Computationally restoring the potency of a clinical antibody against Omicron

https://doi.org/10.1038/s41586-024-07385-1

```
Received: 21 October 2022
```

Accepted: 4 April 2024

Published online: 8 May 2024

Open access

Check for updates

Thomas A. Desautels¹, Kathryn T. Arrildt², Adam T. Zemla³, Edmond Y. Lau², Fangqiang Zhu², Dante Ricci², Stephanie Cronin⁴, Seth J. Zost⁴, Elad Binshtein⁴, Suzanne M. Scheaffer⁵, Bernadeta Dadonaite⁶, Brenden K. Petersen¹, Taylor B. Engdahl⁴, Elaine Chen⁴, Laura S. Handal⁴, Lynn Hall⁴, John W. Goforth³, Denis Vashchenko⁷, Sam Nguyen^{1,20}, Dina R. Weilhammer², Jacky Kai-Yin Lo², Bonnee Rubinfeld², Edwin A. Saada², Tracy Weisenberger², Tek-Hyung Lee², Bradley Whitener^{5,21}, James B. Case⁵, Alexander Ladd¹, Mary S. Silva³, Rebecca M. Haluska⁷, Emilia A. Grzesiak³, Christopher G. Earnhart⁸, Svetlana Hopkins⁹, Thomas W. Bates¹, Larissa B. Thackray⁵, Brent W. Segelke², Tri-lab COVID-19 Consortium^{*}, Antonietta Maria Lillo¹⁰, Shivshankar Sundaram¹¹, Jesse D. Bloom^{6,12}, Michael S. Diamond^{5,13,14}, James E. Crowe Jr^{4,15}, Robert H. Carnahan^{4,15} & Daniel M. Faissol^{1⊠}

The COVID-19 pandemic underscored the promise of monoclonal antibody-based prophylactic and therapeutic drugs¹⁻³ and revealed how quickly viral escape can curtail effective options^{4,5}. When the SARS-CoV-2 Omicron variant emerged in 2021, many antibody drug products lost potency, including Evusheld and its constituent, cilgavimab⁴⁻⁶. Cilgavimab, like its progenitor COV2-2130, is a class 3 antibody that is compatible with other antibodies in combination⁴ and is challenging to replace with existing approaches. Rapidly modifying such high-value antibodies to restore efficacy against emerging variants is a compelling mitigation strategy. We sought to redesign and renew the efficacy of COV2-2130 against Omicron BA.1 and BA.1.1 strains while maintaining efficacy against the dominant Delta variant. Here we show that our computationally redesigned antibody, 2130-1-0114-112, achieves this objective, simultaneously increases neutralization potency against Delta and subsequent variants of concern, and provides protection in vivo against the strains tested: WA1/2020, BA.1.1 and BA.5. Deep mutational scanning of tens of thousands of pseudovirus variants reveals that 2130-1-0114-112 improves broad potency without increasing escape liabilities. Our results suggest that computational approaches can optimize an antibody to target multiple escape variants, while simultaneously enriching potency. Our computational approach does not require experimental iterations or pre-existing binding data, thus enabling rapid response strategies to address escape variants or lessen escape vulnerabilities.

The COVID-19 pandemic has underscored the promise of monoclonal antibody-based drugs as prophylactic and therapeutic treatments for infectious disease. Multiple monoclonal antibody drug products that have demonstrated efficacy in preventing COVID-19 (ref. 1) were developed and authorized for emergency use by the US FDA, reducing deaths, hospitalization rates² and reducing viral load³.

Despite these efforts, the SARS-CoV-2 variant Omicron BA.1 escaped many emergency-use monoclonal antibody and antibody combination drug products^{6,7}. First reported in November 2021, BA.1 outcompeted

all other variants of concern (VOCs) worldwide within weeks⁸. BA.1 contains over 50 substitutions, including 15 in the spike protein receptor-binding domain (RBD), the primary target for therapeutic and prophylactic antibodies. These substitutions reduce or eliminate the neutralization capacity of many authorized prophylactic and therapeutic antibodies^{4,5,7}.

In particular, the antibody combination Evusheld—so far, the only antibody drug approved for pre-exposure prophylaxis in immunocompromised patients for whom vaccination is not always protective¹—was

¹Computational Engineering Division, Lawrence Livermore National Laboratory, Livermore, CA, USA. ²Biosciences and Biotechnology Division, Lawrence Livermore National Laboratory, Livermore, CA, USA. ³Global Security Computing Applications Division, Lawrence Livermore National Laboratory, Livermore, CA, USA. ⁴Vanderbilt Vaccine Center, Vanderbilt University Medical Center, Nashville, TN, USA. ⁵Department of Medicine, Washington University School of Medicine, St. Louis, MO, USA. ⁶Basic Sciences Division and Computational Biology Program, Fred Hutchinson Cancer Center, Seattle, WA, USA. ⁷Applications Simulations and Quality Division, Lawrence Livermore National Laboratory, Livermore, CA, USA. ⁸Joint Program Executive Office for Chemical, Biological, Radiological, and Nuclear Defense, US Department of Defense, Frederick, MD, USA. ⁹Joint Rsearch and Development Inc., Stafford, VA, USA. ¹⁰Los Alamos National Laboratory, Biotechnology, Washington University School of Medicine, St. Louis, MO, USA. ¹⁰Howard Hughes Medical Institute, Seattle, WA, USA. ¹⁰Department of Pediatrics, Vanderbilt University Medical Center, Nashville, TN, USA. ¹⁰Porsent address: Google, Alphabet Inc., Mountain View, CA, USA. ²¹Present address: Vir Biotechnology, San Francisco, CA, USA. *A list of authors and their affiliations appears at the end of the paper. ⁵⁶e-mail: faissol1@llnl.gov



Fig. 1 | **Application of the GUIDE computationally driven drug engineering platform to Omicron.** Given a parental antibody and target antigens, a design space was defined and a collection of co-structures were estimated (left). Within the computational design phase (centre), a sequence generator used predictions of multiple properties to propose multi-point mutant antibody candidates, and a Bayesian optimization agent selected proposed sequences that were then simulated. On the basis of Pareto optimality, mutational distance

and sequence diversity, 376 computationally evaluated sequences were selected and experimentally evaluated for binding in immunoassays (centre right). The best sequences were then evaluated for neutralization of SARS-CoV-2 variants, and the single best sequence was identified (right). See Supplementary Methods for details. FEP, free energy perturbation; MD, molecular dynamics; SFE, structural fluctuation estimation.

overwhelmed by Omicron variants. Evusheld combines tixagevimab plus cilgavimab, which are derived from the progenitor monoclonal antibodies COV2-2196 and COV2-2130, respectively. The two-antibody cocktail exhibits 10–100-fold reduction in neutralizing potency against Omicron BA.1 compared with wild-type SARS-CoV-2 (refs. 4,9), but COV2-2130 lost approximately 1,000-fold neutralization potency against Omicron BA.1.1 compared with strains circulating earlier in the pandemic^{7,10,11}.

COV2-2130 is a class 3 RBD-targeting antibody that blocks interaction between the RBD and human angiotensin-converting enzyme (ACE2) without competing with antibodies targeting the class 1 site on the RBD. Thus, class 1 and class 3 antibodies can be combined or co-administered for simultaneous binding and synergistic neutralization¹². Although antibodies that target the class 3 site of the RBD have clear therapeutic utility in antibody combinations, the emergence of Omicron BA.1 and BA.1.1 undermined many antibodies currently available⁴. Furthermore, potently neutralizing antibodies targeting class 3 sites on the RBD are less frequently identified¹², suggesting that they are more difficult to replace.

Computational redesign of a clinically proven monoclonal antibody shows promise for recovering efficacy against escape variants. especially for antibodies known to complement other monoclonal antibodies as part of a combination antibody drug product¹². Thus, we sought to optimize COV2-2130 to restore potent neutralization of escape variants by introducing a small number of mutations in the paratope, then computationally assessing improvement to binding affinity. Our computational approach-generative unconstrained intelligent drug engineering (GUIDE)-combines high-performance computing, simulation and machine learning to co-optimize binding affinity to multiple antigen targets, such as RBDs from several SARS-CoV-2 strains, along with other critical attributes such as thermostability. The computational platform operates in a 'zero-shot' setting; that is, designs are created without iteration through, or input from, wet laboratory experiments on proposed antibody candidates, relatives or other derivatives of the parental antibody. Although more challenging, this zero-shot approach enables rapid production of antibody candidates optimized for multiple target antigens in response to exigencies presented by escape variants. Over a 3-week period, our computational platform repaired the activity of COV2-2130 against Omicron variants. The best-resulting antibody design introduces just four amino acid substitutions into COV2-2130, which could enable an immunobridging strategy in which the established efficacy and safety profile of the parental antibody is leveraged to enable an accelerated regulatory approval and enter clinical use more rapidly and at lower cost. Furthermore, this strategy may provide a rapid pathway for mitigating the threat of future viruses and their continually evolving mutations.

Computational design

Our antibody design platform leverages simulation and machine learning to generate mutant antibody sequences that are co-optimized for multiple critical properties, without requiring experimental feedback or pre-existing binding data (Fig. 1). The platform comprises three phases: problem formulation, computational design and selection of mutant antibody candidates, and experimental validation of proposed candidates.

We formulated a problem by identifying a parental antibody, a set of target antigens and corresponding co-structures. In this case, we redesigned the COV2-2130 antibody¹² for simultaneous binding improvements to Omicron BA.1 and BA.1.1 while maintaining binding to the Delta variant. We used co-structures that were both experimentally determined and computationally estimated, starting from co-structures including the wild-type antigen¹³. As an experimental structure of the Omicron RBD was not available at the onset of our design process, we estimated the structure of the complex of the RBD with COV2-2130 using template-based structural modelling¹⁴. We incorporated experimentally determined Omicron RBD structures¹⁵ as they became available. We considered 25 paratope residues for mutation, primarily in or near the heavy (H) or light (L) chain complementarity determining regions (CDRs)¹⁶, H2, H3, L1 and L2, and allowed up to 9 amino acid substitutions per mutant sequence, resulting in a search space containing over 10¹⁷ possible mutant sequences.

