

Anti- SARS-CoV-2 Receptor Binding Domain Antibody Evolution after mRNA Vaccination

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Summary

Severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) infection produces B-cell responses that continue to evolve for at least one year. During that time, memory B cells express increasingly broad and potent antibodies that are resistant to mutations found in variants of concern¹. As a result, vaccination of coronavirus disease 2019 (COVID-19) convalescent individuals with currently available mRNA vaccines produces high levels of plasma neutralizing activity against all variants tested^{1,2}. Here, we examine memory B cell evolution 5 months after vaccination with either Moderna (mRNA-1273) or Pfizer-BioNTech (BNT162b2) mRNA vaccines in a cohort of SARS-CoV-2 naïve individuals. Between prime and boost, memory B cells produce antibodies that evolve increased neutralizing activity, but there is no further increase in potency or breadth thereafter. Instead, memory B cells that emerge 5 months after vaccination of naïve individuals express antibodies that are similar to those that dominate the initial response. While individual memory antibodies selected over time by natural infection have greater potency and breadth than antibodies elicited by vaccination, the overall neutralizing potency of plasma is greater following vaccination. These results suggest that boosting vaccinated individuals with currently available mRNA vaccines will increase plasma neutralizing activity but may not produce antibodies with breadth equivalent to those obtained by vaccinating convalescent individuals.

Between January 21 and July 20, 2021, we recruited 32 volunteers with no history of prior SARS-CoV-2 infection receiving either Moderna (mRNA-1273; n=8) or Pfizer-BioNTech (BNT162b2; n=24) mRNA vaccines for sequential blood donation. Matched samples were

obtained at 2 or 3 time points. Individuals indicated as “prime” were sampled an average of 2.5 weeks after receiving their first vaccine dose. Individuals who completed their vaccination regimen were sampled after an average of 1.3 months after the boost (median=35.5 days) which is not statistically different from the 1.3 month sampling in our naturally infected cohort³ (median=38.5 days, $p=0.21$). Individuals sampled at 1.3 months were sampled again approximately 5 months after the second vaccine dose. The volunteers ranged in age from 23-78 years (median=34.5 years), 53% were male and 47% female (for details see Methods and Supplementary Tables 1 and 2).

Plasma binding and neutralization assays

Plasma IgM, IgG, and IgA responses to SARS-CoV-2 receptor binding domain (RBD) were measured by enzyme linked immunosorbent assay (ELISA)³. As reported by others^{2,4-6} there was a significant increase in IgG reactivity to RBD between prime and boost ($p<0.0001$, Fig. 1a). IgM and IgA titers were lower than IgG titers and remained low after the second vaccine dose (Extended data Fig. 1a and b). The magnitude of the response was inversely correlated with age after the prime ($r=-0.54$, $p=0.005$), but in this limited sample set the age difference was no longer significant at 1.3 or 5 months after the second vaccine dose (Fig. 1b, Extended data Fig. 1c). Between 1.3 and 5 months after the boost, anti-RBD titers of IgG and IgA decreased significantly. IgG titers decreased by an average of 4.3-fold (range: 1.7- to 10.2-fold) and the loss of activity was directly correlated to the time after vaccination ($p<0.0001$, Fig. 1a and c and Extended data Fig. 1a and b).

Neutralizing activity was measured using HIV-1 pseudotyped with the SARS-CoV-2 spike^{1,3,7,8}. Naïve individuals showed variable responses to the initial vaccine dose with a geometric mean half-maximal neutralizing titer (NT₅₀) of 171 (Fig. 1d, Supplementary Table 2). The magnitude of the neutralizing responses to the initial vaccine dose in naïve volunteers was inversely correlated with age ($r=-0.39$, $p=0.05$, Fig. 1e). Both binding and neutralizing responses to the second vaccine dose were correlated to the prime ($r=0.46$, $p=0.02$, Extended data Fig. 1d; $r=0.54$, $p=0.003$, Extended data Fig. 1e) and produced a nearly 12-fold increase in the geometric mean neutralizing response that was similar in males and females and eliminated the age-related difference in neutralizing activity in the individuals in this cohort (Fig. 1d, Extended data Fig. 1f and Fig. 1e and Extended data Fig. 1g). 1.3 and 5 months after the boost naïve vaccinees had 4.9- and 3.6 fold higher neutralizing titers than a cohort of infected individuals measured 1.3³- and 6.2⁷-months after symptom onset, respectively ($p<0.0001$, Fig. 1d). Neutralizing responses were directly correlated to IgG anti-RBD titers ($r=0.96$, $p<0.0001$, Fig. 1f). Thus, the data obtained from this cohort agree with prior observations showing a significant increase in plasma neutralizing activity that are correlated with improved vaccine efficacy in naïve individuals that receive the second dose of mRNA vaccine^{2,6,9,10} and higher neutralizing titers in fully vaccinated than infected individuals^{2,6}.

The 28 individuals assayed 5 months after vaccination had a 7.1-fold decrease in geometric mean neutralizing titer from their 1.3-month measurement ($p<0.0001$, Fig. 1d), with a range of 1.4- to 27-fold. Neutralizing activity was inversely correlated with the time from vaccination ($r=-0.82$, $p<0.0001$, Fig. 1g), and directly correlated to IgG anti-RBD binding titers when assessed 5 months after vaccination (Extended data Fig. 1h). As reported by others¹¹, the ratio of binding to

neutralizing serum titers was significantly higher in vaccinated than convalescent individuals at the 1.3-month time point ($p < 0.0001$, Fig. 1h). However, the difference was no longer apparent at the later time point (Fig. 1h).

We and others showed that the neutralizing responses elicited by mRNA vaccination are more potent against the original Wuhan Hu-1 strain than for some of the currently circulating variants of concern^{2,12-14}. To confirm these observations, we measured the neutralizing activity of 15 paired plasmas from naive individuals 1.3 and 5 months after the second vaccine dose against B.1.1.7 (alpha variant), B.1.351 (beta variant), B.1.526 (iota variant), P.1 (gamma variant) and B.1.617.2 (delta variant). Consistent with previous reports^{13,15-17} the neutralizing activity against the variants was lower than against the original Wuhan Hu-1 strain (Fig. 1i, Supplementary Table 3). Initial geometric mean neutralizing titers at 1.3 months against B.1.351, B.1.1.7, B.1.526, P.1 and B.1.617.2 were 5.7, 1.8, 1.1, 1.4 and 2.7-fold lower than against Wuhan-Hu respectively (Fig. 1i). In the months following vaccination there was a decrease in neutralizing activity against Wuhan Hu-1 (R683G) and all the variants with geometric mean neutralizing titers for WT, B.1.351, B.1.1.7, B.1.526, P.1 and B.1.617.2 decreasing by 2.9-, 1.8-, 2.3-, 2.9-, 2.4- and 2.6-fold, respectively (Fig. 1i and Supplementary Table 3).

Monoclonal Antibodies

Circulating antibodies produced by plasma cells can prevent infection if present at sufficiently high concentrations at the time of exposure. In contrast, the memory B cell compartment contains long lived antigen-specific B cells that mediate rapid recall responses that contribute to long term protection¹⁸. To examine the nature of the memory compartment elicited by one or two

mRNA vaccine doses and its evolution after 5 months we used flow cytometry to enumerate B cells expressing receptors that bind to Wuhan Hu-1 (wild type, WT) and the B.1.351 K417N/E484K/N501Y variant RBDs (Fig. 2a and b, and Extended data Fig. 2). Although neutralizing antibodies develop to other parts of the spike (S) protein we focused on RBD because it is the dominant target of the memory antibody neutralizing response^{19,20}. Wuhan-Hu RBD-specific memory B cells developed after the prime in all volunteers examined and their numbers increased for up to 5 months after vaccination (Fig. 2a). Memory B cells binding to the B.1.351 K417N/E484K/N501Y variant RBD were detectable but in lower numbers than wild type RBD-binding B cells in all samples examined (Fig. 2b). Whereas IgG memory cells increased after the boost, IgM-expressing memory B cells that made up 23% of the memory compartment after the prime were nearly absent after boosting (Fig. 2c). Finally, circulating RBD-specific plasmablasts were readily detected after the prime but were infrequent after the boost (Fig. 2d, and Extended data Fig. 2d).

The memory compartment continues to evolve up to one year after natural infection with selective enrichment of cells producing broad and potent neutralizing antibodies¹. To determine how the memory compartment evolves after vaccination, we obtained 2328 paired antibody sequences from 11 individuals sampled at the time points described above (Fig. 2e and f, Extended Data Fig 3, Supplementary Table 4). As expected *IGHV3-30* and *IGHV3-53* were over-represented after the first and second vaccine dose and remained over-represented 5 months after vaccination²¹⁻²³ (Extended data Fig. 4).

All individuals examined showed expanded clones of memory B cells that expressed closely related *IGHV* and *IGHL* genes (Fig. 2e and f, Extended data Fig. 4). Paired prime and 1.3 month post boost samples showed expanded clones of memory B cells some of which were shared across plasmablasts, IgM and IgG prime, and IgG boost memory cells (Extended data Fig. 3 and 5). Thus, the cell fate decision controlling the germinal center versus plasmablast decision is not entirely affinity dependent since cells with the same initial affinity can enter both compartments to produce clonal relatives²⁴.

The relative fraction of memory cells found in expanded clones varied between prime and boost and between individuals and decreased over time (Fig. 2e-g). Overall, clones represented 30%, 21%, and 9.7% of all sequences after prime, 1.3- and 5-month time points respectively (Fig. 2g). Nevertheless, clones of memory B cells continued to evolve for up to 5 months in vaccinated individuals as evidenced by the appearance of unique clones. Notably, unique clones appearing after 1.3 and 5 months represent a greater or equal fraction of the total memory B cell pool than the persisting clones (Fig. 2e-f, 16% vs 9.6% and 5.1% vs 4.7%, respectively, Extended data Fig. 3b). Finally, memory B cells emerging after the boost showed significantly higher levels of somatic mutations than plasmablasts or memory B cells isolated after the prime, and they continue to accumulate mutations up to 5 months post-boost (Fig. 2h, and Extended data Fig. 3d). In conclusion the memory B cell compartment continues to evolve for up to 5 months after mRNA vaccination.

Neutralizing Activity of Monoclonal Antibodies

We performed ELISAs to confirm that the antibodies isolated from memory B cells bind to RBD (Extended data Fig. 6). 458 antibodies were tested by ELISA including: 88 isolated after the first vaccine dose; 210 isolated after the boost; and 160 isolated from individuals that had been fully vaccinated 5 months earlier. Among the 458 antibodies tested 430 (94%) bound to the Wuhan Hu-1 RBD indicating that the method used to isolate RBD-specific memory B cells was highly efficient (Supplementary Table 5-6). The geometric mean ELISA half-maximal concentration (EC_{50}) of the antibodies obtained after prime, and 1.3- and 5-months after the second dose was 3.5, 2.9 and 2.7 ng/ml respectively, suggesting no major change in binding over time after vaccination (Extended data Fig. 6 and Supplementary Table 5-6).

430 RBD-binding antibodies were tested for neutralizing activity using HIV-1 pseudotyped with the SARS-CoV-2 spike^{3,8}. The geometric mean half-maximal inhibitory concentration (IC_{50}) of the RBD-specific memory antibodies improved from 376 ng/ml to 153 ng/ml between the first and second vaccine dose ($p=0.0005$, Fig. 3a). The improvement was reflected in all clones (IC_{50} 377 vs. 171 ng/ml, $p=0.01$ Fig. 3b), persisting clones (IC_{50} 311 vs. 168, Fig. 3c, Supplementary Table 6), unique clones (IC_{50} 418 vs. 165 ng/ml, $p=0.03$ Fig. 3d), and single antibodies (IC_{50} 374 vs. 136 ng/ml, Fig. 3e). The increase in neutralizing activity between the first and second vaccine dose was associated with a decrease in the percentage of non-neutralizing antibodies (defined as $IC_{50} > 1000$ ng/ml) and increased representation of neutralizing antibodies ($p=0.003$, Fig. 3a). In conclusion, memory B cells recruited after the second dose account for most of the improvement in neutralizing activity in this compartment between the 2 vaccine doses. Thus, in addition to the quantitative improvement in serum neutralizing activity there is also an improvement in the neutralizing activity of the antibodies expressed in the memory compartment after boosting.

In contrast, there was no significant improvement in neutralizing activity of the monoclonal antibodies obtained between 1.3 and 5 months after vaccination ($p>0.99$, Fig. 3a). Although there was some improvement among B cell clones, which was accounted for by the small minority of persisting clones, neither was significant ($p=0.58$ and 0.46 , Fig. 3b-e, Supplementary table 6). In contrast, memory antibodies obtained from convalescent individuals showed improved neutralizing activity between 1.3³ and 6.2 months⁷ with IC_{50} of 171 ng/ml to 116 ng/ml (Fig. 3a), which improved further after 1 year¹. This improvement was due to increased neutralizing activity among persisting clones ($p=0.003$, Fig. 3c).

