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Synthetic repertoires derived from convalescent COVID-19 patients enable discovery of SARS-CoV-2 neutralizing antibodies and a novel quaternary binding modality

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41 Abstract

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43 The ongoing evolution of SARS-CoV-2 into more easily transmissible and infectious 44 variants has sparked concern over the continued effectiveness of existing therapeutic antibodies 45 and vaccines. Hence, together with increased genomic surveillance, methods to rapidly develop 46 and assess effective interventions are critically needed. Here we report the discovery of SARS-47 CoV-2 neutralizing antibodies isolated from COVID-19 patients using a high-throughput platform. 48 Antibodies were identified from unpaired donor B-cell and serum repertoires using yeast surface 49 display, proteomics, and public light chain screening. Cryo-EM and functional characterization of the antibodies identified N3-1, an antibody that binds avidly (K_{d,app} = 68 pM) to the receptor binding 50 51 domain (RBD) of the spike protein and robustly neutralizes the virus *in vitro*. This antibody likely 52 binds all three RBDs of the trimeric spike protein with a single IgG. Importantly, N3-1 equivalently 53 binds spike proteins from emerging SARS-CoV-2 variants of concern, neutralizes UK variant 54 B.1.1.7, and binds SARS-CoV spike with nanomolar affinity. Taken together, the strategies 55 described herein will prove broadly applicable in interrogating adaptive immunity and developing 56 rapid response biological countermeasures to emerging pathogens.

57 Introduction

The rapid global dissemination of the severe acute respiratory syndrome coronavirus 2 58 (SARS-CoV-2)¹, the cause of coronavirus disease 19 (COVID-19)², has highlighted our extreme 59 60 vulnerability to novel microbial threats. The speed of SARS-CoV-2 transmission and absence of 61 widespread adaptive immunity created a pandemic that overwhelmed the international medical 62 community. This situation was exacerbated by the scarcity of treatment options, especially early 63 in the pandemic. Functional immune repertoire analysis has the potential to efficiently address 64 this scarcity. By analyzing primary immune responses directed towards emerging pathogens, 65 newly elicited antibodies can be identified and rapidly deployed to treat patients.

The COVID-19 pandemic has stimulated global research efforts to identify SARS-CoV-2neutralizing antibodies for therapeutic and prophylactic applications. Initial attempts to identify neutralizing antibodies focused on screening extant antibodies elicited against previous SARS-CoV strains. These efforts were largely unsuccessful owing to limited cross-reactivity³, primarily due to SARS-CoV-2's sequence divergence in the receptor-binding domain (RBD) of its trimeric spike protein⁴.

72 The ectodomain (ECD) of the SARS-CoV-2 spike (S) protein is essential for initial binding 73 and subsequent entry of the virus into human cells and has been the primary target for 74 therapeutics and vaccine formulations. The ECD consists of the S1 subunit, containing an N-75 terminal domain (NTD) and RBD, and the S2 subunit, containing the fusion machinery that 76 mediates entry into host cells. The RBD initiates attachment through interaction with angiotensin-77 converting enzyme 2 (ACE2)⁵⁻⁷. The functional significance of the ACE2-RBD interaction makes the RBD a prime target for neutralizing antibodies⁸⁻¹⁰. Targeting this domain increases the 78 79 selective pressure on the RBD, which may promote the emergence of escape mutants that

80 maintain virulence. Indeed, neutralizing antibodies targeting a single epitope can induce virus
81 escape in cell culture, quickly rendering antibodies ineffective¹¹.

The FDA approved antibody cocktail of REGN10933 and REGN10987¹² targets distinct 82 83 epitopes of the RBD and sustains some neutralization activity against new SARS-CoV-2 variants 84 B.1.1.7 and B.1.351, originally identified in the UK and South Africa, respectively¹³⁻¹⁵. Recently, 85 a deep mutational scan of the RBD found that the single amino acid mutation E406W increased 86 the IC50 of REGN10933 by more than 300-fold and also detrimentally affected REGN10987 87 binding¹⁶. The scan also identified amino acid changes that compromised the epitope of Eli Lilly's 88 antibody LY-CoV016¹⁶. Moreover, the mass sequencing of 5.085 SARS-CoV-2 genomes from the 89 Houston metropolitan area published in September 2020 already identified numerous spike 90 protein mutations that affect the existing neutralizing antibodies' abilities to bind to their epitopes¹⁷.

91 The B.1.1.7 (UK), B.1.1.248 (Brazil), and B.1.351 (South Africa) variants are of special 92 concern because they have each accumulated multiple spike protein mutations. These mutations 93 are located in the NTD, RBD, and furin cleavage site. NTD-directed antibodies in particular show 94 reduced or abolished binding to these strains¹⁴. More concerning than the loss of binding ability, 95 however, is the reduced neutralizing activity of convalescent plasma from patients infected in the 96 spring of 2020^{14,18}. Furthermore, currently available vaccines are largely based on an earlier 97 prefusion-stabilized spike variant (Wuhan). When assayed, sera from patients vaccinated with the 98 Moderna and Pfizer/BioNTech vaccines showed a significant decrease in neutralization activity 99 towards strain B.1.351, while B.1.1.7 was only mildly affected¹⁴. It is therefore critical to identify 100 neutralizing antibodies to a broad spectrum of non-overlapping epitopes on the spike protein to 101 achieve highly potent and persistent neutralization¹¹.

102 To this end, we developed a multi-pronged antibody discovery and informatics strategy 103 involving both proteomic analysis of donor sera and selection of combinatorially paired heavy and 104 light chain (VH-VL) libraries from donor B-cell receptor repertoires. Specifically, we used Ig-Seq 105 proteomics to identify candidate variable heavy chains (VHs) of serum antibodies binding SARS-106 CoV-2 RBD or ECD. We identified productive light chain pairs for these heavy chains either from a bioinformatically derived set of "public light chains" (PLCs) or through combinatorial pairing with 107 108 donor light chain repertoires via yeast surface display (YSD) selection. In addition to Ig-Seq 109 interrogation of circulating IgGs, we employed YSD for selection of high-affinity antibodies by 110 combinatorial display of all donor VH-VL pairs as Fab fragments through multiple rounds of 111 selection.

112

113 Together, this holistic strategy allowed us to probe the secreted circulating antibody 114 repertoire and the nascent cellular repertoire of a primary immune response. By interrogating the 115 combined repertoire, we gained valuable insight into the antibody response elicited by SARS-116 CoV-2 and discovered neutralizing antibodies to multiple distinct domains of the spike protein. 117 We report the structural characterization of two such neutralizers, including a highly potent 118 neutralizing antibody (N3-1) that binds a quaternary epitope of the trimeric spike protein via a 119 novel binding modality. Finally, we tested binding to emerging spike variants of SARS-CoV-2 and 120 demonstrated N3-1 maintains robust binding to these variants and neutralizes B.1.1.7.

121

122

123 Results

124 Ig-Seq analysis of the serological repertoire reveals candidate heavy chains

125 Ig-Seq provides a proteomic snapshot of the serum antibody repertoire by using mass
 126 spectrometry to identify heavy chain complementarity-determining region 3 (CDR3) peptides of
 127 antigen-enriched antibodies (**Fig. 1**)¹⁹. To identify relevant SARS-CoV-2 antibodies present in the

128 serological repertoire, we first isolated antibodies from the serum of two donors by antigen enrichment chromatography using immobilized SARS-CoV-2 RBD and ECD fragments. We then 129 130 employed Ig-Seq mass spectrometry to identify abundant IgG clonotypes. A total of 15 and 21 131 unique clonotypes were identified with high confidence from donors 1 and 2, respectively 132 (**Extended Data Figs. 1-2**). This is notably lower than what has been observed in previous Ig-133 Seg studies¹⁹⁻²², perhaps because the previous studies involved donors after boost vaccination 134 which is likely to elicit a more robust and diverse response than the primary response to natural 135 SARS-CoV-2 infection observed here. Regardless, the clonotypes identified provided a sufficient 136 set of VH candidates to express and characterize as recombinant anti-SARS-CoV-2 monoclonal 137 antibodies (mAbs) in combination with suitable light chains.

138

139 Public light chains recover productive pairs for most VHs

While Ig-Seq provides valuable information for antibody discovery, it does not identify light chain partners, which usually requires additional laborious techniques. To expedite light chain discovery, we analysed a published dataset of natively paired memory B-cell sequences¹⁹ from three healthy donors and bioinformatically derived a panel of nine light chains that show high frequencies of productive pairings with a diverse set of VH sequences (**Fig. 1c**).

At the time of blood draw, the donors in the dataset were healthy and asymptomatic. This lack of polarization and relative homeostasis potentially affords broader insight into common VH-VL pairings, independent of antigen specificity. Indeed, many of the observed light chains paired promiscuously with a diverse set of VH genes. The V gene usage of undiscerning light chains was remarkably consistent among the three donors and therefore were termed "public light chains" (PLCs). From our secondary analysis of this dataset, we identified six kappa V genes that accounted for 61% of the kappa VH-VL pairs and three lambda V genes representing 22% of

152 lambda pairs (Fig. 2a). Together, these nine public light chain (PLC) V genes accounted for 43%
153 of observed VH-VL pairs in the analyzed repertoires.

154 We hypothesized that this abundance could indicate an enhanced ability to form 155 productive VH-VL pairs independent of VH sequence and antigen specificity. To test this 156 hypothesis, we constructed germline versions of these light chains with their most frequently observed J gene as published on the iRepertoire website²⁰⁻²³ (Extended Data Table 1) and 157 158 obtained full-length light chains for each PLC. SARS-CoV-2 specific Ig-Seg heavy chains were 159 then expressed with each PLC as a full-length IgG1. EC50 determination via ELISA using 160 recombinant spike protein demonstrated that of the 22 VHs tested, all paired productively with 161 one or more of the PLCs, resulting in several highly potent and specific antibodies (Fig. 2b). 162 Through this simple and rapid screening method we identified a panel of functional antibody 163 candidates for SARS-CoV-2 neutralization testing.

