted in GraphPad Prism 9 software and 374 assembled in Adobe Illustrator. FFRNT50 of <20 was treated as 10 for plot purposes and 375 statistical analysis. The above FFRNT₅₀ protocol has been reliably used to support the clinical 376 development of COVID-19 vaccines²⁴. Thus, we applied the same FFRNT protocol to the current 377 study. 378

379 Statistics & Reproducibility

No statistical method was used to predetermine the sample size. The samples were collected based on availability. No data were excluded from the analyses. The experiments were not randomized. Patient information was blinded in the study. The investigators were blinded to sample identity during data collection and/or analysis. The experiments were performed in duplication. All attempts at replication were successful.

Continuous variables were summarized as the geometric mean with 95% confidence intervals or median. Sera with undetectable (<20) antibody titers were assigned an antibody titer of 10, for purposes of GMT calculations or statistical comparisons. Comparison between neutralization titers was performed using a Wilcoxon matched-pairs signed-rank test using GraphPad Prism 9.0. Absolute *P* values were provided. *P*<0.05 was considered statistically significant. Images were assembled using Adobe Illustrator.

391 Data availability

- The raw data that support the findings of this study are shown in the Source data files. The sequence of SARS-CoV-2 variants can be accessed through GISAID (https://gisaid.org) with the following codes: BA.4/5 (BA.4: GISAID EPI_ISL_11 542270; BA.5: GISAID EPI_ISL_11542604; BA.4 and BA.5 have the identical spike sequence), BA.4.6 (GISAID EPI_ISL_15380489), BA.2.75.2 (GISAID EPI_ISL_14458978), BF.7 (GISAID EPI_ISL_14425795), BQ.1.1 (GISAID EPI_ISL_15542649) and XBB.1 (GISAID EPI_ISL_15232105). The sequence of SARS-CoV-2 mNG can be found in the supplementary information of our previous study²⁵.
- 399 Methods-only reference
- 400
 21
 Xie, X. et al. An Infectious cDNA Clone of SARS-CoV-2. Cell Host Microbe 27, 841-848 e843

 401
 (2020). https://doi.org:10.1016/j.chom.2020.04.004
- 402
 22
 Xie, X. *et al.* Engineering SARS-CoV-2 using a reverse genetic system. *Nat Protoc* **16**, 1761-1784

 403
 (2021). https://doi.org:10.1038/s41596-021-00491-8
- 40423Xie, X. et al. Neutralization of SARS-CoV-2 Omicron sublineages by 4 doses of the original mRNA405vaccine. Cell Reports 41, 111729 (2022).
- 406 <u>https://doi.org/10.1016/j.celrep.2022.111729</u>
- 407
 24
 Mulligan, M. J. *et al.* Phase I/II study of COVID-19 RNA vaccine BNT162b1 in adults. *Nature* 586,

 408
 589-593 (2020). <u>https://doi.org:10.1038/s41586-020-2639-4</u>
- 409 25 Muruato, A. E. *et al.* A high-throughput neutralizing antibody assay for COVID-19 diagnosis and
 410 vaccine evaluation. *Nat Commun* **11**, 4059 (2020). <u>https://doi.org:10.1038/s41467-020-17892-0</u>