Affinity, Epitopes and Neutralization Breadth

To examine affinity maturation after vaccination, we performed biolayer interferometry (BLI) experiments using the Wuhan Hu-1 RBD³. 147 antibodies were assayed, 30 obtained after the prime, 74 1.3-months after boosting, and 43 5-months after the boost. Geometric mean IC_{50} s were comparable for the antibodies obtained from the 1.3- and 5-month time points (Extended data Fig. 7a). Overall, there was a 3- and 7.5 fold increase in affinity between the antibodies obtained between the first 2, and second 2 time points respectively (Fig. 4a). After 5 months the affinity of the antibodies obtained from vaccinated individuals was similar to antibodies obtained from naturally infected volunteers (Fig 4a). However, there was no correlation between affinity and neutralizing activity of the antibodies tested at any of the 3 time points (Extended Data Fig. 7b).

We also compared the affinities of pairs of antibodies obtained from persisting clones between 1.3 and 5 months after vaccination. Persisting clones obtained at 1.3 and 5 months from vaccinated individuals showed a median 4.5-fold increase in affinity ($p < 0.0001$, Fig. 4b). In contrast, a comparable group of persisting clonal antibodies obtained from convalescent individuals 1.3 and 6.2 months after infection showed a median 11.2-fold increase in affinity ($p = 0.002$, Fig. 4b).

To determine whether the epitopes targeted by the monoclonal antibodies were changing over time, we performed BLI experiments in which a preformed antibody-RBD complex was exposed to a second monoclonal targeting one of 4 classes of structurally defined epitopes^{1,3} (see schematic in Extended data Fig. 8a). There was no significant change in the distribution of targeted epitopes among 52 randomly selected antibodies with comparable neutralizing activity obtained from the 1.3- and 5-month time points (Extended Data Fig. 8b and c, and Extended Data Fig. 9).

In addition to the increase in potency, the neutralizing breadth of memory antibodies obtained from persisting clones from convalescent individuals increases with time after infection^{1,7,25}. To determine whether there is a similar increase in breadth with time after vaccination, we selected 20 random antibodies from the prime or 1.3 months after boost, with representative levels of activity against the original Wuhan Hu-1 strain, and measured their neutralization potency against a panel of pseudotypes encoding RBD mutations which were selected for resistance to different RBD antibody classes and/or are associated with circulating variants of concern (Extended data. Fig. 10). There was little change in breadth between prime and 1.3 months after

boost, with only a small increase in resistance to K417N and A475V substitutions (Extended data Fig. 10, Supplementary Table 7).

In addition, we assayed 19 pairs of neutralizing antibodies expressed by persisting clones obtained 1.3 and 5 months after vaccination against the same mutant pseudotype viruses (Fig. 4c and Supplementary Table 8). They were compared to 7 previously reported²⁵, plus 9 additional pairs of antibodies obtained from convalescent individuals at 1.3- and 6.2-month time points (Fig. 4d and Supplementary Table 8). Whereas only 36 of 190 (19%) of the vaccinee antibody-mutant combinations showed improved potency, 95 of the 160 (59%) convalescent pairs did so ($p < 0.0001$, Fig. 4c, d and e). Moreover, only 4 of the 19 (21%) vaccine antibody pairs showed improved potency against pseudotypes carrying B.1.617.2 (delta variant)-specific RBD amino acid substitutions (L452R/T478K), while 11 out of 16 (69%) of the convalescent antibody pairs showed improved activity against this virus ($p = 0.007$, Fig. 4c, d and e). We conclude that there is less increase in breadth in the months after mRNA vaccination than in a similar interval in naturally infected individuals.

Circulating antibodies are produced by an initial burst of short-lived plasmablasts²⁶⁻²⁸, and maintained by plasma cells with variable longevity²⁹⁻³². SARS-CoV-2 infection or mRNA vaccination produces an early peak antibody response that decreases by 5-10-fold after 5 months^{7,33-37}. Notably, neutralization titres elicited by vaccination exceed those of COVID-19 recovered individuals at all comparable time points assayed. Nevertheless, neutralizing potency against variants is significantly lower than against Wuhan Hu-1, with up to 5-10-fold reduced activity against the B.1.351 variant^{5,6,13,14,38}. Taken together with the overall decay in

neutralizing activity there can be 1-2 orders of magnitude decrease in serum neutralizing activity after 5 or 6 months against variants when compared to the peak of neutralizing activity against Wuhan Hu-1. Thus, antibody mediated protection against variants is expected to wane significantly over a period of months, consistent with reports of reinfection in convalescent individuals and breakthrough infection by variants in fully vaccinated individuals³⁹⁻⁴².

In contrast to circulating antibodies, memory B cells are responsible for rapid recall responses⁴³⁻⁴⁶, and the number of cells in this compartment is relatively stable over the first 5-6 months after mRNA vaccination or natural infection^{7,47}. In both cases memory B cells continue to evolve as evidenced by increasing levels of somatic mutation and emergence of unique clones.

The memory response would be expected to protect individuals that suffer breakthrough infection from developing serious disease. Both natural infection and mRNA vaccination produce memory antibodies that evolve increased affinity. However, vaccine-elicited memory monoclonal antibodies show more modest neutralizing potency and breadth than those that developed after natural infection^{1,7}. Notably, the difference between the memory compartment that develops in response to natural infection vs mRNA vaccination reported above is consistent with the higher level of protection from variants conferred by natural infection⁴².

There are innumerable differences between natural infection and mRNA vaccination that could account for the differences in antibody evolution over time. These include but are not limited to: 1. Route of antigen delivery, respiratory tract vs. intra-muscular injection^{48,49}; 2. The physical nature of the antigen, intact virus vs. conformationally stabilized prefusion S protein⁵⁰; 3.

Antigen persistence, weeks in the case of natural infection⁷ vs. hours to days for mRNA⁵¹. Each of these could impact on B cell evolution and selection directly, and indirectly through differential T cell recruitment.

The increase in potency and breadth in the memory compartment that develops after natural infection accounts for the exceptional responses to Wuhan Hu-1 and its variants that convalescent individuals develop when boosted with mRNA vaccines^{1,5}. The expanded memory B cell compartment in mRNA vaccinees should also produce high titers of neutralizing antibodies when vaccinees are boosted or when they are re-exposed to the virus⁵². Boosting vaccinated individuals with currently available mRNA vaccines should produce strong responses that mirror or exceed their initial vaccine responses to Wuhan-Hu but with similarly decreased coverage against variants. Whether an additional boost with Wuhan-Hu-based or variant vaccines or re-infection will also elicit development of memory B cells expressing antibodies showing increased breadth remains to be determined. Finally, timing an additional boost for optimal responses depends on whether the objective is to prevent infection or disease⁵³. Given the current rapid emergence of SARS-CoV-2 variants, boosting to prevent infection would likely be needed on a time scale of months. The optimal timing for boosting to prevent serious disease will depend on the stability and further evolution of the memory B cell compartment.

FIGURES

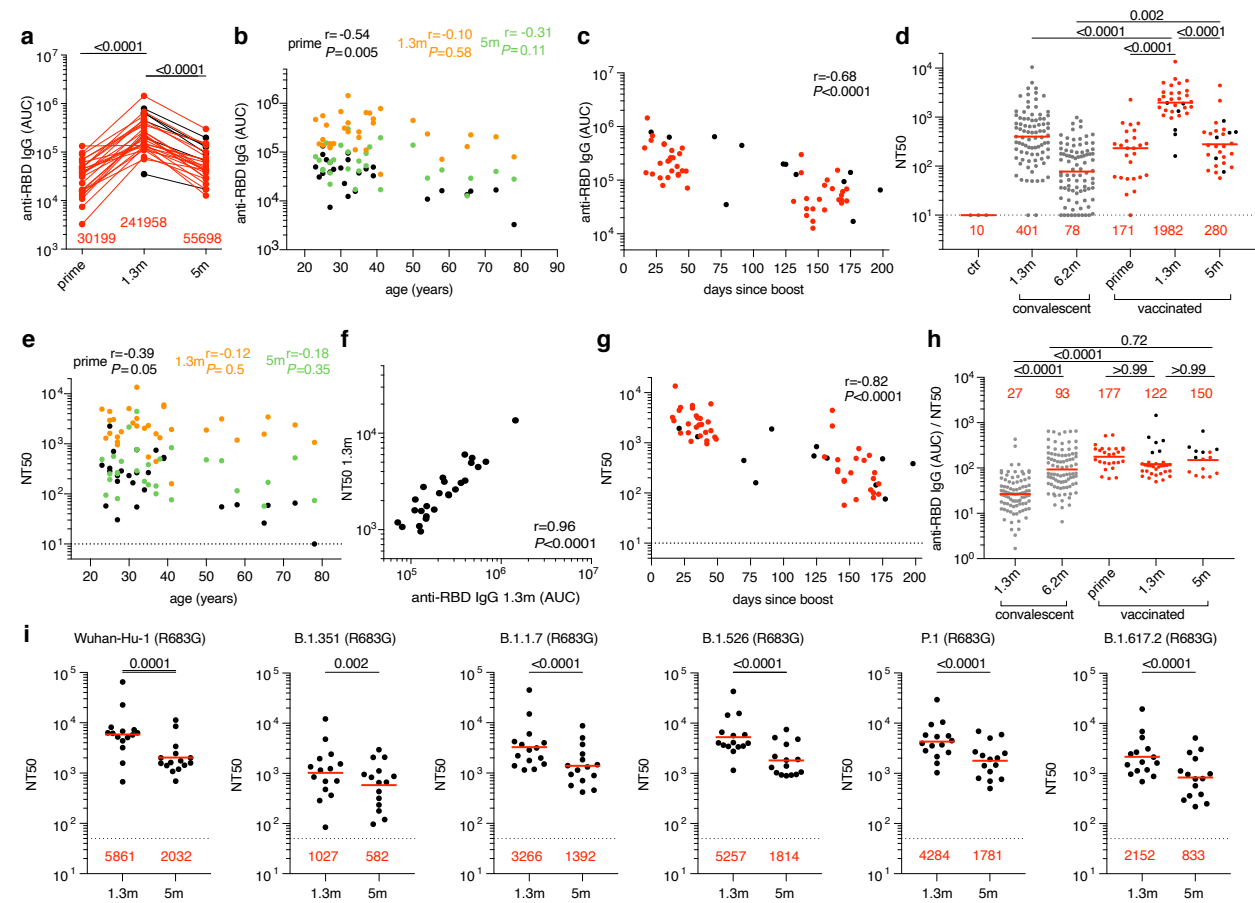


Fig. 1: Plasma ELISAs and neutralizing activity.

a, Graph shows area under the curve (AUC, Y-axis) for plasma IgG antibody binding to SARS-CoV-2 RBD after prime, and 1.3 months (m) and 5 months (m) post-second vaccination for n=32 paired samples. Samples without a prime value are shown in black (n=6). **b**, Graph shows plasma IgG antibody binding (AUC, Y-axis) plotted against age (X-axis) after prime (black), and 1.3 months (orange) and 5 months (green) post-second vaccination. **c**, Graph shows AUC values from **a** (Y-axis) plotted against time after vaccination (X-axis). Samples without a prime value are shown in black. **d**, NT50 values in pre-pandemic plasma (ctr) as well as plasma from convalescent individuals 1.3m³ and 6.2m⁷ after infection (grey) and in n=32 vaccinated individuals after prime, and 1.3- and 5-months after receiving 2 doses of an mRNA vaccine. Samples without a prime

value are shown in black. **e**, NT50 values (Y-axis) vs. age (years, X-axis) in individuals after prime (black), or 1.3 months (1.3m, orange) or 5 months (5m, green) after boosting with an mRNA vaccine. **f**, NT50 values (Y-axis) vs. IgG antibody binding (AUC, X-axis) 1.3 months after 2 doses of an mRNA vaccine (n=26). **g**, Graph shows NT50 values (Y-axis) vs. days after boost (X-axis) in individuals receiving two doses of an mRNA vaccine. Samples without a prime value are shown in black. **h**, Ratio of anti-RBD IgG antibody (AUC) to NT50 values (Y-axis) plotted for convalescent infected individuals (grey) 1.3m³ or 6.2m⁷ after infection, and vaccinated individuals after the prime, and 1.3m and 5m after receiving 2 doses of an mRNA vaccine. Samples without a prime value are shown in black. **i**, Plasma neutralizing activity against indicated SARS-CoV-2 variants of interest/concern (n=15 paired samples at 1.3- and 5-months after full vaccination). Refer to Methods for a list of all substitutions/deletions/insertions in the spike variants. All experiments were performed at least in duplicate. Red bars and values in **a**, **d**, **h** and **i** represent geometric mean values. Statistical significance in **a**, **d**, **h** and **i** was determined by Kruskal-Wallis test with subsequent Dunn's multiple comparisons, and in **b**, **c**, **e**, **f**, and **g** by Spearman correlation test.

