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Prevalent, protective, and convergent IgG recognition of SARS-CoV-2 non-RBD spike epitopes

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The molecular composition and binding epitopes of the immunoglobulin G (IgG) antibodies that circulate in blood plasma after severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) infection are unknown. Proteomic deconvolution of the IgG repertoire to the spike glycoprotein in convalescent subjects revealed that the response is directed predominantly (>80%) against epitopes residing outside the receptor binding domain (RBD). In one subject, just four IgG lineages accounted for 93.5% of the response, including an amino (N)-terminal domain (NTD)-directed antibody that was protective against lethal viral challenge. Genetic, structural, and functional characterization of a multidonor class of "public" antibodies revealed an NTD epitope that is recurrently mutated among emerging SARS-CoV-2 variants of concern. These data show that "public" NTD-directed and other non-RBD plasma antibodies are prevalent and have implications for SARS-CoV-2 protection and antibody escape.

he severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) spike ectodomain (S-ECD) folds into a multidomain architecture (1, 2) and includes the receptor binding domain (RBD), which is essential for viral infectivity, and the structurally adjacent amino (N)-terminal domain (NTD), which plays an uncertain role. Humoral immunity to the spike (S) surface glycoprotein can correlate with protection (3), and it is the primary antigenic target for most vaccines and monoclonal antibodies (mAbs). That the B cell repertoire can recognize multiple spike epitopes is supported by extensive single-cell cloning campaigns (4-9). However, the identity, abundance, and clonality of the immunoglobulin G (IgG) plasma antibody repertoire and the epitopes it may target are not known (10-12). Divergence between the two repertoires is biologically plausible (13-17), and the evidence in COVID-19 includes a paradoxical disconnect between virus-neutralizing IgG titers and RBD-specific B cell immunity (6, 11, 18, 19).

To analyze the IgG repertoire, we collected blood during early convalescence from four seroconverted study subjects (P1 to P4) who experienced mild COVID-19 disease that manifested with plasma virus-neutralization titers in the lowest quartile (P1 and P3), the second highest quartile (P2), or the highest quartile (P4) compared to a larger cohort (table S1 and fig. S1). The lineage composition and relative abundance of constituent IgG antibodies comprising the plasma response to either intact stabilized S-ECD (S-2P (1)) or RBD was determined using the Ig-Seq pipeline (13, 14, 20) that integrates analytical proteomics of affinity-purified IgG fractions with peripheral B cell antibody variable region repertoires (BCR-Seq).

IgG lineages detected by Ig-Seq in the S-ECD fraction but absent from the RBD fraction were deemed to be reactive with spike epitopes outside the RBD. In subject P3, we detected six IgG lineages that bound to S-ECD (Fig. 1A). Four of these (Lin.1 to Lin.4) accounted for 93.5% abundance of the total plasma IgG S-ECD response and exhibited extensive intralineage diversity (fig. S2) indicative of clonal expansion and selection. Notably, the top three lineages (Lin.1 to Lin.3; >85% abundance) all bound to non-RBD epitopes (S2 subunit or NTD). Bulk serology enzyme-linked immunosorbent assays (ELISAs) recapitulated the Ig-Seq result and

demonstrated similarly high levels of non-RBDbinding IgG(P > 0.05) (Fig. 1B), confirming that RBD-binding plasma antibodies constitute only a minor proportion of all spike-binding IgG in naturally infected individuals (21). In all four subjects, the detected plasma IgG repertoire to S-ECD was oligoclonal, comprising only 6 to 22 lineages, with the top-ranked lineage constituting 15 to 50% of the total abundance. On average, 84% of the anti-S-ECD plasma IgG repertoire bound to epitopes outside the RBD (Fig. 1C), a finding consistent with data from single B cell analyses (22), and the most abundant plasma IgG lineage in all donors recognized a non-RBD epitope (Figs. 1A and 2A and fig. S3).