164

165 Yeast Surface Display-IgSeq identifies additional candidate antibodies

166 Our PLC screening strategy provides a method to rapidly generate functionally paired 167 antibodies, but the decreased sequence space of the nine PLCs may limit the ability to efficiently 168 identify potent binders. Thus, to search for additional high-affinity antibodies, we randomly paired 169 24 Ig-Seg heavy chains with donor-derived light chain libraries for yeast surface display (YSD) 170 selections (Fig. 1b), which allowed us to efficiently screen Fabs for each heavy chain/light chain 171 library combination. In YSD, heavy/light chain pairs were displayed as Fabs on the surface of a 172 humanized yeast strain that expresses human protein chaperones to improve antibody 173 expression. We then selected antigen-binding Fabs using two-color fluorescence cell sorting to 174 maximize Fab expression and binding to either SARS-CoV-2 spike (S-2P) or RBD (Extended 175 Data Figs. 3-4). As Fab enrichment over successive rounds of selection indicates antigen

specificity, we performed three rounds and identified and quantified productive VH-VL pairs based on MinION nanopore sequencing of Fab amplicons with sequence error correction and complementary Illumina reads (**Fig. 2c**, **Extended Data Figs. 5-6**). When light chain preference was compared, we observed extensive correspondence between YSD selection and PLC screening.

181

182 Parallel selections *via* yeast surface display yield functionally diverse antibodies

183 Much of the early stage primary antibody response is inaccessible to Ig-Seg discovery, as 184 detection is limited to secreted antibody proteins with sufficient abundance in serum²⁴. To broaden 185 our search for neutralizing antibodies, we employed our YSD platform to directly mine the cellular 186 repertoire for high affinity B-cell receptor sequences (Fig. 1d). We expressed and displayed large 187 VH-VL libraries (>10⁷) created by combinatorial pairing of donor IgG heavy and light chain amplicons. To maximize the functional diversity of selected antibodies, we performed three 188 189 rounds of selection on the initial libraries, using various gating strategies to ensure phenotypic 190 diversity (i.e. a variety of antibody expression and antigen binding strengths). Ultimately, the 191 parallelized selection strategy yielded populations with distinct binding characteristics as 192 confirmed by cytometric phenotyping of single clones (Extended Data Figs. 4, 7). We observed 193 progressive enrichment of specific antibodies with the dominant clone from each course of 194 selections generally approaching 20% of the total population after round three (Extended Data 195 Fig. 8). Individual clones from enriched populations were selected as candidates for further 196 characterization either at random or based on their abundance from the bioinformatic analysis. 197 Importantly, YSD detected enrichment of VH sequences that were not observed by Ig-Seq 198 proteomics, demonstrating the advantages and increased sensitivity of this multi-pronged 199 approach to antibody discovery.

200

201 Non-cognate VH-VL pairs are potent neutralizing antibodies to SARS-CoV-2

202 We initially screened the isolated IgG against the ECD, NTD, and RBD of the spike protein 203 using ELISA to identify the most promising candidates for authentic live virus neutralization of 204 SARS-CoV-2. We prioritized non-RBD antibodies in our screens in order to gain insights into 205 alternative neutralization mechanisms. From binding data of the full-length IgGs, we selected 94 206 candidate VH-VL pairs for live virus neutralization, and 37 pairs successfully neutralized authentic 207 SARS-CoV-2/WA1 (Fig. 3a, Extended Data Table 2). Curve fitting calculated IC50 values as low 208 as 18 pM, with seven antibodies showing sub-nanomolar values (8-32, 18 pM; N3-1, 251.9 pM; 209 8-131, 300 pM; 8-114, 350 pM; 12C8, 836 pM; A7V3, 955 pM; and 7-6, 977 pM). Twenty-three 210 neutralizing antibodies had IC50 values less than 100 nM (Fig. 3). Our high affinity NTD VH-VL 211 pairs (8-32, 8-131, 8-114, 12C8, A7V3, 7-6, 4C7, 4C8, 7A8) showed strong preference for IGHV1-212 24 gene usage. PLC screening successfully identified VL partners for both YSD and Ig-Seg 213 derived VHs (Fig. 3b-e). Seven of the mAbs were shown to be directed to the S2 domain. N3-1 214 possesses a highly potent RBD-directed IGHV4-31 heavy chain and displayed strong 215 neutralization activity with several PLCs as well as YSD VLs (Extended Data Table 2).

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218

Common binding mode of an NTD-directed antibody

A7V3 was of interest because of the different donor origins of the VH and VL. Its IGHV1-24 was discovered in both IgSeq and YSD from donor 1. However, the IGLV1-51 was mapped to 21 donor 2 and is also related to PLC8. The VH of A7V3 showed a strong preference to PLC8 in 222 ELISA screening, and related light chains appear in six other neutralizing VH-VL pairs, commonly 223 pairing with IGHV1-24 VHs. To ascertain its potential binding mode and contributions from both 224 variable regions, we determined a cryo-EM structure of A7V3 Fab complexed with SARS-CoV-2 225 S-ECD. The initial 3D reconstruction from 715,398 particles had a global resolution of 3.0 Å (**Fig.**

226 **4a**). Although the map revealed an A7V3 Fab molecule bound to each NTD of the trimeric spike. 227 only one Fab defined clear density. Further 3D classification was performed to resolve potential 228 heterogeneity in the particle set. One class was obtained consisting of one fourth of the total 229 particles and exhibited defined density for an NTD and its bound Fab. A local refinement of this 230 map focused on the NTD and Fab yielded a reconstruction with a well-resolved binding interface (Fig. 4b). Similar to the first structurally defined NTD-targeted antibody 4A8²⁵, the heavy chain of 231 232 A7V3 plays a dominant role in binding the N3 loop (residues 141-156) and N5 loop (residues 246-233 260) of the NTD. The light chain of A7V3 makes limited contact with the NTD, and is therefore 234 likely to play a structural role in positioning the heavy chain. H-CDR1, H-CDR3, and L-CDR2 235 collectively form a concave surface packed tightly against the N5 loop. Specifically, Trp100d in 236 H-CDR3, and both Tyr49 and Pro55 in L-CDR2, form a hydrophobic cage to enclose Pro251 at 237 the tip of the N5 loop (**Fig. 4c**). Glu31 from H-CDR1 is expected to form a salt bridge with Arg246, 238 and Asp101 from H-CDR3 is expected to form a hydrogen bond with the hydroxyl group of Tyr248. 239 Pro97 and Phe98 from H-CDR3 insert into a groove flanked by the N3 and N5 loops (Fig. 4d). As 240 opposed to the major binding interface on the N5 loop, A7V3 only makes a minor contact with the 241 N3 loop, mainly mediated by H-CDR1 and H-CDR2. Akin to a conserved Phe from H-CDR2 of 242 4A8, Phe51 contacts Lys147 via a π -cation interaction (**Fig. 4d-e**). In stark contrast, H-CDR3 of 243 A7V3 is much shorter than its counterpart in 4A8, which contains a Phe that forms a π - π stacking 244 interaction with Trp152 on the N3 loop. Collectively, the structural characterization of A7V3 245 demonstrates that our semi-synthetic approach is capable of yielding a novel neutralizing antibody 246 that binds to the NTD using an established modality.

247

248 An RBD-targeted antibody exhibits a novel quaternary binding mode

249 Structurally defined RBD-targeted antibodies or nanobodies complexed with SARS-CoV-2 S may be generally grouped into two types based on binding sites. The first type, mainly 250 251 discovered from the IGHV3-53 germline family, recognizes the ACE2-binding site and directly blocks host receptor engagement (e.g. antibodies CC12.1²⁶, C105²⁷ and VHH E²⁸). The second 252 253 type, from a more diverse germline family, recognizes a relatively conserved region on the RBD 254 that is mostly buried and contacts a neighboring RBD in the down conformation (e.g. C135²⁹. S309³⁰ and VHH V²⁸). From negative stain EM analysis of the IgG N3-1-spike complex, we 255 256 observed density for two Fabs, likely derived from the same IgG molecule, bound to multiple 257 RBDs on a single trimeric spike (Extended Data Figs. 12). We also found the affinity of N3-1 IgG 258 to SARS-CoV-2 spike is nearly 1000-fold stronger than N3-1 Fab to spike (Extended Data Fig. 259 13).

260

261 To investigate this unique binding mode, we determined a cryo-EM structure of N3-1 Fab 262 bound to SARS-CoV-2 S-ECD to a global resolution of 2.8 Å (Fig. 5a). We observed that two N3-263 1 Fabs bind to a single trimeric spike with one Fab binding to RBD in the up conformation and the 264 other Fab simultaneously engaging two RBDs: one in the up conformation and one in the down 265 conformation. We performed focused refinement on the Fab bound to the two RBDs (Fig. 5b), 266 which substantially improved the interpretability of the map in this region. The well-resolved Fab-267 RBD binding interface revealed two completely distinct epitopes on RBD-up and RBD-down (Fig. 268 5c). For RBD-up interaction, contacts are made by the Fab H-CDR1, H-CDR3 and L-CDR2, which together bury 862 Å² surface area. For the RBD-down interaction, contacts are made by the Fab 269 H-CDR2, H-CDR3 and L-CDR3, which buries 696 Å² surface area. Notably, the relatively long H-270 271 CDR3 loop (18 a.a.) engages both RBDs via hydrophobic and polar interactions. H-CDR3 272 residues Tyr98. Phe99 and Arg100a pack against a hydrophobic pocket formed by Tyr369. 273 Phe377, Lys378 and Pro384 on RBD-up, which are highly conserved between SARS-CoV and

274 SARS-CoV-2. This pocket is barely exposed when RBD is in the down conformation and is part 275 of the shared epitopes targeted by cross-reactive antibodies CR3022 and COVA1-16³¹. Lys378, 276 from the upper ridge of the pocket, forms a cation- π interaction with Tyr98 of H-CDR3, and its 277 amine group is expected to form a salt bridge with Asp100h of H-CDR3. In addition, main chain 278 atoms of Cys379 and Tyr369 form hydrogen bonds with Phe99 and Arg100a from H-CDR3, 279 strengthening this primary Fab binding interface on RBD-up. Furthermore, the sidechain 280 guanidinium of Arg408 on RBD-up likely has polar interactions with Tyr49 and Glu55 from L-281 CDR2, which along with H-CDR1, constitutes the secondary binding interface on RBD-up.