Forum		Grades		Serum collection	Serum	contract to the contract of the				*FFRNT _{so}			
ID	Age (year)	(F/M)	Ethnicity	time (days post-dose 4 vaccination)	collection date	mRNA Vaccine type	USA- WA1/2020	^s BA.4/5 -spike	BF.7- spike	BA.4.6- spike	BA.2.75.2 -spike	BQ.1.1- spike	XBB.1- spike
1	62	F	White	37	3/16/2022	BTN162b	320	^10	10	10	10	10	10
2	80	M	Hispanic	30	4/7/2022	BTN162b	10240	320	80	80	40	20	20
3	84	F	White	55	5/11/2022	BTN162b	1280	80	14	10	10	10	10
4	92	F	White	27	5/12/2022	BTN162b	2560	113	40	40	28	10	10
5	78	M	White	56	5/17/2022	BTN162b	320	10	10	14	10	10	10
6	71	F	White	27	5/17/2022	mRNA-1273 (dose 1- 3), BTN162b (dose 4)	2560	320	320	320	80	80	40
7	83	F	White	23	5/18/2022	BTN162b	1280	160	113	80	20	20	10
8	87	м	White	24	5/19/2022	BTN162b	1280	113	226	160	10	57	10
9	80	м	White	52	5/23/2022	BTN162b	5120	640	320	226	80	80	40
10	84	м	White	26	5/24/2022	BTN162b	640	20	10	10	10	10	10
11	75	M	Black	47	5/25/2022	BTN162b	1810	57	40	40	226	20	40
12	90	м	Black	34	5/25/2022	BTN162b	2560	160	80	80	40	40	20
13	59	F	Hispanic	27	5/25/2022	BTN162b (dose 1-3), mRNA-1273 (dose 4)	1810	160	113	160	10	28	10
14	72	F	White	52	6/2/2022	BTN162b	1810	40	40	40	20	14	10
15	73	м	White	94	6/3/2022	mRNA-1273	640	40	40	28	10	10	10
16	67	F	Black	50	6/7/2022	BTN162b	1280	80	80	80	20	40	10
17	75	F	White	78	6/8/2022	BTN162b	2560	453	320	320	80	80	40
18	86	м	White	49	6/9/2022	BTN162b	1810	320	226	320	40	20	20
19	66	F	White	48	6/9/2022	BTN162b	453	20	20	14	10	10	10
20	80	м	Black	44	6/10/2022	BTN162b	1280	160	160	160	40	28	20
21	78	M	White	73	6/13/2022	BTN162b	905	40	40	28	20	10	10
22	86	м	White	51	6/16/2022	BTN162b	1280	20	20	20	10	10	10
23	84	F	White	35	6/20/2022	BTN162b	20480	1280	640	453	226	160	80
24	94	F	White	47	6/26/2022	BTN162b	1280	80	40	40	20	20	10
25	87	F	Hispanic	43	6/30/2022	BTN162b	905	160	320	160	28	14	10
Median	80	-	-	47	÷.,		-	-	+	-	-	-	-
#GMT	-		1.72	-			1533	95	69	62	26	22	15
*95% CI	6.50	5.7	1.7	-		(1 7 -)	1036-2268	56-160	41-115	38-103	17-38	16-31	12-20

*Individual FFRNT₅₀ value is the geometric mean of duplicate FFRNT results.

^FFRNT₅₀ of <20 was treated as 10 for plot purposes and statistical analysis.

#Geometric mean neutralizing titers (GMT).

[†]95% confidence interval (95% CI) for the GMT.

^{\$}This data set was reported previously⁵.

Sarum		200 - 200	1990 C	Serum collection	Serum	Last dose of mRNA				FFRNT ₅₀			
ID	Age (year)	Gender (F/M)	Race or Ethnicity	time (days post- BA.5-bivalent booster)	collection date	vaccine before BA.5- bivalent booster	USA- WA1/2020	BA.4/5- spike	BF.7- spike	BA.4.6- spike	BA.2.75 .2-spike	BQ.1.1- spike	XBB.1- spike
1	34	F	Black	15	9/30/2022	Dose 3	5120	640	1280	640	640	226	160
2	78	F	White	17	10/3/2022	Dose 4	5120	640	640	320	320	160	80
3	86	м	White	21	10/4/2002	Dose 4	1280	28	10	10	20	10	10
4	31	м	Asian	20	10/6/2022	Dose 2	5120	1280	1280	905	160	160	80
5	61	F	White	15	10/11/2022	Dose 4	5120	320	453	320	160	80	57
6	58	F	Black	14	10/11/2022	Dose 3	7241	1810	1810	1280	320	640	113
7	69	м	White	22	10/11/2022	Dose 4	2560	80	80	57	57	20	14
8	67	F	Asian	21	10/12/2022	Dose 4	20480	2560	1280	640	160	160	40
9	77	м	Asian	27	10/13/2022	Dose 4	2560	226	320	160	80	40	40
10	39	F	White	15	10/13/2022	Dose 3	5120	320	320	226	80	160	20
11	73	м	White	24	10/13/2022	Dose 4	1810	160	160	80	80	40	28
12	83	F	White	17	10/13/2022	Dose 4	1280	40	80	40	40	10	10
13	79	F	White	26	10/16/2022	Dose 3	3620	320	640	320	226	57	57
14	35	F	Asian	29	10/21/2022	Dose 3	5120	160	320	226	80	80	40
15	73	м	White	26	10/17/2022	Dose 3	7241	320	320	160	80	80	28
16	76	м	White	32	10/17/2022	Dose 4	7241	1280	905	640	226	160	160
17	71	м	White	28	10/18/2022	Dose 4	1280	40	57	40	40	20	10
18	22	м	Hispanic	19	10/18/2022	Dose 3	10240	2560	1280	1280	320	453	80
19	61	F	White	30	10/19/2022	Dose 4	640	80	80	57	20	28	14
20	56	м	White	21	10/19/2022	Dose 3	10240	640	640	453	160	160	80
21	66	F	White	26	10/19/2022	Dose 4	3620	320	320	160	80	80	40
22	76	F	White	30	10/20/2022	Dose 4	3620	226	226	160	10	80	10
23	61	F	Asian	31	10/20/2022	Dose 3	10240	5120	5120	2560	640	453	226
24	77	F	Black	31	10/20/2022	Dose 4	5120	80	80	40	40	28	10
25	59	F	unknown	28	10/21/2022	Dose 4	2560	320	320	160	160	80	28
26	71	м	Hispanic	22	10/21/2022	Dose 4	1280	160	160	80	80	80	14
27	70	F	White	22	10/21/2022	Dose 3	1280	113	57	28	20	20	10
28	79	F	White	25	10/22/2022	Dose 4	2560	226	160	80	57	10	20
29	79	M	White	18	10/22/2022	Dose 4	2560	160	320	160	226	160	80
Median	70	-	-	22	-		-	-	-	-	-	1	
#GMT		-	-				3620	298	305	183	98	73	35
*95% CI					2270		2668-4912	181-490	185-503	111-299	66-146	47-112	24-50