Our computational design approach was implemented as a multiobjective optimization problem defined over this large space of mutations to COV2-2130 paratope residues. We considered five critical antibody properties: (1) binding affinity to the Omicron BA.1 RBD, (2) binding affinity to the BA.1.1 RBD, (3) binding affinity to the Delta RBD, (4) thermostability, and (5) 'humanness'. We expected that restored antibody affinity to each RBD variant would result in restored neutralization because the parental antibody, COV2-2130, competes with ACE2 in SARS-CoV-2 spike binding¹². Four complementary computational tools enabled affinity prediction: atomistic potential of mean force molecular dynamics simulations, structural fluctuation estimation¹⁷, Rosetta Flex¹⁸ and FoldX¹⁹. We estimated thermal stability using the free



Fig. 2 | **Computationally designed IgG antibodies improve Omicron binding and maintain parental thermostability and binding to historical strains. a**, The parental COV2-2130 (orange circles) and computationally designed antibodies (2130-1-0114-112 in purple triangles, 2130-1-0104-024 in blue diamonds and remainder in black) were assayed for thermal shift (n = 3 technical replicates). Melting temperature (T_m) calculated based on the Boltzmann method. Data are mean and s.d. b-e, The parental COV2-2130 antibody and computationally designed antibodies (see symbols in **a**) and cross-reactive positive control antibody S309 (magenta squares) were analysed for relative binding to four SARS-CoV-2 spike RBD variants in the Gyrolab immunoassay: WA1/2020 (**b**), Delta B.1.617.2 (**c**), Omicron BA.1 (**d**) and Omicron BA.1.1 (**e**). Lines represent a four-parameter logistic regression model fit using GraphPad Prism to each titration, executed without technical replicates. mAb, monoclonal antibody.

energy perturbation method²⁰. Humanness was quantified as the score under the AbBERT model²¹, a deep language model trained on a large database of human antibody sequences²². We used these tools to initialize a sequence generator, which proposed multi-residue mutations to the amino acid sequence of COV2-2130. Next, we used distributed software agents, each using Bayesian optimization or rules-based methods, to select a subset of promising candidate sequences to simulate in Rosetta Flex, yielding predicted binding affinities. In less than 3 weeks, we evaluated more than 125,000 antibody candidates.

We calculated the Pareto optimal set²³ based on the outputs of these tools, resulting in 3,809 sequences. Owing to experimental capacity, we further downselected from among the Pareto set based on mutational distance and sequence diversity to ultimately designate 376 antibody sequences for experimental evaluation.

Experimental evaluation

Antibody and antigen production

We experimentally validated the 376 designed candidates. To leverage available resources at multiple experimental sites, we split candidates into partially overlapping sets 1 and 2. Set 1 consisted of 230 designs expressed as IgG in HEK-293 cells (ATUM), and set 2 consisted of 204 designs expressed as IgG via a pVVC-mCisK_hG1 vector (Twist BioScience) in transiently transfected CHO cells. Omicron antigens were produced in Expi293F cells (Thermo Fisher Scientific) and purified on HisTrap Excel columns (Cytiva).

In the following experiments, we selected antigens or viral strains to gauge the success of three goals: (1) improving binding affinity and efficacy to BA.1 and BA.1.1; (2) maintaining efficacy to historical strains, for which design explicitly targeted Delta but experiments often substituted WA1/2020 D614G; and (3) determining robustness to emerging VOCs.

Designed antibodies maintain expression

Because in silico derivatization of antibody sequences can compromise production yield, we measured the concentrations of the 230 COV2-2130-derived recombinant antibodies in set 1 and compared these concentrations to that of the parental antibody. The purified concentrations of 73.9% of redesigned antibodies exceeded that of the parental COV2-2130 antibody (170 of 230 monoclonal antibodies at more than 171.2 mg l^{-1}), reaching as high as 305 mg l^{-1} . Our designed candidates for downstream characterization retained fundamental production properties of the parental antibody, with just 10% of designed antibodies producing poor yields relative to the parental molecule (22 of 230 monoclonal antibodies at less than 135 mg l^{-1} , that is, less than 80% of the parental antibody yield).

Thermostability and binding Omicron

We screened all designed antibodies for binding to RBDs. Set 1 was screened via a single-concentration immunoassay (Gyrolab xPlore) in the contexts of WA1/2020, Delta, BA.1 or BA.1.1 RBDs (Extended Data Fig. 1). For set 2, we used a multi-concentration immunoassay (ELISA: Extended Data Fig. 2) in the context of wild-type, BA.1 or BA.1.1 RBDs. In the single-concentration immunoassay, this value was chosen as a single dilution factor, causing most designed antibody samples to fall in the dynamic range of the positive control. In both cases, we compared the binding of the designed antibodies with a broadly cross-reactive, non-ACE2-competitive control antibody (S309)²⁴ and the parental COV2-2130 antibody. As intended, most antibody designs had altered binding profiles, indicating that the designed mutations were consequential. Approximately 11% of the designs of set 1 retained WA1/2020 antigen binding at the measured concentration; roughly 6% improved binding to BA.1, and 5% did so to BA.1.1. The corresponding numbers for set 2 were 9% to WA1/2020 and 8% to BA.1. Following this initial screen, we downselected both sets of antibody designs to those with improved binding to Omicron subvariants BA.1 and BA.1.1 for further characterization.

These downselected antibodies were re-manufactured at larger scale. We characterized the resulting IgG antibodies by immunoassay and thermal shift (melt temperature) assessments. In agreement with our screens, seven of the eight top-performing antibodies preserved comparable binding with WA1/2020 and Delta RBDs, improving over the parental COV2-2130 antibody with respect to their binding to Omicron BA.1 and BA.1.1 RBDs (Fig. 2). Furthermore, seven of the eight antibodies had melting temperatures and expression properties comparable with those of COV2-2130. One antibody, 2130-1-0114-111, had reduced



Fig. 3 | **Designed antibodies improve pseudoviral neutralization over COV2-2130. a**–**e**, The parental COV2-2130 antibody (orange circles), the crossreactive positive control antibody S2K146 (magenta squares), the negative control antibody DENV-2D22 (grey x) and down-selected computationally designed antibodies (symbols as indicated in the key) were assayed by

melting temperature (Extended Data Table 1). The antibody 2130-1-0114-112 displayed best-in-class binding across all RBD variants and had no substantial difference in thermal stability compared with the parental COV2-2130 antibody.

Restored pseudoviral neutralization

We performed pseudovirus neutralization assays to characterize the functional performance of five selected antibody designs (Fig. 3 and Extended Data Table 2), compared with parental COV2-2130; the positive control S2K146 (ref. 25), which competes with ACE2 binding; and the negative control DENV-2D22 (ref. 26). Our designs maintained neutralization activity against pseudoviruses displaying historical spike proteins (WA1/2020 D614G) and achieved neutralization of those with Omicron BA.1 spikes. The single-best candidate design, 2130-1-0114-112, restored potent neutralization in the context of BA.1.1 and showed a two-order-of-magnitude improvement in the half-maximal inhibitory concentration (IC₅₀) versus parental COV2-2130 for BA.1 and BA.4. These pseudovirus neutralization test results showed that our designs neutralized BA.2 and BA.4 more potently than COV2-2130, despite the emergence of these VOCs after the conception of our designs.

Restored authentic virus neutralization

We evaluated 2130-1-0114-112 (containing four mutations: GH112E, SL32A, SL33A and TL59E) for authentic virus neutralization performance against several strains of SARS-CoV-2 by a focus reduction neutralization test in Vero-TMPRSS2 cells (Extended Data Fig. 3). The strains that we used included several Omicron targets: WA1/2020 D614G, Delta (B.1.617.2), BA.1, BA.1.1, BA.2, BA.2.12.1, BA.4, BA.5 and BA.5.5. In all cases apart from Delta, 2130-1-0114-112 had an IC₅₀ < 10 ng ml⁻¹. Compared with the parental COV2-2130, 2130-1-0114-112 restored potent neutralization activity to both BA.1 (8.08 ng ml⁻¹) and BA.1.1 (7.77 ng ml⁻¹), showed a more than fivefold improvement in IC $_{50}$ to BA.2 (2.4 ng ml $^{-1}$) and BA.2.12.1 (2.27 ng ml⁻¹), and conferred 50-fold or greater improvements in IC₅₀ to BA.4 (3.16 ng ml⁻¹), BA.5 (3.51 ng ml⁻¹) and BA.5.5 (5.29 ng ml⁻¹). We also evaluated 2130-1-0114-112 and a less-mutated alternative design, 2130-1-0104-024 (containing two mutations: SL32W and TL59E), in plaque assays with Vero E6-TMPRSS2-T2A-ACE2 cells (Extended Data Fig. 4). IC₅₀ values for 2130-1-0104-024 were 37.7 ng ml⁻¹, 75.94 ng ml⁻¹ and 781.7 ng ml⁻¹ for Delta, BA.1 and BA.1.1 viruses, respectively.

neutralization with lentiviruses pseudotyped with spike variants of WA1/2020 D614G (**a**), Omicron BA.1 (**b**), Omicron BA.1.1 (**c**), Omicron BA.2 (**d**) and Omicron BA.4 (**e**). Curves are four-parameter logistic regression models fit to two (**a**-**d**) or four (**e**) replicate serial dilutions using GraphPad Prism.

