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COVID-19 vaccines and antibody therapies†

Prediction and mitigation of mutation threats to

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Antibody therapeutics and vaccines are among our last resort to end the raging COVID-19 pandemic. They, however, are prone to over 5000 mutations on the spike (S) protein uncovered by a Mutation Tracker based on over 200 000 genome isolates. It is imperative to understand how mutations will impact vaccines and antibodies in development. In this work, we first study the mechanism, frequency, and ratio of mutations on the S protein which is the common target of most COVID-19 vaccines and antibody therapies. Additionally, we build a library of 56 antibody structures and analyze their 2D and 3D characteristics. Moreover, we predict the mutation-induced binding free energy (BFE) changes for the complexes of S protein and antibodies or ACE2. By integrating genetics, biophysics, deep learning, and algebraic topology, we reveal that most of the 462 mutations on the receptor-binding domain (RBD) will weaken the binding of S protein and antibodies and disrupt the efficacy and reliability of antibody therapies and vaccines. A list of 31 antibody disrupting mutants is identified, while many other disruptive mutations are detailed as well. We also unveil that about 65% of the existing RBD mutations, including those variants recently found in the United Kingdom (UK) and South Africa, will strengthen the binding between the S protein and human angiotensin-converting enzyme 2 (ACE2), resulting in more infectious COVID-19 variants. We discover the disparity between the extreme values of RBD mutation-induced BFE strengthening and weakening of the bindings with antibodies and angiotensin-converting enzyme 2 (ACE2), suggesting that SARS-CoV-2 is at an advanced stage of evolution for human infection, while the human immune system is able to produce optimized antibodies. This discovery, unfortunately, implies the vulnerability of current vaccines and antibody drugs to new mutations. Our predictions were validated by comparison with more than 1400 deep mutations on the S protein RBD. Our results show the urgent need to develop new mutation-resistant vaccines and antibodies and to prepare for seasonal vaccinations

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Introduction

The expeditious spread of the coronavirus disease 2019 (COVID-19) pandemic caused by severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) has led to 95 932 739 confirmed cases and 2 054 853 fatalities as of January 20, 2021. In the 21st century, three major outbreaks of deadly pneumonia have been caused by β-coronaviruses: SARS-CoV (2002), Middle East respiratory syndrome coronavirus (MERS-CoV) (2012), and

SARS-CoV-2 (2019). Similar to SARS-CoV and MERS-CoV, SARS-CoV-2 causes respiratory infections, and the transmission of viruses occurs among family members or in healthcare settings at the early stages of the outbreak. However, SARS-CoV-2 has an unprecedented scale of infection. Considering the high infection rate, high prevalence rate, long incubation period,2 asymptomatic transmission,3,4 and potential seasonal pattern5 of COVID-19, the development of specific antiviral drugs, antibody therapies, and effective vaccines is of paramount importance. Traditional drug discovery takes more than ten years, on average, to bring a new drug to the market.6 However, developing potent SARS-CoV-2 specific antibodies and vaccines is a relatively more efficient and less time-consuming strategy to combat COVID-19 for the ongoing pandemic.7 Antibody therapies and vaccines depend on the host immune system. Recently, studies have been working on the host-pathogen interaction, host immune responses, and the pathogen immune evasion strategies,8-13 which provide insight into understanding the mechanism of antibody therapies and vaccine development.

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[†] Electronic supplementary information (ESI) available: (S1) Methods; (S2) multiple sequence alignments of antibodies and pairwise identity scores; (S3) random coil percentages of antibody paratopes; and (S4) additional analysis of antibody-S protein complexes. See DOI: 10.1039/d1sc01203g

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The immune system is a host defense system that protects the host from pathogenic microbes, eliminates toxic or allergenic substances, and responds to an invading pathogen.14 It has the innate immune system and adaptive immune system as two major subsystems. The innate system provides an immediate but non-specific response, while the adaptive immune system provides a highly specific and effective immune response. Once the pathogen breaches the first physical barriers, such as the epithelial cell layers, secreted mucus layer, and mucous membranes, the innate system will be triggered to identify pathogens by pattern recognition receptors (PRRs), which is expressed on dendritic cells, macrophages, or neutrophils.15 Specifically, PPRs identify pathogen-associated molecular patterns (PAMPs) located on pathogens and then activate complex signaling pathways that introduce inflammatory responses mediated by various cytokines and chemokines, which promote the eradication of the pathogen. 16,17 Notably, the transmission of SARS-CoV-2 even occurs in asymptomatic infected individuals, which may delay the early response of the innate immune response.8 Another important line of host defense is the adaptive immune system. B lymphocytes (B cells) and T lymphocytes (T cells) are special types of leukocyte that are the acknowledged cellular pillars of the adaptive immune system.18 Two major subtypes of T cells are involved in the cellmediated immune response: killer T cells (CD8+ T cells) and helper T cells (CD4+ cells). The killer T cells eradicate cells invaded by pathogens with the help of major histocompatibility complex (MHC) class I. MHC class I molecules are expressed on the surface of all nucleated cells.19 The nucleated cells will firstly degrade foreign proteins via antigen processing when viruses infect them. Then, the peptide fragments will be presented by MHC class I, which will activate killer T cells to eliminate these infected cells by releasing cytotoxins.20 Similarly, helper T cells cooperate with MHC class II, a type of MHC molecule that is constitutively expressed on antigen-presenting cells, such as macrophages, dendritic cells, monocytes, and B cells.21 Helper T cells express T cell receptors (TCR) to recognize antigen bound to MHC class II molecules. However, helper T cells do not have cytotoxic activity. Therefore, they cannot kill infected cells directly. Instead, the activated helper T cells will release cytokines to enhance the microbicidal function of macrophages and the activity of killer T cells.22 Notably, an unbalanced response can result in a "cytokine storm," which is the main cause of the fatality of COVID-19 patients.23 Correspondingly, a B cell gets involved in the humoral immune response and identifies pathogens by binding to foreign antigens with its B cell receptors (BCRs) located on its surface. The antigens that are recognized by antibodies will be degraded to peptides in B cells and displayed by MHC class II molecules. As mentioned above, helper T cells can recognize the signal provided by MHC class II and upregulate the expression of the CD40 ligand, which provides extra stimulation signals to activate antibodyproducing B cells,24 making millions of copies of antibodies (Ab) that recognize the specific antigen. Additionally, when the antigen first enters the body, the T cells and B cells will be activated, and some of them will be differentiated to long-lived

These long-lived memory cells will play a role in quickly and specifically recognizing and eliminating a specific antigen that encountered the host and initiated a corresponding immune response in the future.²⁵ The vaccination mechanism is to stimulate the primary immune response of the human body, which will activate T cells and B cells to generate the antibodies and long-lived memory cells that prevent infectious diseases, which is one of the most effective and economical means of combating COVID-19 at this stage.