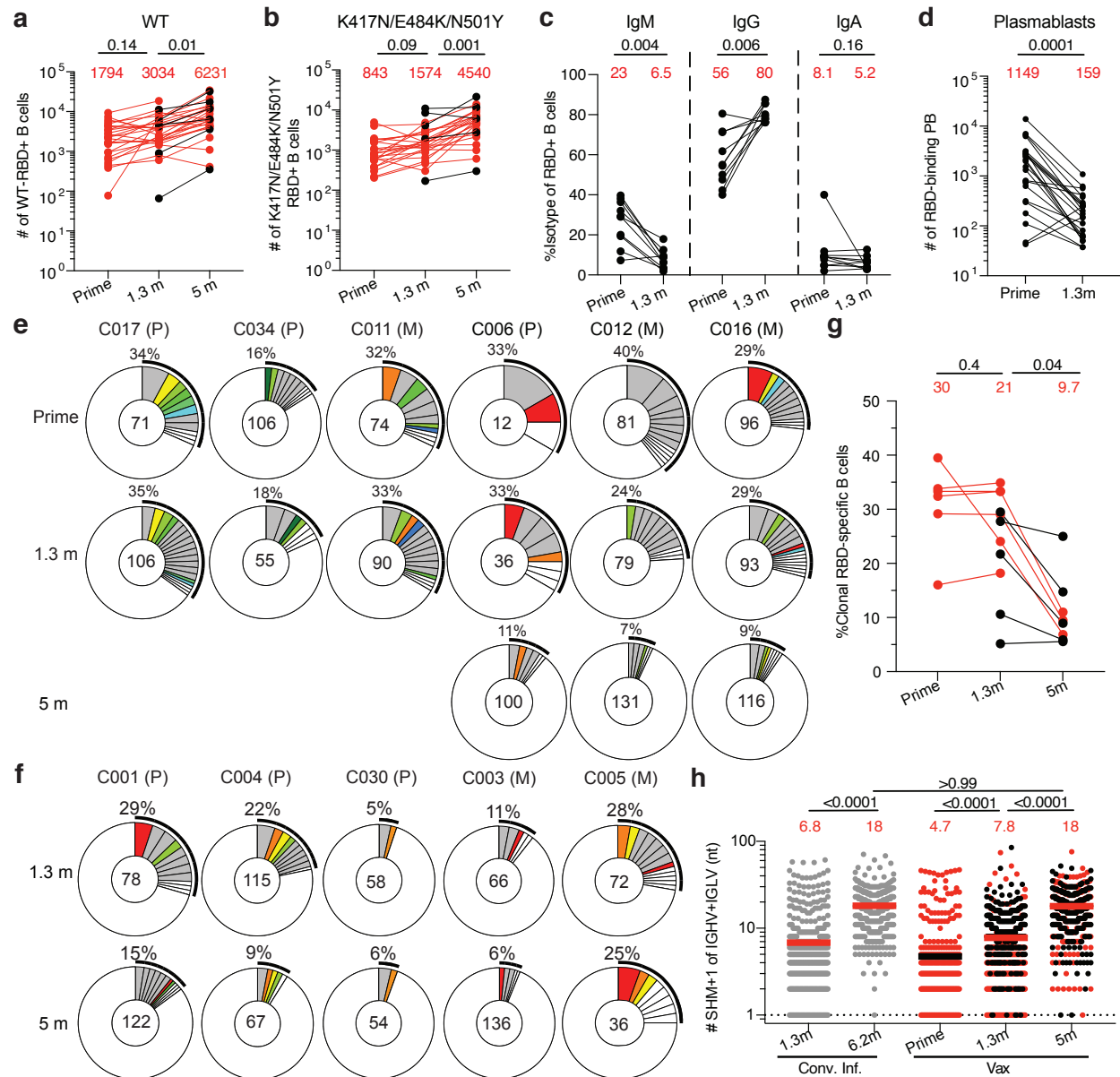


Fig. 2: Anti-SARS-CoV-2 RBD B cells after vaccination. **a-d**, Graphs summarizing **a**, the number of Wuhan-Hu RBD (WT)-specific memory B cells per 10 million B cells for vaccinees and **b**, the number of antigen-specific memory B cells cross-reactive with both WT and K417N/E484K/N501Y RBD mutant per 10 million B cells for n=32 vaccinated individuals after prime, 1.3- and 5-months after 2 doses of vaccination. Samples without a prime value are shown in black. **c**, the frequency of IgM, IgG, or IgA isotype expression by Wuhan-Hu RBD-specific memory B cells after prime or 1.3 months post-boost (n=10), and **d**, number of Wuhan-Hu RBD-

binding plasmablasts per 10 million B cells (n=26) after prime or 1.3 months post-boost. Red numbers indicate geometric means. Gating strategy is in Extended Data Fig. 2. **e, and f**, Pie charts show the distribution of IgG antibody sequences obtained from memory B cells from 11 individuals after **e**, prime and 1.3-months or 5-months post-boost and **f**, 1.3- and 5-months post-boost. The number inside the circle indicates the number of sequences analyzed for the individual denoted above the circle, with Pfizer vaccinees indicated by (P) and Moderna by (M). Pie slice size is proportional to the number of clonally related sequences. The black outline and associated numbers indicate the percentage of clonally expanded sequences detected at each time point. Colored slices indicate persisting clones (same *IGHV* and *IGLV* genes, with highly similar CDR3s) found at more than one timepoint within the same individual. Grey slices indicate clones unique to the timepoint. White slices indicate repeating sequences isolated only once per time point. **g**, Graph shows the relative percentage of clonal sequences at each time point in **e** and **f**. The red numbers indicate the geometric means. Samples without a prime value are shown in black. **h**, Number of nucleotide somatic hypermutations (SHM) in the *IGHV* and *IGLV* combined (also Supplementary Table 4) in the antibodies illustrated in **e** and **f**, compared to the number of mutations obtained after 1.3³ or 6.2⁷ months after infection (grey). Horizontal bars and red numbers indicate mean number of nucleotide mutations at each time point. Samples without a prime value are shown in black. Statistical significance in **a, b, g**, and **h** was determined by Kruskal Wallis test with subsequent Dunn's multiple comparisons, and in **c** and **d** was determined using Wilcoxon matched-pairs signed rank test.

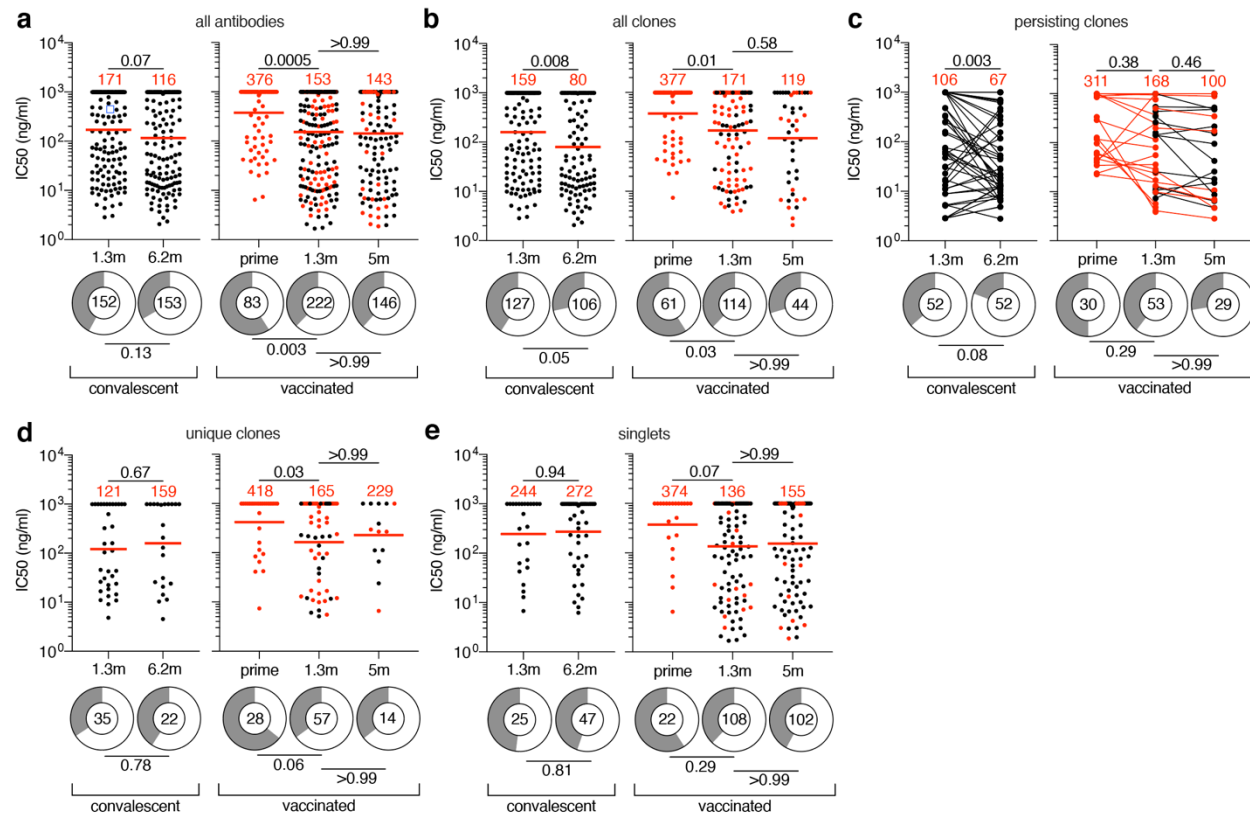


Fig. 3: Anti-SARS-CoV-2 RBD monoclonal antibodies. a-e, Graphs show anti-SARS-CoV-2 neutralizing activity of monoclonal antibodies measured by a SARS-CoV-2 pseudotype virus neutralization assay using wild-type (Wuhan Hu-1⁵⁴) SARS-CoV-2 pseudovirus^{3,8}. Half-maximal inhibitory concentration (IC₅₀) values for all antibodies (a), all clones (b), persisting clones (c), unique clones (d) and singlets (e) isolated from COVID-19 convalescent individuals 1.3³ and 6.2⁷ months after infection or from vaccinated individuals after prime, and 1.3- or 5-months after 2 doses of vaccine. Each dot represents one antibody, where 451 total antibodies were tested including the 430 reported herein (Supplementary Table 5), and 21 previously reported antibodies derived from vaccinees within 8 weeks post-vaccination¹³. Antibodies isolated from samples without a prime value are shown in black. Pie charts illustrate the fraction of non-neutralizing (IC₅₀ > 1000 ng/ml) antibodies (grey slices), inner circle shows the number of antibodies tested

per group. Horizontal bars and red numbers indicate geometric mean values. Statistical significance was determined by Kruskal Wallis test with subsequent Dunn's multiple comparisons. Statistical significance for ring plots was determined using Fisher's exact test with subsequent Bonferroni-correction. All experiments were performed at least twice.