Prophylaxis in vivo

To compare the prophylactic efficacy of 2130-1-0114-112 and the parental COV2-2130 monoclonal antibody in vivo, we administered a single 100 µg (approximately 5 mg kg⁻¹ total) dose to K18-hACE2 transgenic mice 1 day before intranasal inoculation with WA1/2020 D614G, BA.1.1 or BA.5 (88 mice in total, 9-10 for each monoclonal antibody and viral strain). Although Omicron viruses are less pathogenic in mice than in humans, they replicate efficiently in the lungs of K18-hACE2 mice^{27,28}. Viral RNA levels were measured at 4 days post-infection in the nasal washes, nasal turbinates and lungs (Fig. 4). As expected, the parental COV2-2130 monoclonal antibody effectively reduced WA1/2020 D614G infection in the lungs (180,930-fold), nasal turbinates (42-fold) and nasal washes (25-fold) compared with the isotype control monoclonal antibody. However, the COV2-2130 monoclonal antibody lost protective activity to BA.1.1 in all respiratory tract tissues, whereas to BA.5. protection was maintained in the lungs (13.622-fold) but not in the nasal turbinates or nasal washes. Compared with the isotype control monoclonal antibody (Fig. 4), 2130-1-0114-112 protected against lung infection by WA1/2020 D614G (399,945-fold reduction), BA.1.1 (53,468-fold reduction) and BA.5 (160,133-fold reduction). Moreover, in the upper respiratory tract, 2130-1-0114-112 also conferred protection to WA1/2020 D614G, BA.1.1 and BA.5. The differences in protection between the parental COV2-2130 and derivative 2130-1-0114-112 monoclonal antibodies were most apparent in mice infected with BA.1.1, which directly parallels the neutralization data (Fig. 3 and Extended Data Figs. 3 and 4).

Potency without additional liabilities

To understand the neutralization breadth of 2130-1-0114-112 relative to its ancestral antibody, we mapped the epitopes for both antibodies using spike-pseudotyped lentiviral deep mutational scanning (DMS)²⁹. For each antibody, we mapped escape mutations in both the BA.1 and the BA.2 spikes. DMS experiments showed that the escape profile of both COV2-2130 and 2130-1-0114-112 in the context of both BA.1 and BA.2 backgrounds is consistent with the epitope of the antibodies, but with differences in sensitivity to particular mutations (Fig. 5). Consistent with live and pseudovirus neutralization assays (Fig. 3 and Extended Data Figs. 3 and 4), mutations at RBD positions R346 and L452 are sites of substantial escape from both antibodies (Fig. 5). In addition, both antibodies lose potency with mutations at site K444 (such as K444T



Fig. 4 | **2130-1-0114-112 provides in vivo prophylactic protection against SARS-CoV-2 variants. a**–i, Eight-week-old female K18-hACE2 mice were administered 100 µg (approximately 5 mg kg⁻¹) of the indicated monoclonal antibody treatment by intraperitoneal injection 1 day before intranasal inoculation with 10⁴ focus-forming units (FFU) of WA1/2020 D614G (a,d,g), Omicron BA.1.1 (**b,e,h**) or Omicron BA.5 (**c,f,i**). Tissues were collected 4 days after inoculation. Viral RNA levels in the lungs (**a**–**c**), nasal turbinates (**d**–**f**) and nasal washes (**g**–**i**) were determined by RT–qPCR (lines indicate median of log₁₀ values); n = 9 (WA1/2020 D614G and BA.1.1 isotype control groups) or 10 (all others) mice per group, from two experiments. The limit of assay detection is shown as a horizontal dotted line. Statistical comparisons between groups were by Kruskal–Wallis ANOVA with Dunn's multiple comparisons post-test; *P* values are as listed or not significant (NS) if *P* > 0.05. All analyses were conducted in GraphPad Prism.

found in BO.1* variants). The reversion mutation S446G in the BA.1 background increases the neutralization potency of both antibodies (negative escape values in heatmaps) (Fig. 5c) and probably contributes to greater neutralization potency to the BA.2 variant (Fig. 3 and Extended Data Fig. 3), which carries G446. Most mutations at RBD sites K440 and R498 are slightly sensitizing to the COV2-2130 antibody in both BA.1 and BA.2 backgrounds, but provide weak escape for 2130-1-0114-112 in the BA.1 background and have even weaker effect in the BA.2 background. In agreement with pseudovirus neutralization assays (Fig. 3), comparison of mutation-level escape showed that the 2130-1-0114-112 antibody is substantially more potent than COV2-2130 to the BA.1 variant and retains better potency against viruses with additional mutations in both BA.1 and BA.2 backgrounds (Fig. 5a,b). However, even with improved potency, 2130-1-0114-112 is still vulnerable to escape at multiple RBD residues in the 444-452 loop, which is the site of convergent substitutions in several Omicron lineages³⁰. Many of these variants contain multiple substitutions at positions identified by DMS as important for neutralization or in close proximity to the COV2-2130 epitope, including BQ.1.1 (R346T and K444T), XBB (R346T, V445P and G446S) and BN.1 (R346T, K356T and G446S). To understand the impact of these VOCs, we assessed the ability of 2130-1-0114-112 to neutralize BQ.1.1, XBB and BN.1 in pseudoviral neutralization studies. Consistent with the previously known liabilities of COV2-2130 and our DMS results, 2130-1-0114-112 loses neutralizing activity to these VOCs (Extended Data Fig. 5), probably due to substitutions at 444 and combinatorial effects of multiple substitutions within the COV2-2130 epitope present in these variants. Together, these data demonstrate that 2130-1-0114-112 exhibits improved potency against many individual substitutions without incurring additional escape liabilities, although RBD residues such as 444 remain critical for neutralization activity of both 2130-1-0144-112 and COV2-2130.

Structural basis for restored potency. To elucidate the key intermolecular interactions that form the interface and determine Omicron RBD recognition by 2130-1-0114-112, we performed 3D reconstructions of the complex between the SARS-CoV-2 Omicron BA.2 spike and the 2130-1-0114-112 Fab fragment using cryo-electron microscopy (cryo-EM). Reconstruction using refinement of the full complex gave a map with average resolution of 3.26 Å, but the interface region between the BA.2 RBD and the 2130-1-0114-112 Fab was not well resolved, presumably due to the flexibility of the RBD–Fab region in the reconstruction. To resolve details at the intermolecular interface, we performed focused refinement of this portion of the structure. Focused refinement resulted in an effective resolution of approximately 3.6 Å for this region (Electron Microscopy Data Bank EMD-28198 and EMD-28199, and Protein Data Bank 8EKD) (Fig. 6 and Extended Data Fig. 6).

This model shows the binding interface of 2130-1-0114-112–RBD and elucidates how 2130-1-0114-112 regains neutralization potency to Omicron VOCs. The parental COV2-2130 forms extensive interactions with the RBD through CDRH2 and CDRH3, as well as CDRL1 and CDRL2 (ref. 13) with hydrogen bond networks and hydrophobic interactions. To improve binding interactions with Omicron subvariants, 2130-1-0114-112 modifies three CDR loops: G112E in CDRH3, S32A and S33A in CDRL1, and T59E in CDRL2.

The RBD N440K substitution, identified in the DMS as sensitizing for escape from COV2-2130 but less so for 2130-1-0114-112, is on the edge of the interface with the 2130-1-0114-112 CDRH3 loop and does not make direct contact with the CDRH3 substitution G112E. However, N440K introduces a positive charge to a local environment that has substantial hydrophobic-to-hydrophobic contact. The negative charge introduced by the CDRH3 G112E substitution (Fig. 6b) might improve the electrostatic interactions in this region. It is possible that E112 and K440 are interacting by coordinating a water molecule, but the structural resolution is not sufficient to confirm this type of interaction. These experimental structural results are also consistent with our molecular dynamics simulations, which showed this transient interaction between CDRH3 E112 and RBD K440.

The local environment around the CDRL1 loop is mostly hydrophobic (comprising the RBD residues L452, F490 and L492, as well as the Omicron mutation E484A) with a hydrogen bond from LC N34 (Fig. 6c). The hydrophilic-to-hydrophobic CDRL1 substitutions introduced in 2130-1-0114-112, S32A and S33A, may favour the local environment and strengthen hydrophobic interactions with the RBD (Fig. 6c,e). This is supported by the DMS identification of sensitivity to hydrophobic-to-hydrophilic substitutions at RBD position 452 for both 1230-1-0114-112 and the parental COV2-2130. Finally, the T59E mutation in the CDRL2 loop establishes a new salt bridge with the RBD substitution Q498R present in Omicron RBDs. This new salt bridge probably strengthens the interaction with the RBD (Fig. 6d,e).

2130-1-0114-112 distributes four substitutions across three of the four CDR loops comprising the parental COV2-2130 paratope. Mutations to CDRH3 loop were less fruitful than mutations in the L1 and L2 (Extended Data Fig. 7a compared with Extended Data Fig. 7d) when looking across all antibody candidates. Among successful candidates, substitutions at positions 32 and 33 in CDRL1 appear enriched—particularly with hydrophobic residues—consistent with our analysis of this region of the experimentally solved structure of 2130-1-0114-112–BA.2 spike. Another candidate, 2130-1-0104-024, achieves improved affinity and neutralization with only two substitutions: S32W in CDRL1 and T59E in CDRL2. However, full neutralization potency is not reached without the potential charge accommodation mediated by G112E. This suggests



Fig. 5 | COV-2130 and 2130-1-0114-112 escape mapping using DMS. a, b, Comparison between IC50 values measured using DMS for COV-2130 and 2130-1-0114-112 antibodies in BA.1 (a) and BA.2 (b) backgrounds, with key mutations highlighted. Arbitrary units in both plots are on the same scale. Interactive plots that display each mutation can be found at https://dms-vep.org/SARS-CoV-2 Omicron BA.1 spike DMS COV2-2130/compare IC50s.html for the BA.1 background and at https:// dms-vep.org/SARS-CoV-2_Omicron_BA.2_spike_ DMS COV2-2130/compare IC50s.html for the BA.2 background, c.d. Heatmaps of mutation escape scores at key sites for each antibody in BA.1(c) and BA.2(d) backgrounds. Escape scores were calculated relative to the wild-type amino acid in the same virus background. X marks wild-type amino acid in the relevant background. Amino acids not present in the DMS libraries lack squares; grey squares are mutations that strongly impair spike-mediated infection. Mutations identified in **a**.**b** are shown with a heavy line surrounding the corresponding box in **c**, **d**. Interactive heatmaps for full spike can be found for the BA.1 background at https://dms-vep.org/ SARS-CoV-2_Omicron_BA.1_spike_DMS_COV2-2130/COV2-2130_vs_2130-1-0114-112_escape.html and https://dms-vep.org/SARS-CoV-2 Omicron BA.2_spike_DMS_COV2-2130/COV2-2130_vs_2130-1-0114-112_escape.html for the BA.2 background.

that a combination of new bonds and accommodating charge changes optimized the restored affinity and potency of 2130-1-0114-112 with Omicron variants (Extended Data Fig. 8). Altogether, the structural model of 2130-1-0114-112 with the BA.2 RBD helps explain the observed restoration of potency to early SARS-CoV-2 Omicron VOCs.

