1	Combating the SARS-CoV-2 Omicron variant with non-Omicron						
2	neutralizing antibodies						
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23							

24 Abstract

The highly mutated and transmissible Omicron variant has provoked serious 25 concerns over its decreased sensitivity to the current coronavirus disease 2019 26 (COVID-19) vaccines and evasion from most anti-severe acute respiratory 27 syndrome coronavirus 2 (SARS-CoV-2) neutralizing antibodies (NAbs). In this 28 study, we explored the possibility of combatting the Omicron variant by 29 30 constructing bispecific antibodies based on non-Omicron NAbs. We engineered ten IgG-like bispecific antibodies with non-Omicron NAbs named GW01, 16L9, 31 4L12, and REGN10987 by fusing the single-chain variable fragments (scFvs) 32 of two antibodies through a linker and then connecting them to the Fc region of 33 IgG1. Surprisingly, eight out of ten bispecific antibodies showed high binding 34 affinity to the Omicron receptor-binding domain (RBD) and exhibited extreme 35 breadth and potency against pseudotyped SARS-CoV-2 variants of concern 36 (VOCs) including Omicron, as well as authentic Omicron(+R346K) variants. Six 37 38 bispecific antibodies containing the cross-NAb GW01 neutralized Omicron variant and retained their abilities to neutralize other sarbecoviruses. Bispecific 39 antibodies inhibited Omicron infection by binding to the ACE2 binding site. A 40 cryo-electron microscopy (cryo-EM) structure study of the representative 41 bispecific antibody FD01 in complex with the Omicron spike (S) revealed 5 42 distinct trimers and one unique bi-trimer conformation. The structure and 43 mapping analyses of 34 Omicron S variant single mutants elucidated that two 44 scFvs of the bispecific antibody synergistically induced the RBD-down 45 conformation into 3-RBD-up conformation, enlarged the interface area, 46 accommodated the S371L mutation, improved the affinity between a single IgG 47 and the Omicron RBD, and hindered ACE2 binding by forming bi-trimer 48 conformation. Our study offers an important foundation for anti-Omicron NAb 49 design. Engineering bispecific antibodies based on non-Omicron NAbs may 50 provide an efficient solution to combat the Omicron variant. 51

53 INTRODUCTION

Omicron, first identified in South Africa and reported to the WHO at the end of 54 Nov. 2021, became the dominant severe acute respiratory syndrome 55 coronavirus 2 (SARS-CoV-2) variant globally in Jan. 2022. Omicron is the most 56 severely altered version of SARS-CoV-2, with more than 30 mutations in the 57 spike protein, sixteen of which are in the receptor-binding domain (RBD). 58 59 Omicron variants extensively escape neutralization by sera from vaccinated or convalescent individuals ¹⁻¹⁰. Moreover, most neutralizing antibodies (NAbs), 60 including many clinical-stage monoclonal antibodies (mAbs), have completely 61 lost their neutralization potency against Omicron ^{1,3,4,11}. Therefore, there is an 62 urgent need to explore and develop countermeasures against the Omicron 63 variant. 64

In this study, we used four NAbs named GW01, 16L9, 4L12, and REGN10987, 65 which failed to bind or neutralize the Omicron variant, to engineered full-length 66 67 IgG-like bispecific antibodies. We surprisingly found that these bispecific antibodies could neutralize all the VOCs, including Omicron and authentic 68 Omicron(+R346K), while the parental antibody cocktail 69 showed no neutralization against Omicron. Cryo-EM structure study showed six dynamic 70 states of the Omicron S trimer upon bispecific antibody binding, including a 71 novel bi-trimer conformation, within which RBDs were all in "up" conformations. 72 This bi-trimer is critical for inhibiting ACE2 binding and explains the superiority 73 of the bispecific antibody. These novel bispecific antibodies are strong 74 candidates for the treatment and prevention of infection with the Omicron 75 variant and VOCs and other sarbecoviruses that may cause future emerging or 76 reemerging coronavirus diseases. 77

78

79 **RESULTS**

80 Isolation of three non-Omicron neutralizing antibodies from COVID-19

81 convalescent individuals

We sorted and cultured SARS-CoV-2 S-specific memory B cells from two recovered coronavirus disease 2019 (COVID-19) patients and discovered three anti-SARS-CoV-2 NAbs, designated GW01, 4L12, and 16L9. The germlines and CDR3 of these antibodies are listed in Table S1. All three antibodies showed strong binding to the RBD of SARS-CoV-2 (**Fig. 1A**). However, they had no or weak binding to the S trimer or S RBD of the Omicron variant (**Fig. 1A**).

GW01, 4L12, and 16L9 potently neutralized SARS-CoV-2 and the VOCs Alpha, 89 Beta, Gamma, and Delta, but they failed to neutralize the Omicron variant (Fig. 90 **1B**). A panel of control NAbs failed to neutralize the Omicron except S309. S309 91 neutralized Omicron to a similar degree as previous reports^{12,13}. GW01 was a 92 cross-NAb that was able to neutralize SARS-CoV and the SARS-related 93 coronaviruses (SARSr-CoVs) RS3367 and WIV1. GW01 antibodies showed no 94 competition with 4L12, 16L9 or the control antibody REGN10987 in binding the 95 96 RBD (Fig. 1C), indicating that GW01 binds to an epitope different from that bound by 4L12, 16L9, and REGN10987. 97

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Binding and neutralization of the Omicron variant by bispecific antibodies 99 100 We constructed bispecific antibodies targeting different epitopes in the RBD using GW01 in combination with 16L9, 4L12, and REGN10987, and explored 101 their possibilities to neutralize Omicron variant. We linked the single-chain 102 variable fragments (scFvs) of the parental antibodies with a (Gly₄Ser)₄ linker 103 104 and then fused them to a hinge-CH2-CH3 fragment of human immunoglobulin (hlgG1 Fc) to generate a single gene-encoded lgG-like bispecific antibody (Fig. 105 **2A**). For example, the sequence order of the GW01-16L9 (FD01) bispecific 106 antibody was as follows: GW01 VL-(Gly4Ser)3-GW01 VH-(Gly4Ser)4-16L9 VL-107 (Gly₄Ser)₃-16L9 VH-hinge-CH2-CH3. SDS–PAGE results showed that the size 108 of the single chain of two representative bispecific antibodies, GW01-16L9 and 109

110 16L9-GW01, was approximately 100 kDa and that the purity was >95% (Fig.
111 2B). Crosslinking bispecific antibodies with glutaraldehyde revealed that the full
112 size of the bispecific antibodies was approximately 200 kDa, which was 10%
113 larger than that of the parental antibodies (180 kDa, Fig. 2B).

We constructed ten bispecific antibodies and tested their binding abilities to the 114 RBD or S trimer of SARS-CoV-2 and the Omicron variant. Eight bispecific 115 antibodies, FD01 (GW01-16L9), 16L9-GW01, GW01-REGN10987, 116 117 REGN10987-GW01, GW01-4L12, 4L12-GW01, 4L12-REGN10987, and 4L12-16L9, not only strongly bound to the RBD of SARS-CoV-2 and the S trimer and 118 RBD proteins of the Omicron variant (Fig. 2C) but also showed high binding 119 affinity to these proteins (Fig. 2D). These results indicated that the structure of 120 the bispecific antibody increased the parental antibody binding affinity to the 121 Omicron RBD. 122

To understand the breadth of these bispecific antibodies, we performed a 123 124 neutralization assay using SARS-CoV-2 pseudoviruses, including Alpha, Beta, Gamma, Delta, and Omicron variants, and the sarbecoviruses SARS-CoV, 125 WIV1 and RS3367. Surprisingly, these eight bispecific antibodies potently 126 neutralized the Omicron variant with IC50 values from 39.7 to 548 ng/ml. Six 127 bispecific antibodies containing the cross-NAb GW01 strongly neutralized all 128 the tested VOCs and sarbecoviruses (Fig. 2E). FD01 and GW01-REGN10987 129 were the best broadly NAbs, with geometric mean (GM) IC50 values of 25.4 130 and 12.1 ng/ml, respectively. 4L12-REGN10987 and 4L12-16L9 strongly 131 132 neutralized all the tested VOCs with GM IC50 values of 6.9 and 6.8 ng/ml, respectively (Fig. 2E). However, the parental NAb combinations showed no 133 neutralization against the Omicron variant (Fig. 2E). Taken together, these data 134 indicated that bispecific antibodies consisting of non-Omicron NAbs efficiently 135 neutralize the Omicron variant in a way that is different from the antibody 136 cocktail. 137

138 To confirm the neutralization efficacy of the bispecific antibodies, we performed

plaque reduction neutralization assays with an authentic Omicron variant containing the R346K mutation, which escapes more SARS-CoV-2 NAbs than the Omicron variant³. All five representative bispecific antibodies efficiently neutralized the live Omicron variant (**Fig. 2F**), confirming that the bispecific antibodies composed of non-Omicron NAbs are able to neutralize the Omicron variant.