As mentioned above, secreted by B cells of the adaptive immune system, antibodies can recognize and bind to specific antigens. Conventional antibodies (immunoglobulins) are Yshaped molecules that have two light chains and two heavy chains.26 Each light chain is connected to the heavy chain via a disulfide bond, and heavy chains are connected through two disulfide bonds in the mid-region known as the hinge region. Each light and heavy chain contains two distinct regions: the constant region (stem of the Y) and variable region ("arms" of the Y).27 An antibody binds the antigenic determinant (also called epitope) through the variable regions in the tips of the heavy and light chains. There is an enormous amount of diversity in the variable regions. Therefore, different antibodies can recognize many different types of antigenic epitope. To be specific, there are three complementarity determining regions (CDRs) that are arranged non-consecutively in the tips of each variable region. CDRs generate most of the variations between antibodies, which determine the specificity of individual antibodies. In addition to conventional antibodies, camelids also produce heavy-chain-only antibodies (HCAbs). HCAbs, also referred to as nanobodies, or VHHs, contain a single variable domain (VHH) that makes up the equivalent antigen-binding fragment (Fab) of conventional immunoglobulin G (IgG) antibodies.28 This single variable domain can typically acquire affinity and specificity for antigens comparable to conventional antibodies. Nanobodies can easily be constructed into multivalent formats and have higher thermal stability and chemostability than most antibodies do.29 Another advantage of nanobodies is that they are less susceptible to steric hindrances than large conventional antibodies.30

Considering the broad specificity of antibodies, seeking potential antibody therapies has become one of the most feasible strategies to fight against SARS-CoV-2. In general, an antibody therapy is a form of immunotherapy that uses monoclonal antibodies (mAb) to target pathogenic proteins. The binding of an antibody and pathogenic antigen can facilitate an immune response, direct neutralization, radioactive treatment, the release of toxic agents, and cytokine storm inhibition (aka immune checkpoint therapy). The SARS-CoV-2 entry into a human cell is facilitated by the process of a series of interactions between its spike (S) protein and the host receptor angiotensin-converting enzyme 2 (ACE2), primed by host transmembrane protease serine 2 (TMPRSS2).31 As such, most COVID-19 antibody therapeutic developments focus on the SARS-CoV-2 spike protein antibodies that were initially generated from the patient immune response and T-cell pathway inhibitors that block T-cell responses. A large number of antibody therapeutic drugs are in clinical trials.

memory cells, such as memory T cells and memory B cells.

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Fifty-five S protein antibody structures are available in the Protein Data Bank (PDB), offering a great resource for mechanistic analysis and biophysical studies.

Currently, most of the antibody therapy developments focus on the use of antibodies isolated from patients' convalescent plasma to directly neutralize SARS-CoV-2,32-34 although there are efforts to alleviate the cytokine storm. A more effective and economical means to fight against SARS-CoV-2 is the vaccine, 35 which is the most anticipated approach for preventing the COVID-19 pandemic. A vaccine is designed to stimulate effective host immune responses and provide active acquired immunity by exploiting the body's immune system, including the production of antibodies, and is made of an antigenic agent that resembles a disease-causing microorganism, or surface protein, or genetic material that is needed to generate the surface protein. For SARS-CoV-2, the first choice of surface proteins is the spike protein. There are four types of COVID-19 vaccine, as shown in Fig. 1. (1) Virus vaccines use the virus itself in a weakened or inactivated form. (2) Viral-vector vaccines are designed to genetically engineer a weakened virus, such as measles or adenovirus, to produce coronavirus S proteins in the body. Both replicating and non-replicating viral-vector vaccines are being studied now. (3) Nucleic-acid vaccines use DNA or mRNA to produce SARS-CoV-2 S proteins inside host cells to stimulate the immune response. (4) Protein-based vaccines are designed to directly inject coronavirus proteins, such as S protein or membrane (M) protein, or their fragments, into the body. Both protein subunits and viral-like particles (VLPs) are under development for COVID-19.36 Among these technologies, nucleic-acid vaccines are safe and relatively easy to develop.³⁶ However, they have not been approved for any human use before.

However, the general population's safety concerns are the major factors that hinder the rapid approval of vaccines and antibody therapies. A major potential challenge is an antibodydependent enhancement, in which the binding of a virus to suboptimal antibodies enhances its entry into host cells. All vaccine and antibody therapeutic developments are currently based on the reference viral genome reported on January 5, 2020.37 SARS-CoV-2 belongs to the Coronaviridae family and the Nidovirales order, which has been shown to have a genetic proofreading mechanism regulated by non-structure protein 14 (NSP14) in synergy with NSP12, i.e., RNA-dependent RNA polymerase (RdRp).38,39 Therefore, SARS-CoV-2 has a higher fidelity in its transcription and replication process than other singlestranded RNA viruses, such as the flu virus and HIV. However, the S protein of SARS-CoV-2 has been undergoing many mutations, as reported in ref. 40 and 41. As of January 20, 2021, a total of 5003 unique mutations on the S protein have been detected on 203 346 complete SARS-CoV-2 genome sequences. Among them, 462 mutations were on the receptor-binding domain (RBD), the most popular target for antibodies and vaccines. Therefore, it is of paramount importance to establish a reliable paradigm to predict and mitigate the impact of SARS-CoV-2 mutations on vaccines and antibody therapies. Moreover, the efficacy of a given COVID-19 vaccine depends on many factors, including the SARS-CoV-2 biological properties

associated with the vaccine, mutation impacts, vaccination schedule (dose and frequency), idiosyncratic response, and assorted factors such as ethnicity, age, gender, and genetic predisposition. The effect of COVID-19 vaccination also depends on the fraction of the population that accepts vaccines. It is essentially unknown at this moment how these factors will unfold for COVID-19 vaccines.