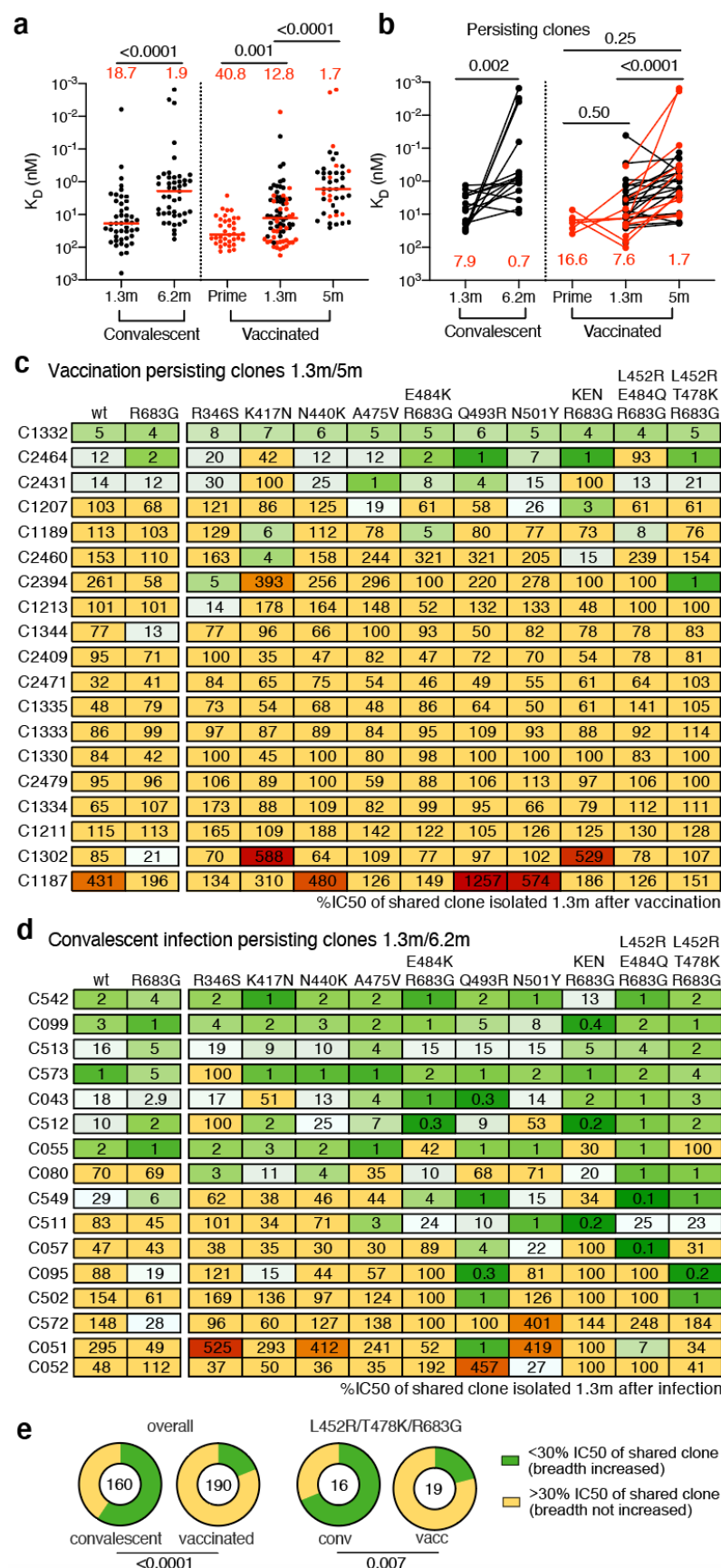


Fig. 4: Affinity and Breadth. a-b, Graphs show antibody K_{DS} for Wuhan-Hu RBD measured by BLI. **a,** antibodies isolated from convalescent individuals 1.3³- (n=42) and 6.2-months⁷ (n=45) after infection or from vaccinees after prime (n=36), and 1.3- (n=74) and 5-months (n=43) post-second vaccination. **b,** Clonally-paired antibodies isolated from convalescent individuals 1.3³- and 6.2⁷-months after infection (n=15) or vaccinated individuals between prime and 1.3 month (n=3), prime and 5 months (n=3), or 1.3- and 5-months after full vaccination (n=26). Antibodies isolated from samples without a prime value are shown in black. Red horizontal bars and numbers indicate median values. Statistical significance was determined using Kruskal Wallis test with subsequent Dunn's multiple comparisons. **c-d,** Heat-maps show inhibitory concentrations of antibodies isolated 5m after vaccination (**c**) or 6.2m⁷ after infection (**d**) normalized to their shared clone isolated 1.3m after vaccination (**c**) or 1.3m³ after infection (**d**), expressed as %IC₅₀, against indicated mutant SARS-CoV-2 pseudoviruses (Supplementary Table 8). Antibodies with improved (<30%) IC₅₀ compared to their clonal relative isolated at an earlier timepoint are colored in shades of green with most improved antibodies in darkest green. Antibodies with worse (>300%) IC₅₀ than their clonal relative isolated at an earlier timepoint are colored in red with the most worsened antibodies in dark red. Antibodies that did not change their IC₅₀ by more than ~3-fold are shown in yellow. **e,** Pie charts illustrate the fraction of antibodies showing improved (<30%, green) vs. not improved (yellow) IC₅₀ compared to their clonal relative isolated at an earlier timepoint, inner circle shows the number of antibody-mutant combinations analyzed per group. Statistical significance for ring plots was determined using Fisher's exact test.

METHODS

Study participants.

Participants were healthy volunteers receiving either the Moderna (mRNA-1273) or Pfizer-BioNTech (BNT162b2) mRNA vaccines against severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) who were recruited for serial blood donations at Rockefeller University Hospital in New York between January 21 and July 20, 2021. The majority of participants (n=28) were *de novo* recruited for this study, while a subgroup of individuals (n=4) were from a long-term study cohort¹³. Eligible participants were healthy adults with no history of infection with SARS-CoV-2, as determined by clinical history and confirmed through serology testing, receiving one of the two Moderna (mRNA-1273) or Pfizer-BioNTech (BNT162b2), according to current dosing and interval guidelines. Exclusion criteria included incomplete vaccination status, presence of clinical signs and symptoms suggestive of acute infection with or a positive reverse transcription polymerase chain reaction (RT-PCR) results for SARS-CoV-2 in saliva, or a positive (coronavirus disease 2019) COVID-19 serology. Seronegativity for COVID-19 was established through the absence of serological activity toward the nucleocapsid protein (N) of SARS-CoV-2. Participants presented to the Rockefeller University Hospital for blood sample collection and were asked to provide details of their vaccination regimen, possible side effects, comorbidities and possible COVID-19 history. All participants provided written informed consent before participation in the study and the study was conducted in accordance with Good Clinical Practice. The study was performed in compliance with all relevant ethical regulations and the protocol (DRO-1006) for studies with human participants was approved by the Institutional Review Board of the Rockefeller University. For detailed participant characteristics see Supplementary Tables 1 and 2.

410

411 **Blood samples processing and storage.**

412 Peripheral Blood Mononuclear Cells (PBMCs) obtained from samples collected at Rockefeller

413 University were purified as previously reported by gradient centrifugation and stored in liquid

414 nitrogen in the presence of Fetal Calf Serum (FCS) and Dimethylsulfoxide (DMSO)^{3,7}.

415 Heparinized plasma and serum samples were aliquoted and stored at -20°C or less. Prior to

416 experiments, aliquots of plasma samples were heat-inactivated (56°C for 1 hour) and then stored

417 at 4°C.

418

419 **ELISAs**

420 Enzyme-Linked Immunosorbent Assays (ELISAs)^{55,56} to evaluate antibodies binding to SARS-

421 CoV-2 RBD were performed by coating of high-binding 96-half-well plates (Corning 3690) with

422 50 µl per well of a 1 µg/ml protein solution in Phosphate-buffered Saline (PBS) overnight at 4°C.

423 Plates were washed 6 times with washing buffer (1× PBS with 0.05% Tween-20 (Sigma-

424 Aldrich)) and incubated with 170 µl per well blocking buffer (1× PBS with 2% BSA and 0.05%

425 Tween-20 (Sigma)) for 1 hour at room temperature. Immediately after blocking, monoclonal

426 antibodies or plasma samples were added in PBS and incubated for 1 hour at room temperature.

427 Plasma samples were assayed at a 1:66 starting dilution and 10 additional threefold serial

428 dilutions. Monoclonal antibodies were tested at 10 µg/ml starting concentration and 10 additional

429 fourfold serial dilutions. Plates were washed 6 times with washing buffer and then incubated

430 with anti-human IgG, IgM or IgA secondary antibody conjugated to horseradish peroxidase

431 (HRP) (Jackson Immuno Research 109-036-088 109-035-129 and Sigma A0295) in blocking

432 buffer at a 1:5,000 dilution (IgM and IgG) or 1:3,000 dilution (IgA). Plates were developed by

addition of the HRP substrate, 3,3',5,5'-Tetramethylbenzidine (TMB) (ThermoFisher) for 10 minutes (plasma samples) or 4 minutes (monoclonal antibodies). The developing reaction was stopped by adding 50 μ l of 1 M H₂SO₄ and absorbance was measured at 450 nm with an ELISA microplate reader (FluoStar Omega, BMG Labtech) with Omega and Omega MARS software for analysis. For plasma samples, a positive control (plasma from participant COV72, diluted 66.6-fold and ten additional threefold serial dilutions in PBS) was added to every assay plate for normalization. The average of its signal was used for normalization of all the other values on the same plate with Excel software before calculating the area under the curve using Prism V9.1(GraphPad). Negative controls of pre-pandemic plasma samples from healthy donors were used for validation (for more details please see³). For monoclonal antibodies, the ELISA half-maximal concentration (EC₅₀) was determined using four-parameter nonlinear regression (GraphPad Prism V9.1). EC₅₀s above 2000 ng/mL were considered non-binders.

Proteins

The mammalian expression vector encoding the Receptor Binding-Domain (RBD) of SARS-CoV-2 (GenBank MN985325.1; Spike (S) protein residues 319-539) was previously described⁵⁷.

SARS-CoV-2 pseudotyped reporter virus

A panel of plasmids expressing RBD-mutant SARS-CoV-2 spike proteins in the context of pSARS-CoV-2-S_{Δ19} has been described^{13,25,58}. Variant pseudoviruses resembling variants of interest/concern B.1.1.7 (first isolated in the UK), B.1.351 (first isolated in South-Africa), B.1.526 (first isolated in New York City), P.1 (first isolated in Brazil) and B.1.617.2 (first isolated in India) were generated by introduction of substitutions using synthetic gene fragments

(IDT) or overlap extension PCR mediated mutagenesis and Gibson assembly. Specifically, the variant-specific deletions and substitutions introduced were:

B.1.1.7: ΔH69/V70, ΔY144, N501Y, A470D, D614G, P681H, T761I, S982A, D118H

B.1.351: D80A, D215G, L242H, R246I, K417N, E484K, N501Y, D614G, A701V

B.1.526: L5F, T95I, D253G, E484K, D614G, A701V.

P.1: L18F, T20N, P26S, D138Y, R190S, K417T, E484K, N501Y, D614G, H655Y, T1027I,

V1167F

B.1.617.2: T19R, Δ156-158, L452R, T478K, D614G, P681R, D950N

The E484K, K417N/E484K/N501Y, L452R/E484Q and L452R/T478K substitution, as well as

the deletions/substitutions corresponding to variants of concern listed above were incorporated

into a spike protein that also includes the R683G substitution, which disrupts the furin cleavage

site and increases particle infectivity. Neutralizing activity against mutant pseudoviruses were

compared to a wildtype (WT) SARS-CoV-2 spike sequence (NC_045512), carrying R683G

where appropriate.

SARS-CoV-2 pseudotyped particles were generated as previously described^{3,8}. Briefly, 293T

cells were transfected with pNL4-3ΔEnv-nanoluc and pSARS-CoV-2-S_{Δ19}, particles were

harvested 48 hours post-transfection, filtered and stored at -80°C.

Pseudotyped virus neutralization assay

Fourfold serially diluted pre-pandemic negative control plasma from healthy donors, plasma

from COVID-19-convalescent individuals or monoclonal antibodies were incubated with SARS-

CoV-2 pseudotyped virus for 1 hour at 37 °C. The mixture was subsequently incubated with

293T_{Ace2} cells³ (for all WT neutralization assays) or HT1080Ace2 cl14 (for all mutant panels and variant neutralization assays) cells¹³ for 48 hours after which cells were washed with PBS and lysed with Luciferase Cell Culture Lysis 5× reagent (Promega). Nanoluc Luciferase activity in lysates was measured using the Nano-Glo Luciferase Assay System (Promega) with the Glomax Navigator (Promega). The relative luminescence units were normalized to those derived from cells infected with SARS-CoV-2 pseudotyped virus in the absence of plasma or monoclonal antibodies. The half-maximal neutralization titers for plasma (NT₅₀) or half-maximal and 90% inhibitory concentrations for monoclonal antibodies (IC₅₀ and IC₉₀) were determined using four-parameter nonlinear regression (least squares regression method without weighting; constraints: top=1, bottom=0) (GraphPad Prism).

Biotinylation of viral protein for use in flow cytometry

Purified and Avi-tagged SARS-CoV-2 RBD or SARS-CoV-2 RBD K417N/E484K/N501Y mutant was biotinylated using the Biotin-Protein Ligase-BIRA kit according to manufacturer's instructions (Avidity) as described before³. Ovalbumin (Sigma, A5503-1G) was biotinylated using the EZ-Link Sulfo-NHS-LC-Biotinylation kit according to the manufacturer's instructions (Thermo Scientific). Biotinylated ovalbumin was conjugated to streptavidin-BV711 (BD biosciences, 563262) and RBD to streptavidin-PE (BD Biosciences, 554061) and streptavidin-AF647 (Biolegend, 405237)³.

Flow cytometry and single cell sorting

Single-cell sorting by flow cytometry was described previously³. Briefly, peripheral blood mononuclear cells were enriched for B cells by negative selection using a pan-B-cell isolation kit

according to the manufacturer's instructions (Miltenyi Biotec, 130-101-638). The enriched B cells were incubated in Fluorescence-Activated Cell-sorting (FACS) buffer (1× PBS, 2% FCS, 1 mM ethylenediaminetetraacetic acid (EDTA)) with the following anti-human antibodies (all at 1:200 dilution): anti-CD20-PECy7 (BD Biosciences, 335793), anti-CD3-APC-eFluor 780 (Invitrogen, 47-0037-41), anti-CD8-APC-eFluor 780 (Invitrogen, 47-0086-42), anti-CD16-APC-eFluor 780 (Invitrogen, 47-0168-41), anti-CD14-APC-eFluor 780 (Invitrogen, 47-0149-42), as well as Zombie NIR (BioLegend, 423105) and fluorophore-labeled RBD and ovalbumin (Ova) for 30 min on ice. Single CD3-CD8-CD14-CD16-CD20+Ova-RBD-PE+RBD-AF647+ B cells were sorted into individual wells of 96-well plates containing 4 µl of lysis buffer (0.5× PBS, 10 mM Dithiothreitol (DTT), 3,000 units/ml RNasin Ribonuclease Inhibitors (Promega, N2615) per well using a FACS Aria III and FACSDiva software (Becton Dickinson) for acquisition and FlowJo for analysis. The sorted cells were frozen on dry ice, and then stored at -80 °C or immediately used for subsequent RNA reverse transcription. For plasmablast single-cell sorting, in addition to above antibodies, B cells were also stained with anti-CD19-BV605 (Biolegend, 302244), and single CD3-CD8-CD14-CD16-CD19+CD20-Ova-RBD-PE+RBD-AF647+ plasmablasts were sorted as described above. For B cell phenotype analysis, in addition to above antibodies, B cells were also stained with following anti-human antibodies: anti-IgD-BV421 (Biolegend, 348226), anti-CD27-FITC (BD biosciences, 555440), anti-CD19-BV605 (Biolegend, 302244), anti-CD71- PerCP-Cy5.5 (Biolegend, 334114), anti- IgG-PECF594 (BD biosciences, 562538), anti-IgM-AF700 (Biolegend, 314538), anti-IgA-Viogreen (Miltenyi Biotec, 130-113-481).