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146 Bispecific antibodies bind to the ACE2-binding site

Competition assays were then performed to evaluate the abilities of the 147 bispecific antibodies to inhibit the binding of SARS-CoV-2 RBD to the 148 recombinant ACE2 protein (Fig. 2G). All five representative bispecific 149 antibodies prevented RBD binding to ACE2 protein, while the control antibody 150 S309 did not affect the S/ACE2 interaction. FD01 showed a strong inhibitory 151 effect against ACE2 binding. These results indicated that bispecific antibodies 152 inhibit Omicron variant infection by occupying the ACE2-binding site on the 153 154 RBD.

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156 Snapshots of Omicron S-FD01 structures determined by cryo-EM

To further investigate the neutralization mechanism of the bispecific antibodies, 157 we chose FD01 (GW01-16L9) as a representative antibody for structural study. 158 Local refinement focused on the RBD and ScFvs improved the interface region 159 to 3.51 Å resolution and allowed us to unambiguously build the RBD and scFvs 160 (Table S2). We determined the cryo-EM structure of the prefusion stabilized 161 SARS-CoV-2 Omicron S ectodomain trimer in complex with the bispecific 162 antibody FD01 (Omicron S-FD01), revealing 6 states of the complex: In the 163 state 1 Omicron S-FD01 structure (~26% of the particles), only one 16L9 binds 164 to the "up" RBD, and the two down RBDs have no antibody binding (up-down-165 down RBDs, 1 scFv, 3.47 Å). In state 2 (~38%), the bispecific antibody FD01 166 (GW01-16L9) binds to a widely open RBD, the so-called "wide up" state, 16L9 167 binds to an "up" RBD, and the third RBD remains in a down state (wide up-up-168

down RBDs, 3 scFvs, 3.70 Å). In both state 3 (~8%) and state 4 (~15%), two 169 FD01 (GW01-16L9) bind separately to two "wide up" state RBDs, and the third 170 RBD represents the half-up conformation in state 3 (wide up-wide up-half up 171 RBDs, 4 scFvs, 3.91 Å) and the up conformation in state 4 (wide up-wide up-172 up RBDs, 4 scFvs, 3.47 Å), without antibody binding. In state 5 (~4%), all RBDs 173 are in the "wide up" state, each bound with an FD01 (GW01-16L9) (all wide up 174 RBDs, 6 scFvs, 3.87 Å). In the final state 6 (~4% of the particles), two state 5 175 trimers are connected by three bispecific FD01 antibodies, forming a bi-trimer 176 structure (bi-trimer, all wide up RBDs, 12 scFvs, 6.11 Å) (Fig. 3, Fig. S2 and 177 S3). 178

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180 Collaborative binding mechanism of FD01 bispecific antibody

These six cryo-EM structures represent the conformational transitions of the 181 Omicron S trimer upon FD01 binding. To simplify the presentation, three 182 protomers of a spike trimer are clockwisely defined as 1, 2, and 3 (Fig. 3, Fig. 183 4). The apo spike trimer (state 0) includes one regular up RBD and two down 184 RBDs. First, 16L9 binds to the "up" RBD (RBD-1) of the apo spike trimers and 185 forms the state 1 confirmation. After that, GW01, connected with 16L9, binds to 186 RBD-1, inducing it into a wide up state (via an ~13 Å outward motion) and 187 pushing RBD-2 to flip from the "down" to the "up" state, making enough space 188 to accommodate the first bispecific antibody FD01 (16L9 and GW01) on RBD-189 1 and the 16L9 of the second FD01 on RBD-2. A slight ~3 Å inward motion of 190 RBD-3 is induced by the neighboring RBD-2 to stabilize this state (state 2). 191 Then, in state 3, the "up" RBD-2 opens up further to the "wide up" state, 192 allowing both 16L9 and GW01 of the second FD01to bind RBD-2. In addition, 193 RBD-3 is pushed up to a "half up" state. In state 4, the "half up" RBD-3 opens 194 up to the regular "up" state and is ready for 16L9 binding. Following that, in state 195 5, the third FD01 binds to RBD-3 and induces RBD-3 to adopt the "wide up" 196

state. Finally, two trimers in state 5 form a bi-trimer induced by three pairs of Fc
 regions from six antibodies.

Thus, the six states represent the continuous conformational transitions starting from the first 16L9 binding to three FD01 binding and the final bi-trimer formation, which inhibits the ACE2 binding by aggregating virions. In addition to the motion of RBDs, the N-terminal domains (NTDs) of the trimer are also moved or rotated following the motion of their neighboring RBDs.

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FD01 targets two conserved epitopes

16L9 and GW01 bind to two different sites of one RBD (Fig. 5A). The epitope 206 of 16L9 almost overlaps with the receptor-binding motif (RBM), while GW01 207 binds outside the RBM. The binding of 16L9 and the RBD buries a 1061 $Å^2$ 208 surface area, and a total of 20 residues from RBD are involved. The interaction 209 between 16L9 and RBD is largely driven by extensive hydrophilic and 210 hydrophobic interactions between CDRH1, CDRH2, CDRH3 and CDRL1 of 211 16L9 and RBD (Fig. 5C). Residues D405, T415, D420, Y421, Y453, L455, F456, 212 Y473, A475, N477, Y489, R493, P500, Y501, and H505 of the RBD are involved 213 in this interaction, forming 14 pairs of hydrogen bonds and 3 patches of 214 hydrophobic interactions (Fig. 5C). In addition, the hydrogen bond between S96 215 of CDRL3 and R403 from the RBD and the salt bridge between E52 of CDRL2 216 and R493 from the RBD further enhance the interaction (Fig. 5C). 217 Coincidentally, residues Y453, A475, Y489, R493, T500, Y501, and H505 of the 218 Omicron RBD are important for ACE2 recognition and binding¹⁴. The 219 neutralization activity test showed that 16L9 alone was able to broadly 220 neutralize SARS-CoV-2 and SARS-CoV-2 variants. This result implies that 221 16L9 targets the conserved residues of the RBM, which is needed for receptor 222 223 binding.

GW01 interacts with another novel conserved epitope beyond the binding site of 16L9. The binding site of GW01 and the RBD has a buried surface area of 668.2 Å. The interaction between GW01 and the RBD is mainly contributed by 227 CDRH3. The long loop (226-YGPPDVFNY-234) of CDRH3 engages with Y369,

F374, T376, F377, Y508, and V503 from the RBD, forming 3 patches of
hydrophobic interactions and 2 pairs of hydrogen bonds. D155 on CDRH1 and
N53 on CDRL2 are also involved in the interaction by forming hydrogen bonds
between V503 and N370, respectively (Fig. 5D).

Interestingly, the simultaneous binding of 16L9 and GW01 with the RBD
introduces additional interactions. Hydrogen bonds are formed between N178
and N196 of GW01 and S26, Y93, S96 and N98 of 16L9 (Fig. 5B), which further
enhances the interaction between FD01 and S.

Structural alignment of the 16L9-GW01-RBD complex with the ACE2–RBD
complex indicated that both 16L9 and GW01 were able to compete with ACE2
when binding to the RBD (Fig. 5E), which is consistent to the competition assay.

Bispecific antibodies accommodated the mutations in the Omicronvariant

242 We constructed 34 single mutants of the Omicron variant to identify the key residues that mediate resistance to GW01, 16L9, 4L12, and REGN0987. The 243 S371L mutation, which was found to stabilize the Omicron into a single-RBD-244 down conformation¹², greatly decreased the neutralization activities of GW01, 245 4L12, and REGN10987 (Table 1). The S375F mutation decreased the 246 neutralization activity of GW01 by 16-fold and resulted in resistance to 247 REGN10987. The K417N mutation resulted in complete resistance to 16L9 248 (>1000-fold). All six tested bispecific antibodies showed only a slight decrease 249 250 (12.4- to 25.5-fold) or no change in neutralization activity against the S371L mutant. Therefore, bispecific antibodies bind to the ACE2-binding site of the 251 RBD and accommodate the S371L mutation of the Omicron variant, resulting 252 in extraordinary breadth for these bispecific antibodies. 253

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255 **DISCUSSION**

The recently emerged SARS-CoV-2 Omicron strain raised unprecedented 256 global concern about invalidation of most FDA-approved antibody drugs, 257 including LY-CoV555, LY-CoV016, REGN10933, REGN10987, AZD8895 and 258 AZD1061¹⁵. Previous studies have shown that combining two NAbs that target 259 different neutralizing epitopes in the SARS-CoV-2 S protein increased 260 therapeutic and prophylactic efficacy. However, the most potent anti-SARS-261 CoV-2 antibody cocktail, RENG10933/REGN10987, failed to neutralize the 262 263 Omicron variant. Zhou tested 10 antibody combinations against Omicron and three antibody combinations with increasing neutralization contained a VH1-58 264 derived anti-Omicron NAb that bound RBD in the "up" position. Cyro-EM 265 structure showed that the antibody combination with improved neutralization 266 (B1-182.1/A19-46.1) synergistically induced the 3-RBD-up conformation¹². 267 However, this antibody combination approach is not an ideal solution because 268 few anti-Omicron NAbs are available. 269

Using the NAbs that failed to neutralize the Omicron variant, we constructed a 270 271 serial of novel bispecific antibodies that capable of neutralizing all SARA-CoV-2 variants of concern (VOCs), including the Omicron strain. Interestingly, single 272 IgG parental antibodies or the combination of parental antibodies failed to 273 neutralize Omicron, although the neutralization activity against SARS-CoV-2 or 274 SARS-CoV-2 Alpha, Beta, Gamma, and Delta was remarkable. Thus, the 275 effective neutralization of Omicron by bispecific antibodies may be due to the 276 277 construction of this bispecific antibody.