There is no doubt that any preparation that leads to an improvement in the COVID-19 vaccination effect will be of tremendous significance to human health and the world economy. Therefore, in this work, we integrate genetic analysis and computational biophysics, including artificial intelligence (AI), as well as additional enhancement from advanced mathematics to predict and mitigate mutation threats to COVID-19 vaccines and antibody therapies. We perform single nucleotide polymorphism (SNP) calling41 to identify SARS-CoV-2 mutations. For mutations on the S protein, we analyze their mechanism,42 frequency, ratio, and secondary structural traits. We construct a library of 56 existing antibody structures by January 1, 2021 from the PDB and analyze their twodimensional (2D) and three-dimensional (3D) characteristics. We further predict the mutation-induced binding free energy (BFE) changes of antibody and S protein complexes using a topology-based network tree (TopNetTree),43 which is a stateof-the-art model that integrates deep learning and algebraic topology.44-46 In this work, TopNetTree is trained with newly available deep mutation datasets on the S protein, ACE2, and some antibodies and its predictions are validated with thousands of experimental data points. Our studies indicate that most mutations will significantly disrupt the binding of essentially all known antibodies to the S protein. Therefore, vaccines and antibody drugs that were developed based on the early SARS-CoV-2 genome will be seriously compromised by mutations. Additionally, we show that most known mutations will strengthen the binding between the S protein and ACE2, which gives rise to more infectious variants. Our studies also reveal that SARS-CoV-2 is at an advanced stage of evolution with respect to its ability to infect humans. Although the human immune system is able to produce antibodies that are optimized with respect to a pathogen, the antibodies, once produced, are very vulnerable to attack by mutants.

Mutations on the spike protein 2

As a fundamental biological process, mutagenesis changes the organism's genetic information and serves as a primary source for many kinds of cancer and heritable diseases, which is a driving force for evolution. 47,48 Generally speaking, virus mutations are introduced by natural selection, replication mechanism, cellular environment, polymerase fidelity, gene editing, random genetic drift, recent epidemiological features, host immune responses, etc. 49,50 Notably, understanding how mutations have changed the SARS-CoV-2 structure, function, infectivity, activity, and virulence is of great importance for coming up with life-saving strategies in virus control, containment, prevention, and medication, especially in the development of antibodies and vaccines. Genome sequencing, SNP

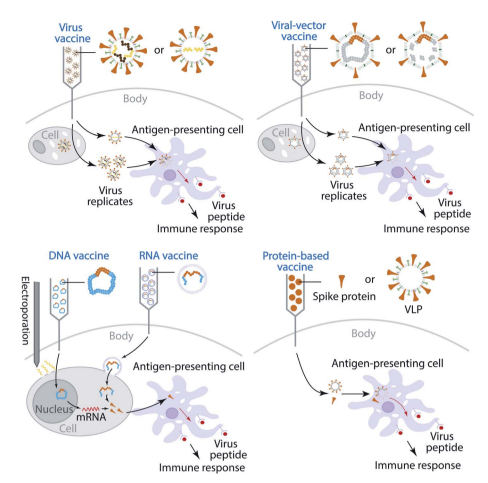


Fig. 1 Illustration of four types of COVID-19 vaccine that are currently in development.

calling, and phenotyping provide an efficient means to parse mutations from a large number of viral samples⁴⁰ (see the ESI (S1†)). In this work, we retrieved more than 200 000 complete SARS-CoV-2 genome sequences from the GISAID database⁵¹ and

created a real-time interactive SARS-CoV-2 Mutation Tracker to report more than 26 000 unique single mutations along with their mutation frequency on SARS-CoV-2 as of January 20, 2021. Fig. 2 is a screenshot of our online Mutation Tracker. It

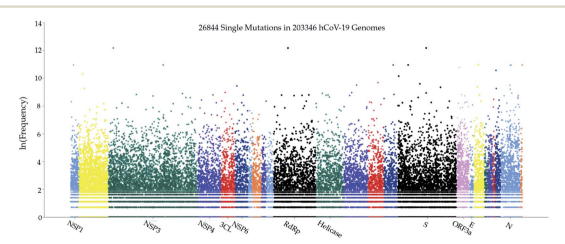


Fig. 2 The distribution of genome-wide SARS-CoV-2 mutations on 26 proteins. The y-axis represents the natural log frequency for each mutation on a specific position of the complete SARS-CoV-2 genome. While only a few landmark positions are labeled with gene (protein) names, the relative positions of other genes (proteins) can be found in our Mutation Tracker (https://users.math.msu.edu/users/weig/SARS-CoV-2_Mutation_Tracker.html).

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Table 1 The distribution of 12 SNP types among 5003 unique mutations and 467 604 non-unique mutations on the S gene of SARS-CoV-2 worldwide. $N_{\rm U}$ is the number of unique mutations and $N_{\rm NU}$ is the number of non-unique mutations. $R_{\rm U}$ and $R_{\rm NU}$ represent the ratios of 12 SNP types among unique and non-unique mutations

SNP type	Mutation type	$N_{ m U}$	$N_{ m NU}$	$R_{ m U}$	$R_{ m NU}$
A > T	Transversion	454	5236	9.07%	1.12%
A > C	Transversion	341	2571	6.82%	0.55%
A > G	Transition	700	199 015	13.99%	42.56%
T > A	Transversion	356	1614	7.12%	0.35%
T > C	Transition	779	19 313	15.57%	4.13%
T > G	Transversion	277	1940	5.54%	0.41%
C > T	Transition	542	158 898	10.83%	33.98%
C > A	Transversion	313	10 301	6.26%	2.20%
C > G	Transversion	156	968	3.12%	0.21%
G > T	Transversion	435	34 421	8.69%	7.36%
G > C	Transversion	225	6090	4.50%	1.30%
G > A	Transition	425	27 237	8.49%	5.82%