Antibody sequencing, cloning and expression

Antibodies were identified and sequenced as described previously^{3,59}. In brief, RNA from single cells was reverse-transcribed (SuperScript III Reverse Transcriptase, Invitrogen, 18080-044) and the cDNA was stored at -20°C or used for subsequent amplification of the variable IGH, IGL and IGK genes by nested PCR and Sanger sequencing. Sequence analysis was performed using MacVector. Amplicons from the first PCR reaction were used as templates for sequence- and ligation-independent cloning into antibody expression vectors. Recombinant monoclonal antibodies were produced and purified as previously described³.

Biolayer interferometry

Biolayer interferometry assays were performed as previously described³. Briefly, we used the Octet Red instrument (ForteBio) at 30°C with shaking at 1,000 r.p.m. Affinity measurement of anti-SARS-CoV-2 IgGs binding were corrected by subtracting the signal obtained from traces performed with IgGs in the absence of WT RBD. The kinetic analysis using protein A biosensor (ForteBio 18-5010) was performed as follows: (1) baseline: 60sec immersion in buffer. (2) loading: 200sec immersion in a solution with IgGs 10 $\mu\text{g}/\text{ml}$. (3) baseline: 200sec immersion in buffer. (4) Association: 300sec immersion in solution with WT RBD at 20, 10 or 5 $\mu\text{g}/\text{ml}$ (5) dissociation: 600sec immersion in buffer. Curve fitting was performed using a fast 1:1 binding model and the Data analysis software (ForteBio). Mean equilibrium dissociation constant (K_D) values were determined by averaging all binding curves that matched the theoretical fit with an R^2 value ≥ 0.8 .

Computational analyses of antibody sequences

Antibody sequences were trimmed based on quality and annotated using Igblastn v.1.14. with IMGT domain delineation system. Annotation was performed systematically using Change-O toolkit v.0.4.540⁶⁰. Heavy and light chains derived from the same cell were paired, and clonotypes were assigned based on their V and J genes using in-house R and Perl scripts. All scripts and the data used to process antibody sequences are publicly available on GitHub (https://github.com/stratust/igpipeline/tree/igpipeline2_timepoint_v2).

The frequency distributions of human V genes in anti-SARS-CoV-2 antibodies from this study was compared to 131,284,220 IgH and IgL sequences generated by⁶¹ and downloaded from cAb-Rep⁶², a database of human shared BCR clonotypes available at <https://cab-rep.c2b2.columbia.edu/>. Based on the 112 distinct V genes that make up the 7936 analyzed sequences from Ig repertoire of the 11 participants present in this study, we selected the IgH and IgL sequences from the database that are partially coded by the same V genes and counted them according to the constant region. The frequencies shown in Extended Data Fig. 4 are relative to the source and isotype analyzed. We used the two-sided binomial test to check whether the number of sequences belonging to a specific *IGHV* or *IGLV* gene in the repertoire is different according to the frequency of the same IgV gene in the database. Adjusted p-values were calculated using the false discovery rate (FDR) correction. Significant differences are denoted with stars.

Nucleotide somatic hypermutation and Complementarity-Determining Region (CDR3) length were determined using in-house R and Perl scripts. For somatic hypermutations, *IGHV* and *IGLV* nucleotide sequences were aligned against their closest germlines using Igblastn and the number

of differences were considered nucleotide mutations. The average number of mutations for V genes was calculated by dividing the sum of all nucleotide mutations across all participants by the number of sequences used for the analysis.

Data availability statement: Data are provided in Supplementary Tables 1-8. The raw sequencing data and computer scripts associated with Figure 2 and Extended Data Fig. 3 have been deposited at Github (https://github.com/stratust/igpipeline/tree/igpipeline2_timepoint_v2). This study also uses data from “A Public Database of Memory and Naive B-Cell Receptor Sequences” (<https://doi.org/10.5061/dryad.35ks2>), PDB (6VYB and 6NB6) and from “High frequency of shared clonotypes in human B cell receptor repertoires” (<https://doi.org/10.1038/s41586-019-0934-8>).

Code availability statement: Computer code to process the antibody sequences is available at GitHub (https://github.com/stratust/igpipeline/tree/igpipeline2_timepoint_v2).

Data presentation

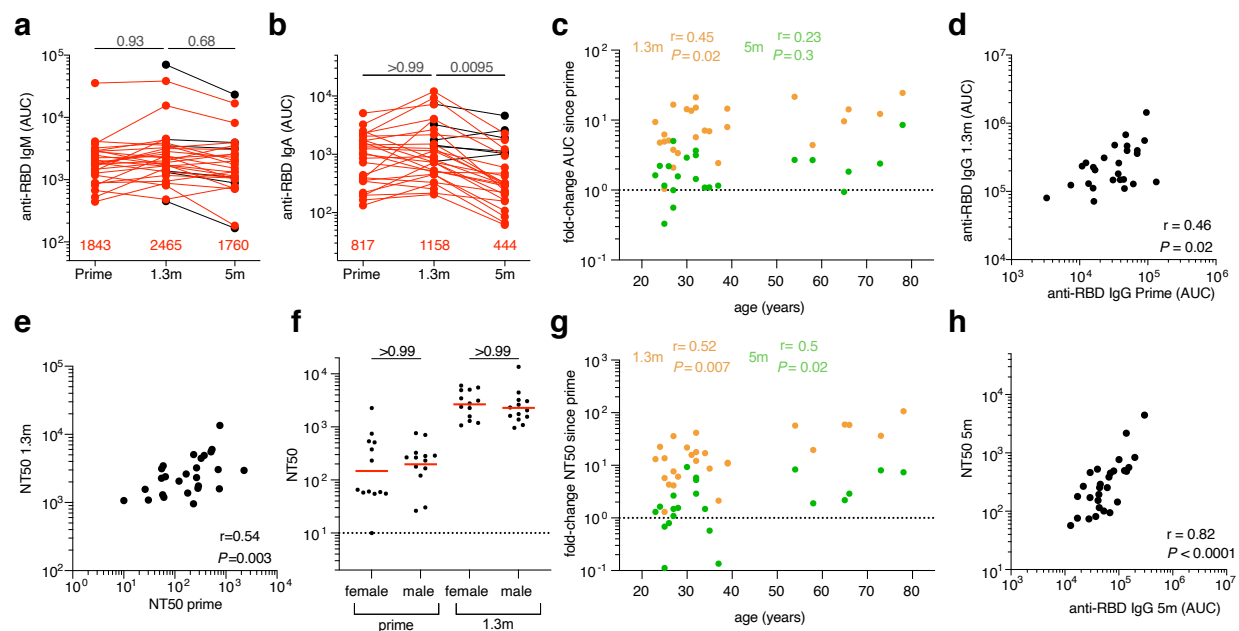
Figures arranged in Adobe Illustrator 2020.

Competing interests: The Rockefeller University has filed a provisional patent application in connection with this work on which M.C.N. is an inventor (US patent 63/021,387). The patent has been licensed by Rockefeller University to Bristol Meyers Squib.

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Author Contributions: P.D.B., T.H., and M.C.N. conceived, designed and analyzed the experiments. M. Caskey and C.G. designed clinical protocols. A.C, F.M., D.S.B., Z.W., S.F., P.M., M.A., E.B., J.D.S., I.S., J.D. F.S., F.Z., and T.B.T. carried out experiments. A.G. and M. Cipolla produced antibodies. D.S.B., M.D., M.T., K.G.M., C.G. and M. Caskey recruited participants, executed clinical protocols. T.Y.O. and V.R. performed bioinformatic analysis. A.C., F.M, D.S.B., Z.W., S.F., and M.C.N. wrote the manuscript with input from all co-authors.

615 **EXTENDED FIGURES**



616

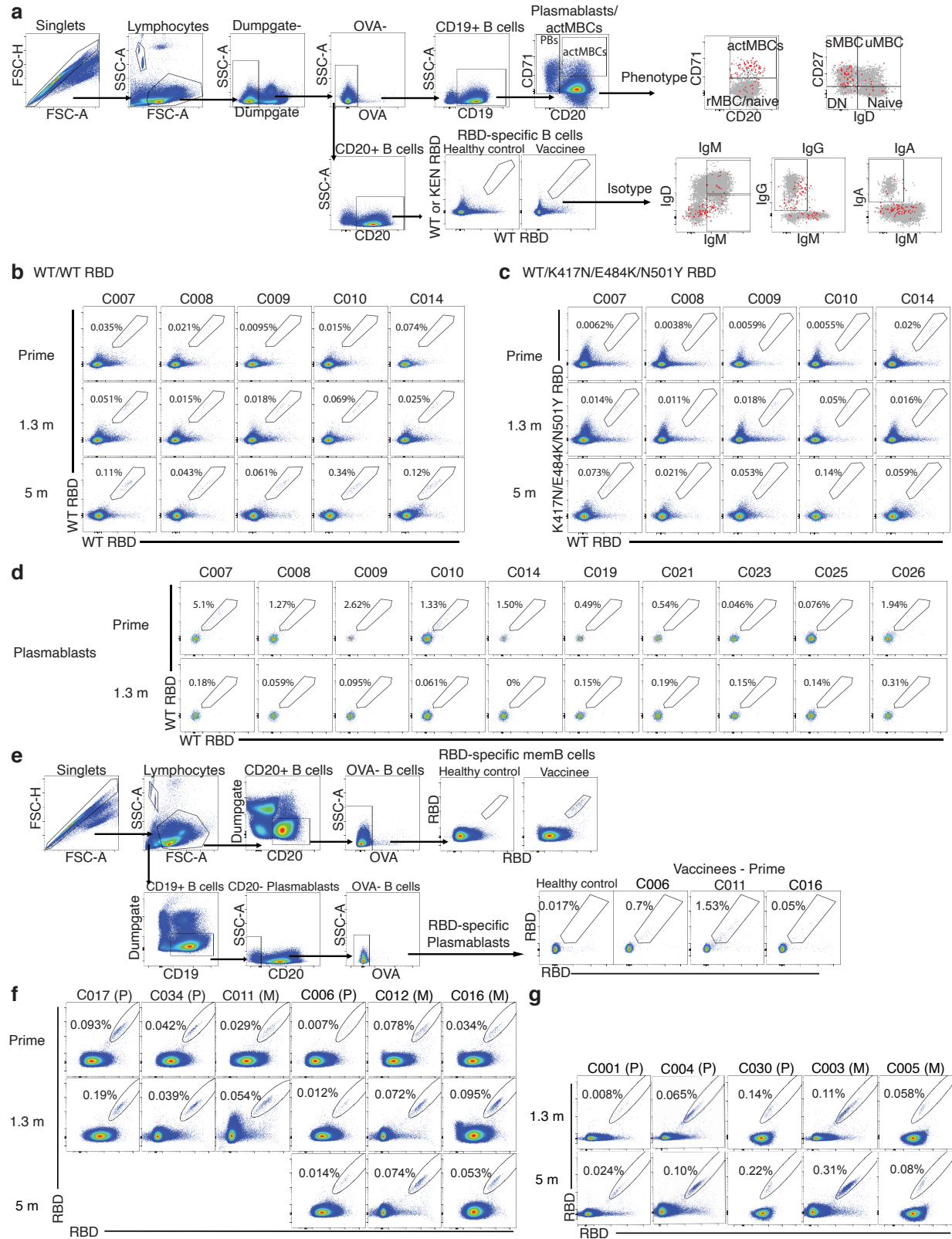
617 **Extended Data Fig 1: Plasma ELISA and neutralization.**