The structure of FD01 (GW01-16L9) bispecific antibody gives a hit at this 278 279 collaborative binding mechanism. First, one 16L9 scFv binds to the exposed epitope of the "up" state RBD-1 in the apo Omicron S trimer as a trigger. Then, 280 the 20 aa GS linker between 16L9 and GW01 guides GW01 to its targeting 281 epitope of RBD-1, pushing RBD-1 more open, which unlocks the "down" state 282 of RBD-2 and induces it to adopt the up state. Thus, another 16L9 scFv could 283 easily catch the "up" state RBD-2. The same triggering process would occur on 284 RBD-2 and RBD-3, allowing the binding of the second and the third FD01 to 285

RBD-2 and RBD-3, respectively. GW01 Fab alone could not trigger binding to 286 the apo state of the Omicron trimer without the help of 16L9 and the GS linker 287 guider. Although 16L9 can bind to the first "up" state RBD, it lacks the ability to 288 release other "down" state RBDs for further binding (Fig. S5). In summary, the 289 two scFvs of the GW01-16L9 bispecific antibody have collaborative roles in the 290 neutralization process. The neutralization mechanism of FD01 may be 291 mediated by the unique engineering of the combination of two antibodies into 292 293 one, which induces RBD-up conformation, enlarges the interface area, improves the affinity of a single IgG and the RBD, stabilizes the interaction by 294 additional interactions between the two antibodies, forms bi-trimer which 295 hinders the ACE2 binding by aggregating virions, and therefore blocks the 296 infection of SARS-CoV-2 Omicron variant. 297

Taken together, the unique construction of bispecific antibodies enables non-Omicron NAbs to neutralize Omicron variant. Our approach can rescue the majority of the SARS-CoV-2 antibodies, such as REGN10987, to overcome the resistance of Omicron and prepare for future SARS-CoV-2 variants.

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312 AUTHOR CONTRIBUTIONS

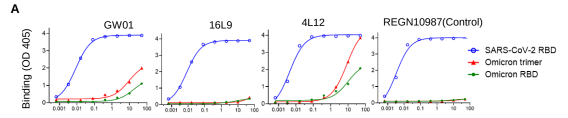
313 JH, LS, and FW conceived and designed the experiments. JH, ML, and FW

performed B cell sorting and antibody cloning. YDW and YJ constructed the 314 bispecific antibodies and performed neutralization assay, ELISA, bilayer 315 interferometry experiments. YDW, YJ, and QW constructed and expressed 316 SARS-CoV-2 pseudovirus mutants and purification antibodies. LS, XZ, WZ, and 317 ZC performed the structural studies. JZ and YQW were responsible for the 318 authentic virus experiments. JH, YDW, YJ, FW, LS, XZ, WZ, JZ, and YQW 319 analyzed the data. YW and TZ supervised the project. JH, LS, FW, YDW, and 320 321 XZ wrote the manuscript.

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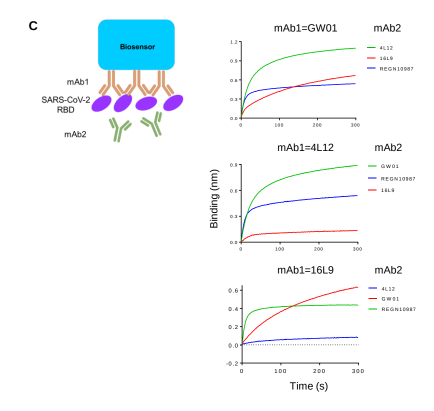
323 COMPETING INTERESTS

324 Patents about the bispecific antibodies in this study are pending.



Antibody concentration (µg/mL)

Ab / Virus ID	IC50 (ng/ml)								% of	GM	Mediar	
	WT	Alpha	Beta	Gamma	Delta	Omicron	SARS-CoV	WIVI	RS3367	Neut	IC50	IC50
GW01	23.1	15.4	51.0	89.1	50.2	>10,000	56.8	2.3	1.4	89	18.7	36.6
16L9	4.1	9.3	412	5.4	7.0	>10,000	>10,000	>10,000	>10,000	56	14.2	7.0
4L12	4.5	6.4	12.0	1.3	19.2	>10,000	>10,000	>10,000	>10,000	56	6.1	6.4
S309	30.8	86.6	41.1	60.0	63.5	192	24.1	113	295	100	73.7	63.5
REGN10987	2.3	1.7	5.7	38.5	32.9	>10,000	>10,000	>10,000	>10,000	56	7.8	5.7
CC12.1	17.3	301	>10,000	24.5	20.3	>10,000	>10,000	>10,000	>10,000	44	40.1	22.4
REGN10989	5.7	0.4	>10,000	>10,000	2.9	>10,000	>10,000	>10,000	>10,000	33	1.8	2.9
4A8	196	2269	>10,000	>10,000	1224	>10,000	>10,000	>10,000	>10,000	33	817	1224



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Figure 1. Isolation of three non-Omicron neutralizing antibodies from COVID-19 convalescent individuals.

(A) Binding of GW01, 16L9, and 4L12 to the SARS-CoV-2 RBD, Omicron RBD
and trimer in an ELISA. REGN10987 was used as a control. (B) Neutralizing
activities of GW01, 16L9, and 4L12 were determined against pseudotyped
SARS-CoV-2 and its variants Alpha, Beta, Gamma, Delta, and Omicron, as well
as sarbecoviruses. REGN10987 was used as a control. (C) Binding of 4L12,
16L9, and RGN10987 to the SARS-CoV-2 RBD in competition with GW01, as
measured by bilayer interferometry experiments.

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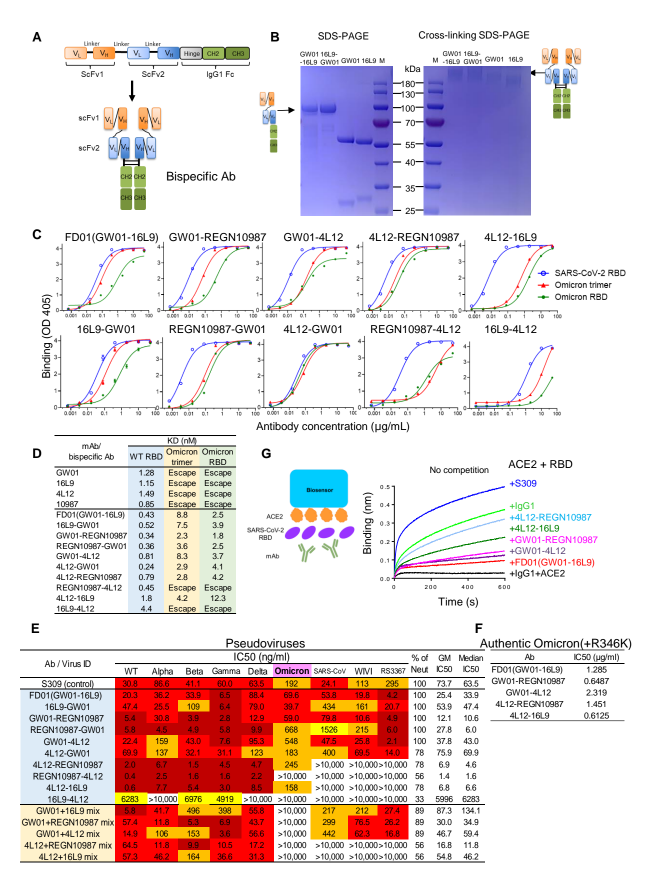
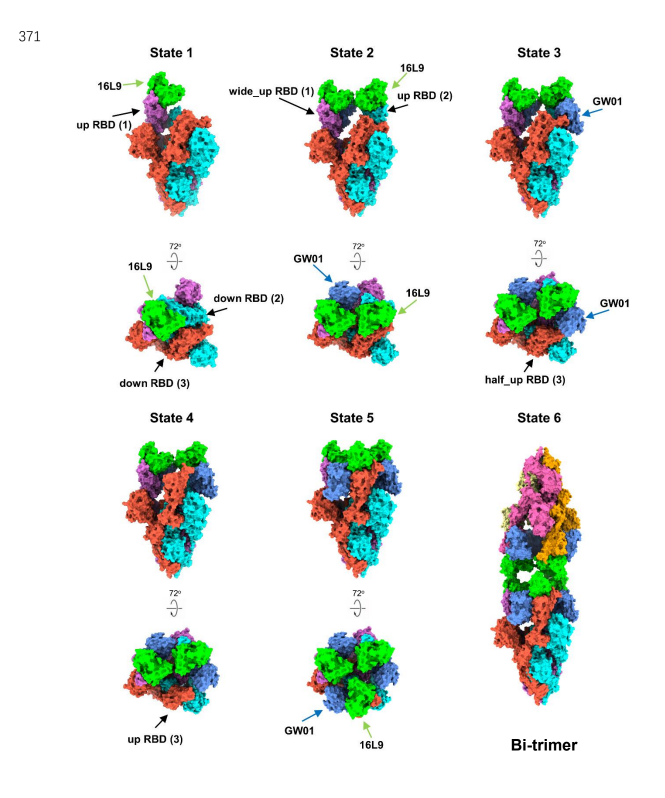


Figure 2. Binding and neutralization of the Omicron variant by bispecific antibodies.