Table 2 The statistics of non-degenerate mutations on the secondary structure of SARS-CoV-2 S protein. The unique and non-unique mutations are considered in the calculation. N_U , N_{NU} , AR_U , and AR_{NU} represent the number of unique mutations, the number of non-unique mutations, the average rate of unique mutations, and the average rate of non-unique mutations on the secondary structure of S protein, respectively. Here, the secondary structure is mostly extracted from the crystal structure of 7C2L; the missing residues are predicted by RaptorX-Property

Secondary structure	Length	$N_{ m U}$	$N_{ m NU}$	AR_U	AR_{NU}
Helix	249	516	9535	2.07	38.29
Sheet	276	711	20 422	2.58	73.99
Random coils	748	2100	350 659	2.81	468.80
Whole spike	1273	3327	380 616	2.61	298.99

describes the distribution of mutations on the complete coding region of SARS-CoV-2. The y-axis shows the natural log frequency for each mutation at a specific position. A reader can download the detailed mutation SNP information from our Mutation Tracker website.

As mentioned before, the S protein has become the first choice for antibody and vaccine development. Among the 203 346 complete genome sequences, 5003 unique single mutations are detected on the S protein. The number of unique mutations (N_U) is determined by counting the same type of mutation in different genome isolates only once, while the number of non-unique mutations (N_{NU} , i.e., frequency) is calculated by counting the same type of mutation in different genome isolates repeatedly. Table 1 lists the distribution of 12 SNP types among unique and non-unique mutations on the S protein of SARS-CoV-2 worldwide. It can be seen that C > T and A > G are the two dominant SNP types, which may be due to the innate host immune response via APOBEC and ADAR gene editing.42

Moreover, 144 non-degenerate mutations occurred on the S protein RBD, which are relevant to the binding of SARS-CoV-2 S protein and most antibodies as well as ACE2. Additionally, the

218 mutations that occurred on the S protein N-terminal domain (NTD) (residue id: 14 to 226) are relevant to the binding of another two antibodies (4A8 and FC05) and SARS-CoV-2 S protein.

Furthermore, since antibody CDRs are random coils, the complementary antigen-binding domains must involve random coils as well. Table 2 lists the statistics of non-degenerate mutations on the secondary structures of SARS-CoV-2 S protein. Here, the secondary structures are mostly extracted from the crystal structure of 7C2L,52 and the missing residues are predicted by RaptorX-Property.⁵³ We can see that for both unique and non-unique cases, the average mutation rates on the random coils of the S protein have the highest values. Particularly, the 23 403 A > G-(D614G) mutation on the random coils has the highest frequency of 192 284. If we do not consider the 23 403 A > G-(D614G) mutations, then the unique and nonunique average rates on the random coils of S protein still have the highest values (2.81 and 212.01), indicating that mutations are more likely to occur on the random coils. Consequently, the natural selection of mutations may tend to disrupt antibodies.

SARS-CoV-2 antibodies 3

In this work, we consider 56 3D structures available from the PDB (https://www.rcsb.org) before January 1, 2021. These 56 structures include 51 structures of antibodies binding to S protein RBD, 4 structures of antibodies having binding domains outside the S protein RBD, and an ACE2-S protein complex. Among the four structures having binding domains outside the RBD, there are three distinct antibodies not binding to the RBD, namely 4A8,52 FC05,⁵⁴ and 2G12.⁵⁵ This is because FC05 has two sets of structures (PBD IDs 7CWU and 7CWS) that differ from each other by their components on the RBD (i.e., H014 and P17). Some antibodies are given as combinations of other unique ones. Among the 51 antibodies on the RBD, there are only 42 unique ones, including MR17-K99Y as a mutant of MR17.56

3.1 3D antibody structure alignment on the S protein

We present the 3D alignment of 45 structures of SARS-CoV-2 S protein with ACE2 and antibodies (excluding the mutant MR17-K99Y of MR17) in Fig. 3. ACE2 in Fig. 3(a) is a reference. Fig. 3(a)-(j) list 42 single antibodies binding to the RBD, and Fig. 3(k) includes the other 3 alignments of 4A8, FC05, and 2G12 whose binding domains are outside the RBD. Fig. 3(m) presents a 3D structure of a single chain of S protein. The PDB IDs of these complexes can be found in Fig. 4.

Fig. 3 reveals, except for Fab 52,62 S309,57 CR3022,63 EY6A,67 4A8,52 FC05,54 and 2G12,55 all the other 38 antibodies have their binding sites spatially clashing with that of ACE2. Notably, the paratopes of H014 (ref. 69) and S304 (ref. 60) do not overlap with that of ACE2 directly, but in terms of 3D structures, their binding sites still overlap. This suggests that the bindings of 39 antibodies are in direct competition with that of ACE2. Theoretically, this direct competition reduces the viral infection rate. Such antibodies with strong binding ability will directly neutralize SARS-CoV-2 without the need for antibody-