Discussion

We set out to rapidly design and validate derivatives of the COV2-2130 antibody that restore potent in vitro neutralization to BA.1 and BA.1.1 Omicron subvariants while maintaining binding and neutralization to previous strains of SARS-CoV-2. In addition, we sought to retain favourable thermostability properties and maintain the humanness of the sequences, a data-driven measure of similarity to known human sequences. Despite multiple mutations in the COV2-2130 epitope present in Omicron BA.1 and BA.1.1, we achieved these design objectives by applying a computationally driven, multi-objective approach. We chose to take a risk-seeking approach that increased the chance of obtaining at least one highly potent design, ideally several, by choosing diverse sequences predicted to have substantial effects on binding.

Several designed antibody candidates successfully restored neutralization potency to Omicron subvariants. In our top antibody design, 2130-1-0114-112, four substitutions accommodate Omicron escape mutations in BA.1 and BA.1.1 without sacrificing potency against Delta. This engineered antibody is thermostable; potently neutralizes Omicron BA.2, BA.4, BA.5 and BA.5.5; and restores prophylactic efficacy in vivo. Our approach for extending the utility of a high-value antibody complements state-of-the-art ex vivo discovery of high-value antibodies with responsive computational modification.

The distributed nature of the improvements in 2130-1-0114-112, with four mutations spanning three CDRs, plausibly makes this antibody comparatively robust to subsequent escape, insofar as DMS results demonstrate an improvement in binding over the parental COV2-2130 where escape substitutions present in the design targets, BA.1 and BA.1.1, are mitigated without new vulnerabilities. 2130-1-0114-112 does not mitigate the reliance of the parental COV2-2130 antibody on the RBD residue K444 and sensitivity to substitutions at this position. Although our top candidate does not neutralize the BQ.1.1 and XBB variants, which contain multiple substitutions within the COV2-2130 epitope, DMS results indicate that 2130-1-0114-112 reduces the impact of some of the mutations of the variants.

Our design approach shows potential for expediting the path of new drug products to clinical use, including lower development costs and risks versus identifying wholly new drug products of comparable breadth and efficacy. Our top-performing antibody restores in vivo efficacy and achieves potent and broad neutralization of many SARS-CoV-2



Fig. 6 | **Cryo-EM structure of neutralizing antibodies 2130-1-0114-112 in complex with Omicron BA.2 RBD. a**, Atomic model of the RBD–Fab complex, coloured by chain: BA.2 RBD in red, 2130-1-0114-112 HC in yellow and 2130-1-0114-112 LC in green. BA.2 RBD mutations are in orange, and 2130-1-0114-112 mutations are in cyan and blue (HC and LC) (left). A close-up view of the RBD–Fab interface, showing WA1 RBD (Protein Data Bank 7L7E, light brown shading) aligned with the BA.2 RBD (right). **b**–**d**, Details showing the 2130-1-0114-112 modified residues and their interaction with BA.2 RBD, coloured as in **a**. Residue labels are shown in black for the BA.2 complex and brown for the overlaid WA1-2130 complex. The orange and green dashed lines indicate hydrogen bond and hydrophobic interactions, respectively; the yellow dashed lines are labelled with distances. CDRH3 residue Glu112 (left) and with the

surface coloured by electrostatic potential (right), showing the positive and negative charges of RBD Lys444 and CDRH3 Glu112 (**b**). CDRL1 Ala32 and Ala33 hydrophobic network (left) and with the nearby RBD surface coloured by hydrophobicity (right; orange to cyan indicates hydrophobic to hydrophilic) (**c**). CDRL2 Glu59 salt bridge with RBD residue Arg498 (**d**). **e**, 2D diagram of Fab 2130-1-0114-112 paratope and epitope residues involved in hydrogen bonds and salt bridges (yellow and red dashed lines, respectively; distances in Å) and hydrophobic interactions (curved lines with rays). Atoms are shown as circles, with oxygen, carbon and nitrogen in red, black and blue, respectively. Interacting residues that belong to CDR loops are coloured in corresponding shades. The asterisks indicate mutated residues. Image created with Ligplot+³⁴.

VOCs by substituting just four amino acids into a parental antibody that has been extensively tested for safety, manufacturability and clinical efficacy¹. Given increasing evidence that neutralization is a correlate of protection from severe COVID-19 in patients treated with monoclonal antibody therapies^{31,32}, an immunobridging strategy has been proposed as a response to rapidly evolving SARS-CoV-2 variants to shorten the pathway of improved monoclonal antibodies to clinical use³³. Rapid computational rescue of high-value, potentially rare, antibodies in clinical use presents a high-impact, real-world application of our work that could be made more impactful with such an immunobridging strategy. We demonstrate successful re-targeting without requiring major sequence changes or acquisition of new liabilities. The urgency for a design approach like ours is clear given that existing antibody drug development approaches are struggling to match the rapid pace of SARS-CoV-2 evolution.

Although the individual components comprising our approach are built on existing computational approaches, we integrate them into a novel framework that demonstrates (1) a computational approach to antibody optimization that gains neutralization to a new target, (2) successful optimization of an antibody to achieve high potency to multiple targets (for example, multiple escape variants) without requiring experimental iterations, and (3) computationally restoring or improving efficacy with in vivo validation. The computational approach that we used in this work did not require iterative improvement based on feedback from experimental evaluations, nor did it require availability of data on antibody candidates tested against the target antigens, either of which would result in further delays when responding to an emergent variant. Furthermore, our fundamental approach is adaptable to more modest or decentralized computing resources than those used in this study.

Future work seeks to extend our computational approach to include additional predictive models, such as models predicting antibody expression, protein aggregation and polyreactivity. Our models for predicting antibody-antigen binding heavily depend on performing simulations with sufficiently accurate models of antibody-antigen co-structures, which is an important limitation. Consequently, we are developing experimental datasets to advance machine learning-based approaches for predicting binding directly from sequence, as well as incorporating emerging artificial intelligence-based approaches for determining and refining structural models.

In this study, we demonstrate an innovative computational methodology capable of creating an array of antibody designs targeting the initial subvariants of Omicron SARS-CoV-2. A subset of these designed antibodies display enhancements over the parental COV2-2130 antibody, including superior binding, broad and potent neutralization, and in vivo protection against Omicron BA.1.1. Our approach demonstrates an adaptable antibody-based therapeutic discovery strategy, enabling rapid deployment in response to emerging viral threats or evolutionary shifts. Furthermore, the limited number of amino acid substitutions in our redesigned antibodies suggests the feasibility of an immunobridging strategy for accelerated regulatory approval, especially if the parental antibody has received regulatory clearance for use in humans. Our computational method can also proactively mitigate liabilities identified via DMS, potentially limiting the impact of escape variants and thereby extending the therapeutic utility of the designed antibody in a clinical setting.

Online content

Any methods, additional references, Nature Portfolio reporting summaries, source data, extended data, supplementary information, acknowledgements, peer review information; details of author contributions and competing interests; and statements of data and code availability are available at https://doi.org/10.1038/s41586-024-07385-1.

- Levin, M. J. et al. Intramuscular AZD7442 (tixagevimab–cilgavimab) for prevention of Covid-19. N. Engl. J. Med. 386, 2188–2200 (2022).
- Dougan, M. et al. Bamlanivimab plus etesevimab in mild or moderate Covid-19. N. Engl. J. Med. 385, 1382–1392 (2021).
- Weinreich, D. M. et al. REGN-COV2, a neutralizing antibody cocktail, in outpatients with Covid-19. N. Engl. J. Med. 384, 238–251 (2020).
- 4. VanBlargan, L. A. et al. An infectious SARS-CoV-2 B.1.1.529 Omicron virus escapes
- neutralization by therapeutic monoclonal antibodies. *Nat. Med.* 28, 490–495 (2022).
 Iketani, S. et al. Antibody evasion properties of SARS-CoV-2 Omicron sublineages. *Nature* 604, 553–556 (2022).
- Case J B et al. Resilience of \$309 and A7D7442 monoclonal antibody treatments
- against infection by SARS-CoV-2 Omicron lineage strains. *Nat. Commun.* **13**, 3824 (2022). 7. Wang, Q. et al. Alarming antibody evasion properties of rising SARS-CoV-2 BQ and XBB
- subvariants. *Cell* **186**, 279–286.e8 (2023). 8. Viana, R. et al. Rapid epidemic expansion of the SARS-CoV-2 Omicron variant in southern
- Africa. Nature **603**, 679–686 (2022). 9. Cameroni, E. et al. Broadly neutralizing antibodies overcome SARS-CoV-2 Omicron
- antigenic shift. Nature 602, 664–670 (2022).
- Tuekprakhon, A. et al. Antibody escape of SARS-CoV-2 Omicron BA.4 and BA.5 from vaccine and BA.1 serum. Cell 185, 2422–2433.e13 (2022).
- Takashita, E. et al. Efficacy of antibodies and antiviral drugs against Omicron BA.2.12.1, BA.4, and BA.5 subvariants. N. Engl. J. Med. 387, 468–470 (2022).
- Zost, S. J. et al. Potently neutralizing and protective human antibodies against SARS-CoV-2. Nature 584, 443–449 (2020).
- Dong, J. et al. Genetic and structural basis for SARS-CoV-2 variant neutralization by a two-antibody cocktail. Nat. Microbiol. 6, 1233–1244 (2021).