*Individual FFRNT₅₀ value is the geometric mean of duplicate FFRNT results. #Geometric mean neutralizing titers (GMT). [†]95% confidence interval (95% CI) for the GMT.

				Serum collection	Ferry	Last dose of				*FFRNT ₅₀				
Serum ID	Age (year)	Gender (F/M)	Race or Ethnicity	Race or Ethnicity	time (days post- BA.5-bivalent booster)	collection date	mRNA vaccine before BA.5- bivalent booster	USA- WA1/2020	BA.4/5- spike	BF.7- spike	BA.4.6- spike	BA.2.75.2 -spike	BQ.1.1- spike	XBB.1- spike
1	19	F	White	19	10/4/2022	Dose 3	10240	1280	1280	640	640	160	113	
2	69	F	White	15	10/6/2022	Dose 4	10240	5120	5120	2560	905	1280	320	
3	59	F	White	15	10/7/2022	Dose 3	20480	10240	10240	7241	2560	2560	453	
4	46	F	White	18	10/11/2022	Dose 2	5120	2560	2560	640	320	320	160	
\$5	80	F	Black	22	10/11/2022	Dose 3	10240	7241	5120	3620	2560	640	320	
6	76	F	Hispanic	21	10/12/2022	Dose 3	5120	1280	1280	905	453	160	113	
7	67	м	White	15	10/12/2022	Dose 4	5120	2560	1280	1280	320	320	80	
8	52	F	White	30	10/13/2022	Dose 4	5120	1280	1280	905	160	640	160	
9	48	F	Asian	24	10/13/2022	Dose 3	14482	7241	5120	2560	1280	640	320	
10	58	F	White	28	10/13/2022	Dose 2	5120	640	640	320	160	226	40	
11	67	F	Black	25	10/14/2022	Dose 2	5120	2560	2560	1280	640	640	160	
12	20	F	Hispanic	21	10/14/2022	Dose 2	1810	320	453	160	80	160	28	
^{\$} 13	75	м	White	21	10/14/2022	Dose 3	20480	2560	2560	1280	1280	453	320	
14	64	F	White	19	10/4/2022	Dose 4	7241	2560	2560	1280	2560	453	320	
15	90	F	White	22	10/18/2022	Dose 3	10240	1280	1280	1280	320	640	160	
16	35	F	Hispanic	32	10/18/2022	Dose 2	1280	320	226	160	80	80	28	
\$17	39	м	Asian	31	10/19/2022	Dose 3	10240	5120	2560	2560	640	320	80	
^{\$} 18	67	м	White	17	10/19/2022	Dose 4	10240	2560	1810	1280	640	640	226	
\$19	68	м	Hispanic	14	10/19/2022	Dose 3	5120	2560	1280	905	320	160	80	
20	51	M	White	28	10/21/2022	Dose 3	7241	905	320	320	320	160	57	
21	65	м	Asian	29	10/20/2022	Dose 4	2560	453	160	80	113	57	40	
22	64	м	White	16	10/21/2022	Dose 3	2560	640	640	320	40	40	20	
\$23	73	M	Black	31	10/22/2022	Dose 4	640	57	40	28	20	10	10	
Median	64	-	-	21		-	-	-	-	-	-	-	-	
#GMT	-	<u></u>	20			540 C	5776	1558	1223	744	367	267	103	
*95% CI		2	12	2	-		3994-8352	922-2631	704-2125	425-1301	209-644	158-452	66-162	

*Individual FFRNT₅₀ value is the geometric mean of duplicate FFRNT results.

*Geometric mean neutralizing titers (GMT).

[†]95% confidence interval (95% CI) for the GMT.