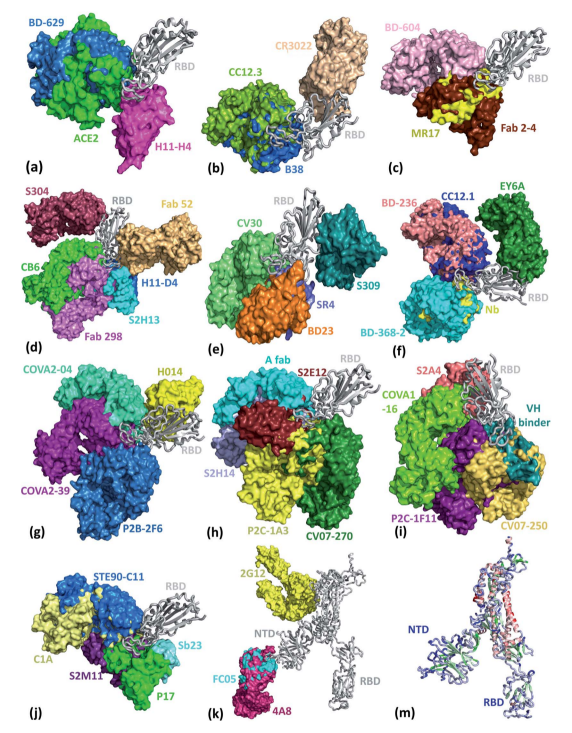


Fig. 3 Aligned structures of 46 complexes of the S protein and ACE2 and single antibodies. (a) – (j) The 3D alignment of the available unique 3D structures of SARS-CoV-2 S protein RBD in binding complexes with 42 antibodies (MR17-K99Y is excluded because its binding mode is the same as that of MR17). (k) The 3D alignment of the three antibodies binding outside RBD. (m) The 3D structure of S protein RBD. The red, green, and blue colors represent helix, sheet, and random coils of RBD, respectively. The darker color represents the higher mutation frequency on a specific residue. The structures are (a) ACE2 (6M0J),⁵⁷ BD-629 (7CH5), H11-H4 (6ZBP); (b) CC12.3 (6XC4),⁵⁸ B38 (7BZ5),⁵⁹ CR3022 (6XC3),⁵⁸ (c) BD-604 $(7CH4), MR17 (7C8W), ^{56} Fab 2-4 (6XEY); ^{56} (d) S304 (7JW0), ^{60} CB6 (7C01), ^{61} Fab 52 (7K9Z), ^{62} S2H13 (7JV6), ^{60} H11-D4 (6YZ5), ^{63} Fab 298 (7K9Z); ^{62} (e) Fab (7C01), ^{61} Fab (7C01),$ $\text{CV30 (6XE1)}, ^{64} \text{ BD23 (7BYR)}, ^{65} \text{ SR4 (7C8V)}, ^{56} \text{ S309 (6WPS)}; ^{66} \text{ (f) CC12.1 (6XC2)}, ^{58} \text{ EY6A (6ZCZ)}, ^{67} \text{ BD-236 and nanobody (Nb) (7CHE)}, ^{68} \text{ BD-368-2}, ^{68$ (7CHH);⁶⁸ (g) H014 (7CAH),⁶⁹ COVA2-04 (7JMO),⁷⁰ COVA2-39 (7JMP),⁷⁰ P2B-2F6 (7BWJ);⁷¹ (h) P2C-1A3 (7CDJ), CV07-270 (6XKP),⁷² S2H14 (7JX3),60 A fab (7CJF), S2E12 (7K45);75 (i) CV07-250 (6XKQ),72 P2C-1F11 (7CDI), VH binder (7JWB),74 S2A4 (7JVA),60 COVA1-16 (7JMW);75, (j) C1A (7KFV),76 STE90-C11 (7B3O),77 Sb23 (7A29),78 S2M11 (7K43),73 P17 (7CWM);79; and (k) 4A8 (7C2L),52 FC05 (7CWU),54 and 2G12 (7L06).55

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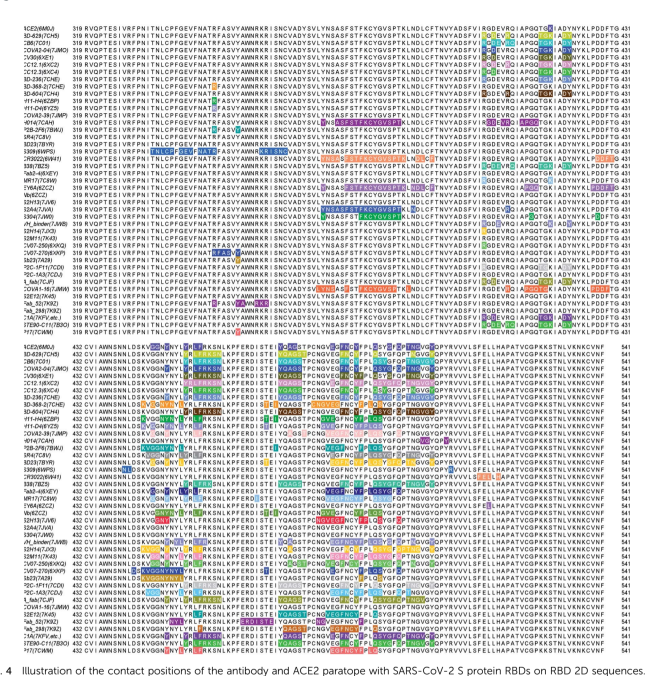


Fig. 4 Illustration of the contact positions of the antibody and ACE2 paratope with SARS-CoV-2 S protein RBDs on RBD 2D sequences. The corresponding PDB IDs are given in parentheses.

dependent cell cytotoxicity (ADCC), antibody-dependent cellular phagocytosis (ADCP), or other immune mechanisms.

The paratopes of S309, Fab 52, CR3022, and EY6A on the RBD are away from that of ACE2, leading to the absence of binding competition.66,67,80 One study shows that the ADCC and ADCP mechanisms contribute to the viral control conducted by S309 in infected individuals.66 For Fab 52, it was suggested that its mechanism could involve S protein destabilization. 62 For CR3022, one research indicates that it neutralizes the virus in a synergistic fashion.81 For EY6A, the hypothesis is that glycosylation of ACE2 accounts for at least part of the observed crosstalk between ACE2 and EY6A.67 More radical examples are 4A8, FC05, and 2G12. 4A8 binds to the NTD of the S protein (Fig. 3(h)), which is quite far from the RBD. It is speculated that 4A8 may neutralize SARS-CoV-2 by restraining the conformational changes of the S protein, which is very important for the SARS-CoV-2 cell entry.⁵² FC05 is combined with P17 or H014 to form a cocktail.54 2G12 binds to the S protein S2 domain.55 Any antibody or drug that can inhibit the serine protease TMPRSS2 priming of the S protein priming can effectively stop the viral cell entry.31

3.2 2D residue contacts between antibodies and the S protein RBD

Fig. 3 provides a visual illustration of antibody and ACE2 competitions. It remains to be known in the residue detail what has happened to these competitions. To better understand the antibody and S protein interactions, we study the residue contacts between antibodies and the S protein. We include the ACE2 as a reference but excluding antibodies MR17-K99Y as well as 4A8, FC05, and 2G12 that bind to other domains.