- Zemla, A. et al. AS2TS system for protein structure modeling and analysis. Nucleic Acids Res. 33, W111–W115 (2005).
- Mannar, D. et al. SARS-CoV-2 Omicron variant: antibody evasion and cryo-EM structure of spike protein–ACE2 complex. Science 375, 760–764 (2022).
- Sela-Culang, I., Kunik, V. & Ofran, Y. The structural basis of antibody-antigen recognition. Front. Immunol. https://doi.org/10.3389/fimmu.2013.00302 (2013).
- Zemla, A. et al. SARS-COV-2 Omicron variant predicted to exhibit higher affinity to ACE-2 receptor and lower affinity to a large range of neutralizing antibodies, using a rapid computational platform. Preprint at *bioRxiv* https://doi.org/10.1101/2021.12.16.472843v1 (2021).
- Barlow, K. A. et al. Flex ddG: Rosetta Ensemble-based estimation of changes in protein-protein binding affinity upon mutation. J. Phys. Chem. B 122, 5389–5399 (2018).
- Schymkowitz, J. et al. The FoldX web server: an online force field. Nucleic Acids Res. 33, W382-W388 (2005).
- Zhu, F. et al. Large-scale application of free energy perturbation calculations for antibody design. Sci. Rep. 12, 12489 (2022).
- 21. Vashchenko, D. et al. AbBERT: learning antibody humanness via masked language modeling. Preprint at *bioRxiv* https://doi.org/10.1101/2022.08.02.502236v1 (2022).
- Olsen, T. H., Boyles, F. & Deane, C. M. Observed Antibody Space: a diverse database of cleaned, annotated, and translated unpaired and paired antibody sequences. *Protein Sci.* 31, 141–146 (2022).
- Ehrgott, M. Multicriteria Optimization Vol. 491 (Springer Science & Business Media, 2005).
 Pinto, D. et al. Cross-neutralization of SARS-CoV-2 by a human monoclonal SARS-CoV
- antibody. Nature 583, 290–295 (2020).
 Park, Y.-J. et al. Antibody-mediated broad sarbecovirus neutralization through ACE2 molecular mimicry. Science 375, 449–454 (2022).
- de Alwis, R. et al. Identification of human neutralizing antibodies that bind to complex epitopes on dengue virions. Proc. Natl Acad. Sci. USA 109, 7439–7444 (2012).
- 27. Uraki, R. et al. Characterization and antiviral susceptibility of SARS-CoV-2 Omicron BA.2. Nature **607**, 119–127 (2022).
- Ying, B. et al. Boosting with variant-matched or historical mRNA vaccines protects against Omicron infection in mice. Cell 185, 1572–1587.e11 (2022).
- Dadonaite, B. et al. A pseudovirus system enables deep mutational scanning of the full SARS-CoV-2 spike. Cell 186, 1263–1278.e20 (2023).
- Cao, Y. et al. Imprinted SARS-CoV-2 humoral immunity induces convergent Omicron RBD evolution. Nature 614, 521–529 (2023).
- Stadler, E. et al. Determinants of passive antibody efficacy in SARS-CoV-2 infection: a systematic review and meta-analysis. *Lancet Microbe* 4, e883–e892 (2023).
- Schmidt, P. et al. Antibody-mediated protection against symptomatic COVID-19 can be achieved at low serum neutralizing titers. *Sci. Transl. Med.* 15, eadg2783 (2023).
- US Food and Drug Administration and European Medicines Agency. Summary report of the joint EMA–FDA workshop on the efficacy of monoclonal antibodies in the context of rapidly evolving SARS-CoV-2 variants. FDA https://www.fda.gov/media/165344/download (2023).
- Laskowski, R. A. & Swindells, M. B. LigPlot+: multiple ligand-protein interaction diagrams for drug discovery. J. Chem. Inf. Model. 51, 2778–2786 (2011).

Publisher's note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.



Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate

credit to the original author(s) and the source, provide a link to the Creative Commons licence, and indicate if changes were made. The images or other third party material in this article are included in the article's Creative Commons licence, unless indicated otherwise in a credit line to the material. If material is not included in the article's Creative Commons licence and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this licence, visit http://creativecommons.org/licenses/by/4.0/.

© The Author(s) 2024

Tri-lab COVID-19 Consortium

Emily Z. Alipio Lyon¹⁰, Penelope S. Anderson¹⁰, Kathryn T. Arrildt², Aram Avila-Herrera³, Thomas W. Bates¹, William F. Bennett², Feliza A. Bourguet¹⁶, Julian C. Chen¹⁰, Matthew A. Coleman², Nicole M. Collette², Anastasiia Davis¹⁰, Thomas A. Desautels¹, Byron D. Vannest², Daniel M. Faissol¹, Erika J. Fong¹¹, Sean Gilmore², John W. Goforth³, Andre R. Goncalves¹, Emilia A. Grzesiak³, Sara B. Hall², Rebecca M. Haluska⁷, Brooke Harmon¹⁷, Wei He², Steven A. Hoang-Phou², Alexander Ladd¹, Mikel Landajuela¹, Edmond Y. Lau², Ted A. Laurence¹⁸, Tek Hyung Lee², Felipe Leno Da Silva¹, Chao Liu², Jacky Kai-Vin Lo², Terrel N. Mundhenk¹, Mariam V. Mohagheghi², Peter R. McIlroy¹⁷, Brenden K. Petersen¹, Le Thanh Mai Pham¹⁰, Dante Ricci², Bonnee Rubinfeld², Edwin A. Saada², Joseph C. Sanchez¹⁰, Nicholas Watkins¹¹, Dina R. Weilhammer², Tracy Weisenberger², Jiachen Yang¹, Congwang Ye¹¹, Adam T. Zemla³, Boya Zhang¹ & Fangqiang Zhu²

¹⁶Global Security Principal Directorate, Lawrence Livermore National Laboratory, Livermore, CA, USA. ¹⁷Biotechnology and Bioengineering, Sandia National Laboratories, Livermore, CA, USA. ¹⁸Materials Science Division, Lawrence Livermore National Laboratory, Livermore, CA, USA. ¹⁹Bioresource and Environmental Security, Sandia National Laboratories, Livermore, CA, USA.

Reporting summary

Further information on research design is available in the Nature Portfolio Reporting Summary linked to this article.

Data availability

The EM map and model have been deposited to the Electron Microscopy Data Bank (EMD-28198 and EMD-28199) and the Protein Data Bank (PDB;8EKD). Other protein structural data were used in this work (PDB 7L7E and 7T9K) and in analysis (PDB 7X66, 7XAZ, 8IOS, 8IF2 and 8GB8). Sequence data that support the findings of this study have been deposited in GenBank under accessions PP474664–PP474679 and are available in the Supplementary Information. Source data for Fig. 4 are provided with the paper. DMS library variant data and antibody per replicate DMS selection data can be accessed at https://github. com/dms-vep/SARS-CoV-2_Omicron_BA.2_spike_DMS_COV2-2130 and https://github.com/dms-vep/SARS-CoV-2_Omicron_BA.1_spike_DMS_ COV2-2130 GitHub repositories. Source data are provided with this paper.

Code availability

The ImprovWF dynamic workflows code is available at https://github. com/LLNL/improvwf. Selected GUIDE workflow components are available at https://github.com/LLNL/CRPCA. All DMS data analysis was performed using dms-vep-pipeline version 1.8, which can be accessed at https://github.com/dms-vep/dms-vep-pipeline/tree/51e73d601 bd770eb6e9abd21f57fb4365699c984. Code and notebooks related to DMS runs are available at https://dms-vep.github.io/SARS-CoV-2_ Omicron_BA.1_spike_DMS_COV2-2130/ and https://dms-vep.github. io/SARS-CoV-2_Omicron_BA.2_spike_DMS_COV2-2130/.

Acknowledgements This work was performed under the auspices of the US Department of Energy by Lawrence Livermore National Laboratory (LLNL) under contract DE-AC52-07NA27344. The work was supported by the US Department of Defense's Joint Program Executive Office for Chemical, Biological, Radiological and Nuclear Defense, in collaboration with the Defense Health Agency COVID funding initiative under agreement 11647302 in support of the GUIDE program, the Defense Advanced Research Projects Agency (DARPA) agreement numbers HR0011154580 and HR0011-18-2-0001 (A. Jenkins), Laboratory Directed Research and Development programs (20-ERD-032 and 20-ERD-064) at LLNL, and grants and contracts from the US NIH (R01 AI157155, NIAID Centers of Excellence for Influenza Research and Response (CEIRR) contract 75N93019C00051, and NIAID B-cell Epitope Discovery and Mechanisms of Antibody Protection Contract 75N93019C00074). The following reagents were obtained through BEI Resources, NIAID, NIH: SARS-related coronavirus 2, Wuhan-Hu-1 spike D614Gpseudotyped lentiviral kit (NR-53817) and the HEK-293T-hACE2 cell line (NR-52511), B.D. and J.D.B. were supported in part by the NIH/NIAID grant R01A1141707. J.D.B. is an Investigator of the Howard Hughes Medical Institute. EM data collections were conducted at the Center for Structural Biology Cryo-EM Facility at Vanderbilt University. We thank A. Jenkins and J. Brase for technical and programmatic contributions and for facilitating this collaborative research effort; M. Landajuela and J. Yang for technical feedback; E. Jaffe, A. Fountain, A. Avila-Herrera, A. Diaz and L. Consolati for contributions to the editing and final preparation of the manuscript and files; and laboratories of S. Subramaniam and X. Wang for their cryo-EM structures of Omicron ahead of release (LLNL-JRNL-839587).