Binding analysis of P3 mAbs CM29 to CM32 representing the most expanded clones within each of lineages Lin.1 to Lin.4 showed that CM29 (Lin.1) recognizes the S2 subunit [dissociation constant $(K_D) = 6.6$ nM], CM30 and CM31 (Lin.2 and Lin.3 with $K_D = 0.8$ and 37.7 nM, respectively) were specific for the NTD, and CM32 (Lin.4) bound the RBD (K_D = 6.0 nM), as expected from the Ig-Seq differential affinity purifications (Fig. 1A and table S2). CM30 potently neutralized authentic SARS-CoV-2 in vitro [median inhibitory concentration (IC₅₀) = $0.83 \,\mu\text{g/ml}$] and CM32 was slightly less potent (2.1 µg/ml), whereas CM29 and CM31 showed minimal neutralization activity (Fig. 1D).

We then determined the capacity of mAbs CM29 to CM32, singly and in combination, to confer prophylactic protection in vivo to virus challenge using the MA10 mouse model of SARS-CoV-2 infection (23, 24). Even though the RBD-directed mAb CM32 could neutralize authentic virus in vitro and had relatively high antibody-dependent cellular phagocytosis (ADCP) activity (fig. S4), it did not protect in vivo (fig. S5), possibly because of amino acid changes in the MA10 virus. Similarly, no protection was observed for the non-neutralizing S2-directed mAb CM29 or non-neutralizing NTD-directed mAb CM31. The neutralizing mAb CM30, derived from the top-ranking NTDtargeting IgG lineage (21% abundance), was the sole plasma antibody that conferred complete protection to MA10 viral challenge (Fig. 1, E and F, and fig. S5). Administration of a cocktail comprising the top non-RBD plasma mAbs CM29 to CM31 (>85% of the IgG plasma lineages to S-ECD; Fig. 1A) showed the most

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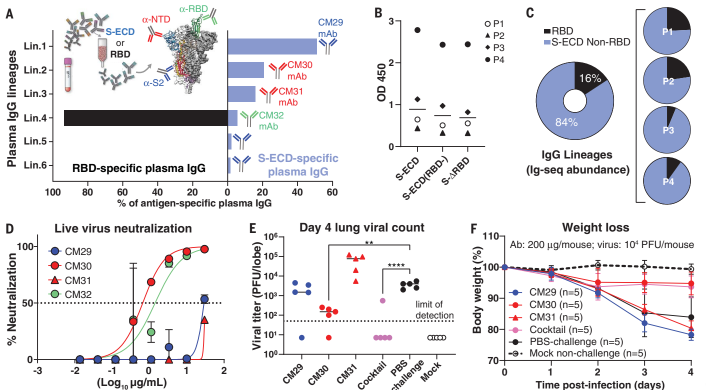


Fig. 1. Most plasma IgG antibodies bind non-RBD spike epitopes such as the NTD. (A) Affinity purification using spike S-ECD (1) or RBD for subject P3. Plasma IgG lineage identities, binding specificity, and relative abundance were mapped by means of Ig-seq proteomics (14), facilitating recombinant plasma mAb characterization; anti-RBD (green); anti-S2 (blue); and anti-NTD (red). (B) IgG ELISA binding (1:150 plasma dilution) to S-ECD alone or in the presence of RBD (50 µg/ml) [S-ECD(RBD-)] or S-ΔRBD deletion mutant. (**C**) Quantitative Ig-seq determination of anti-RBD and non-RBD IgG mAb abundance in early convalescent

plasmas across four subjects. **(D)** Authentic virus neutralization (in duplicate) of the four most abundant plasma IgGs (CM29, CM30, CM31, CM32) from plasma lineages Lin.1, Lin.2, Lin.3, and Lin.4 in subject P3. **(E** and **F)** Prophylactic protection of 12-month-old BALB/c mice (n = 5 per group) against lethal challenge with a high dose (10^4 PFU) of mouse-adapted (MA10) SARS-CoV-2. Cocktail of non-RBD mAbs ($200 \mu g$ per mouse) at 2:1:1 ratio reflecting their relative plasma abundance. **P < 0.005; ****P < 0.0001, determined by one-way analysis of variance (ANOVA) with Dunnett's multiple comparisons test.

robust protection and lung viral titers below the limit of detection in high-viral load challenge [10⁴ plaque-forming units (PFU)].