282

283 In contrast, the N3-1 binding site on RBD-down overlaps with the ACE2-binding site, 284 where 11 of 19 N3-1 epitope residues are also involved in ACE2 binding (Fig. 5d). Phe486 on 285 RBD-down inserts into a hydrophobic pocket formed by Trp96 (L-CDR3), Tyr52, Tyr58 (H-CDR2), 286 Arg100e and Val100g (H-CDR3). The sidechain guanidinium of Arg100e not only forms a 287 hydrogen bond with Tyr489, but also contacts Phe486 through cation- π interactions. In addition, 288 Asn487 and Gln 493 on RBD-down are expected to form polar interactions with Ser93 of L-CDR3 289 and Thr57 of H-CDR2. The angle of approach of N3-1 prevents ACE2 binding by both trapping 290 one RBD in the down position, thereby preventing exposure of the ACE2 binding site, as well as 291 by sterically inhibiting ACE2 access to the bound 'up' RBD. (Fig. 5e). Collectively, the N3-1 292 antibody engages an extensive quaternary epitope on neighboring RBDs through a novel binding 293 modality.

294

295 mAb N3-1 binds robustly to spike proteins from emerging variants of concern

296 Next, we tested whether binding of A7V3 (NTD-targeting) or N3-1 (RBD-targeting) was 297 detrimentally affected by mutations emerging in SARS-CoV-2 variants of concern (VOC). We 298 assessed binding to the 12 most abundant SARS-CoV-2 variants from Houston¹⁷ and did not

299 observe detrimental losses in binding avidity (Extended Data Table 5). Next, we used 300 mammalian surface display of spike variants to test binding of N3-1 to mutations that were 301 previously shown to reduce or abrogate binding to one or both of the Regeneron antibodies 302 (REGN10987 and REG10933) as well as a commonly found mutation in Australia (S477N). 303 Strikingly, N3-1 was able to bind each of the mutant spike variants. N3-1 binding was slightly 304 enhanced by 10 of the 12 mutant spike proteins tested (L455A, G476D, E406W, Q493F, K444A, 305 K444Q, V445A, G446A, G446V, and S477N) using this assay. Variant Y453A was 306 indistinguishable from binding to the prefusion stabilized SARS-CoV-2 S D614G, and F486K was 307 the only mutation shown to marginally reduce binding ($\sim 6\%$) (**Extended Data Fig. 15**). We then 308 tested binding to B.1.1.7, B.1.1.248, and B.1.351 spike protein variants using the same 309 mammalian surface display assay. Binding of A7V3 was reduced by all three variants, a result 310 predicted from our structural work. In contrast, N3-1 binding was largely unperturbed by the 311 accumulated mutations (Fig. 6a) of B.1.1.7, B.1.1.248, and B.1.351 spike proteins in this assay 312 (Fig. 6b). To validate the mammalian display screening results, we examined the binding of the 313 SARS-CoV-2 S Wuhan-Hu-1, and its variants B.1.1.7, and B.1.351 to N3-1 Fab or N3-1 IgG using 314 surface plasmon resonance (SPR). N3-1 Fab binding to each of the variants was only able to be 315 fit to a heterogenous binding model (Extended Data Fig. 13), consistent with the structural 316 studies. N3-1 IgG exhibited comparable binding to Wuhan-Hu-1 and its variants, with 67.9 pM 317 apparent affinity to Wuhan-Hu-1, 296 pM apparent affinity to B.1.1.7, and 300 pM apparent affinity 318 to B.1.351 (Fig. 6c-e). We also tested binding via SPR to SARS-CoV spike protein and discovered 319 that N3-1 IgG cross-reacts with SARS-CoV spike with 28 nM affinity (Extended Data Figure 13). 320 Finally, we corroborated these findings with live virus neutralization assays with circulating VOC 321 B.1.1.7. We found that the wildtype WA/1 strain was fully neutralized by both N3-1 and A7V3, but 322 B.1.1.7 was only neutralized by N3-1 at the same titer as WA/1 (Fig. 6f), consistent with our 323 structure-based predictions.

324

325 Discussion

326 SARS-CoV-2 and its evolved progeny represent a moving target for therapeutic 327 development. While the rapid response of the global research community has led to the discovery 328 of many neutralizing mAbs³²,^{25,33} the emergence and widespread dissemination of viral variants 329 such as the U.K. variant B.1.1.7, Brazilian variant B.1.1.248, and South African variant B.1.351 330 stresses the need for continued efforts to rapidly identify and develop antibodies that can bind to 331 diverse epitopes. By combining proteomic analyses of antibody repertoires from patient samples 332 and yeast-display with public light chain pairing, we developed a high-throughput antibody 333 discovery strategy that identified 37 neutralizing antibodies that target multiple regions of the 334 SARS-CoV-2 spike protein, including the RBD, NTD, and the S2 domain, and that broadly bind 335 multiple viral variants.

To accelerate the discovery of neutralizing antibodies, we first performed Ig-Seq proteomic analyses on patient samples. However, because Ig-Seq does not provide information on relative affinities, epitope targeting, and light chain pairings, we adopted two strategies for light chain discovery: a rapid screening method utilizing public light chains (PLCs) and a YSD light chain selection. Ultimately, we found that both methods independently yielded multiple neutralizing antibodies, with several of the most potently neutralizing mAbs converging in two or more of our approaches.

The neutralizing antibodies identified relied on eight IGHV genes, including IGHV3-30 and IGHV1-2, consistent with studies that show these genes are overrepresented in SARS-CoV-2 neutralizing antibodies³². In addition, IGHV1-24 was previously found to contribute to a number of NTD-directed neutralizing antibodies^{25,34} and was observed in 10 of the neutralizing antibodies herein. We found representative neutralizers from each of our discovery strategies, although each

348 method yielded one or more unique IGVH or IGVL genes, validating the high coverage afforded349 by this multi-pronged approach.

350 The rapid PLC screening method appears particularly useful not only for expediting 351 functional testing of Ig-Seg-derived heavy chains, but also to generate high-affinity antibodies. 352 For instance, N3-1 exhibits a sub-nanomolar IC-50 (252 pM) and binding affinity (68 pM), while 353 A7V3 has an IC-50 of 950 pM. A7V3 is a particularly interesting demonstration of the utility of 354 the strategy, because its heavy chain was first identified from one donor, while the light chain was 355 mapped to another donor. Initial PLC screening indicated a nearly identical IGLC VJ gene (PLC8) 356 as a productive partner, which further confirmed the utility of PLC screening for productive 357 antibody VH-VL pairings.

358 We resolved high-resolution cryo-EM structures of these two newly discovered 359 neutralizing antibodies, and found that N3-1 targets the RBD, whereas A7V3 targets the NTD. 360 Structural analysis of N3-1 revealed that the L-CDR2 and L-CDR3s were both engaged in binding, 361 which suggests that our bioinformatically derived PLCs may directly contribute to neutralization. 362 A7V3 bound to a common neutralization site on the NTD³⁴, and ultimately, PLC8 was found to 363 productively pair with three different neutralizing antibodies that were most commonly associated 364 with NTD-binding heavy chains. Our work not only establishes the first empirical demonstration 365 that public light chains can form functional pairings across heavy chains, but also debuts their 366 utility as a prolific method for high affinity antibody discovery, potentially improving rapid-response 367 antibody discovery for neo-antigens in future epidemics. Our results further suggest that naturally 368 paired repertoires may be unnecessary for efficient antibody discovery, in contrast to a recently 369 published work where cognate pairing was deemed necessary for effective neutralization³⁵.

370 The cryo-EM structure of N3-1 is of particular importance because it reveals a novel 371 binding mechanism at a quaternary epitope of the spike trimer, thus allowing this antibody to bind 372 the RBD in both the 'up' and 'down' conformations. This greatly enhances the binding affinity of 373 the full mAb as it enables binding to all three RBD subunits in the spike protein trimer. As a 374 consequence of this unique mechanism of action, N3-1 is minimally perturbed by spike mutations 375 in the newly emerged variants B.1.1.7, B.1.1.248, and B.1.351, where it was shown to still have 376 low picomolar affinities and neutralized B.1.1.7 with a similar titer to WA/1. N3-1 is also capable 377 of binding the distantly related SARS-CoV spike protein with low nanomolar affinity (28 nM). Given 378 the mechanism of binding and the fact that the N3-1 binding epitope for the RBD-down 379 conformation corresponds to the ACE2-binding site, we fully expect N3-1 to show neutralization 380 of VOCs not tested here.

381 The strategy we described here will aid in the rapid identification and development of 382 monoclonal antibodies to extant SARS-CoV-2 variants, and can be readily utilized for identifying 383 antibody therapeutics for variants yet to be discovered and for other explosive pathogen 384 zoonoses. Candidate mAbs may be produced in less than 10 days, and have desirable characteristics - presence in immune responses, public light chain robustness, validation of 385 386 expression via yeast display – that may enable production at scale. Because of the modular 387 nature of the synthetic biology-like discovery process, the overall strategy could also be readily 388 extended to affinity maturation or directed evolution of tighter binding mAbs to distinct antigens or 389 epitopes. In the long term, the ability to store, combine, and continually reassess repertoires 390 should prove broadly useful in not only discovering novel antibodies with potential therapeutic 391 applications, but also for studying the evolution of adaptive immunity.