618 **a,b**, Graph shows area under the curve (AUC, Y-axis) for plasma IgM (**a**) or IgA (**b**) antibody
619 binding to SARS-CoV-2 RBD after prime, and 1.3- and 5-months post-boost for paired samples
620 from $n=32$ vaccinated individuals. Samples without a prime value are shown in black. **c**, Graph
621 shows age (years, X-axis) vs. fold-change of IgG-binding titers (AUC, Y-Axis) between prime
622 and 1.3m (orange) or 5m (green) post-boost. **d**, IgG antibody binding after prime (AUC, X-axis)
623 vs. IgG antibody binding after 1.3 months post-boost (AUC, Y-axis) and **e**, NT50 values after
624 prime (X-axis) vs. NT50 values after 1.3 months post-boost (Y-axis) in individuals receiving two
625 doses of an mRNA vaccine ($n=26$). **f**, NT50 values after prime and 1.3 months post-boost in
626 females and males receiving 2 doses of an mRNA vaccine. **g**, Graph shows age (years, X-axis) vs
627 fold-change of NT50 (X-axis) between prime and 1.3m (orange) or 5m (green) post-boost. **h**, NT50
628 values (Y-axis) vs. IgG antibody binding (AUC, X-axis) 5 months after boost in individuals

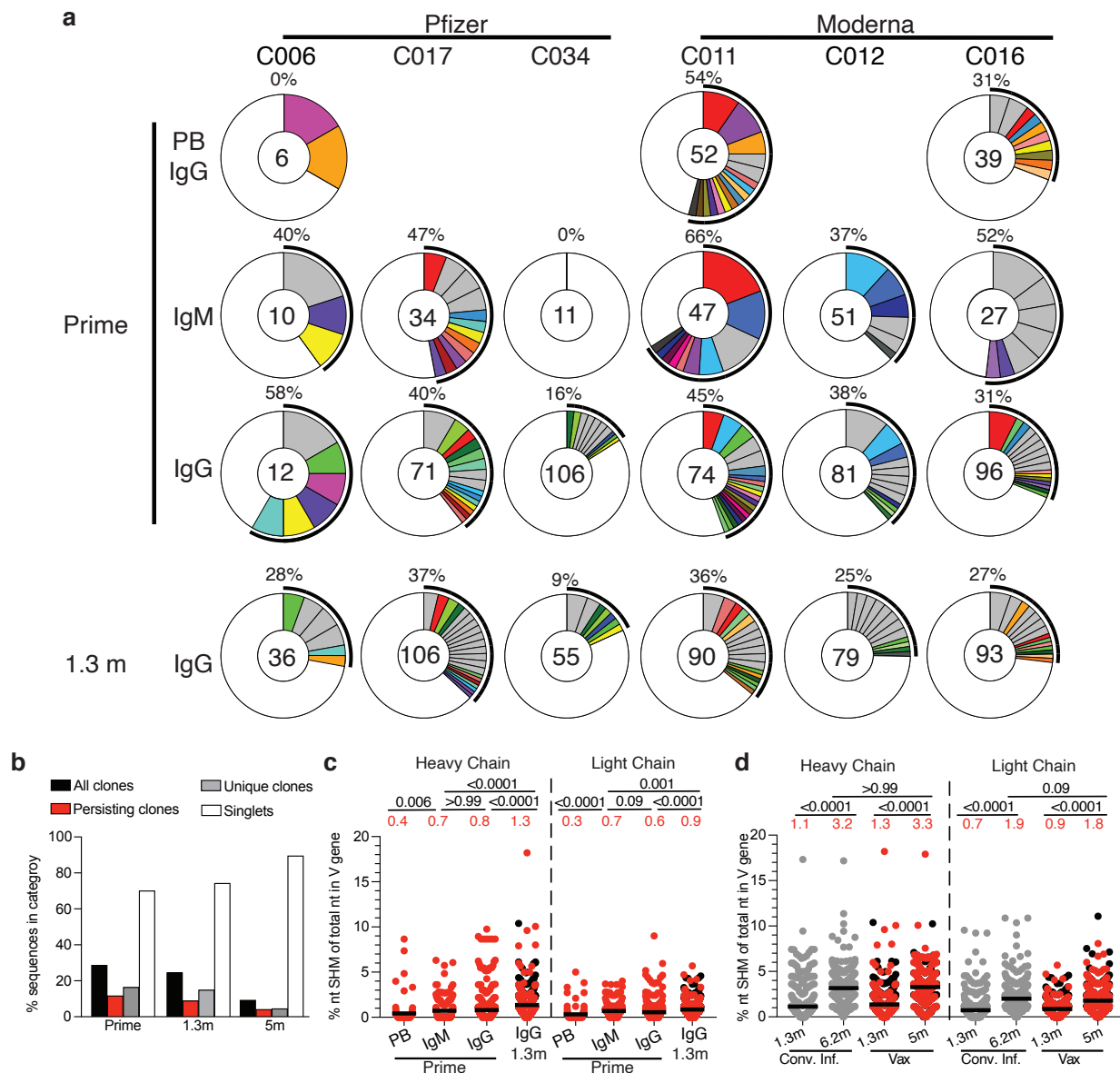
629 receiving two doses of an mRNA vaccine (n=28). All experiments were performed at least in
630 duplicate. Red values or bar in **a**, **b** and **f** represent geometric mean values. Statistical significance
631 in **a**, **b**, and **f** was determined by Kruskal-Wallis test with subsequent Dunn's multiple
632 comparisons, or by Spearman correlation test in **c**, **d**, **e**, **g**, and **h**.

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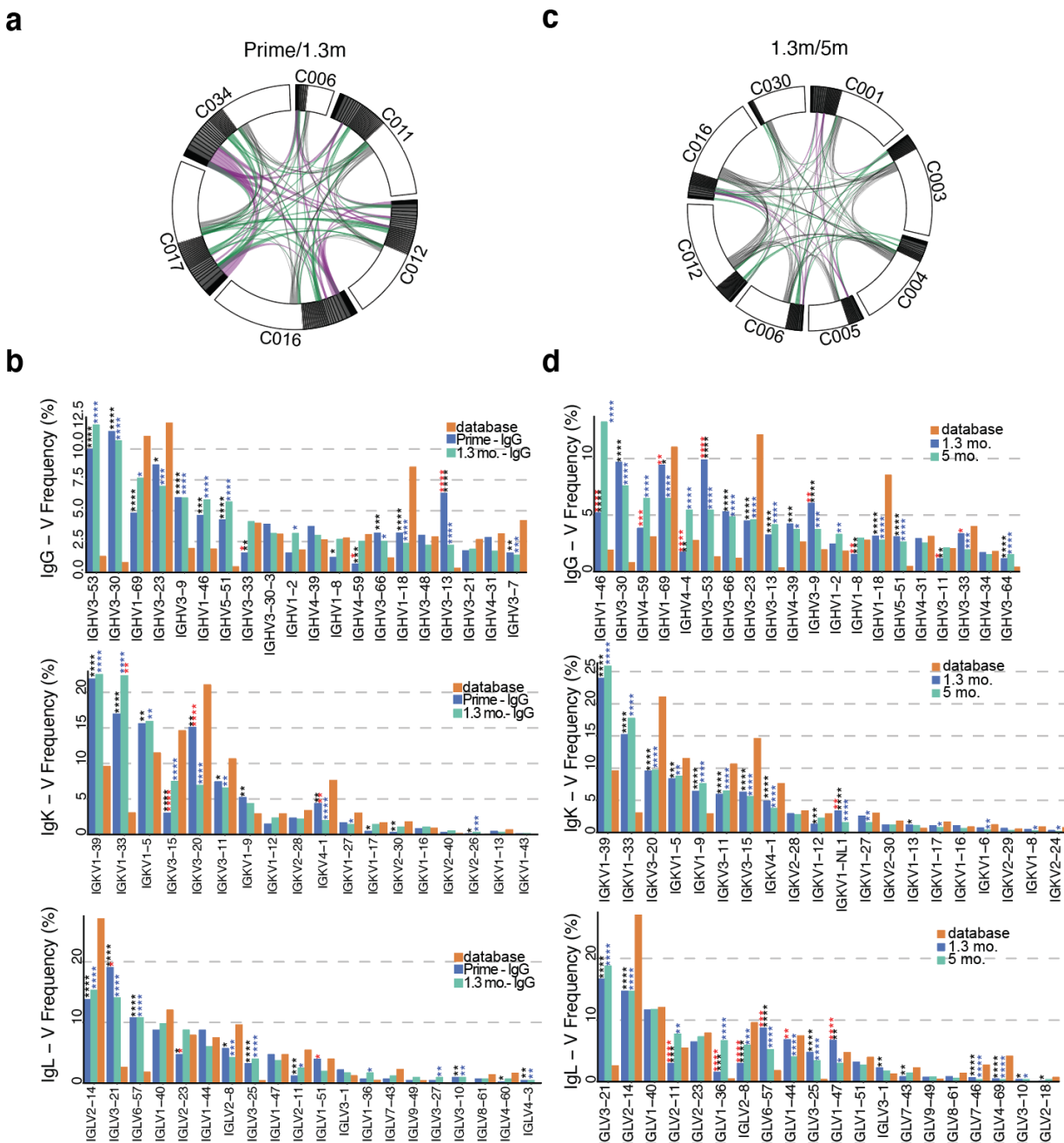


Extended Data Fig. 2: Flow Cytometry. a, Gating strategy for phenotyping. Gating was on singlets that were CD19⁺ or CD20⁺ and CD3-CD8-CD16-Ova-. Anti-IgG, IgM, IgA, IgD, CD71 and CD27 antibodies were used for B cell phenotype analysis. Antigen-specific cells were detected based on binding to RBD WT-PE⁺ and RBD WT/KEN (K417N/E484K/N501Y)-AF647⁺. **b-c,** Flow cytometry plots showing the frequency of **b**, RBD WT-binding memory B cells, and **c**, RBD-binding memory B cells cross-reactive with WT and K417N/E484K/N501Y mutant RBD in 5 selected individuals, after prime, 1.3 months, and 5 months post-second vaccination. **d,** Flow cytometry plots showing frequency of RBD-binding plasmablasts, in 10 selected vaccinees after prime or 1.3 months post-boost. **e,** Gating strategy for single-cell sorting for CD20⁺ memory B cells (top panel) or CD19⁺CD20⁻ plasmablasts (bottom panel) which were double positive for RBD-PE and RBD-AF647. **f-g,** Representative flow cytometry plots showing dual AlexaFluor-647-RBD and PE-RBD-binding, single-cell sorted B cells from **f**, 6 individuals after prime and 1.3 months or 5 months post-boost and **g**, 5 individuals from 1.3- or 5-months post-boost. Percentage of RBD-specific B cells is indicated.



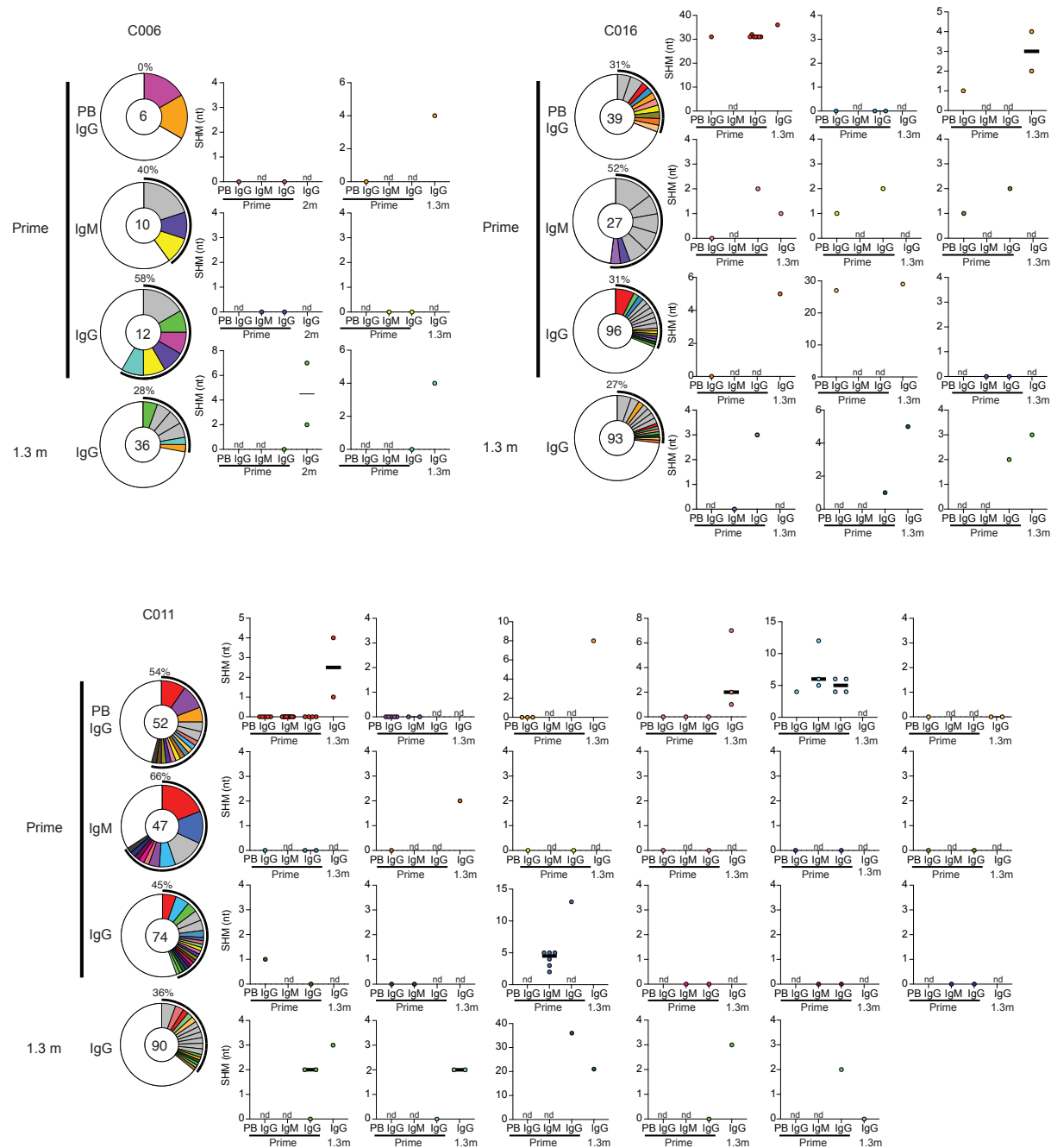
Extended Data Fig 3: anti-SARS-CoV-2 RBD-specific plasmablast and memory B cells responses after vaccination. **a**, Pie charts show the distribution of antibody sequences from 6 individuals after prime (upper panel) or 1.3 months post-boost (lower panel). Sequences derived from IgG plasmablast (PB), IgM memory B cells (MBC), and IgG MBC compartments were analyzed after prime, while only IgG MBCs were analyzed at 1.3 months after boost, as indicated to the left of the plots. The number inside the circle indicates the number of sequences analyzed for the individual denoted above the circle. Pie slice size is proportional to the number of clonally