Subject P2, with ~10-fold higher neutralizing titer compared with subject P3 (fig. S1 and table S1), displayed a more polyclonal IgG response (Fig. 2A), with 12 out of 15 lineages (>80% total abundance) in the anti-S-ECD repertoire recognizing non-RBD epitopes. Conspicuously, as with P3, the most abundant S-ECD-directed plasma antibodies target the S2 subunit, with the four topmost lineages (68% total abundance) binding to S2. mAbs CM25 and CM17, representative of two NTDtargeting lineages each constituting ~2.5% of the response at day 56 (Ig-Seq Lin.6 and Lin.9) (Fig. 2A), were both encoded by unmutated or near-germline immunoglobulin G heavychain variable region 1-24 (IGHV1-24). We found an additional NTD-targeting unmutated IGHV1-24 plasma mAb (CM58) in subject P4. CM17, CM25, and CM58 bound S-ECD with similar single-digit nanomolar affinity (Fig. 2B and table S2), and all three potently neutralized SARS-CoV-2 virus, with IC₅₀ values of 0.01 to 0.81 µg/ml comparable to those of S309 anti-RBD control (25) (Fig. 2C, fig. S6, and table S2). For all three mAbs, preadministration in the MA10 mouse model resulted in significantly reduced lung viral titers after infection with 10^5 PFU (Fig. 2D; P < 0.001), resulting in 100% survival, compared with just 40% in the control group (Fig. 2E). CM17-and CM25-treated cohorts exhibited only minimal weight loss (Fig. 2F). Thus, IGHV1-24 is intrinsically suited for potent and protective targeting of the NTD.

B cell expression of IGHV1-24 in COVID-19 (\sim 5 to 8%) (5, 7, 26) is \sim 10-fold higher than in healthy individuals (0.4 to 0.8%) (27). Moreover, we could detect IGHV1-24 plasma antibodies only in S-ECD fractions (mean 3.7%) but not among anti-RBD IgGs (Fig. 3, A and B). Alignment of CM17, CM25, and CM58 with four neutralizing IGHV1-24 anti-NTD mAbs cloned from peripheral B cells [4A8 (4), 1-68 (5), 1-87 (5), COVA2-37 (7)] and an additional antibody [COV2-2199 (8)] identified a class of convergent heavy-chain variable ($V_{\rm H}$) immune receptor sequences (Fig. 3C). In

all cases, three glutamate (Glu) residues (Glu³⁶ Glu⁵⁹, and Glu⁸⁰) located in complementaritydetermining region-H1 (CDR-H1), CDR-H2, and framework H3 (FWR-H3), respectively, as well as a phenylalanine (Phe) residue (Phe⁵⁶) in CDR-H2, were invariably unmutated and are specific to the electronegative IGHV1-24 [isoelectric point (pI) = 4.6]. The convergent V_H genes paired promiscuously with six distinct light-chain variable (V_L) genes, yet CDR-H3 peptide lengths were restricted (14 or 21 amino acids) (table S3). A "checkerboard" bindingcompetition experiment (Fig. 3D) indicated the presence of at least two epitope clusters on the NTD, including one targeted by all of the tested IGHV1-24 mAbs (4A8, CM25, CM17, CM58, and 1-68) and the IGHV3-11 mAb CM30. Another NTD epitope was identified by CM31 (IGHV2-5, 6.4% mutation), which overlapped with CM30 (IGHV3-11; 3.1% mutation), CM58, and 1-68 but did not compete with the other three IGHV1-24 NTD mAbs.