In Fig. 4, the paratopes of 42 individual antibodies (excluding MR17-K99Y) and ACE2 were aligned on the S protein RBD 2D sequence, and their contact regions are highlighted. From the figure, one can see that, except for Fab 52, S309, CR3022, EY6A, H014, and S304, all the other 36 antibodies have their antigenic epitopes overlapping with the ACE2, especially on the residues from 486 to 505 of the RBD. Although the paratopes of H014 and S304 do not overlap with that of ACE2 directly, their binding sites still overlap in 3D structures. Therefore, these 38 antibodies competitively bind against ACE2 as revealed in Fig. 3.

3.3 Antibody sequence alignment and similarity analysis

The next question is whether there is any connection or similarity between the antibody paratopes in our library, particularly for those antibodies that share the same binding sites. To better understand this perspective, we carry out multiple sequence alignment (MSA) to further study the similarities and differences among existing antibodies. Many antibodies are very similar to each other and can be classified into several clusters using the CD-HIT suite.82 The first and largest cluster includes COVA2-04, CC12.1, BD-236, BD-604, B38, EY6A, S304, P2C-1A3, A fab, C1A, STE90-C11, and CB6. Their identity scores to CB6 are 90.48%, 94.74%, 93.59%, 93.35%, 94.77%, 92.52%, 90.62%, 90.51%, 91.18%, 94.08%, and 93.00%, respectively. The second cluster contains BD-629, CC12.3, P2C-1F11, and CV30. Their identity scores to CV30 are 95.41%, 96.32%, and 97.68%, respectively. The third cluster has CV07-270 and COVA2-39, and the pairwise identity score is 90.18%. The fourth cluster is composed of H11-H4, H11-D4, and Nb, and their identity scores to Nb are 99.25% and 95.52%, respectively. They are all nanobodies. The fifth cluster has Fab 298 and COVA1-16, and the pairwise identity score is 90.80%. Their alignment plots are given in the ESI (Fig. S1-S5†).

The above similarity indicates that the adaptive immune systems of individuals have a common way to generate antibodies. On the other hand, the existence of five distinct clusters, as well as antibodies 4A8, FC05, and 2G12 suggests the diversity

in the immune response. Note that we have also included ACE2 in our MSA as a reference, but none of the existing antibodies are similar to ACE2 because they were created from entirely different mechanisms.

4 Mutation impacts on SARS-CoV-2 antibodies

To investigate the influences of existing S protein mutations on the binding free energy (BFE) of S protein and antibodies, we consider 462 mutations that occurred on the S protein RBD, which are relevant to the binding of SARS-CoV-2 S protein and antibodies as well as ACE2. Additionally, 540 mutations occurred on the NTD of the S protein (residue id: 14 to 226) which are relevant to the binding of SARS-COV-2 S protein and antibody 4A8 (PDB: 7C2L). We predict the free energy changes following existing mutations using our TopNetTree model.43 The mutations on the RBD are considered for the predictions of BFE changes. Our predictions are built from the X-ray crystal structure of SARS-CoV-2 S protein and ACE2 (PDB 6M0J),57 and various antibodies (PDBs 6WPS, 66 6XC2, 58 6XC3, 58 6XC4, 58 6XC7, 58 6XE1, 64 6XEY, 83 6XKP, 72 6XKQ, 72 6YLA, 63 6YZ5, 6Z2M, 6ZBP, 6ZCZ, 67 6ZER, 67 7A29,⁷⁸ 7B3O, 7BWJ,⁷¹ 7BYR,⁶⁵ 7BZ5,⁵⁹ 7C01,⁶¹ 7C2L,⁵² 7C8V,⁵⁶ 7C8W,⁵⁶ 7CAH,⁶⁹ 7CAH,⁶⁹ 7CAN,⁵⁶ 7CDI, 7CDJ, 7CH4,⁶⁸ 7CH5,⁶⁸ 7CHB,⁶⁸ 7CHE,⁶⁸ 7CHF,⁶⁸ 7CHH,⁶⁸ 7CJF, 7CWM,⁷⁹ 7CWN⁷⁹ 7JMO,⁷⁰ 7JMP,⁷⁰ 7JMW,⁷⁵ 7JV6,⁶⁰ 7JVA,⁶⁰ 7JVC,⁶⁰ 7JW0,⁶⁰ 7JWB,⁷⁴ 7JX3,⁶⁰ 7K43,⁷³ 7K45,⁷³ 7K9Z,⁶² 7KFV,⁷⁶ 7KFW,⁷⁶ 7KFX,⁷⁶ and 7KFY⁷⁶). The BFE change following mutation ($\Delta \Delta G$) is defined as the subtraction of the BFE of the mutant type from the BFE of the wild type: $\Delta \Delta G = \Delta G_W - \Delta G_M$, where ΔG_W is the BFE of the wild type and $\Delta G_{\rm M}$ is the BFE of the mutant. Therefore, a negative BFE change means that the mutation decreases affinities, making the protein-protein interaction less stable.

Four antibody–S protein complexes are examined in this section. Next, we present a library of mutation-induced BFE changes for all mutations and 51 antibodies, as well as ACE2. The statistical analysis of mutation impacts on antibodies is discussed.

4.1 Single antibody-S protein complex analysis

For four antibody-S protein complexes, since there are too many mutations, we only consider those mutations whose

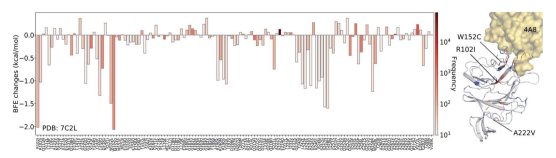


Fig. 5 Illustration of SARS-CoV-2 mutation-induced binding free energy changes for the complexes of S protein and 4A8 (PDB: 7C2L). The blue color in the structure plot indicates a positive BFE change while the red color indicates a negative BFE change, and toning indicates the strength. Here, mutations R102I, W152C, W152L, S247N, and Y248H could potentially disrupt the binding of antibody 4A8 and S protein.