(A) Schematic diagrams showing the structures of bispecific antibodies. (B) SDS-PAGE and cross-linking SDS-PAGE gels showing the sizes of the representative bispecific antibodies and their parental antibodies. (C) Binding specificities of the bispecific antibodies to the SARS-CoV-2 RBD-his, Omicron trimer-his, or RBD-his protein. (D) Binding affinities of the bispecific antibodies to the SARS-CoV-2 RBD-his, Omicron trimer-his, or RBD-his protein were measured by bilayer interferometry experiments. (E) Neutralization by bispecific antibodies and combinations of parental antibodies against the VOCs, including Omicron variant, and sarbecoviruses. (F) Neutralization of five representative bispecific antibodies against authentic Omicron(+R346K) variants. (G) Five representative bispecific antibodies bind to the ACE2-binding site and block RBD binding to ACE2. Binding of ACE2 to the SARS-CoV-2 RBD in competition with bispecific antibodies (red), S309 (blue), control IgG1 (green), and IgG1+ACE2 (black).



372 Figure 3. Cryo-EM structures of the Omicron S trimer in complex with

373 the bispecific antibody FD01.

- 374 The bispecific antibody FD01 binds to Omicron S trimers in six states. Two
- perpendicular views of Omicron S-FD01 are shown in surface representation,
- with 16L9 ScFv in lime and GW01 ScFv in cornflower blue.
- 377

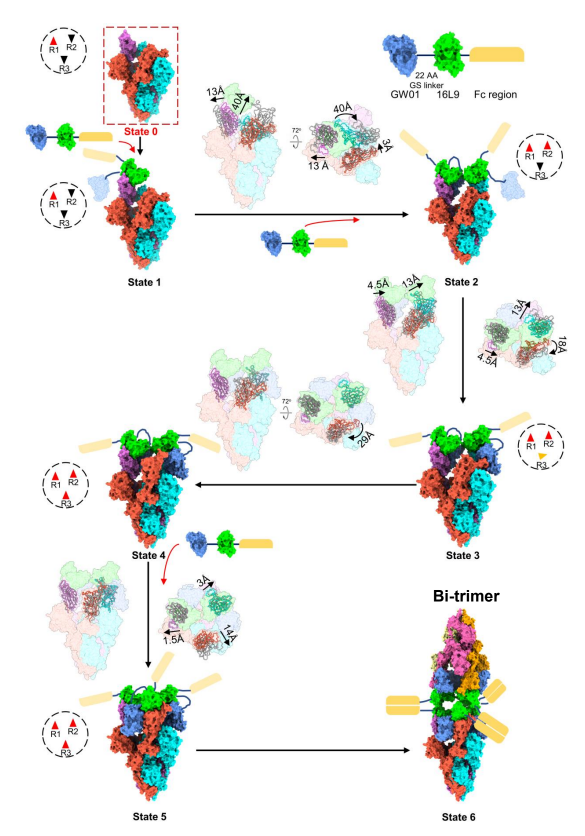


Figure 4. Conformation transitions of Omicron S-FD01 in all states.

- 380 State 0 inside the red dashed box is a hypothetical apo state structure. Small
- 381 triangles inside the black dashed circle indicate the up/half_up/down states of
- 382 three RBDs in the trimer.

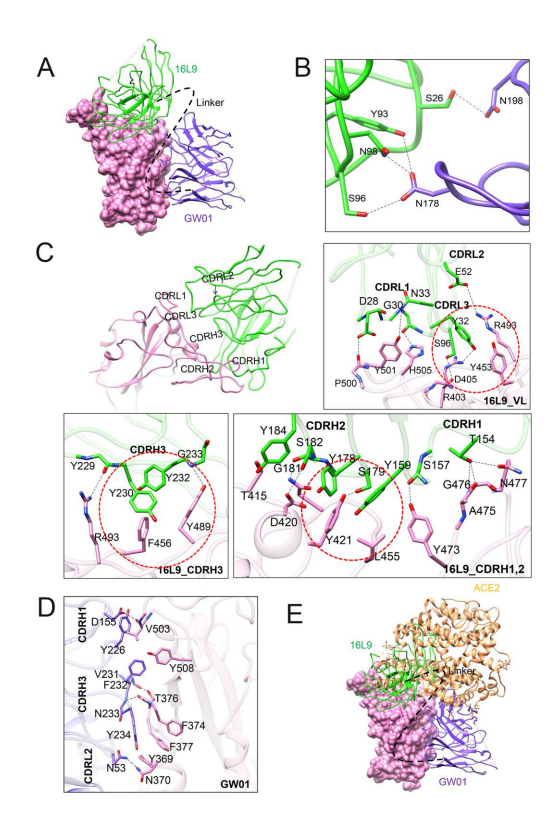


Figure 5. Two conserved epitopes recognized by FD01. (A) Close-up view 385 of the interaction between FD01 and Omicron; the Omicron RBD is displayed 386 in pink in surface representation. 16L9 and GW01 are shown as cartoons 387 colored green and medium-blue, respectively. (B) The interface between 16L9 388 and GW01. (C-D) The interaction of 16L9 (C) and GW01 (D). The residues 389 involved in interactions are represented as sticks. Polar interactions are 390 indicated as dotted lines. (E) Ribbon diagrams of FD01 and ACE2 (PDBID: 391 392 7T9L) bound to the Omicron RBD.

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Table 1. Neutralization by GW01, 16L9, 4L12, REGN10987, and six
bispecific antibodies against 34 strains containing single mutations
present within the Omicron variant.