Reported as SARS-CoV-2 RT-PCR positive. Nucleocapsid testing was not performed.

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\boxtimes		For Bayesian analysis, information on the choice of priors and Markov chain Monte Carlo settings
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		Our web collection on <u>statistics for biologists</u> contains articles on many of the points above.

Software and code

Policy information about availability of computer code				
Data collection	Cytation 7 with Gen5 software			
Data analysis	GraphPad Prism 9.0, Adobe Illustrator			

For manuscripts utilizing custom algorithms or software that are central to the research but not yet described in published literature, software must be made available to editors and reviewers. We strongly encourage code deposition in a community repository (e.g. GitHub). See the Nature Portfolio guidelines for submitting code & software for further information.

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All manuscripts must include a data availability statement. This statement should provide the following information, where applicable:

- Accession codes, unique identifiers, or web links for publicly available datasets
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The raw data that support the findings of this study are shown in the Source data files. The sequence of SARS-CoV-2 variants can be accessed through GISAID (https://gisaid.org) with the following codes: BA.4/5 (BA.4: GISAID EPI_ISL_11 542270; BA.5: GISAID EPI_ISL_11542604; BA.4 and BA.5 have the identical spike sequence), BA.4.6 (GISAID EPI_ISL_15380489), BA.2.75.2 (GISAID EPI_ISL_14458978), BF.7 (GISAID EPI_ISL_14425795), BQ.1.1 (GISAID EPI_ISL_15542649) and XBB.1 (GISAID EPI_ISL_15232105). The sequence of SARS-CoV-2 mNG can be found in the supplementary information of our previous study.

Human research participants

Policy information about studies involving human research participants and Sex and Gender in Research.

Reporting on sex and gender	Human serum samples were collected based on availability. Both gender are included.
Population characteristics	Samples were collected based on availability. Varied ages with both genders are included. The population contains varied races or ethnicity, including white, Hispanic, black, and Asian. Subjects have received at least two doses of the COVID-19 vaccine with or without evidence of SARS-CoV-2 infection.
Recruitment	No patients were recruited in this study. No informed consent was required, because these deidentified sera were leftover specimens before being discarded. No diagnoses or treatment was involved either.
Ethics oversight	The research protocol regarding the use of human serum specimens was reviewed and approved by the University of Texas Medical Branch (UTMB) Institutional Review Board (IRB#: 20-0070).

Note that full information on the approval of the study protocol must also be provided in the manuscript.

Field-specific reporting

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Life sciences study design

All studies must disclose on these points even when the disclosure is negative.

Sample size	No sample size calculation was performed. Samples were collected based on the availability.
Data exclusions	No data was excluded in the study.
Replication	Each human serum sample was analyzed in duplication. The averaged results from the duplication were reported in this study. All attempts at replication were successful.
Randomization	This is no randomization in this study. All samples available were analyzed for the neutralizing activities against WT SARS-CoV-2 and variants in the same experimental settings.
Blinding	Patient information was blinded in the study. The investigators were blinded to sample identity during data collection and/or analysis.

Reporting for specific materials, systems and methods

We require information from authors about some types of materials, experimental systems and methods used in many studies. Here, indicate whether each material, system or method listed is relevant to your study. If you are not sure if a list item applies to your research, read the appropriate section before selecting a response.

Materials & experimental systems			thods	
n/a	Involved in the study	n/a	Involved in the study	
\boxtimes	Antibodies	\boxtimes	ChIP-seq	
	Eukaryotic cell lines	\boxtimes	Flow cytometry	
\ge	Palaeontology and archaeology	\boxtimes	MRI-based neuroimaging	
\ge	Animals and other organisms			
\boxtimes	Clinical data			
\times	Dual use research of concern			

Eukaryotic cell lines

Policy information about <u>cell lines</u>	s and Sex and Gender in Research
Cell line source(s)	Vero E6 cells (ATCC [®] CRL-1586) were obtained from ATCC; VeroE6-TMPRSS2 cells were Vero E6 cells expressing TMPRSS2 purchased from SEKISUI XenoTech, LLC.
Authentication	VeroE6 Cells have been authenticated by ATCC using morphologies and other groups using STR profiling (reference: Almeida JL, Hill CR, Cole KD. VeroE6-TMPRSS2 cells have been authenticated by the vendor. Authentication of African green monkey cell lines using human short tandem repeat markers. BMC Biotechnol. 2011;11:102. Published 2011 Nov 7. doi:10.1186/1472-6750-11-102).
Mycoplasma contamination	All cell lines were tested negative for mycoplasma.
Commonly misidentified lines (See <u>ICLAC</u> register)	None.