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518 Online Methods

519 Strains and media. Yeast strain EBY100 (MATa AGA1::GAL1-AGA1::URA3 ura3-52 trp1 leu2-520 delta200 his3-delta200 pep4::HIS3 prbd1.6R can1 GAL) was acquired from ATCC (cat. no. MYA-521 4941) and used for antibody expression and selection. To improve antibody expression the 522 human chaperones BIP (binding immunoglobulin protein) and PDI (protein disulfide isomerase) 523 were genomically integrated as an expression cassette in the HO locus. Yeast were grown in rich 524 medium (YPD; Takara, cat. no. 630409) or in selective medium for leucine prototrophs after library 525 transformation (Takara cat. no. 630310). YEP-galactose was used for expression of displayed 526 antibody libraries (1% yeast extract, 1% bacto-peptone, 0.5% NaCl, 2% galactose, 0.2% glucose).

527 **Antigens and antibodies.** Spike antigen was biotinylated using the EZ-link kit (Thermo Scientific, 528 cat. no. 21435) and labeled with streptavidin-AF647 (Invitrogen, cat. no. S32357). RBD was 529 labeled with a mouse anti-human Fc-AF647 (Southern Biotech, cat. no. 9042-31). Fab library light 530 chains were labeled with anti-FLAG M2-FITC (Sigma, cat. no. F4049).

531 **Donors.** Blood was collected from 3 PCR-confirmed, symptomatic patients. Donors 1 and 2 were 532 collected at day 12 post-onset of symptoms. Donor 3 was collected on day 11. None of the donors 533 were hospitalized or experienced severe disease. PBMCs and plasma were both collected by 534 density gradient centrifugation using Histopaque-1077 (Sigma-Aldrich).

Preparation of serum antibodies for Ig-Seq proteomic analysis. Serum samples were prepared for Ig-Seq analysis as previously described³⁴. Briefly, total IgG was isolated from plasma using Pierce Protein G Plus Agarose (Pierce Thermo Fisher Scientific) and cleaved into F(ab')2 fragments with IdeS protease. Antigen-specific F(ab')2 was enriched by affinity chromatography against recombinant SARS-CoV-2 S-2P or RBD protein cross-linked to NHS-activated agarose resin (Thermo Fisher Scientific). Eluted F(ab')2 fractions were concentrated by vacuum

541 centrifugation and prepared for mass spectrometry-based proteomic analysis as previously 542 described.²⁴

543 LC-MS/MS analysis of antigen-enriched antibodies. Liquid chromatography-tandem mass 544 spectrometry analysis was carried out on a Dionex Ultimate 3000 RSLCnano system coupled to 545 an Orbitrap Fusion Lumos Mass Spectrometer (Thermo Scientific). Samples were loaded onto an 546 Acclaim PepMap 100 trap column (75 µm × 2 cm; Thermo Scientific) and separated on an Acclaim 547 PepMap RSLC C18 column (75 µm × 25 cm; Thermo Scientific) with a 3%-40% acetonitrile 548 gradient over 60 min at a flow-rate of 300 nl/min. Peptides were eluted directly into the Lumos 549 mass spectrometer using a nano-electrospray source. Mass spectra were acquired in data-550 dependent mode with a 3 sec. cycle time. Full (MS1) scans were collected by FTMS at 120,000 551 resolution (375-1600 m/z, AGC target = 5E5). Parent ions with positive charge state of 2-6 and 552 minimum intensity of 3.4E4 were isolated by quadrupole (1 m/z isolation window) and fragmented 553 by HCD (stepped collision energy = 30 + (-3%)). Fragmentation (MS2) scans collected by ITMS 554 (rapid scan rate, AGC target = 1E4). Selected ions and related isotopes were dynamically 555 excluded for 20 sec (mass tolerance = +/-10 ppm).

Antibody variable chain sequencing. Peripheral blood mononuclear cells (PBMCs) from processed donor samples were provided in Trizol as a kind gift from Dr. Gregory C. Ippolito³⁴. IgG and IgM VH and VL cDNA libraries were separately amplified from PBMC RNA of three donors and sequenced to create donor specific reference databases, from which the complete amino acid sequences of serum IgG proteins could subsequently be determined based on their mass spectral identifications. **Extended data Figure 1** shows the distribution and diversity of clonotypes and V-gene usage for the variable heavy chain repertoires of donors 1 and 2.

563 Ig-Seg MS data analysis. Mass spectra were analyzed using Proteome Discoverer 2.2 software 564 (Thermo Scientific). Precursor masses were first recalibrated with the Spectrum File RC node 565 using a consensus human reference proteome database (UniProt) with common contaminants 566 (MaxQuant) and precursor mass tolerance of 20 ppm. Recalibrated mass spectra were searched 567 against a custom database for each donor consisting of donor-derived VH sequences, VL 568 sequences, and the human and contaminant sequences using the Sequest HT node. Mass 569 tolerances of 5 ppm (precursor) and 0.6 Da (fragment) were used. Static carbamidomethylation 570 of cysteine (+57.021 Da) and dynamic oxidation of methionine (+15.995 Da) were considered. 571 False discovery rates for peptide-spectrum matches (PSMs) were estimated by decoy-based 572 error modelling through the Percolator node. Label-free guantitation (LFQ) abundances were 573 calculated from precursor areas using the Minora Feature Detector and Precursor lons Quantifier 574 nodes.

575 Resulting PSMs were filtered according to methods described (Boutz, 2014). Briefly, peptide 576 sequences differing only by isoleucine/leucine substitution were considered equivalent and 577 combined into a single PSM. PSMs were re-ranked by posterior error probability, q-value, and 578 Xscore. Only top-ranked, high-confidence PSMs (FDR < 1%) were retained for each scan. If two 579 or more PSMs had identical top-ranked scores, they were considered ambiguous and removed. 580 PSMs for the same peptide sequence were summed and the average mass deviation (AMD) was 581 calculated for each peptide. Peptides with AMD greater than 2 ppm were filtered out. Peptides 582 mapping to VH sequences from a single clono-group were considered clono-specific. Clono-583 specific peptides overlapping the CDR3 sequence by four amino acids or more were considered 584 CDR3-informative.

585 For each clono-group, PSMs and LFQ abundances of clono-specific CDR3-informative peptides 586 were summed. Ratios of elution:flow-through PSMs and LFQ abundances were calculated; only 587 clono-groups with both ratios > 5 were considered elution-specific.

588 Library assembly and bacterial transformation. Donor B-cell VH and VL amplicons were 589 amplified via PCR to include adapters for cloning into yeast expression vectors. Assembly into 590 the yeast kappa and lambda expression vectors was done via Golden Gate assembly. Library 591 assemblies were prepared in 20 µL reactions as follows: 2 µL 10X AARI buffer (ThermoFisher 592 Scientific, cat. no. B27), 0.4 µL 50X oligo buffer (ThermoFisher Scientific, cat. no. ER1582), 0.2 593 uL 100 mM ATP (ThermoFisher Scientific, cat, no. R0441), 20 fmol backbone DNA, 40 fmol VH 594 and VL amplicons, 0.5 µL (2 U/µL) AARI endonuclease (ThermoFisher Scientific, cat. no. 595 ER1582), and 0.5 µL T7 ligase (3000 U/µL) (NEB, cat. no. M0318). Each assembly was scaled 596 up to 16 total reactions in 8-well strips. Thermocycling consisted of the following protocol: 37°C, 597 15 minutes; 37°C, 2 minutes, 16°C, 1 minute; go to step 2, x74; 37°C, 60 minutes; 80°C, 15 598 minutes; hold at 4°C. Assemblies were consolidated and column purified using Promega Binding 599 Solution (Promega, cat. no. A9303) to bind DNA to a Zymo-spin II column (Zymo Research, cat. 600 no. C1008). The column was washed twice with DNA Wash Buffer (Zvmo Research, cat, no. 601 D4003) and eluted in 30 µL nuclease-free water. For library transformations, DH10B cells were 602 diluted 1:100 from confluent culture into 50 mL Superior broth (AthenaES, cat. no. 0105). When 603 cells reached an OD_{600} of 0.4-0.6, they were washed 3X with cold 10% glycerol and resuspended 604 to a final volume of 600 µL. The purified library was added to cells and electroporated at 1.8 kV 605 in an E. coli Pulser electroporator (Bio-Rad) using Genepulser 0.2 cm cuvettes (Bio-Rad, cat. no. 606 1652086) at 200 µL per transformation.

607 **Library transformation into yeast and protein expression.** Purified libraries were linearized 608 for integration into the yeast genome via homologous recombination at the Leu2 locus. For each

609 1 µg library plasmid, 0.5 µL Notl (10 units/µL) (NEB, cat. no. R0189) was used with the supplied 610 Buffer 3.1 in 10 µL. Reactions were incubated at 37°C overnight and heat inactivated at 80°C for 611 20 minutes. Digests were pooled and column purified as described in previous sections and 612 eluted in 25 µL nuclease-free water. Our strain was electroporated as described elsewhere¹. We 613 found that 10 µg of linearized DNA was sufficient for library sizes of 10⁶, and that library sizes 614 could reach >10⁷ with 20 µg DNA. Transformed yeast were recovered in SD -Leu medium (see 615 "strains and media" section). Libraries were passaged once at 1:100 before protein expression to 616 reduce contamination from untransformed cells. To express Fab libraries, yeast were washed in 617 YEP-galactose (see "strains and media") and diluted 1:10 into 10 mL final volume. Cells were 618 induced for 48 hours at 20°C with shaking.