To better understand the IGHV1-24 interactions with the spike NTD, we determined a cryo-electron microscopy (EM) structure of

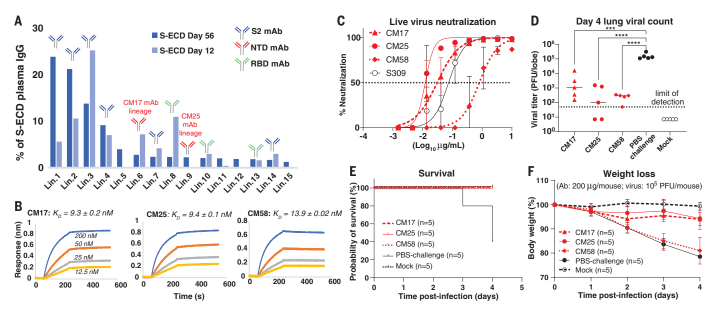


Fig. 2. Protective spike NTD-targeting antibodies are prevalent in COVID-19 convalescent plasma. (**A**) Temporal Ig-seq dynamics of the anti-S-ECD IgG repertoire at days 12 and 56 after symptom onset. (**B**) Biolayer interferometry (BLI) sensorgrams to S-ECD ligand of anti-NTD mAbs CM17, CM25 (subject P2), and CM58 (subject P4). (**C**) In vitro live virus neutralization (performed in duplicate). (**D** to **F**) In vivo prophylactic protection of 12-month-old BALB/c mice (*n* = 5 per group) against high-dose intranasal challenge (10⁵ PFU) of mouse-adapted (MA10) SARS-CoV-2. ***P < 0.0007; ****P < 0.0001, determined by one-way ANOVA with Dunnett's multiple comparisons test.

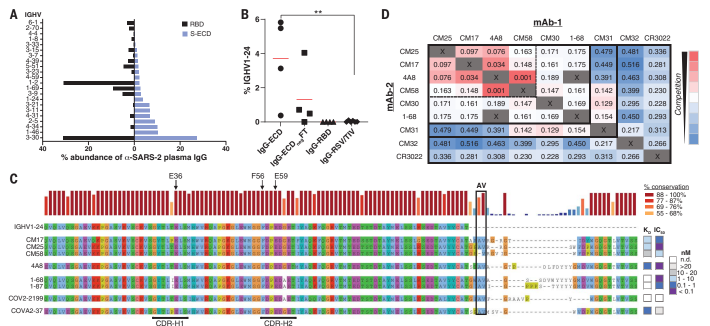


Fig. 3. Genetic basis of a shared, or public, class of IGHV1-24 plasma antibodies targeting the spike NTD. (A) IGHV usage of plasma antibodies in all subjects (n=4). **(B)** Comparative IGHV1-24 usage of anti-S-ECD (IgG-ECD) and anti-RBD (IgG-RBD) plasma antibodies, or in depleted S-ECD affinity column flow-through (IgG-ECD_{neg}FT) in all subjects (n=4). IgG-RSV/TIV: IgG specific to respiratory syncytial virus (RSV) or trivalent influenza vaccine hemagglutinin HA1 (TIV) in healthy controls after vaccination (n=6). **P<0.01, determined by Mann–Whitney U test. **(C)** Sequence alignment of IGHV1-24 neutralizing anti-NTD IgGs from plasma (CM17, CM25, and CM58) or from peripheral B cells [4A8 (4), 1-68 and 1-87 from a subject with ARDS (5), COV2-2199 (13), and COVA2-37 (mild disease subject)] (7). Arrows point to

specific IGHV1-24 residues. Heatmap shows recombinant mAb affinity (K_D) and live-virus neutralization (IC_{50}) for individual antibodies. (**D**) Competitive BLI binding assay ("checkerboard competition") of NTD-binding mAbs found in this study (CM17, CM25, CM58, CM30, and CM31) and others (4A8 and 1-68). RBD-binding mAbs CM32 and CR3022 included for comparison. Numbers refer to the shift, in nanometers, after second mAb binding to the preformed mAb–NTD complex. Dashed box drawn to highlight strong competition (<0.1 nm shift) among 4A8 and three IGHV1-24 mAbs examined in this study. Abbreviations for the amino acid residues are as follows: A, Ala; C, Cys; D, Asp; E, Glu; F, Phe; G, Gly; H, His; I, Ile; K, Lys; L, Leu; M, Met; N, Asn; P, Pro; Q, Gln; R, Arg; S, Ser; T, Thr; V, Val; W, Trp; and Y, Tyr.