related sequences. The black outline indicates the frequency of clonally sequences detected in each patient. Colored slices indicate persisting clones (same *IGHV* and *IGLV* genes, with highly similar CDR3s) found in multiple compartments and/or timepoints within the same patient. Grey slices indicate clones unique to the compartment. White indicates sequences isolated once. **b**, Graph shows the percentage of total paired-sequences analyzed at either prime, 1.3- or 5-months post-boost, that can be found as part of all clones (black bars), persisting clones (red bars), unique clones (grey bars), or singlets (white bar). **c-d**, Ratio of the number of somatic nucleotide mutations over the nucleotide length of the V gene in the Ig heavy and light chains, separately, in antibodies detected in **c**, different B cell compartments after prime or 1.3 months post-boost and **d**, 1.3 or 5 months post-boost compared to convalescent infected (grey) individuals after 1.3³ and 6.2⁷ months post-infection (also Supplementary Table 4). Horizontal bars and red numbers indicate mean ratio in each compartment at each time point. Sequences derived from samples without a prime value are shown in black. Statistical significance in **c** and **d** was determined using a Kruskal Wallis test with subsequent Dunn's multiple comparisons.



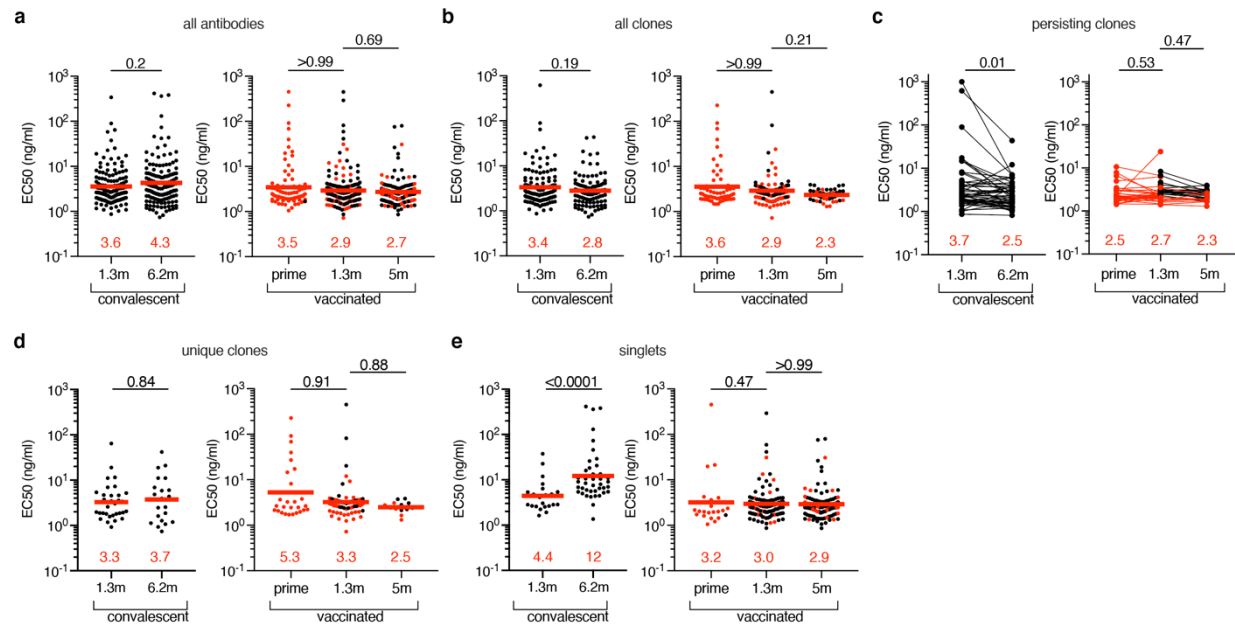
Extended Data Fig. 4: Frequency distribution of human V genes. **a**, Circos plot depicting relationship between antibodies that share V and J gene usage in both IgH and IgL when comparing prime/1.3m IgG MBC sequences. Purple, green, and grey lines connect related clones, clones and singlets, and singlets to each other, respectively. **b**, Graph shows relative abundance of human heavy chain *IGHV* (top), light chain *IGKV* (middle) or *IGLV* (bottom) genes comparing Sequence

Read Archive accession SRP010970 (orange), and IgG MBCs after prime (blue) or 1.3 months post-boost (green). Statistical significance was determined by two-sided binomial test. * = $p \leq 0.05$, ** = $p \leq 0.01$, *** = $p \leq 0.001$, **** = $p \leq 0.0001$. Color of stars indicates: black - comparing Database versus Prime; blue - comparing Database versus 1.3m; red - comparing Prime versus 1.3m. **c**, Circos plot depicting relationship between antibodies that share V and J gene usage in both IgH and IgL when comparing 1.3 m/5 m IgG MBC sequences. Purple, green, and grey lines connect related clones, clones and singlets, and singlets to each other, respectively. **d**, Graph shows relative abundance of human heavy chain *IGHV* (top), light chain *IGKV* (middle) or *IGLV* (bottom) genes comparing Sequence Read Archive accession SRP010970 (orange), and IgG MBCs after 1.3 months (blue) or 5 months (green) post-vaccination. Statistical significance was determined by two-sided binomial test. * = $p \leq 0.05$, ** = $p \leq 0.01$, *** = $p \leq 0.001$, **** = $p \leq 0.0001$. Color of stars indicates: black - comparing Database versus 1.3 months; blue - comparing Database versus 5 months; red - comparing 1.3 months versus 5 months.

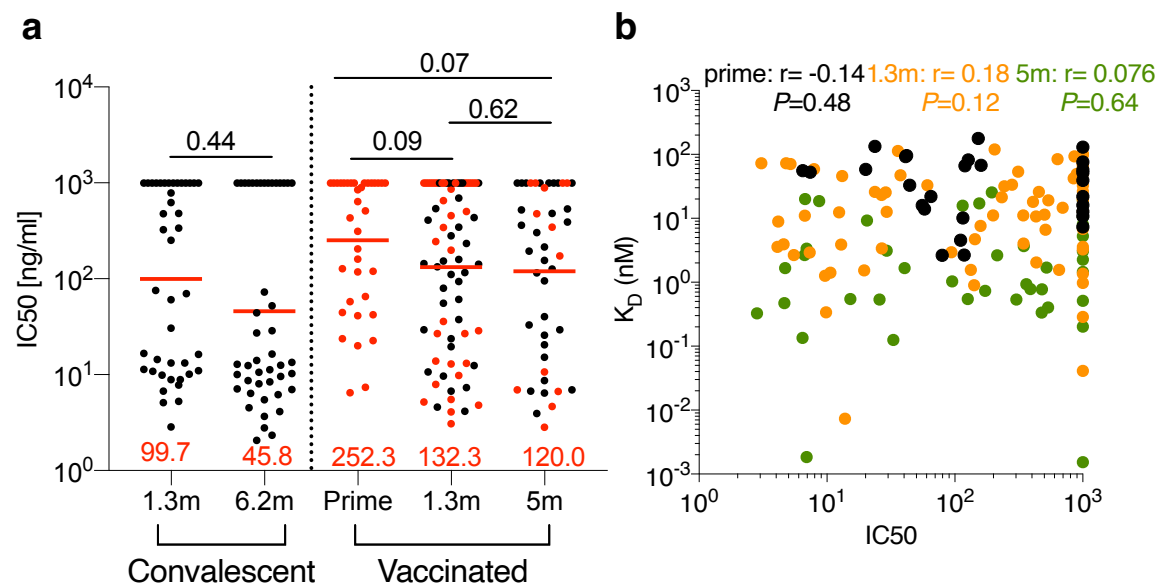


Extended Data Fig. 5: Somatic hypermutation of anti-SARS-CoV-2 RBD antibody clones after prime or boost. Clonal evolution of RBD-binding B cells from 3 individuals for which plasmablasts, IgM memory B cells, and IgG memory B cells were analyzed after prime, and IgG memory B cells were analyzed after 1.3 months post-boost (as described in Extended Data Fig. 3). The number of somatic nucleotide mutations found in shared clonal families found in at least 2

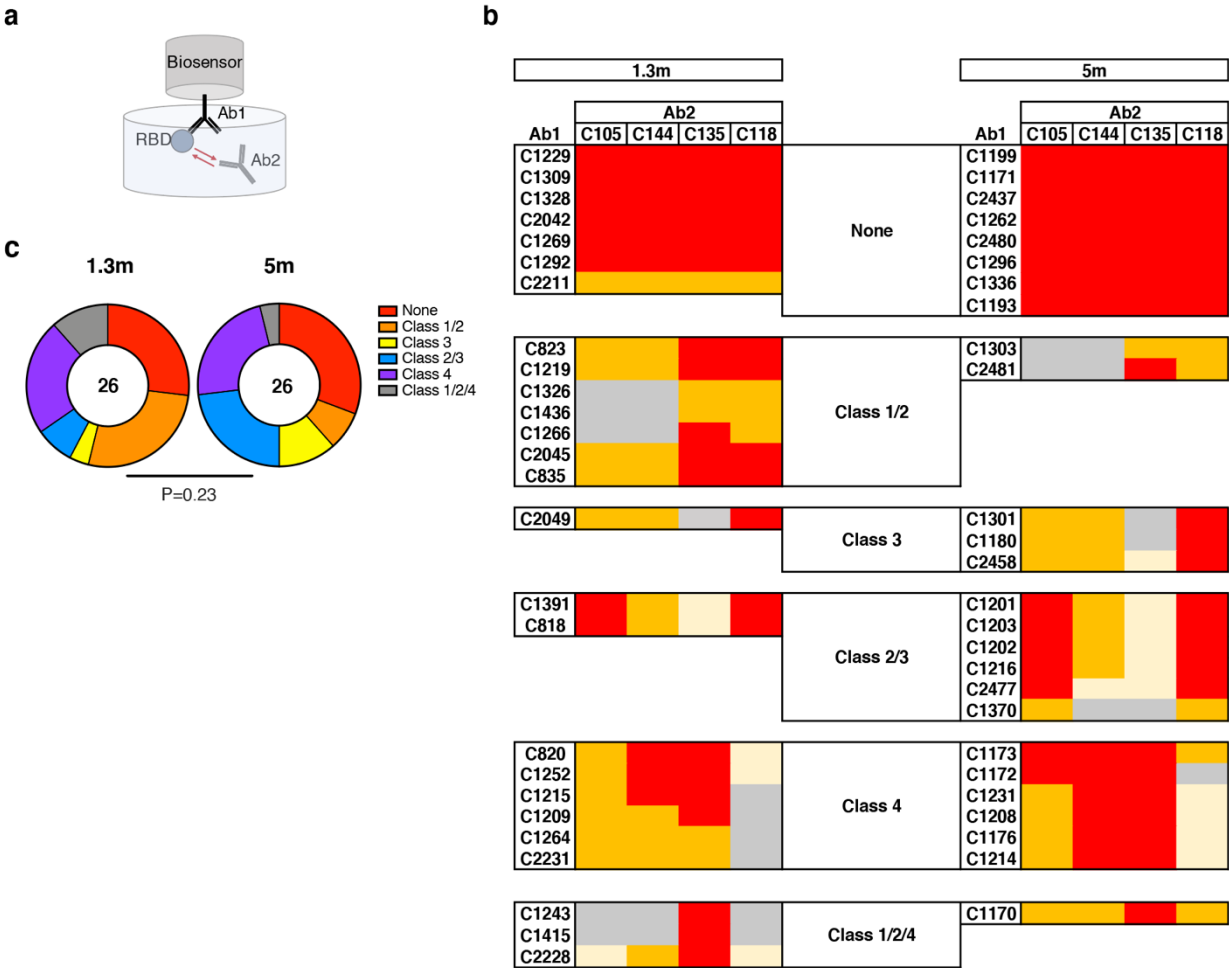
704 different compartments is graphed to the right of each donut plot. Color of dot plots match the
705 color of pie slices within the donut plot, which indicate persisting clones. nd – clone was Not
706 Detected in the indicated compartment. Black horizontal line indicates median number of SHM.
707
708



Extended Data Fig. 6: Anti-SARS-CoV-2 RBD monoclonal antibodies ELISAs. a-e, Graphs show anti-SARS-CoV-2 binding activity of monoclonal antibodies measured by ELISA against RBD. ELISA half-maximal concentration (EC₅₀) values for all antibodies (a), all clones (b), persisting clones (c), unique clones (d) and singlets (e) isolated from COVID-19 convalescent individuals 1.3³ and 6.2⁷ months after infection (left panel) or from vaccinated individuals after prime, or 1.3m or 5m after receiving the second dose of mRNA vaccination (right panel). Each dot represents one antibody. Antibodies isolated from samples without a prime value are shown in black. Red horizontal bars and numbers indicate geometric mean values. Statistical significance was determined by Mann-Whitney test (left panels of a, b, d and e), Kruskal-Wallis test with subsequent Dunn's multiple comparisons (right panels of a-e) or by Wilcoxon test (left panel of c). All experiments were performed at least twice.