619 Fab library labeling and selection. Expressed yeast libraries were harvested at 100 µL 620 (representing approximately 10⁷ cells) and washed with PBSA buffer (1X PBS, 2 mM EDTA, 0.1% 621 Tween-20, 1% BSA, pH 7.4). Antigen was incubated with cells in 1 mL in PBSA at 200 nM at RT 622 for one hour, washed with PBSA at 4C, and labeled with secondary antibodies (mouse anti-human 623 FITC, 1:100; streptavidin-AF647, 1:100; mouse anti-human Fc-AF647, 1:50). Cells were washed 624 2X and resuspended in 2 mL in cold PBSA for sorting. Cell sorting was performed using a Sony SH800 fluorescent cell sorter. For first round libraries, 10⁷ events were sorted into 2 mL SD-Leu 625 626 medium supplemented with penicillin/streptomycin (Gibco, cat. no. 15140122). Cells were 627 recovered by shaking incubation for 1-2 days for further rounds of selection or plated directly for 628 phenotyping clones.

Next generation sequencing. Genome extraction was performed on yeast cultures of libraries and sorted rounds underwent genome extraction using a commercial kit (Promega, cat. no. A1120) with zymolyase (Zymo Research, cat. no. E1004). 100 ng genomic template was used to amplify the heavy and light chains separately or as one amplicon for short or long-read

sequencing, respectively. For amplification of heavy chain genes only, primers JG.VHVLK.F and
JG.VH.R were used. For amplification of light chain genes only, primers JG.VL.F and JG.VHVK.R
or JG.VHVL.R were used for kappa and lambda vectors, respectively. For amplification of paired
genes, primers JG.VHVLK.F and JG.VHVK.R or JG.VHVL.R were used. Amplicons were column
purified and deep sequenced with an iSeq. In parallel, we obtained ~1.8 kb sequences spanning
the entire VH and VL using MinION nanopore sequencing (Oxford Nanopore Technologies Ltd.,
MinION R10.3).

640 Colony PCR and Sanger sequencing. Sorted yeast populations were plated on SD -Leu and 8-641 32 colonies per plate were picked into 2 mL microplates either by hand or using a QPIX 420 642 (Molecular Devices) automatic colony picker. Cultures were grown at 1000 rpm at 3 mm orbit at 643 30°C overnight. Cells (20 µL) were transferred to a fresh microplate and washed with 1 mL TE 644 buffer (10 mM Tris, 1 mM EDTA). Cells were incubated with 20 µL zymolyase solution (5 mg/mL 645 zymolyase, 100T in TE) at 37°C for 1 hour. Cells (5 µL) were then used in colony PCR to amplify 646 the paired heavy and light chains. Amplicons were column purified with the Wizard SV 96 PCR 647 Clean-Up System (Promega, cat. no. A9342) and yields were quantified with a Nanodrop 648 spectrophotometer or the Quant-it Broad-Range dsDNA kit (Invitrogen, cat. no. Q33130). 649 Approximately 10 ng (2.5-5 µL) of purified PCR products were then subjected to Sanger 650 sequencing.

Long-read sequencing (donors 1 & 2) Sequencing libraries were prepared from 18 amplicon samples using the Native Barcoding Kit (Oxford Nanopore Technologies; cat. no. EXP-NBD103) paired with the Ligation Sequencing Kit (Oxford Nanopore Technologies; cat. no. SQK-LSK109) according to the manufacturer's directions. Between four and eight sequencing libraries per flow cell were pooled for sequencing on three MinION flow cells (Oxford Nanopore Technologies; R9.4.1) for 72 hours on an Oxford Nanopore Technologies MinION Mk1B device (Oxford

Nanopore Technologies). Raw data was basecalled using the high accuracy model in Guppy(v.3.5.2).

659

Short-read sequencing (phenotyping plates) Sequencing libraries were prepared from 308 amplicon samples using the Nextera DNA Flex Library Preparation kit (Illumina; cat. no. 20018705) according to the manufacturer's directions. Sequencing libraries were pooled and sequenced (2x151bp) on an iSeq 100 (Illumina; California, USA) using iSeq 100 i1 Reagents v.1 (Illumina; cat. no. 20021533).

665

Long-read sequencing (donor 3) Sequencing libraries were prepared from 32 amplicon samples using the Native Barcoding Kit (Oxford Nanopore Technologies; cat. no. EXP-NBD104) paired with the Ligation Sequencing Kit (Oxford Nanopore Technologies; cat. no. SQK-LSK109) according to the manufacturer's directions. Between five and eight sequencing libraries per flow cell were pooled for sequencing on five GridION flow cells (Oxford Nanopore Technologies; R9.4.1) for 72 hours on a GridION Mk1 device (Oxford Nanopore Technologies; Oxford, England, UK). Raw data was live basecalled using the high accuracy model in Guppy (v.3.2.10).

673

574 **Short-read sequencing (donor 3)** Sequencing libraries were prepared from 32 samples using 575 the Nextera DNA Flex Library Preparation kit (Illumina; cat. no. 20018705) according to the 576 manufacturer's directions. Sequencing libraries were pooled and sequenced (2x151bp) on an 577 iSeq 100 (Illumina; California, USA) using the iSeq 100 i1 Reagents v2 (Illumina; cat. no. 578 2009584).

679

Long-read sequencing (YSD-IgSeq) Sequencing libraries were prepared from 16 amplicon
 samples using the Native Barcoding Kit (Oxford Nanopore Technologies; Cat. No. EXP-NBD104)

paired with the Ligation Sequencing Kit (Oxford Nanopore Technologies; Cat. No. SQK-LSK109)
according to the manufacturer's directions. Four sequencing libraries were pooled per flow cell
and sequenced on four GridION flow cells (Oxford Nanopore Technologies; R9.4.1) for 72 hours
on a GridION Mk1 device (Oxford Nanopore Technologies; Oxford, England, UK). Raw data was
live basecalled using the high accuracy model in Guppy (v.4.0.11).

687

Short-read sequencing (YSD-IgSeq) Sequencing libraries were prepared from 16 samples using the Nextera DNA Flex Library Preparation kit (Illumina; cat. no. 20018705) according to the manufacturer's directions. Sequencing libraries were pooled and sequenced (2x150 bp) on an iSeq 100 (Illumina; California, USA) using the iSeq 100 i1 Reagents v2 (Illumina; Cat. No. 2009584).

693 Sequence processing and consolidation into VHVL clones Individual reads after Guppy base 694 calling typically average more than 10% error per base, and numerous tools exist to align and 695 reduce reads into consensus sequences with substantially improved accuracy³⁶. However, such 696 tools were not designed for antibody library sequencing and its huge populations of subtly different 697 sequences which, even assuming successful alignment, group into a myriad of very short, 698 disconnected assemblies. We therefore implemented a bioinformatic pipeline to obtain accurate 699 VHVL sequences from the MinION and iSeq data and to estimate, within each YSD round, the 700 relative abundance of individual VHVL pairs.

Our methods proceed through antibody V(D)J annotation of raw MinION reads using MiXCR (v3.0.13)³⁷; iteratively growing and shrinking sequence clusters based on annotated features from each read; sequence error correction and consolidation within each cluster, optionally including high quality Illumina reads; and finally, VHVL clone definition within each sample and quantitation by number of reads mapped to each clone. For enumeration, we only include counts for reads

with a length of 1700 to 2100 base pairs. This points to a secondary advantage of MinION sequencing, as shorter reads proliferate during PCR and inflate apparent abundance of particular species. Without length-filtering, relative VHVL abundance calculated from both MinION and iSeq reads are strongly correlated.

710 Tissue culture and transient transfection of ECD and RBD. Spike ECD protein and RBD 711 proteins were expressed in Expi293F cells using the manufacturer provided guidelines with slight 712 modifications. In short, a 1 ml frozen working cell bank of Expi293F cells at 1 × 10⁷ viable cells/mL 713 were thawed in a bead bath at 37°C for 2-3 mins. The vial was sprayed with 70% isopropyl alcohol 714 and transferred into a biosafety cabinet. The thawed cells were transferred to a 125 mL non-715 baffled vented shake flask containing 29 mL of fresh pre-warmed ExpiExpression medium at 716 37°C. Cells were incubated in at 37°C with \geq 80% humidity and 8% CO₂ on an orbital shaker at 717 120 rpm and grown until they reached a cell density of 3 × 10⁶ viable cells/ml. Fresh pre-warmed 718 ExpiExpression medium was added to 1 L non-baffled vented shake flask and the 30 mL cell 719 suspension was carefully introduced, making the seeding density of 0.4 x 10⁶ viable cells/mL in a 720 final culture volume of 225 mL. After the cell density reached 3×10^6 viable cells/ml, the culture 721 was expanded to a final volume of 2.25 L in two 2.8 L Thomson Optimum Growth flasks with a 722 seeding density of 0.3×10^6 viable cells/mL. After the cell density reached 3×10^6 viable cells/mL, 723 2 L of the culture was-split into four Thompson flasks with each flask containing 500 mL of culture 724 medium and 500 mL of fresh pre-warmed ExpiExpression medium. The final culture volume in 725 each of the four flasks was 1 L. The cells were incubated at 37°C with ≥80% humidity and 8%

726 CO₂ on an orbital shaker at 100 rpm. When the cells reached a density of 3 × 10⁶ viable cells/mL,

the culture was transferred to two 500 mL sterile centrifuge bottles and the cells were spun down at 100 x g for 10 min. The supernatant was removed, and the cells were resuspended in 4 L of fresh, prewarmed medium. The cells were allowed to re-stabilize in the incubator for 24 h and were transfected at 3×10^6 viable cells/mL in 4 L.

731 Transfection was performed by diluting 4 mg of plasmid DNA (pDNA) in 240 mL of OptiMEM 732 medium in a sterile bottle and gently inverting 3 - 4 times before incubating at room temperature 733 (RT) for 5 min. ExpiFectamine 293 reagent (13 mL) was then diluted in 225 mL OptiMEM in a 734 sterile bottle and inverted 3 - 4 times before incubating for 5 min at RT. The diluted pDNA and 735 ExpiFectamine reagent were carefully mixed and incubated at RT for 15 min. One-fourth of the 736 combined complex was then slowly transferred to each of the flasks while gently swirling the cells 737 during addition. The cultures were again placed in the incubator with shaking at 100 rpm for 18 h. 738 Post-transfection, ExpiFectamine 293 Transfection Enhancer 1 (6 mL) and ExpiFectamine 293 739 Transfection Enhancer 2 (60 mL) were added to each flask. The cell viability was monitored every 740 24 h and the cells were harvested when the viability dropped below 70% or after approximately 3 741 d. Harvesting was done by centrifugation at 15,900 x g for 45 min at 4°C and the supernatant was 742 transferred into sterile bottles.