						IC	50 fold	change	vs. WT			
Sn	oike	Virus ID	GW01	16L9	4L12	REGN	FD01	16L9-	GW01-	GW01-	4L12-	4L12-
						10987		GW01	REGN10987	4L12	REGN10987	16L9
		WT	1.0	1.0	1.0	1.0	1.0	1.0	1.0	1.0	1.0	1.0
		A67V	0.7	0.8	4.4	1.1	2.8	2.1	0.8	3.2	3.3	31.3
		Δ69-70	0.6	1.3	2.5	1.9	2.3	2.6	4.9	3.9	4.5	2.8
		T95I	1.4	1.9	2.5	1.1	1.3	2.0	1.9	2.4	3.3	0.8
		G142D	3.2	4.8	0.7	3.4	2.5	1.2	21.1	1.6	0.8	5.0
N	TD	Δ143-145	0.1	2.4	0.9	0.7	1.3	0.2	0.2	1.3	0.6	1.6
		Δ211	0.5	0.8	9.5	5.2	3.8	0.5	3.3	0.8	0.4	1.4
		L212I	0.6	1.1	1.3	2.1	3.7	2.1	1.5	7.0	5.3	5.0
		ins214EPE	6.4	3.2	6.7	0.9	3.9	1.1	1.9	3.1	5.3	0.3
		G339D	1.0	0.7	4.1	0.5	1.6	0.7	1.1	0.8	32.8	3.1
		S371I	80.0	1.3	116.5	74.1	12.4	14.4	2.6	1.2	25.4	15.6
		S373P	4.8	6.4	2.8	5.9	0.9	0.1	6.0	10.7	0.3	2.8
		S375F	16.0	3.2	n.t.	>1000	1.2	n.t.	n.t.	n.t.	n.t.	n.t.
		K417N	3.9	>1000	0.3	1.7	1.7	1.6	4.3	3.8	5.8	4.4
		N440K	2.3	7.3	3.5	>1000	3.8	1.1	1.1	2.6	0.9	1.0
		G446S	1.1	1.1	9.1	>1000	1.8	1.6	0.5	2.5	0.9	0.8
RBD		S477N	4.1	1.4	16.0	0.9	2.2	2.1	3.7	8.9	3.1	4.1
		T478K	1.5	2.3	4.5	1.0	2.2	2.7	2.8	5.5	2.4	1.6
	RBM	E484A	0.5	3.0	2.5	1.4	4.2	1.3	1.4	3.9	1.0	0.9
		Q493R	1.0	0.6	8.1	0.9	1.9	1.9	4.6	8.3	0.7	1.1
		G496S	1.6	3.5	9.4	4.7	0.8	2.7	2.4	5.8	2.7	1.4
		Q498R	0.4	0.8	4.8	0.9	2.5	0.5	2.2	0.6	0.4	1.4
		N501Y	0.6	2.5	20.3	3.2	2.9	2.0	3.3	6.9	1.0	2.1
		Y505H	0.2	1.0	19.3	0.2	7.9	2.9	6.4	1.7	0.7	0.9
S	D1	T547K	0.6	6.4	4.8	0.2	2.5	0.8	0.6	2.1	0.9	0.4
		D614G	2.4	1.8	4.6	1.5	2.3	1.1	2.3	3.6	3.4	1.2
S	D2	H655Y	4.0	2.1	6.2	0.6	0.7	3.1	4.6	9.1	5.6	2.7
		N679K	3.2	1.9	8.6	0.8	2.5	6.8	6.6	14.0	3.5	4.3
		P681H	4.8	3.5	3.8	1.4	3.7	0.9	0.5	0.6	0.7	1.0
.		N764K	1.5	1.3	7.2	1.1	2.0	5.0	11.4	10.3	4.0	3.3
	=P	D796Y	6.4	0.7	8.2	0.5	1.9	5.0	2.2	8.4	4.0	3.3
		N856K	9.5	1.9	8.1	0.7	0.7	1.3	1.5	4.9	1.6	1.5
		Q954H	0.7	0.8	3.0	1.1	2.8	2.2	2.7	4.5	1.5	1.2
H	R1	N969K	9.5	8.7	n.t.	5.9	2.4	n.t.	n.t.	n.t.	n.t.	n.t.
		L981F	0.5	1.7	0.7	0.2	2.5	0.2	1.6	0.4	1.7	5.3

Fold change is defined as the IC50 of the mutant/the IC50 of the WT. Mutants that resulted in fold change values between 10-50 are highlighted in yellow, and those with values >50 are highlighted in red. The S375F and N969K pseudoviruses were not available to some of the antibodies and are labeled as not tested "n.t.".

405

406

408 Materials and Methods

409 Cell lines, proteins, viruses and plasmids

The human primary embryonic kidney cell lines (HEK293T) and 293T-hACE2 410 cells were cultured in DMEM medium with 10% fetal bovine serum (FBS). RBD-411 his proteins of SARS-CoV-2 and Omicron variant were purchased from Sino 412 Biological. Genes of bispecific antibodies were synthesized by Genscript. The 413 authentic Omicron (B.1.1.529) with R346K mutation used in this study were 414 isolated from COVID-19 patients in Guangzhou, passaged, and titered on Vero 415 E6 cells. African green monkey kidney-derived Vero E6 cells were grown in 416 Dulbecco's modified Eagle's medium (DMEM; Gibco, Grand Island, NY, USA) 417 supplemented with 10% fetal bovine serum (FBS). All work with authentic 418 SARS-CoV-2 was conducted at the Guangzhou Customs Technology Center 419 Biosafety Level 3 (BSL-3) Laboratory. 420

421 **Production of Pseudoviruses**

S genes of SARS-CoV-2 (NC 045512), Alpha (containing 69-70 and 144 422 deletions and N501Y, A570D, D614G, P681H, T716I, S982A, and D1118H 423 substitutions), Beta (containing D80A, D215G, 241-243 deletions and K417N, 424 E484K, N501Y, D614G and A701V substitutions), Gamma (containing 425 L18F,T20N,P26S,D138Y,R190S,K417T,E484K,N501Y,D614G.H655Y,T1027I, 426 and V1176F substitutions), Delta (containing T19R, 157-158 deletions and 427 L452R, T478K, D614G, P681R and D950N substitutions), and Omicron 428 (containing A67V, 69-70del, T95I, G142D, 143-145del, N211I, 212del, 429 ins215EPE, G339D, S371L, S373P, S375F, K417N, N440K, G446S, S477N, 430 T478K, E484A, Q493R, G496S, Q498R, N501Y, Y505H, T547K, D614G, 431 H655Y, N679K, P681H, N764K, D796Y, N856K, Q954H, N969K, L981F 432 substitutions), SARS-CoV, bat SARSr-CoVs (WIV1 and Rs3367) were 433 synthesized by BGI and constructed in pcDNA3.1 vector. Pseudoviruses were 434 generated by co-transfection of 293T cells with an env-deficient HIV backbone 435 pNL4-3.Luc.R-E- backbone and a spike expressing vector^{16,17}. Fifty additional 436

437 spike variants carrying currently circulating single-point mutations were
 438 constructed by site-directed mutagenesis.

439 **Neutralization assay**

The neutralization activities of mAbs and bispecific antibodies were determined 440 using a single-round pseudovirus infection of 293T-hACE2 cells. 10 µl of 5-fold 441 serially diluted antibody was incubated with 40 µl of pseudovirus in 96-well plate 442 at 37 °C for 1 h. 10⁴ 293T-hACE2 cells were then added to the mixture and 443 cultured for 48 h at 37 °C. Cells were lysed and firefly luciferase activity were 444 developed with a luciferase assay system (Promega) and detected on a 445 luminometer (Perkin Elmer). The IC50s of NAbs 446 were 447 calculated using the GraphPad Prism 7.04 software (La Jolla, CA, USA).

448 Memory B-cell staining, sorting and antibody cloning

CD19+IqA-IqD-IqM- primary B cells were sorted out from peripheral blood 449 mononuclear cells (PBMC) of recovered patients of COVID-19 and expanded 450 in vitro in MEM medium with 10% FBS in the presence of irradiated 3T3-451 msCD40L feeder cells, IL-2 and IL-21 as previously described ¹⁸. After fifteen 452 days of incubation, supernatants were screened for neutralization against 453 SARS-CoV-2. From the wells with SARS-CoV-2 neutralization activities, the 454 variable regions of the antibody (VH and VL) genes were amplified by RT–PCR. 455 mAbs were expressed as human IgG1 by HEK293F cells and purified using a 456 457 protein G column (Smart-Lifesciences).

458 **ELISA**

2 μg/ml of SARS-CoV-2 RBD-his, Omicron trimer-his, and Omicron RBD-his protein were coated overnight at 4 °C in a 96-well plate (MaxiSorp Nuncimmuno, Thermo Scientific, USA). Wells were blocked with 5% non-fat milk (Biofroxx, Germany) in PBS for 1 hour at room temperature, followed by incubation with 5-fold serially diluted mAb in disruption buffer (PBS, 5% FBS, 2% BSA, and 1% Tween-20) for 1 hour at room temperature. After 3 washing steps with PBS 0.05% Tween 20 (PBS-T), 1:2500 diluted HRP-conjugated goat
anti-human IgG antibody (Jackson Immuno Research Laboratories, USA) was
added for 1 hour at room temperature. Plates were washed three times with
0.2% Tween-20 in PBS and developed using ABST (Thermo Scientific, USA)
for 30 minutes. Absorbance at 405 nm was read on a Multiskan FC plate reader
(Thermo Scientific, USA).

471 Biolayer interferometry (BLI) binding assay and competition assay

Experiments were carried out on a FortéBio OctetRED96 instrument. The 472 kinetics of monoclonal antibody binding to SARS-CoV-2 RBD-his, Omicron 473 RBD-his, or trimer-his proteins was measured using anti-human IgG (AHC) 474 biosensors. 10 µg/ml of mAbs were immobilized on biosensors for 200s. After 475 a 120 sec stabilization step with 0.02% PBST (PBS with 0.02% Tween), 476 biosensors were moved into 6 µg/ml of RBD-his or trimer-his proteins for the 477 300 sec association step. Then biosensors were moved into 0.02% PBST to 478 479 detect dissociation for 300 sec. The buffer control binding was subtracted to deduct nonspecific binding. Kon, Koff, and KD were calculated by FortéBio Data 480 Analysis software (Version 8.1) using 1:1 binding and a global fitting model. 481

For the Ab competition assay, 10 μg/ml of mAbs 1 were immobilized on the antihuman IgG (AHC) biosensors for 200s. After wash with 0.02% PBST for 120s
to reach baseline, biosensors were moved into 50 μg/ml of IgG1 isotype control
for 200s and then moved into SARS-CoV-2 RBD at 6 μg/ml for 300s. After wash
with 0.02% PBST for 120s, biosensors were moved into 10 μg/ml of mAb2 for
600s to detect the association between mAb2 and SARS-CoV-2 RBD.