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frequencies are greater than 10. We first present the BFE changes $(\Delta \Delta G)$ of the SARS-CoV-2 S protein binding domain with antibody 4A8 in Fig. 5, which is one of the three complexes that are not on the RBD in our collections of S protein and antibody complexes. A total 141 of 540 mutations on residue ID from 14 to 226 have frequencies larger than 10. Most mutations have small BFE changes (from -0.5 kcal mol^{-1} 0.5 kcal mol⁻¹) in their binding free energies, while 28 mutations have negative BFE changes less than -0.5 kcal mol⁻¹. Notably, 53 out of 141 mutations on the binding domain have positive BFE changes, which means that the mutations increase affinities and would make the S protein-4A8 interactions more stable. However, the majority (63%) of mutations have negative BFE changes, including high-frequency mutations R102I and W152C with frequencies of 89 and 356, respectively. Since the largest positive and negative BFE changes are 0.37 and -2.06 kcal mol⁻¹ (-3.1 if low frequency mutations are counted), respectively, the prediction indicates that antibody 4A8 isolated from 10 convalescent patients at the early stage of the pandemic⁵² is an optimized product of the human immune system with respect to the original S protein. It is also noted that many mutations on the binding domain, such as W152L, S247N, and Y248H, have significant negative free energy changes. The mutations on the binding domain with large

negative BFE changes reveal that the binding of antibody 4A8 and S protein will be potentially disrupted.

Next, we study the BFE changes $(\Delta \Delta G)$ induced by 80 mutations on the SARS-CoV-2 S protein RBD for the antibody Fab 2-4 (PDB: 6XEY) in Fig. 6. Antibody Fab 2-4 shares a similar binding domain with ACE2 and thus is a potential candidate for the direct neutralization of SARS-CoV-2. Most mutations induce small changes in the binding free energies, while mutations E484K, E484Q, F486L, and F490S have large negative BFE changes. Overall, 38 out of 80 mutations on the RBD lead to negative BFE changes, which means 48% of mutations will potentially weaken the binding between antibody Fab 2-4 and S protein. For positive BFE changes, the largest value is only 0.55 kcal mol⁻¹ and the average of positive BFE changes is 0.16 kcal mol⁻¹. However, many mutations with negative BFE changes have a very large magnitude, indicating that antibody Fab 2-4 was an immune product optimized with respect to the original un-mutated S protein. In general, the mutations on S protein weaken the Fab 2-4 binding with S protein and make it less competitive with ACE2 as most mutations strengthen the S protein and ACE2 binding. It is interesting to note that mutation E484K is the so-called South Africa variant. It indeed has a strong vaccine-escape effect.

In Fig. 7, we illustrate the mutation-induced BFE changes for antibody MR17 (PDB: 7C8W), which shares the binding domain

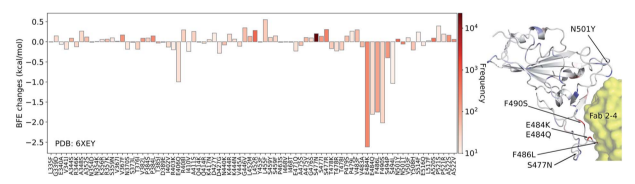


Fig. 6 Illustration of SARS-CoV-2 mutation-induced binding free energy changes for the complexes of S protein and Fab 2-4 (PDB: 6XEY). The blue color in the structure plot indicates a positive BFE change while the red color indicates a negative BFE change, and toning indicates the strength. Here, mutations E484K, E484Q, F486L, and F490S could potentially disrupt the binding of antibody Fab 2-4 and the S protein.

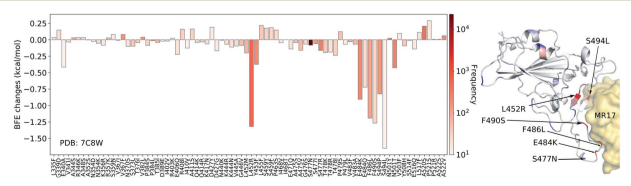


Fig. 7 Illustration of SARS-CoV-2 mutation-induced binding free energy changes for the complexes of S protein and MR17 (PDB: 7C8W). Blue in the structure plot indicates a positive BFE change while red indicates a negative BFE change, and toning indicates the strength. Here, mutations L452R, E484K, F486L, F490S, and S494L could potentially disrupt the binding of antibody MR17 and the S protein.

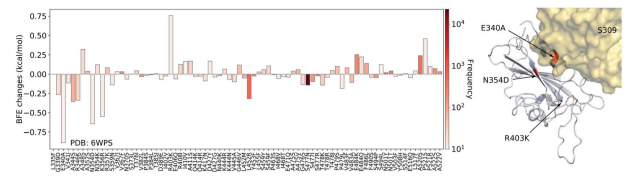


Fig. 8 Illustration of SARS-CoV-2 mutation-induced binding free energy changes for the complexes of S protein and S309 (PDB: 6WPS). The blue color in the structure plot indicates a positive BFE change while the red color indicates a negative BFE change, and toning indicates the strength. Here, mutations E340A, N354D, and K356R could potentially weaken the binding of antibody S309 and the S protein.

with ACE2 as well. One can notice that five mutations, L452R, E484K, F486L, F490S, and S494L, have BFE changes less than -1 kcal mol^{-1} as well as high frequencies. The rest of the mutations have a small magnitude of changes. 27 out of 80 mutations have positive BFE changes with the largest value less than 0.25 kcal mol^{-1} . Our results indicate that antibody MR37 is likely to be isolated from patients at the early stage and thus, it was optimized based on an early version of the SARS-CoV-2 virus. Mutations L452R, E484K, F486L, F490S, and S494L will reduce its competitiveness with ACE2 (Fig. 7).