Author contributions T.A.D., K.T.A., A.T.Z., E.Y.L., F.Z., S.C., S.J.Z., E.B., C.G.E., S.H., L.B.T., B.W.S., A.M.L., S.S., M.S.D., J.E.C. Jr, R.H.C. and D.M.F. contributed to the conception or design of the study. T.A.D., K.T.A., D.R., S.C., S.J.Z., E.B., S.M.S., B.D., T.B.E., E.C., L.S.H., L.H., D.R.W., J.K.-Y.L., B.R., E.A.S., T.W., T.-H.L., B.W., J.B.C., E.A.G., B.K.P., L.B.T., B.W.S., J.D.B., M.S.D., J.E.C. Jr, R.H.C. and D.M.F. acquired, analysed or interpreted data. T.A.D., A.T.Z., E.Y.L., F.Z., J.W.G., D.V., S.N., A.L., M.S.S., R.M.H., E.A.G., B.K.P. and D.M.F. created new software. T.A.D., K.T.A., A.T.Z., E.Y.L., F.Z., D.R., S.C., S.J.Z., E.B., B.D., T.B.E., TW.B., B.K.P., B.W.S., J.D.B., M.S.D., J.E.C. Jr, R.H.C. and D.M.F. drafted or substantively revised the manuscript. Tri-lab COVID-19 Consortium members who directly contributed to this work are also listed as contributing authors. Tri-lab COVID-19 Consortium guthors and/or via contributed indirectly as members of groups led by contributing authors and/or via contributions to general infrastructure and capabilities.

Competing interests M.S.D. is a consultant for Inbios, Vir Biotechnology, Ocugen, Moderna and Immunome. The Diamond laboratory has received unrelated funding support in sponsored research agreements from Moderna, Vir Biotechnology and Emergent BioSolutions. J.E.C. Jr has served as a consultant for Luna Labs USA, Merck Sharp & Dohme Corporation, Emergent Biosolutions and GlaxoSmithKline, is a member of the scientific advisory board of Meissa Vaccines, a former member of the scientific advisory board of Gigagen (Grifols) and is founder of IDBiologics. The laboratory of J.E.C. Jr received unrelated sponsored research agreements from AstraZeneca, Takeda and IDBiologics during the conduct of the study. J.D.B. is on the scientific advisory boards of Apriori Bio, Aerium Therapuetics, Invivod and the Vaccine Company. The LLNL, Los Alamos National Laboratory and Vanderbilt University have applied for patents for some of the antibodies in this paper, for which T.A.D., K.T.A., A.T.Z., E.Y.L., F.Z., A.M.L., R.H.C., J.E.C. Jr and D.M.F. are inventors. Vanderbilt University has licensed certain rights to antibodies in this paper to AstraZeneca. J.D.B. and B.D. are inventors on Fred Hutch licensed patents related to the DMS of viral proteins. All other authors declare no competing interests.

Additional information

Supplementary information The online version contains supplementary material available at https://doi.org/10.1038/s41586-024-07385-1.

Correspondence and requests for materials should be addressed to Daniel M. Faissol. **Peer review information** *Nature* thanks Charlotte Deane and the other, anonymous, reviewer(s) for their contribution to the peer review of this work.

Reprints and permissions information is available at http://www.nature.com/reprints.



Extended Data Fig. 1 | Optimized, single-concentration screening data in Gyrolab immunoassays allow selection of candidates from Set 1 (n = 230) for down-stream characterization. a-d, Parental mAb COV2-2130 (orange circles) and positive control mAb S309 24 (magenta squares) serve as references for computationally designed mAbs in single-concentration immunoassays. Computationally designed antibodies are shown as gray diamonds; selected computationally designed antibodies are highlighted with other colors and symbols as shown in the legend. Target antigens are (a) wild type WA1/2020, (**b**) Delta, (**c**) Omicron BA.1 and (**d**) Omicron BA.1.1. Each screened antibody and antigen combination was evaluated in two replicate assays, shown as individual points in the plot. Each of the controls was replicated in two replicate assays for each of three groups of antibodies, resulting in six replicate assays for each point on the control curves. Control curve points and error bars indicate mean and standard deviation. Curves represent a four-parameter logistic curve fit to the control data. All analysis performed in GraphPad Prism.



Extended Data Fig. 2 | ELISA screening allows set for down selection of candidates from Set 2 (n = 204), for down-stream characterization. a-d, Parental mAb COV2-2130 (orange circles; 3 technical replicates) and positive control mAb S309 24 (magenta squares; 3 technical replicates) serve as references for computationally designed mAbs (black curves). Purple triangles are 2130-1-114-112; blue diamonds are 2130-1-0104-024. Each designed antibody had a single measurement (n = 1) at each concentration. All curves are 4-parameter logistic fits, produced in GraphPad Prism. Target antigens are RBD from (a) wild type WA1/2020, (b) Omicron BA.1 (Acro), which is biotinylated, (c) Omicron BA.1, and (d) Omicron BA.1.1. For the biotinylated antigen in (b) (Acro Biosystems, cat. SPD-C82E4), an additional coat and wash cycle was required to prepare the ELISA plate with streptavidin.



Extended Data Fig. 3 | 2130-1-0114-112 is potent in focus reduction neutralization tests with authentic virus in Vero-TMPRSS2 cells. a-i, 2130-1-0114-112 potently neutralizes (a) WA1/2020 D614G (b) Delta B.1.617.2, (c) Omicron BA.1, (d) Omicron BA.1.1, (e) Omicron BA.2, (f) Omicron BA.2.12.1, (g) Omicron BA.4, (h) Omicron BA.5, and (i) Omicron BA.5.5 authentic viruses in focus reduction neutralization assays in Vero-TMPRSS2 cells. Symbols indicate the values of two technical replicates; curves are 4-parameter logistic regression models fit of normalized data. **j**, IC50 values and 95% confidence intervals corresponding to **a**-**i**. ">" indicates IC50 values > 10,000; "NC" indicates fits that were unconverged, unstable, or with positive hill slope. Analyses were performed in GraphPad Prism.



Extended Data Fig. 4 | Authentic virus neutralization in plaque assays using Vero E6-TMPRSS2-T2A-ACE2 (VAT) cells. a-c, Plaque assay neutralization of (a) Delta, (b) Omicron BA.1 and (c) Omicron BA.1.1 viruses. Data are represented as the normalized infection of mAb-treated virus to virus treated with control human IgG (Invitrogen). For S309, each point shows one of four technical replicates; all other points show one of two technical replicates at each concentration. Curves are two-parameter (IC50, hill-slope) logistic fits to normalized response. **d**, IC50 values (ng/ml, ">" indicates IC50 greater than 10,000 ng/mL) show that 2130-1-0104-024, while having only two mutations from COV2-2130, remains potent against BA.1 and suffers a 20-times loss in potency against BA.1.1. 2130-1-0114-112 is strongly potent against all three tested variants. All analysis performed in GraphPad Prism.



Extended Data Fig. 5 | Normalized pseudoviral neutralization of SARS-CoV-2 VOCs by 2130-1-0114-112. a, f, COV2-2130 and 2130-1-0114-112 potently neutralize WA1/2020 D614G. b, COV2-2130 does not potently neutralize BA.1, whereas 2130-1-0114-112 does, consistent with other pseudoviral neutralization assays. c, 2130-1-0114-112 potently neutralizes BA.2.75, outperforming COV2-2130 by 90-fold. d, e, 2130-1-0114-112 loses substantial potency in the context of BA.4.6 and artificially-produced BA.2.75 + R346T but retains measurable neutralization, demonstrating mitigation of this critical weakness of COV2-2130. g-j, 2130-1-0114-112 maintains weak neutralization of BA.2.75 R346T F486S
(g), but loses detectable neutralization activity against BQ.1.1 (h) and XBB
(i) and exhibits a near-complete loss of neutralization activity against BN.1 (j). Symbols indicate two independent technical replicates at each concentration. Curves represent a four-parameter logistic curve fit to the data with fixed minimum and maximum values (0 and 100, respectively). k, Neutralization IC50 values in ng/ml. ">" indicates IC50 greater than 10,000 ng/mL.
"NC" indicates failure to converge. All analysis conducted in GraphPad Prism.



Extended Data Fig. 6 | See next page for caption.

Extended Data Fig. 6 | Details and VOC comparisons for Cryo-EM structure of neutralizing antibody 2130-1-0114-112 in complex with Omicron BA.2 RBD. a, RBD residues within 7 Å of 2130-1-0114-112. RBD shown in red, with BA.2 mutated residues in orange, and 2130-1-0114-112 in yellow/green. b, CDRH3 Glu112 and RBD Lys440, shown with the EM map and distance between the side chains. c, d, 3D representation of the interaction plot between RBD and 2130-1-0114-112 HC (c, yellow) and LC (d, green). 2130-1-0114-112 is shown as stick and RBD as gray spheres with the contact residues in red. Contact residues are labelled and numbered. e, Fab COV-2130 paratope and epitope residues involved in hydrogen bonding (dashed lines; distances in Å) and hydrophobic interactions with WA1/2020 RBD; compare with Fig. 6d showing BA.2/2130-1-0114-112 interactions. Residues forming hydrophobic interactions are shown as curved lines with rays. Atoms are shown as circles, with oxygen, carbon, and nitrogen in red, black, and blue respectively. Image created with Ligplot + . **f**, Atomic model of the RBD-Fab complex superimposed with WA1-RBD (light brown PDB: 7L7E), XBB1.1-RBD (pink PDB: 8IOS), and BQ1.1 (gray PDB 8IF2). BA.2 RBD is shown in red, with BA.2 mutations in orange. 2130-1-0114-112 HC and LC are yellow and green, with mutations in cyan and blue. Hydrogen bonds are shown as dashed lines.



Extended Data Fig. 7 | **Sequence logos of candidate antibody designs. a**, The set of 376 designed IgG includes mutation at 16 positions in the heavy chain (blue; mutations in green) and 9 positions in the light chain (magenta; mutations in pink). Height of each letter is proportional to the frequency of the amino acid in the group. b, c. This set of 376 sequences is divided into two overlapping sets,

Set 1 (**b**; n = 230) and Set 2 (**c**; n = 204). **d**, From these two sets, a set of eight sequences was selected for production at larger scale and further evaluation including assessment of their thermostability and binding performance (Fig. 2). Selected sequences show reduction in mutations throughout the CDRH3 residues (103-118) mutated in (**a**), especially in residues 103-108.