For ACE2 competition, biosensors were moved into 20 µg/ml of ACE2-Fc for 600s. After baseline, wash, and blocking steps, biosensors were moved into pre-mix of 600 nM of mAb and 100 nM SARS-CoV-2 RBD for 600s. A mixture of ACE2-Fc and SARS-CoV-2 RBD was used as a positive control, while the 492 mixture of IgG1 isotype control and SARS-CoV-2 RBD was used as a negative493 control.

494 Focus reduction neutralization test

SARS-CoV-2 Omicron(+R346K) focus reduction neutralization test (FRNT) was 495 performed in a certified Biosafety level 3 lab. Fifty microliters antibody were 496 serially diluted, mixed with 50 µl of SARS-CoV-2 (100 focus forming unit, FFU) 497 in 96-well microwell plates and incubated for 1 hour at 37°C. Mixtures were then 498 transferred to 96-well plates seeded with Vero E6 cells (ATCC, Manassas, VA) 499 for 1 hour at 37°C to allow virus entry. Inoculums were then removed before 500 media (100)MEM 501 adding the overlay μl containing 1.2% 502 Carboxymethylcellulose, CMC). After 24-hour post infection, the overlay was discarded and the cell monolayer was fixed with 4% paraformaldehyde solution 503 for 2 hours at RT. After permeabilized with 0.2% Triton X-100 for 20 min at room 504 temperature, the plates were sequentially stained with cross-reactive rabbit 505 506 anti-SARS-CoV-2 N IgG (Cat. No.: 40143-T62, Sino Biological Inc) as the primary antibody and HRP-conjugated goat anti-rabbit IgG(H+L) (No.: 109-035-507 088, Jackson ImmunoResearch) as the secondary antibody in 37°C for 1 hour 508 respectively. The reactions were developed with KPL TrueBlue Peroxidase 509 substrates. The numbers of SARS-CoV-2 foci were calculated using CTL 510 ImmunoSpot S6 Ultra reader (Cellular Technology Ltd). Neutralizing activity 511 512 was defined as the ratio of inhibition of SARS-CoV-2 focus comparing diluted antibody to control. 513

514 **Construction and expression of bispecific NAbs**

515 Genes of a bispecific Ab consisting of the scFv of GW01 and scFv of 16L9, 516 REGN10987 or 4L12 were synthesized and codon-optimized by GenScript. The 517 bispecific antibody sequence alignment was as follows: variable light chain (VL) 518 and variable heavy chain (VH) of mAb 1 or mAb 2 were linked with a (Gly₄Ser)₃ 519 linker. VL-VH of mAb 1 and VL-VH of mAb 2 were linked with a (Gly₄Ser)₄ linker 520 and then fused to the expression vector with hinge-CH2-CH3 fragment of human immunoglobulin (hlgG1 Fc). FD01 bispecific antibody sequence order
was as follows: GW01 VL-(Gly4Ser)3-GW01 VH-(Gly4Ser)4-16L9 VL-(Gly4Ser)316L9 VH-hinge-CH2-CH3.

293F cells were transiently transfected with bispecific Abs plasmid. After 6 days
of culture at 37°C in a 5% CO2 incubator, supernatant was collected and filtered.
Bispecific antibodies were purified with protein G colume (Smart-Lifesciences)
and stored in PBS at -80°C.

528 SDS-PAGE and cross-linking SDS-PAGE of bispecific antibodies

The purity and molecular weight of bispecific antibodies were then analyzed by SDS-PAGE and cross-linking SDS-PAGE. Briefly, 5 μ g of bispecific antibodies were mixed with 5x SDS-loading sample buffer containing 10% β mercaptoethanol. The samples were heated for 10 min at 100°C and were then loaded on an SDS gradient gel (4–20% Precast Protein Improve Gels, Genscript Biotech Corporation). The gel was run for 120 min at 120 V, and Coomassie staining was performed.

Extent of dimer was investigated by cross-linking of bispecific antibodies with 536 glutaraldehyde (Sigma-Aldrich). Briefly, 5 µg of antibodies were diluted in 25 µl 537 of PBS in the presence of a 2.7 µM of glutaraldehyde cross-linker. The mixture 538 was incubated at RT for 5 minutes, and then glutaraldehyde was guenched by 539 adding 1 M Tris-HCl buffer (pH 8.0) to a final concentration of 40 mM. After 540 mixing with 5x SDS-loading sample, the protein samples were loaded on a 4-541 20% SDS gradient gel. The gel was run for 180 min at 120 V and confirmed by 542 Coomassie staining. 543

544 Expression and purification of SARS-CoV-2 Omicron Spike

The Human codon gene encoding SARS-CoV-2 Omicron S ectodomain was purchased from GeneScript. The expression plasmid of Omicron S 6P substitution¹⁹ was constructed and transfected into suspension HEK293F using polyethlenimine. After 72 hours, the supernatants were harvested and filtered

for affinity purification by Histrap HP (GE). The protein was then further purified
by gel filtration using Superose 6 increase 10/300 column (GE Healthcare) in
20 mM Tris pH 8.0, 200 mM NaCl.

552 Cryo-EM sample preparation

553 Purified SARS-CoV-2 Omicron S at 1.554 mg/mL was mixed with FD01 554 antibody by a molar ratio of 1:1.5 incubated for 10 min on ice before application 555 onto a freshly glow-discharged holey amorphous nickel-titanium alloy film 556 supported by 400 mesh gold grids ²⁰. The sample was plunged freezing in liquid 557 ethane using Vitrobot IV (FEI/Thermo Fisher Scientific), with 2 s blot time and -558 3 blot force and 10 s wait time.

559 Cryo-EM data collection and image processing

560 Cryo-EM data were collected on a Titan Krios microscope (Thermo Fisher) 561 operated at 300 kV, equipped with a K3 summit direct detector (Gatan) and a 562 GIF quantum energy filter (Gatan) setting to a slit width of 20 eV. Automated 563 data acquisition was carried out with SerialEM software²¹ through beam-image 564 shift method²².

565 Movies were taken in the super-resolution mode at a nominal magnification 566 81,000×, corresponding to a physical pixel size of 1.064 Å, and a defocus range 567 from $-1.2 \mu m$ to $-2.5 \mu m$. Each movie stack was dose-fractionated to 40 frames 568 with a total exposure dose of about 58 e⁻/Å² and exposure time of 3s.

All the data processing was carried out using either modules on, or through, RELION v3.0²³ and cryoSPARC²⁴. A total of 4,363 movie stacks was binned 2 × 2, dose weighted, and motion corrected using MotionCor2 ²⁵ within RELION. Parameters of contrast transfer function (CTF) were estimated by using Gctf ²⁶. All micrographs then were manually selected for further particle picking upon ice condition, defocus range and estimated resolution.

575 Remaining 3,817 good images were imported into cryoSPARC for further 576 patched CTF-estimating, blob-picking and 2D classification. From 2D 577 classification, bi-trimer and trimer particles were observed. Several good 2D 578 classes of these two kind particles were used as templates for template-picking separately. After 2D classification of particles from template-picking was
finished, all good particles from blob-picking and template-picking were merged
and deduplicated, subsequently being exported back to RELION through pyem
package ²⁷.

583 For bi-trimer map, 1,003,956 particles were extracted at a box-size of 540 and 584 rescaled to 180, then carried on 2 round of 3D classification with a soft circular 585 mask of 480 Å in diameter in RELION. Only good classes were selected, 586 yielding 166,441 clean particles. These particles were re-extracted unbinned 587 (1.064 Å/pixel) and auto-refined without applying symmetry, yielding a map at 588 6.11 Å.

For trimer map, 1,003,956 particles were extracted at a box-size of 320 and 589 rescaled to 160, then carried on 1 round of 3D classification with a soft circular 590 mask of 220 Å in diameter in RELION. Three classes with different conformation 591 change on trimer RBDs were selected separately for another round of 3D 592 classification. Particles in different states were auto-refined, CTF-refined and 593 594 polished separately. Some density of RBDs and Fabs in some state were not well-resolved, so we carried out no-alignment 3D classification with NTD-RBD-595 Fabs (NRF) mask to improve those regions. Finally, we got 5 states of Omicron 596 S-FD01 trimer. 597

To get clear interfaces of RBD with Fabs, we did local-refinement focused on 598 that region. We first selected all good 3D-classes with relatively complete RBD 599 and FD01 density within all states. We auto-refined these particles with a C3-600 aligned reference, but the auto-refinement procedure was not applied any 601 symmetry. Then particles were expanded with C3 symmetry and further 602 subtracted with one NRF mask. After no-alignment 3D-classification, 249,122 603 particles with complete NRF density were selected out, exported to cryoSPARC 604 and carried out local refinement, yielding a local-refined map at 3.51 Å. 605

The reported resolutions above are based on the gold-standard Fourier shell correlation (FSC) 0.143 criterion. All the visualization and evaluation of 3D density maps were performed with UCSF Chimera ²⁸ and ChimeraX²⁹. The above procedures of data processing are summarized in Fig. S3 and Fig. S4.