Antigen purification. The feed was prepared adding 0.2 M NaCl and 10 mM imidazole to the supernatant while mixing. The feed material was filtered using a 0.45 μm PES filter membrane pre-wetted with PBS before loading on a prepared Ni-IMAC column. A metal affinity column was prepared by packing IMAC FF beads (Cytiva) into an AxiChrom 70 column housing to a bed height of 9.5 cm and then charging with 50% column volume (CV) of a 0.2 M nickel sulfate solution, washing with water and then 50% CV of 100% "B" buffer (50 mM sodium phosphate buffer

containing 300 mM NaCl and 250 mM imidazole, pH 7.8) to remove weakly bound nickel ions.
The column was then washed with 100% "A" buffer (50 mM sodium phosphate buffer containing
300 mM NaCl and 20 mM imidazole, pH 7.2) prior to sample loading. The prepared (4.8 L) feed
was then loaded onto the column at a linear flow rate of 90 cm/h. After loading, the column was
washed with two CVs of 100% A buffer and two CVs of a 13% B buffer (containing 50 mM
imidazole) before eluting the protein using 100% B (250 mM imidazole) for 3 CVs. All steps except
for the loading step were done at 150 cm/h.

Fractions from the elution peak were pooled and then concentrated 4- to 8-fold by ultrafiltration (UF) using a 115 cm²hollow fiber cartridge with either a 50 kDa (S-2P) or a 10 kDa (RBD) molecular weight cutoff membrane (Repligen) and then diafiltered after concentration by exchanging with 5 volumes of PBS.

The concentration of diafiltered protein was determined by measuring the absorbance at 280 nm versus a PBS blank. The protein concentration in mg/mL was obtained using a divisor of 1.03 mL/mg-cm for S-2P and 1.19 mL/mg-cm for RBD. A qualitative assessment of protein quality was made using SDS-PAGE with SYPRO Ruby staining (BioRad) for reduced and non-reduced samples. Only those preparations showing predominantly full-length S-2P (160 kDa subunits) or RBD (70 kDa) were used in ELISAs for assessing neutralizing antibodies.

Monoclonal antibody expression and purification. VHVL candidates were cloned into custom
Golden Gate compatible pCDNA3.4 vectors for IgG1 expression. For transfections, VL was mixed
3:1 with a corresponding VH. Plasmids were transfected into Expi293F (Invitrogen) cells using
the recommended protocol. Monoclonal antibodies were harvested at 5-7 days post-transfection.
Expi293F cells were centrifuged at 300 x g for 5 min, supernatants were collected and centrifuged
at 3000 x g for 20 min at 4°Cand diluted to 1X PBS final concentration. Each supernatant was

passed through a Protein G or A agarose affinity column (Thermo Scientific). Flow through was
collected and passed through the column three times. Columns were washed with 10 CV of PBS
and antibodies were eluted with 5 mL 100 mM glycine, pH 2.7 directly in neutralization buffer
containing 500 µL 1 M Tris-HCl, pH 8.0.

Phenotyping assays. Sorted clones from rounds two or three of selection were picked into microplates as described previously. After antibody expression, 10 µL of cells were transferred to a fresh 2-mL microplate and washed 2X with 200 µL cold PBSA buffer. Cells were labeled with 200 nM spike or RBD antigen in 50-100 µL PBSA at RT for 1 h with shaking at 1000 rpm, 3 mm orbit. Labeled cells were washed 2X, and secondary labels were applied as previously described. Cells were resuspended in 200 µL ice-cold PBSA just before analysis. Samples were analyzed on a Sony SA3800 Spectral Cell Analyzer.

783 Analysis of mammalian cell surface displayed spike proteins with flow cytometry. The assay has been described in detail in the recent paper by Javanmardi et al³⁸. Briefly, plasmids 784 785 expressing full-length SARS-CoV-2 spike proteins (Spike-Linker-3XFLAG-TM), WT (HexaPro-786 D614G) and variants, were transfected into HEK293T cells using Lipofectamine 2000 according 787 to manufacturer's instructions. After 48 h, cells were collected and resuspended in PBS-BSA and 788 incubated with anti-FLAG (mouse) and anti-spike (human) mAbs for 1 h. shaking at RT. Cells 789 were washed 3X with PBS-BSA, resuspended in mL and incubated with Alexa Fluor 488 (anti-790 mouse) and Alexa Fluor 647 (anti-human) antibodies for 30 min, shaking at 4°C. Cells were 791 washed again and resuspended in PBS-BSA prior to flow cytometry analysis (SA3900 Spectral 792 Analyzer, Sony Biotechnology). All data was analyzed with FlowJo (BD Bioscience).

ELISA. Antigen ELISA plates were made using high-binding plates (Corning, cat. no. 3366) with
 antigen diluted in PBS to a final concentration of 2 μg/mL. Antigen solution (50 μL) was added to

795 microplates and incubated overnight at 4°C with shaking at 100 rpm, 3 mm orbit. Plates were blocked with PBSM (2% milk in PBS) at RT for 1 h. Plates were washed 3X with 300 µL PBS-T 796 797 (0.1% Tween-20). Purified antibodies were prepared to 10 µg/mL in PBSM and serially diluted. 798 Antibodies were incubated for 1 h at RT. Plates were washed 3X with PBS-T, and secondary goat 799 anti-human Fab-HRP (Sigma-Aldrich, cat. no. A0293) was applied at 1:5000 in PBSM in 50 µL 800 and incubated at RT for 45 min. HRP substrate (50 µL) was added to wells and the reaction 801 proceeded for 5-15 min until quenched with 50 µL 4M H₂SO₄and analyzed for absorbance at 450 802 nm in a plate reader.

803 Live virus neutralization assays. A SARS-CoV-2 microneutralization assay was adapted from an assay used to study Ebola virus³⁹. This assay also used SARS-CoV-2 strain WA1. Antibodies 804 805 were diluted in cell culture medium in triplicate. A SARS-CoV-2 monoclonal antibody was used 806 as a positive control. An antibody that does not bind SARS-CoV-2A was used as negative control. 807 Diluted antibodies were mixed with the SARS-CoV-2 WA1 strain, incubated at 37°C for 1 h, then 808 added to Vero-E6 cells at target MOI of 0.4. Unbound virus was removed after 1 h incubation at 809 37°C, and culture medium was added. Cells were fixed 24 h post-infection, and the number of 810 infected cells was determined using SARS-CoV-S specific mAb (Sino Biological, cat. no. 401430-811 R001) and fluorescently-labeled secondary antibody. The percent of infected cells was 812 determined with an Operetta high-content imaging system (PerkinElmer) and Harmonia software. Percent neutralization for each monoclonal antibody at each dilution was determined relative to 813 814 untreated, virus-only control wells.

Live virus neutralization (VN) for variant of concern B.1.1.7 and WA/1 were performed in an orthogonal assay. The ability of the monoclonal antibodies to neutralize SARS-CoV-2 was determined with a traditional VN assay using SARS-CoV-2 strain USA-WA1/2020 (NR-52281-BEI resources), as previously described^{40,41,42,43}. All experiments with SARS-CoV-2 were performed

819 in the Eva J Pell BSL-3 laboratory at Penn State and were approved by the Penn State Institutional 820 Biosafety Committee (IBC # 48625). For each mAb a series of 12 two-fold serial dilutions were 821 assessed from a stock concentration of 1 mg/ml. Triplicate wells were used for each antibody 822 dilution. 100 tissue culture infective dose 50 (TCID50) units of SARS-CoV-2 were added to 2-fold 823 dilutions of the diluted mAb. After incubating for 1 hour at 37°C, the virus and mAb mixture was 824 then added to Vero E6 cells (ATCC CRL-1586) in a 96-well microtiter plate and incubated at 37°C. 825 After 3 days, the cells were stained for 1 hour with crystal violet-formaldehyde stain (0.013% 826 crystal violet, 2.5% ethanol, and 10% formaldehyde in 0.01 M PBS). The endpoint of the 827 microneutralization assay was determined as the highest mAb dilution, at which all 3, or 2 of 3, 828 wells are not protected from virus infection. Percent neutralization ability of each dilution of the 829 mAb was calculated based on the number of wells protected, 3, 2, 1, 0 of 3 wells protected was 830 expressed as 100%, 66.6%, 33.3%, or 0%.

831 Surface plasmon resonance. To investigate the binding kinetics of mAb N3-1 binding to the 832 spikes, purified His-tagged spike variants (SARS-CoV Tor2 S-2P, SARS-CoV-2 Wuhan-Hu-1 S-833 HexaPro, SARS-CoV-2 B.1.1.7 S-Hexapro and SARS-CoV-2 B.1.351 S-HexaPro) were 834 immobilized on a Ni-NTA sensor chip (GE Healthcare) using a Biacore X100 (GE Healthcare). 835 For Fab binding experiments, we immobilized spike proteins to a level of ~450 response units 836 (RUs). Serial dilutions of purified Fab N3-1 were injected at concentrations ranging from 400 to 837 6.25 nM over spike-immobilized flow cell and the control flow cell in a running buffer composed 838 of 10 mM HEPES pH 8.0, 150 mM NaCl and 0.05% Tween 20 (HBS-T). Between each cycle, the 839 sensor chip was regenerated with 0.35 M EDTA, 50 mM NaOH and followed by 0.5 mM NiCl₂. 840 For IgG binding experiments, spike immobilization of 200 RUs was used instead to avoid mass 841 transport effect. Serial dilutions of purified IgG N3-1 were injected at concentrations ranging from 842 25 to 1.56 nM over a spike-immobilized flow cell and the control flow cell. For the SARS-CoV Tor2

S-2P binding experiments, IgG N3-1 concentrations ranging from 100 to 6.25 nM were used.
Response curves were double-reference subtracted and fit to a 1:1 binding model or
heterogeneous ligand binding model using Biacore X100 Evaluation Software (GE Healthcare).