CM25 Fabs bound to trimeric S-ECD (Fig. 4A and figs. S7 and S8). Focused refinement of the CM25-NTD interface resulted in a 3.5-Šreconstruction that revealed a heavy-chain-dominant mode of binding, with substantial contacts mediated by interactions among the three CDRs and the N3 and N5 loops of the NTD (Fig. 4B). The light chain contributes only 11% (86 Ų) of the total CM25 binding

interface, mainly through a stacked hydrophobic interaction between CDR-L2 $\rm Tyr^{55}$ and $\rm Pro^{251}$ within the N5 loop. Distinctive germline IGHV1-24 residues contribute 20% (149 Ų) of the total binding interface. CDR-H1 interacts extensively through hydrogen bonds and contacts between hydrophobic residues, including a salt bridge formed between the conserved Glu 36 residue and the N5 loop residue $\rm Arg^{246}$

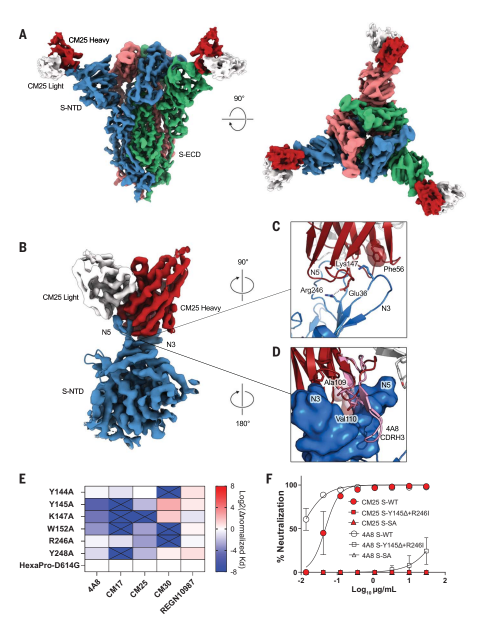


Fig. 4. Structural basis of public IGHV1-24 plasma antibodies, NTD mutations, and antibody escape. (A) Side and top views of the structure of CM25 Fab bound to S-ECD shown as cryo-EM density. (B) Focused refinement density revealing a V_H -dominant mode of binding, with substantial contacts mediated by interactions among the three CDRs and the N3 and N5 loops of the NTD. (C) CDR-H1 interaction includes a salt bridge formed between the distinctively encoded Glu^{36} residue and the N5 loop residue Arg^{246} ; the distinctive Phe^{56} residue in CDR-H2 forms a pi-cation interaction with Lys¹⁴⁷ in the N3 loop. (D) The AV dipeptide interaction with the N3 and N5 loops of the NTD is structurally conserved between mAbs CM25 (red) and 4A8 (pink). (E) Normalized shift (log_2) in binding K_D , as measured by differential BLI affinities for single Ala mutants and parental D614G spike protein. (F) Authentic virus neutralization of CM25 and 4A8 against wild-type (WT), double S-N3/N5 loop mutants, and South Africa (SA) B.1.351 viral variant.

(Fig. 4C). The common IGHV1-24 Phe^{56} residue in CDR-H2 forms a pi-cation interaction with Lys1⁴⁷ in the N3 loop (Fig. 4C). CM25 contains a 14-amino acid CDR-H3 loop that contributes 35% (261 Å²) of the total interface, including the AV aliphatic motif found in all but one of the convergent IGHV1-24 NTD-binding mAbs. Ala¹⁰⁹ and Val¹¹⁰ are buried at the interface in a binding pocket framed by the N3 and N5 loops. A comparison of CM25 with an extant structure of an IGHV1-24 NTD-binding antibody isolated by B cell cloning, 4A8 (4), revealed that the AV dipeptide interaction is structurally conserved, and the 21-amino acid CDR-H3 of 4A8 extends along the outside of the NTD, contributing three additional contacts and 46% $(415 \, \text{Å}^2)$ of the total binding interface (Fig. 4D). Both structures show extensive contacts between the heavy chain of the Fabs and the NTD N3 and N5 loops. The $\mathrm{Glu}^{36}\text{-Arg}^{246}$ salt bridge and an identical CDR-H2 contact between Phe⁵⁶ and Lys¹⁴⁷ are conserved in the 4A8-NTD interface.