Extended Data Fig. 8 | See next page for caption.

Extended Data Fig. 8 | Strains with mutated epitopes are differentially neutralized by COV2-2130 and 2130-1-0114-112. a, b, Starting from the 7L7E structure (a), shown after being separated and rotated to show the contact surfaces (b), complexes of COV2-2130 and 2130-1-0114-112 were naively composed by rigid body superposition of the highest resolution structures available of the VOC RBDs onto the WA1/2020 RBD of 7L7E. Epitope and paratope are outlined (solid black) and colored according to charge (blue for positive, red for negative). Residues are correspondingly outlined according to charge or, if uncharged, hydrophilicity (cyan) or hydrophobicity (yellow). The models were used to infer possible intermolecular clashes and loss of key interactions that could account for loss of affinity and, conversely, relief of clashes or new favorable interactions that could account for gain of affinity. For putative interactions that differ among the antibody and RBD combinations displayed, lines show salt bridges (purple) and hydrogen bonds (cyan). c, The interface between COV2-2130 and WA1/2020 RBD shows a number of favorable electrostatic interactions, including between RBD R346 and HC D56 and RBD K444 and HC D107. d-f, In the modeled interaction between 2130-1-0114-112 and BA.1 (d. RBD from PDB 7X66), separated and rotated (e), and with interactions identified (f) the RBD G446S substitution changes the center of the epitope, resulting in a loss of binding with COV2-2130, perhaps by means of clashes. 2130-1-0114-112 may rescue lost affinity by introduction of LC 59E that may form a favorable interaction with R498 and S446. Also 2130-1-0114-112 may

introduce a favorable interaction with all of the Omicron variants by forming a salt bridge between HC 112E and K440. Further, the designed mutations LC S32A and S33A in 2130-1-0114-112 could enhance favorable hydrophobic interaction across the interface in the A484 region. g, BA.1.1 RBD (RBD from PDB 7XAZ) has all of the mutations of BA.1, but additionally contains the mutation R346K, which may disrupt the favorable interaction with HC D56. h, BA.2 RBD (RBD from PDB 8GB8) lacks the particularly unfavorable G446S mutation present in BA.1 and BA.1.1, and the R346K mutation of BA.1.1. BA.2 also has the K440 and A484 residues that are the interaction partners of 2130-1-0114-112's designed mutations HC E112, LC A32 and LC A33. This combination of preservation of WA1/2020 interactions and the addition of newly exploited interaction partners results in strong neutralization by 2130-1-0114-112. i, j, BQ.1.1 (i, RBD from 8IF2), and XBB.1 (j, RBD from 8IOS) which are not neutralized by either COV2-2130 or 2130-1-0114-112, contain substitutions in the epitope at positions 346 and either 444 or 445 as compared to WA1/2020. All three of R346, K444, and V445 are among the most highly buried residues in the epitope, and the mutation of R346 and K444 removes the two salt bridges formed with these residues. Consequently, substitutions at these positions heavily impact the shape and charge complementarity of both COV2-2130 and 2130-1-0114-112 to the RBD, likely explaining much of the impact to their affinities.

Extended Data Table 1 | Thermal stability and expression yield of selected IgGs

mAb	Tm-Boltzmann (C; mean +/- std)	Yield (mg) per 100 mL culture
COV2-2130	68.54 ± 0.27	28.31
2130-1-0111-002	68.41 ± 0.06	31.64
2130-1-0104-015	68.35 ± 0.22	27.77
2130-1-1231-017	68.52 ± 0.42	19.34
2130-1-0104-024	67.81 ± 0.35	26.93
2130-1-0114-111	65.53 ± 0.25	26.54
2130-1-0114-112	68.23 ± 0.26	48.12
2130-1-1231-174	69.02 ± 0.33	32.63
2130-1-1231-200	68.07 ± 0.27	47.72

Melting temperature (Tm) was determined using a fluorescence-based protein thermal shift assay (GloMeltTM, Biotium). Yield was determined by measuring optical density at 280 nm and deriving antibody quantity using the calculated extinction coefficient.

Extended Data Table 2 | Pseudoviral neutralization potency (IC50) of selected IgGs, corresponding to Fig. 3

		IC5	0 (ng/ml) [95% CI]		
Antibody ID	D614G	BA.1	BA.1.1	BA.2	BA.4
rDENV-2D22	NC	NC	>	NC	>
rS2K146	6.56 [3.29–12.1]	4.38 [2.89–6.55]	2.09 [1.47-2.99]	3.36 [2.49-4.51]	36.0 [16.6–72.3]
rCOV2-2130	3.60 [2.51–5.11]	158 [50.9–392.]	NC	2.53 [1.89–3.34]	64.3 [38.5–106.]
2130-1-0111-002	4.92 [3.62–6.66]	34.8 [25.1–49.5]	>	1.47 [1.21–1.78]	4.69 [1.35–13.2]
2130-1-1231-017	2.43 [1.24–4.26]	61.3 [35.8–111.]	>	0.569 [0.502-0.644]	17.1 [5.50–42.6]
2130-1-0104-024	2.99 [2.11-4.08]	13.3 [9.06–19.7]	78.5 [26.4–227.]	0.220 [0.136-0.342]	2.27 [0.923-4.96]
2130-1-0114-111	3.02 [2.45–3.76]	32.0 [19.3–58.1]	4027 [230–9450]	0.162 [0.108-0.235]	34.3 [17.6–76.0]
2130-1-0114-112	4.08 [3.67–4.55]	2.56 [1.79–3.70]	6.25 [4.51-8.75]	0.985 [0.747–1.29]	0.779 [0.604–0.98

Pseudovirus neutralization IC50 values and 95% confidence intervals corresponding to Fig. 3. ">" indicates a value>10,000 ng/mL; NC indicates positive hill slope or failure to converge.

nature portfolio

Corresponding author(s): Thomas A. Desautels, Daniel M. Faissol

Last updated by author(s): Mar 13, 2024

Reporting Summary

Nature Portfolio wishes to improve the reproducibility of the work that we publish. This form provides structure for consistency and transparency in reporting. For further information on Nature Portfolio policies, see our <u>Editorial Policies</u> and the <u>Editorial Policy Checklist</u>.

Statistics

For all statistical analyses, confirm that the following items are present in the figure legend, table legend, main text, or Methods section.

n/a	Cor	firmed
	\boxtimes	The exact sample size (n) for each experimental group/condition, given as a discrete number and unit of measurement
	\square	A statement on whether measurements were taken from distinct samples or whether the same sample was measured repeatedly
	\boxtimes	The statistical test(s) used AND whether they are one- or two-sided Only common tests should be described solely by name; describe more complex techniques in the Methods section.
\boxtimes		A description of all covariates tested
	\boxtimes	A description of any assumptions or corrections, such as tests of normality and adjustment for multiple comparisons
	\boxtimes	A full description of the statistical parameters including central tendency (e.g. means) or other basic estimates (e.g. regression coefficient) AND variation (e.g. standard deviation) or associated estimates of uncertainty (e.g. confidence intervals)
	\boxtimes	For null hypothesis testing, the test statistic (e.g. <i>F</i> , <i>t</i> , <i>r</i>) with confidence intervals, effect sizes, degrees of freedom and <i>P</i> value noted Give <i>P</i> values as exact values whenever suitable.
\boxtimes		For Bayesian analysis, information on the choice of priors and Markov chain Monte Carlo settings
\boxtimes		For hierarchical and complex designs, identification of the appropriate level for tests and full reporting of outcomes
\boxtimes		Estimates of effect sizes (e.g. Cohen's d, Pearson's r), indicating how they were calculated
		Our web collection on statistics for biologists contains articles on many of the points above.

Software and code

Policy information	about <u>availability of computer code</u>		
Data collection	On Biological data collection was conducted with appropriate, instrument specific software.		
Data analysis	Analysis of individual experiments was conducted in GraphPad Prism v.8.1, 9.0, or 10.2, as noted in individual figure legends.		
	Protein Thermal Shift [™] software version 1.4 (ThermoFisher) was used to generate the melting temperature and fit data.		
	Antibody-antigen structures were minimized and relaxed using standard minimization procedures from Rosetta, Chimera, and GROMACS.		
	The ImprovWF dynamic workflows code is available at: https://github.com/LLNL/improvwf		
	Selected GUIDE workflow components are available at: https://github.com/LLNL/CRPCA		
	Antibody-antigen minimization procedures used Rosetta 3.13, Chimera v1.15, or GROMACS v2021.4		
	Rosetta FlexDDG. Script collected from github in 2020: https://github.com/Kortemme-Lab/flex_ddG_tutorial/blob/master/analyze_flex_ddG.py		
	SFE v1.0		

ddG estimates calculated with FoldX v4 or Rosetta 3.13

All DMS data analysis was performed using dms-vep-pipeline version 1.8, which can be accessed at https://github.com/dms-vep/dms-vep-pipeline/tree/51e73d601bd770eb6e9abd21f57fb4365699c984

Code and notebooks related to DMS runs are available at: https://dms-vep.github.io/SARS-CoV-2_Omicron_BA.1_spike_DMS_COV2-2130/ https://dms-vep.github.io/SARS-CoV-2_Omicron_BA.2_spike_DMS_COV2-2130/

Pseudocode for additional components is available in the Supplemental Material.

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

Data

Policy information about availability of data

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

- Accession codes, unique identifiers, or web links for publicly available datasets
- A description of any restrictions on data availability
- For clinical datasets or third party data, please ensure that the statement adheres to our policy

The EM map and model has been deposited into EMDB (EMD-28198, EMD-28199) and PDB (8EKD). Other protein structural data were employed in this work (PDB: 7L7E, 7T9K) and in analysis (PDB: 7X66, 7XAZ, 8IOS, 8IF2, 8GB8). Sequence data that support the findings of this study have been deposited in GenBank under accessions PP474664 – PP474679 and are available in the Supplementary Information. Source data for Figure 4 are provided with the paper. DMS library variant data and antibody per replicate DMS selection data can be accessed at https://github.com/dms-vep/SARS-CoV-2_Omicron_BA.2_spike_DMS_COV2-2130 and https://github.com/dms-vep/SARS-CoV-2_Omicron_BA.1_spike_DMS_COV2-2130 GitHub repositories.