⁶¹⁰ These sharpened maps were generated by DeepEMhancer ³⁰ and then "vop

⁶¹¹ zflip" to get the correct handedness in UCSF Chimera for subsequent model

612 building and analysis.

613 Model building and refinement

For model building of SARS-CoV-2 Omicron S FD01 complex, the SARS-CoV-614 2 Omicron S trimer model and the antibody model generated by swiss-model ³¹ 615 were fitted into the map using UCSF Chimera and then manually adjusted with 616 COOT ³². Several iterative rounds of real-space refinement were further carried 617 out in PHENIX ³³. The RBD bounded with 16L9 and GW01 was refined against 618 the local refinement map and then docked back into global refinement trimer 619 and bi-trimer maps. Model validation was performed using phenix.MolProbity. 620 Figures were prepared using UCSF Chimera and UCSF ChimeraX²⁹. 621

Data and materials availability: The cryo-EM map and the coordinates of 622 SARS-CoV-2 Omicron S complexed with FD01 have been deposited to the 623 Electron Microscopy Data Bank (EMDB) and Protein Data Bank (PDB) with 624 accession numbers EMD-32655 and PDB 7WOQ (state 1), EMD-32656 and 625 PDB 7WOR (state 2), EMD-32657 and PDB 7WOS (state 3), EMD-32659 and 626 PDB 7WOU (state 4), EMD-32660 and PDB 7WOV (state 5), EMD-32661 and 627 PDB 7WOW (state 6), EMD-32654 and PDB 7WOP (NTD-RBD-GW01-16L9 628 629 local refinement).

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631 **REFERENCES**

632 Cameroni, E. et al. Broadly neutralizing antibodies overcome SARS-CoV-2 Omicron 1 antigenic shift. Nature, doi:10.1038/s41586-021-04386-2 (2021). 633 634 2 Cele, S. et al. Omicron extensively but incompletely escapes Pfizer BNT162b2 635 neutralization. Nature, doi:10.1038/s41586-021-04387-1 (2021). 636 Liu, L. et al. Striking Antibody Evasion Manifested by the Omicron Variant of SARS-CoV-3 637 2. Nature, doi:10.1038/s41586-021-04388-0 (2021). 638 Planas, D. et al. Considerable escape of SARS-CoV-2 Omicron to antibody neutralization. 4

639 Nature, doi:10.1038/s41586-021-04389-z (2021). 640 5 Rössler, A. et al. SARS-CoV-2 Omicron Variant Neutralization in Serum from Vaccinated 641 and Convalescent Persons. doi:10.1056/NEJMc2119236 (2022). 642 6 Carreño, J. M. et al. Activity of convalescent and vaccine serum against SARS-CoV-2 643 Omicron. Nature, doi:10.1038/s41586-022-04399-5 (2021). 644 7 Garcia-Beltran, W. F. et al. mRNA-based COVID-19 vaccine boosters induce neutralizing 645 immunity against SARS-CoV-2 Omicron variant. Cell, doi:10.1016/j.cell.2021.12.033 646 (2022). 647 Deinirattisai, W. et al. Reduced neutralisation of SARS-CoV-2 omicron B.1.1.529 variant 8 648 by post-immunisation serum. Lancet 399, 234-236, doi:10.1016/s0140-6736(21)02844-0 649 (2022). 650 9 Wang, Y. et al. Resistance of SARS-CoV-2 Omicron Variant to Convalescent and 1-8, 651 CoronaVac Vaccine Plasma. Emerg Microbes Infect, 652 doi:10.1080/22221751.2022.2027219 (2022). 653 Wang, Y. et al. The significant immune escape of pseudotyped SARS-CoV-2 variant 10 Omicron. Emerg Microbes Infect 11, 1-5, doi:10.1080/22221751.2021.2017757 (2022). 654 655 11 Dejnirattisai, W. et al. Omicron-B.1.1.529 leads to widespread escape from neutralizing 656 antibody responses. bioRxiv, doi:10.1101/2021.12.03.471045 (2021). 657 Zhou, T. et al. Structural basis for potent antibody neutralization of SARS-CoV-2 variants 12 658 including B.1.1.529. 2021.2012.2027.474307, doi:10.1101/2021.12.27.474307 %J bioRxiv 659 (2021). 660 VanBlargan, L. A. et al. An infectious SARS-CoV-2 B.1.1.529 Omicron virus escapes 13 661 neutralization by therapeutic monoclonal antibodies. Nature Medicine. 662 doi:10.1038/s41591-021-01678-y (2022). 663 Mannar, D. et al. Structural analysis of receptor binding domain mutations in SARS-CoV-14 664 2 variants of concern that modulate ACE2 and antibody binding. Cell Rep 37, 110156, 665 doi:10.1016/j.celrep.2021.110156 (2021). 666 Cao, Y. et al. Omicron escapes the majority of existing SARS-CoV-2 neutralizing 15 667 antibodies. Nature, doi:10.1038/s41586-021-04385-3 (2021). 668 16 Ju, B. et al. Human neutralizing antibodies elicited by SARS-CoV-2 infection. Nature 584, 669 115-119, doi:10.1038/s41586-020-2380-z (2020). 670 Wu, F. et al. Evaluating the Association of Clinical Characteristics With Neutralizing 17 671 Antibody Levels in Patients Who Have Recovered From Mild COVID-19 in Shanghai, China. 672 JAMA internal medicine 180, 1356-1362, doi:10.1001/jamainternmed.2020.4616 (2020). 673 Huang, J. et al. Isolation of human monoclonal antibodies from peripheral blood B cells. 18 Nat Protoc 8, 1907-1915, doi:10.1038/nprot.2013.117 (2013). 674 675 19 Hsieh, C. L. et al. Structure-based design of prefusion-stabilized SARS-CoV-2 spikes. 676 Science 369, 1501-1505, doi:10.1126/science.abd0826 (2020). 677 Huang, X. et al. Amorphous nickel titanium alloy film: A new choice for cryo electron 20 678 microscopy sample preparation. Prog **Biophys** Mol Biol 156, 3-13. 679 doi:10.1016/j.pbiomolbio.2020.07.009 (2020). 680 21 Mastronarde, D. N. Automated electron microscope tomography using robust prediction 681 of specimen movements. J Struct Biol 152, 36-51, doi:10.1016/j.jsb.2005.07.007 (2005). 682 22 Wu, C. et al. High-quality, high-throughput cryo-electron microscopy data collection via

683		beam tilt and astigmatism-free beam-image shift. Journal of structural biology 208,
684		107396, doi:10.1016/j.jsb.2019.09.013 (2019).
685	23	Zivanov, J. et al. New tools for automated high-resolution cryo-EM structure
686		determination in RELION-3. <i>Elife</i> 7, doi:10.7554/eLife.42166 (2018).
687	24	Punjani, A. et al. M. A. cryoSPARC: algorithms for rapid unsupervised cryo-EM structure
688		determination. <i>Nature methods</i> 14, 290-296, doi:10.1038/nmeth.4169 (2017).
689	25	Zheng, S. Q. et al. MotionCor2: anisotropic correction of beam-induced motion for
690		improved cryo-electron microscopy. Nat Methods 14, 331-332, doi:10.1038/nmeth.4193
691		(2017).
692	26	Zhang, K. Gctf: Real-time CTF determination and correction. J Struct Biol 193, 1-12,
693		doi:10.1016/j.jsb.2015.11.003 (2016).
694	27	Asarnow, D., Palovcak, E., Cheng, Y. UCSF pyem v0.5. Zenodo
695		https://doi.org/10.5281/zenodo.3576630 (2019).
696		doi: <u>https://doi.org/10.5281/zenodo.3576630</u> (2019).
697	28	Pettersen, E. F. et al. UCSF Chimera a visualization system for exploratory research and
698		analysis. <i>J Comput Chem</i> 25 , 1605-1612, doi:10.1002/jcc.20084 (2004).
699	29	Pettersen, E. F. et al. UCSF ChimeraX: Structure visualization for researchers, educators,
700		and developers. <i>Protein Sci</i> 30 , 70-82, doi:10.1002/pro.3943 (2021).
701	30	Sanchez-Garcia, R. et al. DeepEMhancer: a deep learning solution for cryo-EM volume
702		post-processing. <i>Commun Biol</i> 4 , 874, doi:10.1038/s42003-021-02399-1 (2021).
703	31	Waterhouse, A. et al. SWISS-MODEL: homology modelling of protein structures and
704		complexes. <i>Nucleic Acids Res</i> 46 , W296-W303, doi:10.1093/nar/gky427 (2018).
705	32	Emsley, P., Lohkamp, B., Scott, W. G. & Cowtan, K. Features and development of Coot.
706		Acta Crystallogr D Biol Crystallogr 66, 486-501, doi:10.1107/S0907444910007493 (2010).
707	33	Afonine, P. V. et al. Real-space refinement in PHENIX for cryo-EM and crystallography.
708		Acta Crystallogr D Struct Biol 74 , 531-544, doi:10.1107/S2059798318006551 (2018).