846 Negative stain EM for spike-lgG complexes. To investigate mAb N3-1 binding to spike proteins, 847 purified SARS-CoV-2 Wuhan-Hu-1 S-HexaPro was incubated with 1.2-fold molar excess of IgG 848 N3-1 in 2 mM Tris pH 8.0, 200 mM NaCl and 0.02% NaN₃ on ice for 10 min. The spike-IgG 849 complexes were at a concentration of 0.05 mg/mL in 2 mM Tris pH 8.0, 200 mM NaCl and 0.02% 850 NaN₃ prior to deposition on a CF-400-CU grid (Electron Microscopy Sciences) that was plasma 851 cleaned for 30 sec in a Solarus 950 plasma cleaner (Gatan) with a 4:1 ratio of O₂/H₂ and stained 852 using methylamine tungstate (Nanoprobes). Grids were imaged at a magnification of 92,000X 853 (corresponding to a calibrated pixel size of 1.63 Å/pix) in a Talos F200C TEM microscope 854 equipped with a Ceta 16M detector. The CTF-estimation, particle picking and 2D classification 855 were all performed in cisTEM⁴⁴.

856 Cryo-EM sample preparation and data collection. Purified SARS-CoV-2 S (HexaPro variant) 857 at 0.2 mg/mL was incubated with 5-fold molar excess of Fab N3-1 in 2 mM Tris pH 8.0, 200 mM 858 NaCl and 0.02% NaN₃ at RT for 30 min. The sample was then deposited on plasma-cleaned 859 UltrAuFoil 1.2/1.3 grids before being blotted for 4 sec with -3 force in a Vitrobot Mark IV and 860 plunge-frozen into liquid ethane. Purified SARS-CoV-2 S with three RBDs covalently trapped in 861 the down conformations (HexaPro-RBD-down variant, S383C/D985C⁴⁵⁻⁴⁷) at 0.2 mg/mL, 862 complexed with 2-fold molar excess of Fab A7V3, was deposited on plasma-cleaned UltrAuFoil 863 1.2/1.3 grids before being blotted for 3 sec with -4 force in a Vitrobot Mark IV and plunge-frozen 864 into liquid ethane. For the HexaPro-N3-1 sample, 3,203 micrographs were collected from a single 865 grid. For the HexaPro-RBD-down-A7V3 sample, 3,636 micrographs were collected from a single 866 grid. FEI Titan Krios equipped with a K3 direct electron detector (Gatan) was used for imaging.

Data were collected at a magnification of 22,500x, corresponding to a calibrated pixel size of 1.07
Å/pix. A full description of the data collection parameters can be found in Extended Data Tables
3-4.

870 Cryo-EM data processing. Gain reference- and motion-corrected micrographs processed by 871 Warp⁴⁸ were imported into cryoSPARC v2.15.0⁴⁹, which was used to perform CTF correction, 872 micrograph curation, particle picking, and particle curation via iterative rounds of 2D classification. 873 The final global reconstructions were then obtained via ab initio reconstruction, iterative rounds 874 of heterogeneous refinement, and subsequently non-uniform homogeneous refinement of final 875 classes with C1 symmetry. For the HexaPro-RBD-down-A7V3 sample, C3 symmetry was 876 attempted in the initial refinement process. Given the low occupancy of the Fabs on a trimeric 877 spike, C1 symmetry was used for the final runs of heterogeneous and homogeneous refinement. 878 To better resolve the Fab-spike interfaces, both datasets were subjected to particle subtraction 879 and focused refinement. Finally, both global and focused maps were sharpened using 880 DeepEMhancer⁵⁰. For A7V3-NTD model building, we used an NTD from the 4A8 complexed spike 881 structure (PDB ID: 7C2L²⁵) and a homologous Fab structure (PDB ID: 6IEK) as an initial model to build into map density using UCSF ChimeraX⁵¹. For N3-1-RBDs model building, we used one 882 RBD-up and one RBD-down from S-HexaPro (PDB ID: 6XKL⁵²) and two homologous Fab 883 884 structures (PDB ID: 5BV7 and 5ITB) as an initial model to build into map density via UCSF 885 ChimeraX. Both models were built further and iteratively refined using a combination of Coot⁵³, 886 Phenix⁵⁴, and ISOLDE⁵⁵. The detailed workflows of cryo-EM data processing and data validation 887 can be found in Extended Data Figures 10, 11 and 14.

888

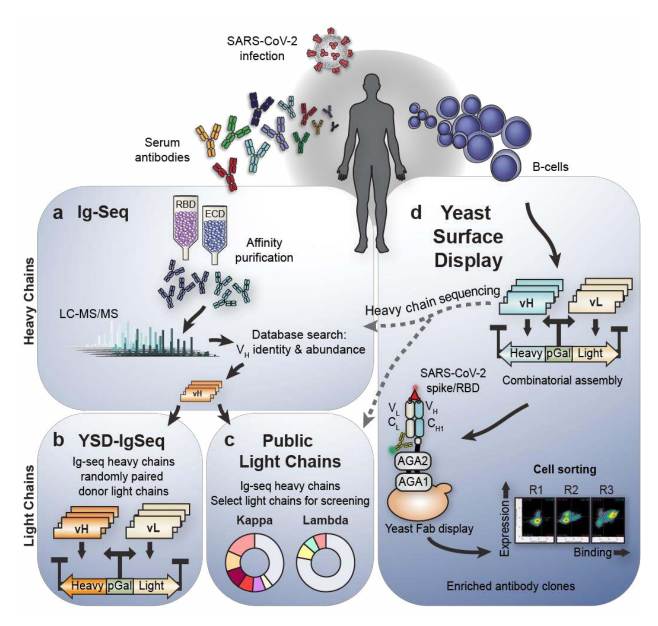
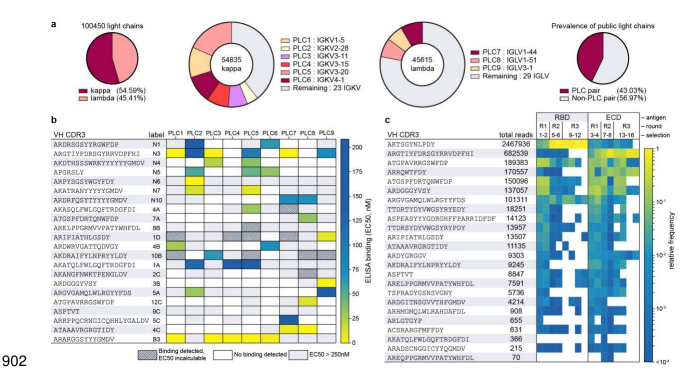


Figure 1. Overview of complementary strategies for discovering convalescent patient-derived anti SARS-CoV-2 neutralizing antibodies.

889

892 Serum antibodies and B-cells are isolated from patient blood post infection. a, IgG is antigen-enriched, digested, and 893 analyzed with tandem mass spectrometry. Searching against a database of donor B cell receptor sequences yields 894 antigen-specific heavy chain candidates. b, To find productive VH-VL pairs, the proteomically identified VH are 895 randomly combined with donor VL, generating a large library of Fab amplicons for selection in yeast. c, Additional 896 productive VH-VL are recovered by matching each proteomic VH candidate against a fixed set of nine public light 897 chains significantly overrepresented in prior repertoire studies. All combinations are screened for binding. d, Yeast 898 surface display provides an independent method to recover VH or VHVL candidates. VH and VL sequences are 899 amplified from donor B-cells, and pairs are randomly assembled into large Fab libraries and selected using 900 fluorescent cell sorting. Yeast pools are then sequenced after each selection to track individual clone dynamics round 901 over round. Promising candidates are expressed as IgG for further characterization.



903 Figure 2. Public light chain and YSD-IgSeq abundance and screening.

a, Nine PLCs emerge from analysis of 100,450 previously published paired VH-VL sequences. b, Screening with VHs
 (H-CDR3 depicted) identified by Ig-Seq and YSD against the panel of nine PLCs to determine productive VHVL
 pairings. IgG mAbs ELISA EC50s revealed that partnering VHs with PLCs can create low nM affinity binders. Grey
 boxes indicate that binding was detected but at an affinity too low to determine an EC50 in the concentration range
 tested. White boxes, no binding was detected. c, VH CDR3 read counts and relative abundance from the YSD-IgSeq
 experiment shown for MinION reads of 1.7 kb to 2.1 kb. Heatmap values are CDR3 frequencies, or read counts
 normalized within each respective sample.

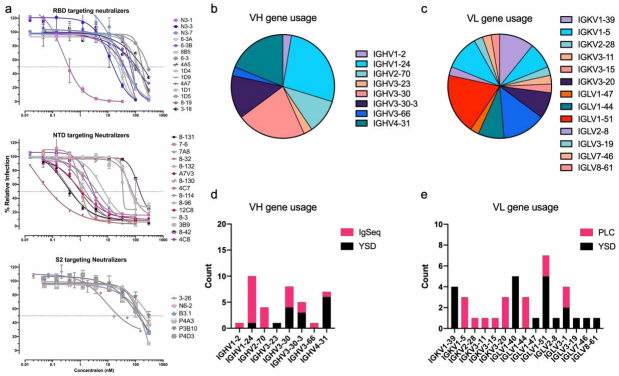


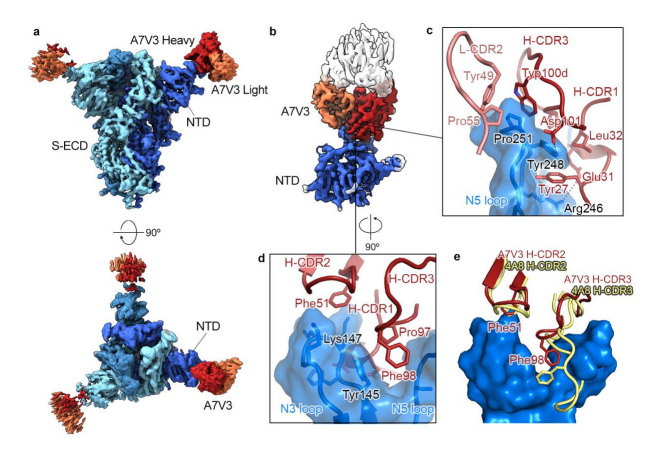
Figure 3. Non-cognate paired antibodies neutralize SARS-CoV-2 live virus.