SARS-CoV-2 variants of concern contain mutations in the NTD N3 and N5 loops, including Y144/Y145Δ and K147E (UK lineage B.1.1.7), W152C (California B.1.429), and $242-244\Delta$ or R246I (South Africa B.1.351). Alanine substitutions at several of these positions ablated binding or reduced affinity more than fivefold by public IGHV1-24 antibodies, as exemplified by 4A8, CM17, and CM25 (Fig. 4E and fig. S9), a result consistent with the CM25-NTD and 4A8-NTD structures. Additionally, we confirmed that an engineered N3-N5 double mutant and native B.1.351 (28) both evade neutralization by mAbs CM25 and 4A8 (Fig. 4F). Thus, mutations in SARS-CoV-2 variants confer escape from public neutralizing anti-NTD antibodies.

In conclusion, we find that the convalescent plasma IgG response to SARS-CoV-2 is oligoclonal and directed overwhelmingly toward non-RBD epitopes in the S-ECD. This includes public, near-germline, and potently neutralizing antibodies against the NTD. The extent to which public anti-NTD antibodies contribute to protection is likely related to their relative concentrations in plasma, which can be dominant in some individuals. Our finding that mutations present in circulating SARS-CoV-2 variants can impair or ablate binding and neutralization by public anti-NTD antibodies may constitute a mechanism of viral escape in a subset of the population. Numerous other NTD mutations-which overlap with the structural epitope recognized by the public IGHV1-24 antibody class-have been described in additional circulating variants, in laboratory escape mutants, and in immunocompromised patients (12, 29-33).

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Writing: Original Draft, W.N.V., N.J.V., J.J.L., and G.C.I.; Writing: Review & Editing: W.N.V., Y.J.H., N.J.V., B.L.I., R.S.B., J.S.M., G.G., J.J.L., and G.C.I.; Funding: J.D.G., R.S.B., J.S.M., G.G., and G.C.I. Competing interests: A patent application submitted by The University of Texas Board of Regents is pending for anti-SARS-CoV-2 monoclonal antibodies described in the manuscript (W.N.V., J.D.G., J.S.M., G.G., J.J.L., and G.C.I.). Data and materials availability: FASTQ VH and VH:VL sequence files have been deposited in the NCBI Sequence Read Archive with accession numbers PRJNA729513. The monoclonal antibodies have been deposited in GenBank (www.ncbi.nlm.nih.gov/genbank/) with accession numbers MZ049539 to MZ049552. Coordinates for the CM25 Fab in complex with trimeric spike ectodomain have been deposited to the Protein Data Bank as PDBID:7M8J. Cryo-EM maps have been deposited to the Electron Microscopy Data Bank under accession code EMD-23717. These structural data are presented in Fig. 4, table S4, and figs. S7 and S8. All other data are available in the main text or the supplementary materials. This work is licensed under a Creative Commons Attribution 4.0 International (CC BY 4.0) license, which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited. To view a copy of this license, visit https://creativecommons.org/licenses/by/4.0/. This license does not apply to figures/photos/artwork or other content included in the article that is credited to a third party; obtain authorization from the rights holder before using such material.

SUPPLEMENTARY MATERIALS

science.sciencemag.org/content/372/6546/1108/suppl/DC1 Materials and Methods Figs. S1 to S9 Tables S1 to S5 References (34–54) MDAR Reproducibility Checklist

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Prevalent, protective, and convergent IgG recognition of SARS-CoV-2 non-RBD spike epitopes

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A public anti-COVID antibody repertoire

Most analyses of the antibody responses induced by severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) infection have focused on antibodies cloned from memory B cells. This approach has led researchers to conclude that neutralizing antibodies (nAbs) primarily target the receptor-binding domain (RBD) of the virus's spike protein. Voss *et al.* took a different approach, using proteomic deconvolution of the serum immunoglobulin G antibody repertoire from four COVID-19 convalescent patients. They found that the nAb response was largely directed against epitopes such as the N-terminal domain (NTD), which lie outside the RBD. Several of these nAbs were shared among donors and targeted an NTD epitope that is frequently mutated by variants of concern.

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