709	Table S1. The germline and CDRH3 sequences of GW01, 4L12, and 16L9.

Donor ID mAb ID	VH	CDRH3 sequence	VL	CDRL3 sequence
Donor 1 GW01	IGHV3-43	AKDRSYGPPDVFNYEYGMDV	IGLV1-44	AAWDDSLNWV
Donor 1 4L12	IGHV3-66	ARDLITYGMDV	IGKV1-9	QQLNSYPPLT
Donor 2 16L9	IGHV3-53	ARGEIQPYYYYGMDV	IGLV2-8	SSYAGSSNFDV

Table S2. Cryo-EM data collection and refinement statistics.

	State	State	State	State 4	State	State	Local
	1	2	3		5	6	refine NRF
Data collection							
and processing							
Magnification				81,000			
Voltage (kV)				300			
Electron exposure				58			
(e–/Ų)				-1.2 to -2	.5		
Defocus range (µm)				1.064			
Pixel size (Å)				1,003,95	6		
Initial particles (no.)				C1			
Symmetry imposed	194,02	62,04	74,41	141,577	39,29	71,568	249,122
Final particles (no.)	6	0	5		4		
Map resolution (Å)	3.47	3.70	3.91	3.47	3.87	6.11	3.51
Refinement							
R.m.s. deviations	0.003	0.003	0.003	0.003	0.003	0.002	0.002
Bond lengths	0.539	0.506	0.499	0.557	0.527	0.437	0.524
(Å)							
Bond angles (°)							
Validation							
MolProbity	2.52	2.48	2.46	2.50	2.53	2.41	2.88
score	9.18	8.06	8.38	8.44	8.14	7.67	10.97
Clashscore	5.14	5.34	5.30	5.39	5.97	5.33	9.21
Rotamer outlier							
(%)							
Ramachandran							
plot	91.75	92.00	92.74	91.84	91.67	93.20	88.62
Favored (%)	7.95	7.90	7.14	8.13	8.17	6.80	11.38
Allowed (%)	0.30	0.11	0.13	0.03	0.16	0.00	0.00
Disallowed (%)							
EMDB	32655	3265	3265	32659	3266	32661	32654
		6	7		0		
PDB	7WOQ	7WO	7WO	7WOU	7WO	7WOW	7WOP
		R	S		V		

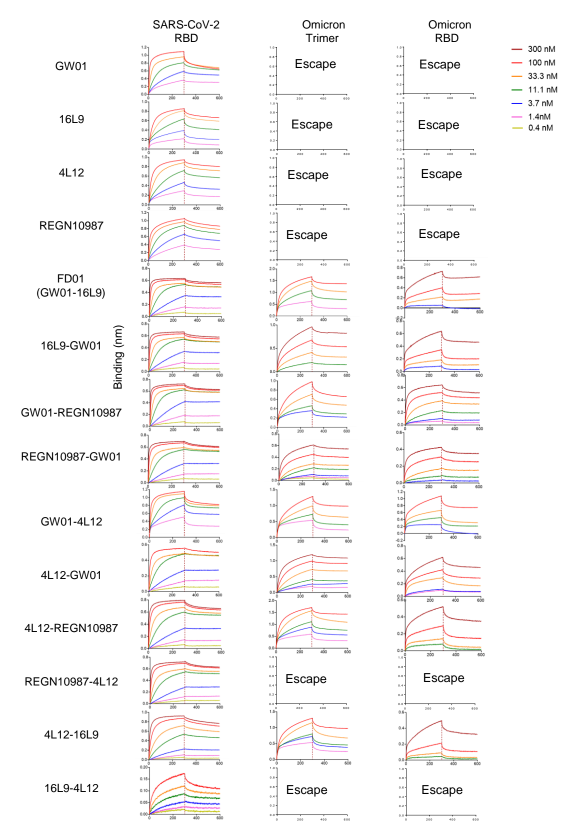
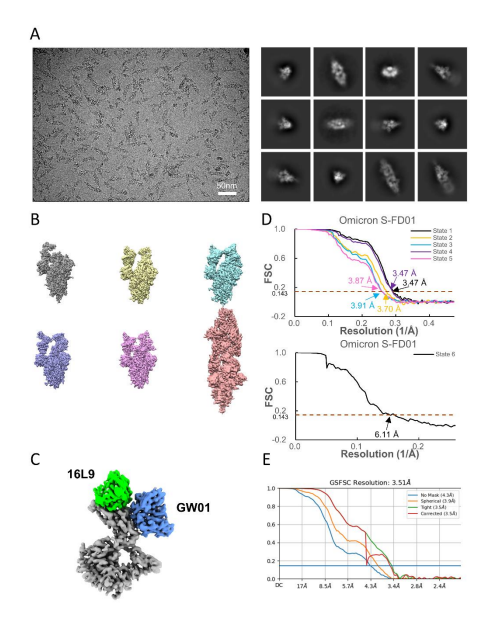
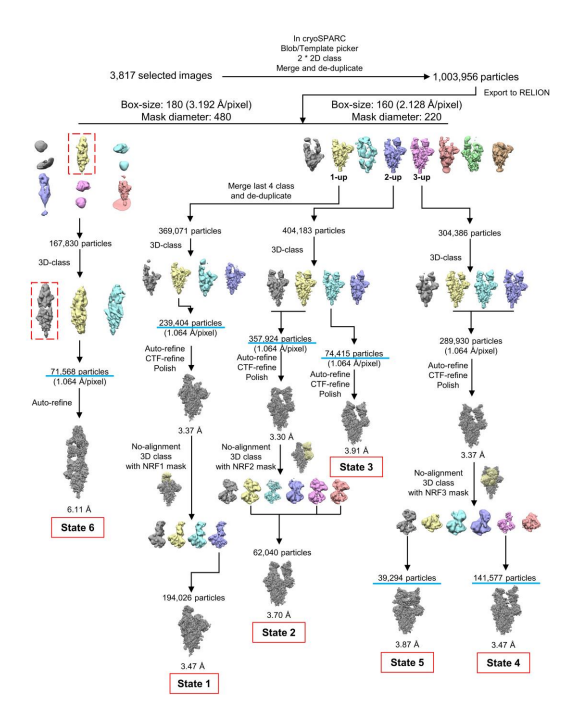


Figure S1. Binding affinities of GW01, 16L9, 4L12, REGN10987, and ten
bispecific antibodies to SARS-CoV-2 RBD-his, Omicron trimer-his and
Omicron RBD-his measured by bilayer interferometry experiments.
Antibodies were immobilized on anti-human IgG (AHC) biosensors and then
tested for their binding abilities to the target proteins.

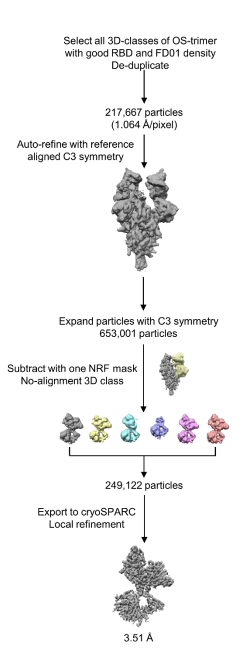


718

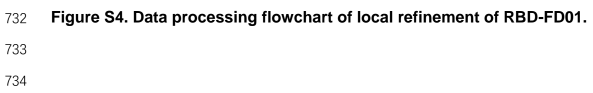
Figure S2. Cryo-EM data collection and processing of FD01 bound SARS-719 CoV-2 Omicron S. (A) Representative electron micrograph and 2D 720 classification results of FD01 bound SARS-CoV-2 S. (B) The reconstruction 721 map of the complex structures at six states. (C) The local-refined map of the 722 NRF region. (D) Gold-standard Fourier shell correlation curves generated in 723 RELION for structures of six states. The 0.143 cut-off is indicated by a 724 horizontal dashed line. (E) Gold-standard Fourier shell correlation curves 725 generated in cryoSPARC for local-refined map. 726



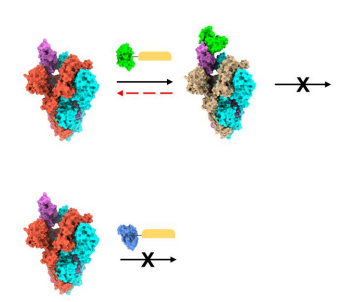
- 727 Figure S3. Data processing flowchart of FD01 bound SARS-CoV-2
- 728 **Omicron S trimer.** Particles number above cyan line is used for particle
- 729 counting statistics.







740



- 741 Figure S5. Hypothesis of binding features when Omicron S trimer meets
- 742 with mAbs of 16L9 or GW01.

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