Human research participants

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

Reporting on sex and gender	N/A
Population characteristics	N/A
Recruitment	N/A
Ethics oversight	N/A

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

Field-specific reporting

Please select the one below that is the best fit for your research. If you are not sure, read the appropriate sections before making your selection.

Life sciences Behavioural & social sciences

nces Ecological, evolutionary & environmental sciences

For a reference copy of the document with all sections, see nature.com/documents/nr-reporting-summary-flat.pdf

Life sciences study design

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

Sample size	For animal studies, no sample sizes were chosen a priori but instead estimated based on prior knowledge of anticipated experimental differences among groups. All experiments with statistical analysis were repeated at least two independent times, each with multiple technical replicates. Experimental size of animal cohorts was determined based on prior experience performing studies in mice.
Data exclusions	Representative data were chosen where appropriate in lieu of combining independent replicates. Data were excluded only for technical reasons (e.g., splash into wells). No data was excluded from animal studies.
Replication	All replications executed were consistent with the representative data presented in the manuscript. Each DMS experiment was done using two independently produced deep-mutational scanning libraries. For animals studies, all experiments had multiple biological and/or technical replicates and are indicated the Figure legends.
Randomization	For animal studies, mice were randomly assigned from large batches obtained from the vendor to different experimental groups in an age-

Randomization	matched distribution.
Blinding	For animal studies, no blinding was performed as handling of BSL3 virus requires exact tracking of infected mice and samples.

Behavioural & social sciences study design

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

Study description Briefly describe the study type including whether data are quantitative, qualitative, or mixed-methods (e.g. quali quantitative experimental, mixed-methods case study).		
Research sample	State the research sample (e.g. Harvard university undergraduates, villagers in rural India) and provide relevant demographic information (e.g. age, sex) and indicate whether the sample is representative. Provide a rationale for the study sample chosen. For studies involving existing datasets, please describe the dataset and source.	
Sampling strategy	Describe the sampling procedure (e.g. random, snowball, stratified, convenience). Describe the statistical methods that were used to predetermine sample size OR if no sample-size calculation was performed, describe how sample sizes were chosen and provide a rationale for why these sample sizes are sufficient. For qualitative data, please indicate whether data saturation was considered, and what criteria were used to decide that no further sampling was needed.	
Data collection	Provide details about the data collection procedure, including the instruments or devices used to record the data (e.g. pen and paper, computer, eye tracker, video or audio equipment) whether anyone was present besides the participant(s) and the researcher, and whether the researcher was blind to experimental condition and/or the study hypothesis during data collection.	
Timing	Indicate the start and stop dates of data collection. If there is a gap between collection periods, state the dates for each sample cohort.	
Data exclusions	If no data were excluded from the analyses, state so OR if data were excluded, provide the exact number of exclusions and the rationale behind them, indicating whether exclusion criteria were pre-established.	
Non-participation	State how many participants dropped out/declined participation and the reason(s) given OR provide response rate OR state that no participants dropped out/declined participation.	
Randomization	If participants were not allocated into experimental groups, state so OR describe how participants were allocated to groups, and if allocation was not random, describe how covariates were controlled.	

Ecological, evolutionary & environmental sciences study design

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

Study description Briefly describe the study. For quantitative data include treatment factors and interactions, design structure (e.g. factoria hierarchical), nature and number of experimental units and replicates.	
Research sample	Describe the research sample (e.g. a group of tagged Passer domesticus, all Stenocereus thurberi within Organ Pipe Cactus National Monument), and provide a rationale for the sample choice. When relevant, describe the organism taxa, source, sex, age range and any manipulations. State what population the sample is meant to represent when applicable. For studies involving existing datasets, describe the data and its source.
Sampling strategy	Note the sampling procedure. Describe the statistical methods that were used to predetermine sample size OR if no sample-size calculation was performed, describe how sample sizes were chosen and provide a rationale for why these sample sizes are sufficient.
Data collection	Describe the data collection procedure, including who recorded the data and how.
Timing and spatial scale	Indicate the start and stop dates of data collection, noting the frequency and periodicity of sampling and providing a rationale for these choices. If there is a gap between collection periods, state the dates for each sample cohort. Specify the spatial scale from which the data are taken
Data exclusions	If no data were excluded from the analyses, state so OR if data were excluded, describe the exclusions and the rationale behind them, indicating whether exclusion criteria were pre-established.
Reproducibility	Describe the measures taken to verify the reproducibility of experimental findings. For each experiment, note whether any attempts to repeat the experiment failed OR state that all attempts to repeat the experiment were successful.
Randomization	Describe how samples/organisms/participants were allocated into groups. If allocation was not random, describe how covariates were controlled. If this is not relevant to your study, explain why.
Blinding	Describe the extent of blinding used during data acquisition and analysis. If blinding was not possible, describe why OR explain why blinding was not relevant to your study.

Field work, collection and transport

Field conditions	Describe the study conditions for field work, providing relevant parameters (e.g. temperature, rainfall).
Location	State the location of the sampling or experiment, providing relevant parameters (e.g. latitude and longitude, elevation, water depth).
Access & import/export	Describe the efforts you have made to access habitats and to collect and import/export your samples in a responsible manner and in compliance with local, national and international laws, noting any permits that were obtained (give the name of the issuing authority, the date of issue, and any identifying information).
Disturbance	Describe any disturbance caused by the study and how it was minimized.

Reporting for specific materials, systems and methods

No

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

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

Antibodies

Antibodies used	Gyrolab immunoassays: A secondary detection antibody served as a fluorescent reporter: Alexa Fluor 647 AffiniPure Fab Fragment Goat Anti-Human IgG, Fcy fragment specific (Jackson ImmunoResearch). The detection Fab was used at 100 nM Our bridging control antibody was s309 (Biovision, A2266 -100) - https://www.abcam.com/products/primary-antibodies/sars-cov-2- spike-glycoprotein-s1-antibody-s309-bsa-and-azide-free-ab289796.pdf Positive control for pseudovirus neutralization assay: S2K146 (DOI: 10.1126/science.abm8143) Negative control for pseudovirus and authentic virus neutralization assays: DENV-2D22 (doi:10.1073/pnas.1200566109) ELISA: 1:5000 dilution goat anti-human IgG conjugated with horseradish peroxidase (HRP) (Southern Biotech, cat. 2014-05, lot L2118- VG00B) FRNT: Anti-S murine antibodies (PMID: 34481543) Dilution: N/A This pool of antibodies was used as pooled hybridoma supernatant FRNT: HRP-conjugated goat anti-mouse IgG (Sigma Cat # A8924, RRID: AB_258426) 1:1000 dilution Mouse studies isotype control antibody: anti-West Nile hE16 mAb (Oliphant2005, PMID: 15852016) Reagent antibodies were obtained from Jackson Immunoresearch 109-607-008
Validation	Validation of all primary antibodies tested for binding antigen by ELISA or with infected cells. For validation data on commercial antibodies used, refer to specification documents from the manufacturers as follows: HRP-conjugated Goat anti-mouse IgG - https://www.sigmaaldrich.com/specification-sheets/638/581/A8924-BULK.pdf HRP-conjugated Goat anti-human IgG, Southern Biotech, cat. 2014-05, lot L2118-VG00B, https://resources.southernbiotech.com/ techbul/2014.pdf AlexaFluor647-conjugated Goat Anti-Human IgG, F(ab') ₂ - https://www.jacksonimmuno.com/lots/000000168261

Eukaryotic cell lines

Cell line source(s) HEK293 cells at ATUM (Newark, CA, USA)	Policy information about <u>cell lines</u>	and Sex and Gender in Research
ExpiCHO-S™ Cells (Thermofisher scientific Cat. No. A29127) Expi293F™ Cells (Thermofisher scientific Cat. No. A14527 HEK293T (ATCC: CRL-11268) Vero-hACE2-TMPRSS2 cells in FRNT were generated at the NIH (VRC, Barney Graham lab) now BEI (BEI NR-54970) - (PMI 32404436)	Cell line source(s)	HEK293 cells at ATUM (Newark, CA, USA) ExpiCHO-S [™] Cells (Thermofisher scientific Cat. No. A29127) Expi293F [™] Cells (Thermofisher scientific Cat. No. A14527 HEK293T (ATCC: CRL-11268) Vero-hACE2-TMPRSS2 cells in FRNT were generated at the NIH (VRC, Barney Graham lab) now BEI (BEI NR-54970) - (PMID: 32404436)

	VAT cells used in Plaque assays: BEI Resources, NIAID, NIH DMS experiments were performed using 293T-hACE2 cells available from BEI:NR-52511
Authentication	None authenticated.
Mycoplasma contamination	All cell lines were confirmed to be negative for mycoplasma during a regular basis.
Commonly misidentified lines (See ICLAC register)	None.

Animals and other research organisms

Policy information about studies involving animals; ARRIVE guidelines recommended for reporting animal research, and Sex and Gender in Research

Laboratory animals	Seven to nine-week-old female heterozygous 18-hACE2 C57BL/6J mice (strain: 2B6.Cg-Tg(K18-ACE2)2Prlmn/J, Cat # 34860) were obtained from The Jackson Laboratory. Mice were housed in groups of 3 to 5. Photoperiod = 12 hr on:12 hr off dark/light cycle. Ambient animal room temperature is 70° F, controlled within ±2° and room humidity is 50%, controlled within ±5%.
Wild animals	No wild animals were used in this study.
Reporting on sex	Only female mice were used in the study.
Field-collected samples	No field collected samples were used in this study.
Ethics oversight	The protocols were approved by the Institutional Animal Care and Use Committee at the Washington University School of Medicine (assurance number A3381–01).

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