911 912 913 914 **a**, SARS-CoV-2 WA1 live virus neutralization assays. The graphs depict the neutralizers binned by their target spike subdomain. **b**, VH gene usage of the neutralizing antibodies as a fraction of the whole. **c**, Stacked bar chart showing 915 neutralizer VH gene usage broken down by discovery method. d-e, Same analysis described in b-c but by VL gene.

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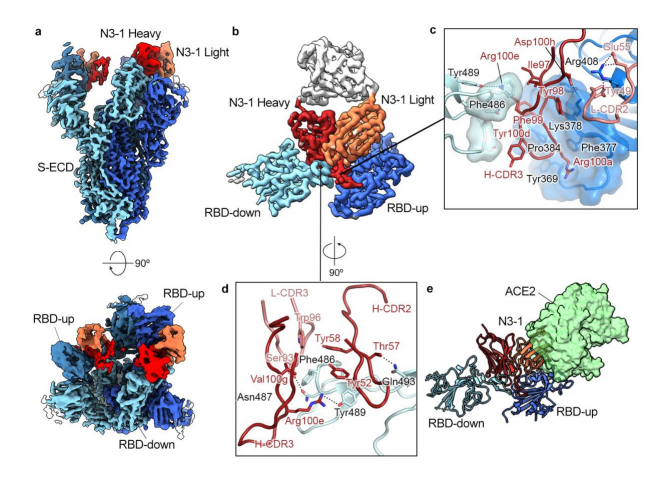


921

922 Figure 4. NTD-directed mAb A7V3 exhibits a common binding mode.

923 a, Cryo-EM structure of A7V3 bound to SARS-CoV-2 S at a global resolution of 3.0 Å. Side view and top-down views 924 of the complex are shown in the upper and the lower panel, respectively. Each protomer is depicted in steel blue, royal 925 blue and sky blue. The heavy chain of A7V3 is colored firebrick, and the light chain is colored coral. b, Focused map 926 of A7V3 bound to NTD reveals a major binding site on the N3 and N5 loops. c, The tip of the N5 loop is surrounded by 927 the hydrophobic residues from L-CDR2 and H-CDR3. Arg246 and Tyr248 contact Glu31 and Asp101 via polar 928 interactions. d, Pro97 and Phe98 in H-CDR3 insert into a groove walled by the N3 and N5 loops. Conserved Phe51 in 929 H-CDR2 forms a pi-cation interaction with Lys147 e, Superimposed structure of 4A8-NTD complex (PDB ID: 7C2L) 930 with A7V3-bound NTD. The molecular surface of the N3 and N5 loops is shown in blue. Unlike 4A8, the relatively short 931 H-CDR3 from A7V3 barely contacts the N3 loop.

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- 933



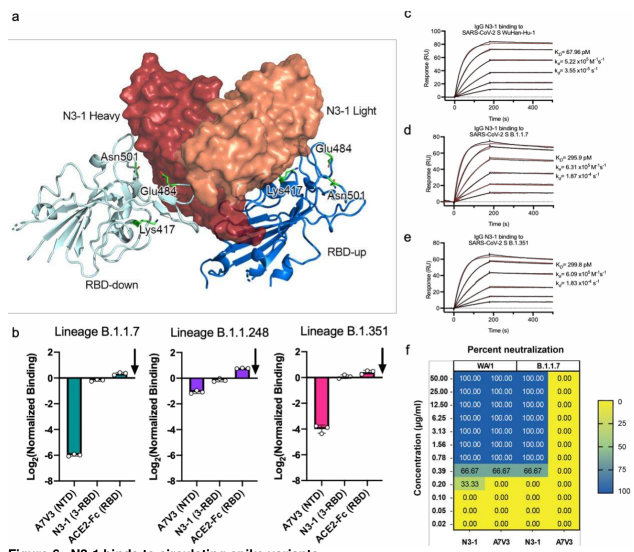
934

935 Figure 5. RBD-directed mAb N3-1 exhibits a unique binding mode by recognizing two distinct epitopes.

936 937 938 a, Cryo-EM structure of N3-1-bound SARS-CoV-2 spike at a global resolution of 2.8 Å. Side view and top-down views of the complex are shown in the upper and the lower panel, respectively. Each protomer is depicted in steel blue, royal 939 blue and sky blue. The heavy chain of N3-1 is colored firebrick, and the light chain is colored coral. b, Focused map of 940 N3-1 bound to RBDs in the up and down conformations with five CDRs involved in the binding interface. c. One face 941 of H-CDR3 contacts a conserved hydrophobic pocket (transparent royal blue surface) on RBD-up. Arg408 forms 942 multiple polar interactions with Tyr49 and Glu55. The other face of H-CDR3 contacts the ACE2-binding site on RBD-943 down. H-CDR1 contacts the epitopes on RBD-up, but it is omitted for clarity. d, The epitope on RBD-down is centered 944 on Phe486, which fits into a hydrophobic surface formed by Trp96, Tyr58, Tyr52, Arg100e and Val100g (clockwise). 945 Arg100e also forms a cation-pi interaction with Phe486 and a hydrogen bond with Tyr489. e, Superimposed crystal 946 structure of RBD-ACE2 complex (PDB ID: 6M0J) with N3-1 bound RBDs. The molecular surface of ACE2 is shown in 947 transparent pale green. The light chain of N3-1 heavily clashes with ACE2. The ACE2-binding site on RBD-down is 948 completely blocked by H-CDR2, H-CDR3 and L-CDR3 [d]. 949

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54 Figure 6. N3-1 binds to circulating spike variants.

955 956 a, Model of N3-1 bound to RBD-up and RBD-down. Mutation sites in B.1.1.7, B.1.1.248, and B.1.351 are highlighted in green. b, The B.1.1.7, B1.1.248, and B.1.351 spike variants were tested with A7V3, N3-1, and chimeric human 957 ACE2-Fc using mammalian display of spike variants and compared to SARS-CoV-2 Wuhan-Hu-1 S-HexaPro-D614G 958 (baseline arrow). c-e, Binding of IgG N3-1 to SARS-CoV-2-S WuHan-Hu-1 [c], variants B.1.1.7 [d] and B.1.351 [e] 959 were assessed by surface plasmon resonance using an NTA sensor chip. Binding data are shown as black lines. For 960 [c-e], the best fit was achieved using a 1:1 binding model and shown as red lines. f, Neutralization of B.1.1.7 and 961 WA/1 by N3-1 and A7V3, where neutralization is measured as the percent of replicates showing complete 962 neutralization at a given concentration (see methods). 963

965 Ethics statement

- 966 The acquisition of blood specimens from convalescent individuals was approved by the
- 967 University of Texas at Austin Institutional Review Board (protocol 2020-03-085; Breadth of
- serum antibody immune responses prior to, or following, patient recovery in asymptomatic and
- 969 non-severe COVID-19). Informed consent was obtained from all participants.
- 970

971 Data availability

- 972 The sequences of neutralizing antibodies have been deposited in GenBank with accession
- 973 numbers. Molecular coordinates for A7V3 and N3-1 Fab complexes with SARS-CoV-2 trimeric
- 974 spike protein have been deposited to the Protein Data Bank. Structural data are presented in
- 975 Figs. 4-5, Extended Data Tables 3-4, and Extended Data Figs. 10-12 and 14.

976

977 Author Contributions

- 978 Conceptualization: JG, CH, AH, ECG, DRB, JSM, and JDG; Methodology: JG, CH, AH, ECG,
- 979 FB, NW, KJ, AH, RR, MJJ, SLW, ZLN, JL, TSS, SVK, VK, RAH, IF, DRB, JSM, and JDG;
- 980 Investigation: JG, CH, AH, ECG, FB, NW, AA, KJ, AH, WNV, JAC, ACM, RR, MJJ, SLW, ZLN,
- JL, RAH, SVK, VK, IJF, JMM, DRB, JSM, and JDG; Data Analysis and Interpretation: JG, CH,
- AH, DRB, NW, AH, JMD, IJF, EMM, JSM, and JDG; Data Curation: JG, CH, AH, ECG, NW,
- 983 DRB, JSM, AH, JDM, JSM, and JDG; Original Draft: JG, CH, AH, ECG, DRB, JMM, JSM, VK,
- EMM, and JDG; Review & Editing: JG, CH, AH, ECG, DRB, ADE, EMM, JMM, IJF, GG, JSM,
- and JDG; Funding: JMD, IJF, GCI, VK, SVK, GG, ADE, JSM and JDG.

986

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- 1014 Experiment Station (to VK and SK); an NIH grant (R01 AI158177-01) to SK; and an NIH grant
- 1015 (R35 GM122480) to EMM. GCI expended discretionary funds from his MBS-20 account.
- 1016
- 1017

1018 Competing Interest Statement

- 1019 JL, ADE, EMM, GG, and DRB declare competing financial interests in the form of provisional
- and granted patent applications relevant to Ig-Seq. JG, CH, ECG, AH, DRB, EMM, JSM, GCI,
- 1021 ADE, GG, and JDG have filed provisional applications for the discovery of neutralizing
- 1022 antibodies. JG, DRB, ECG, AH, and JDG have filed applications for additional methods relevant
- 1023 to this work.