Extended Data Fig. 7. Affinity. Biolayer interferometry measurements. **a**, IC₅₀ values for randomly selected antibodies isolated from convalescents 1.3m³ (n=42) and 6.2m⁷ (n=45) after infection or from vaccinees after prime (n=36), and 1.3m (n=74) and 5m (n=43). Red horizontal lines and numbers indicate geometric mean. Antibodies isolated from samples without a prime value are shown in black. **b**, Graphs show affinities (K_D, Y-axis) plotted against neutralization activity (IC₅₀, X-axis) for antibodies isolated after prime (black), or 1.3m (orange) or 5m (green) post-boost vaccination. Statistical significance was determined using Kruskal Wallis test with subsequent Dunn's multiple comparisons in **a** and Spearman correlation test in **b**.

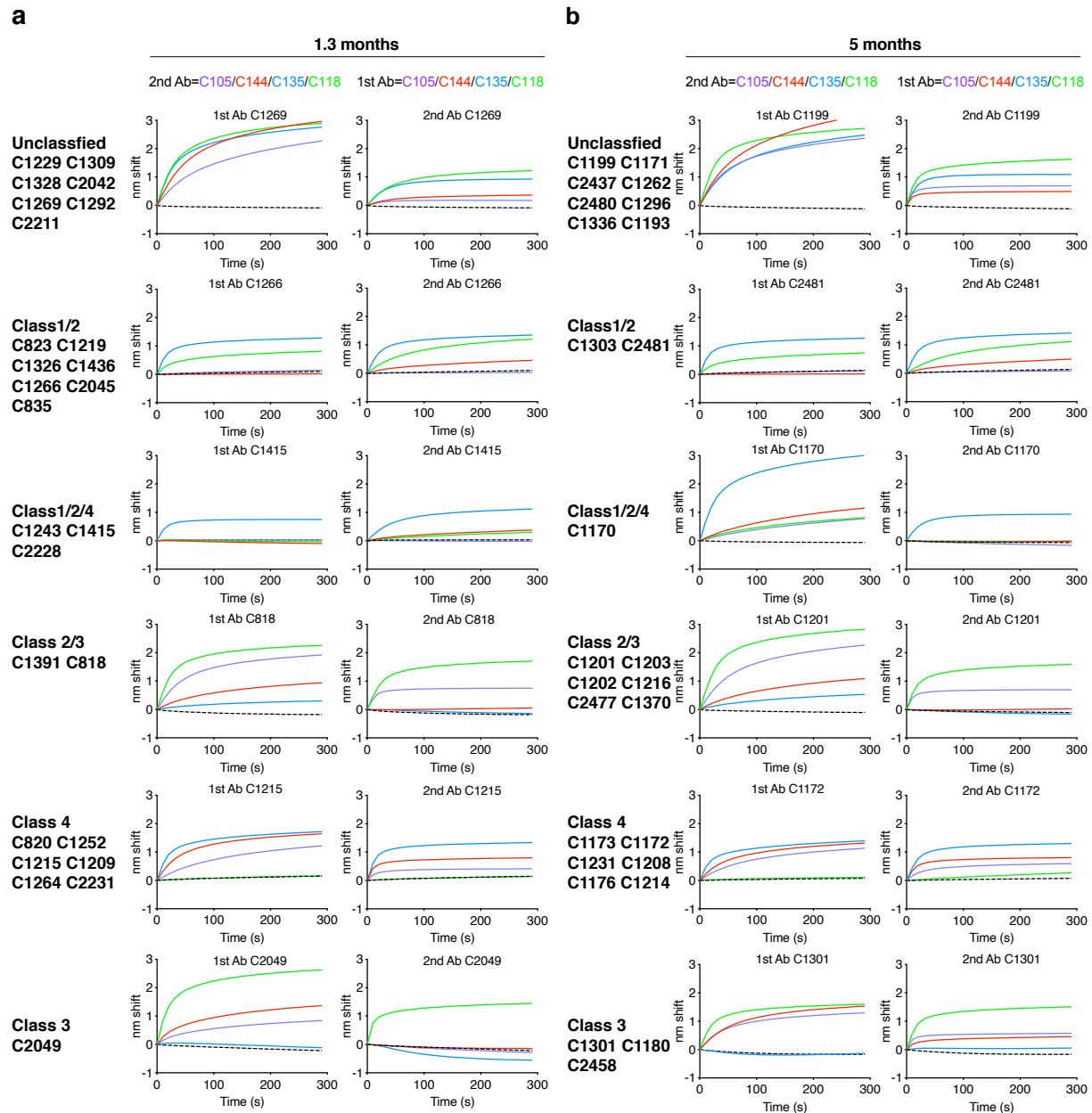


Extended Data Fig. 8. Epitope targeting of anti-SARS-CoV-2 RBD antibodies. **a**, Schematic representation of the BLI experiment for randomly selected antibodies isolated from vaccinees 1.3- and 5 months after full vaccination (each presented group shows n=26 antibodies). **b**, Heat-map of relative inhibition of Ab2 binding to the preformed Ab1-RBD complexes (grey=no binding, yellow=low binding, orange=intermediate binding, red=high binding). Values are normalized through the subtraction of the autologous antibody control. BLI traces can be found in Extended Data Fig. 9. **c**, Pie charts indicate the fraction of antibodies that are assigned to different classes according to their binding pattern as shown in **b** and Extended data Fig. 9. Number in inner

744 circle shows number of antibodies tested. Statistical significance was determined using the Chi-
745 square test in **c**.

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Extended Data Fig. 9. BLI traces from epitope mapping of anti-SARS-CoV-2 RBD antibodies. a-b, BLI traces from competition experiments used to determine epitope targets of anti-SARS-CoV-2 RBD antibodies isolated from vaccinees at 1.3m (a) or 5m (b) post-boost, as illustrated in Extended data Fig. 8.

a

	wt	R683G	R346S	K417N	N440K	A475V	E484K R683G	Q493R	N501Y	KEN R683G	L452R E484Q R683G
C2159	12.2	6.2	9.1	3.3	11.2	8.8	627.1	17.1	20.6	212.0	36.9
C2029	25.1	7.6	>1000	5.9	23.0	19.5	1.8	14.6	57.7	1.6	>1000
C2139	46.0	25.4	33.0	>1000	48.7	>1000	153.6	>1000	>1000	>1000	105.1
C2033	46.2	25.8	32.0	>1000	41.4	868.4	79.5	183.4	668.2	>1000	52.1
C2209	52.6	26.7	40.7	12.5	48.2	214.0	107.3	341.9	214.9	58.8	44.0
C2020	65.8	37.8	164.0	>1000	53.5	71.5	131.3	>1000	119.0	>1000	83.7
C2221	69.3	26.4	47.6	>1000	58.8	>1000	355.2	>1000	46.3	>1000	124.0
C2019	88.5	96.3	>1000	25.5	90.8	70.1	726.4	92.3	125.5	380.3	>1000
C2110	118.4	106.6	92.6	>1000	109.9	797.4	254.9	198.0	301.5	>1000	218.3
C2018	118.9	37.2	55.4	33.4	110.8	98.0	284.5	123.8	135.8	140.0	>1000
C2022	153.4	61.6	114.5	>1000	130.5	>1000	247.7	358.7	162.8	>1000	139.2
C2113	348.4	127.8	242.4	166.8	267.1	347.9	>1000	922.3	339.4	>1000	>1000
C2149	376.8	178.6	259.3	>1000	331.2	>1000	724.8	>1000	>1000	>1000	608.0
C2026	433.1	25.6	258.1	400.1	350.5	925.7	188.0	368.0	387.1	190.1	147.3
C2150	591.0	57.5	672.9	496.9	413.7	783.8	240.8	406.4	543.2	202.1	199.1
C2013	593.3	204.6	391.0	>1000	484.3	117.9	>1000	>1000	>1000	>1000	>1000
C2185	670.6	116.1	440.0	239.1	494.6	818.2	412.8	759.7	485.4	246.7	251.1
C2004	722.5	117.4	529.5	521.1	468.7	>1000	400.1	496.2	928.0	318.2	345.1
C2140	840.9	124.4	706.9	839.5	778.2	>1000	648.7	866.7	815.5	481.7	497.8
C2109	1000.0	198.7	572.6	825.0	336.9	>1000	960.0	762.2	620.2	691.7	464.7

IC50 (ng/ml)

b

	wt	R683G	R346S	K417N	N440K	A475V	E484K R683G	Q493R	N501Y	KEN R683G	L452R E484Q R683G
C2039	1.9	0.5	1.0	0.7	1.2	1.0	>1000	1.4	2.3	>1000	>1000
C2237	6.7	0.7	3.8	2.3	4.7	3.8	342.4	9.1	4.5	815.8	>1000
C2049	10.0	5.2	7.1	319.9	9.6	65.5	10.9	17.5	12.8	>1000	7.2
C2065	11.6	9.8	>1000	4.0	11.1	6.4	387.7	9.6	11.1	123.4	>1000
C2319	13.3	6.3	8.7	>1000	10.2	131.2	7.7	28.4	297.3	>1000	8.5
C2175	17.6	4.9	12.1	5.1	14.2	8.0	506.1	>1000	17.5	347.0	23.5
C2219	20.7	9.8	13.0	5.2	19.4	35.1	>1000	369.6	13.7	>1000	>1000
C2227	48.3	28.5	94.8	20.6	45.0	36.8	>1000	8.7	>1000	>1000	14.3
C2047	49.1	40.4	145.2	>1000	48.8	53.2	168.6	>1000	96.8	>1000	123.4
C2045	52.0	41.2	46.4	>1000	61.1	375.8	62.9	>1000	60.3	>1000	76.4
C2188	90.8	45.7	64.3	>1000	56.6	743.4	134.8	>1000	>1000	>1000	119.6
C2037	148.1	74.1	53.1	32.7	88.0	103.4	378.3	147.5	157.4	246.4	>1000
C2228	178.4	145.9	124.1	70.2	132.2	>1000	770.5	>1000	197.9	886.3	785.3
C2167	200.4	140.1	156.4	13.6	144.2	143.0	287.9	233.5	183.3	28.4	243.6
C2318	351.9	126.5	262.1	113.7	241.7	335.9	286.4	311.0	477.8	244.8	231.4
C2210	366.2	145.7	236.9	188.6	270.5	297.1	382.6	333.2	276.1	381.7	363.0
C2317	429.2	549.9	>1000	105.5	296.6	282.1	>1000	305.6	387.5	>1000	>1000
C2172	451.6	246.0	324.0	214.0	257.3	>1000	363.5	>1000	486.9	>1000	199.7
C2070	584.0	532.7	856.2	260.7	529.0	709.6	884.3	629.7	802.5	838.8	578.2
C2321	843.9	254.3	>1000	648.9	>1000	627.1	400.1	316.5	693.4	445.2	>1000

IC50 (ng/ml)

Extended Data Fig. 10: Breadth of anti-SARS-CoV-2 RBD antibodies elicited after prime and 2 doses of vaccination. a-b, IC₅₀ values for n=40 neutralizing antibodies isolated after prime (a) or 1.3 months post-boost (b) against indicated mutant SARS-CoV-2 pseudoviruses. Color gradient indicates IC₅₀ values ranging from 0 (white) to 1000 ng/ml (red).

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