Finally, we consider the BFE change predictions for the antibody S309 and S protein complex, whose receptor binding motif (RBM) does not overlap with the RBM of ACE2 (see

Fig. 3(e)). The BFE changes induced by 80 mutations are predicted. Among them, 38 changes are positive. Similar to the aforementioned antibodies, most of the mutations lead to small changes in their binding affinity magnitude but three mutations, E340A, N354D, and K356R, induce moderate negative changes. Interestingly, none of the 80 RBD mutations have a major impact on S309. Although mutation R403K might disrupt S309, it does weaken many other antibody bindings with the S protein. While antibodies play a variety of functions in the human immune system, such as neutralization of infection, phagocytosis, antibody-dependent cellular cytotoxicity, etc., their binding with antigens is crucial for these functions. Our analysis of BFE changes following mutations on the S

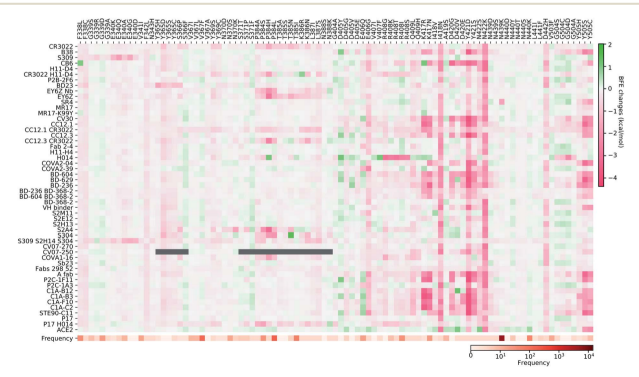


Fig. 9 Illustration of the SARS-CoV-2 helix-residue mutation induced BFE changes for the complexes of S protein and 51 antibodies or ACE2. Positive changes strengthen the binding while negative changes weaken the binding. Mutation frequency is given for each mutation. Grey color indicates that PDB structures do not include residues induced by those mutations.

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protein suggests that some antibodies will be less affected by mutations, which is important for developing vaccines and antibody therapies.

4.2 Mutation impact library

In this section, we build a library of mutation-induced BFE changes for all mutations and all antibodies as well as ACE2. In principle, we could create a library of all possible mutations for all antibodies, as we did for ACE2.⁸⁴ Here, we limit our effort to all existing mutations. Antibody 4A8 on the NTD has been discussed above. We consider antibodies on the RBD.

Based on our earlier analysis, three types of SARS-CoV-2 S protein secondary structural residue have different mutation rates. Among them, the random coils are major components of the RDB and the NTD, as shown in Fig. 3. Most RBD mutations (287 of 462) occur on the residues whose secondary structures are coil, while 93 out of 462 mutations are on the helix, and 82 out of 462 mutations are on the sheet. Therefore, mutations on the RBD are split into three categories based on their locations in secondary structures of helix, sheet, and coil. In Fig. 9, we present the BFE changes for the complexes of the S protein and antibodies or ACE2 induced by mutations on the helix residues of the S protein RBD. The frequency for each mutation is also presented. Most mutations on helix residues lead to negative BFE changes (pink squares), which weaken the bindings, while some mutations induce positive BFE changes (green squares). It

is noted that most mutations lead to the strengthening of the S protein and ACE2 binding, which is consistent with the natural selection rule. Mutations N406G, I418N, N422K, D442H, Y505S, and Y505C give rise to a strong weakening effect on most antibodies. The N439K mutation having the highest frequency, shows a positive BFE change on ACE2 but negative changes on most antibodies. Mutation D405Y appears to strengthen most antibodies.

In Fig. 10, we present the BFE changes for the S protein and antibody (ACE2) complexes following sheet residue mutations of the S protein RBD. Like the last case, most mutations lead to positive BFE changes for ACE2, indicating infectivity strengthening. There are many disruptive mutations, such as R355W, F401I, F401C, I402F, C432G, I434K, A435P, O493P, V510E, V512G, and L513P, that will weaken most antibody and S protein complexes. On the other hand, most mutations strengthen certain antibodies but weaken other ones, which allows the effectiveness of antibody cocktails for better protection. The binding of antibody H014 and the S protein is strengthened by many mutations, particularly S375F, K378O, R403K, and Y453F. Among them, Y453F is an infectivity-strengthening mutation with a relatively high frequency.

Fig. 11–13 present the BFE changes for the S protein and antibody (ACE2) complexes following coil residue mutations of the S protein RBD. Overall, most mutations on coil residues lead to mild negative BFE changes. However, mutations V350F, W353R, I401N, G416V, G431V, Y449D, Y449S, C480R, P491R,

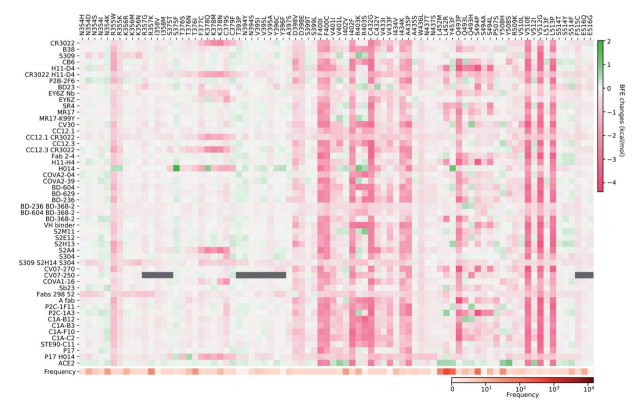


Fig. 10 Illustration of SARS-CoV-2 sheet-residue mutation induced BFE changes for the complexes of S protein and 51 antibodies or ACE2. Positive changes strengthen the binding while negative changes weaken the binding. Mutation frequency is presented for each mutation. Grey color indicates that PDB structures does not include residues induced by those mutations.

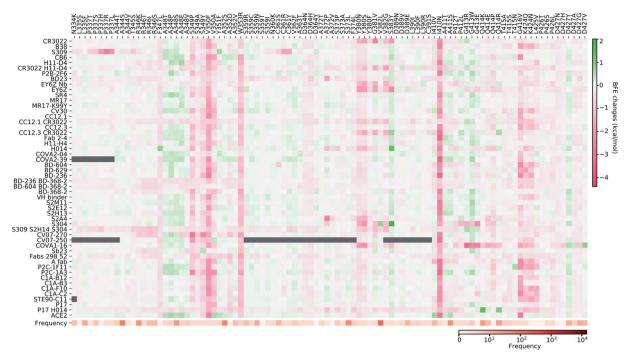


Fig. 11 Illustration of SARS-CoV-2 coil-residue mutation induced BFE changes for the complexes of S protein and 51 antibodies or ACE2. Positive changes strengthen the binding while negative changes weaken the binding. Mutation frequency is presented for each mutation. Grey color indicates that PDB structures do not include residues induced by those mutations.

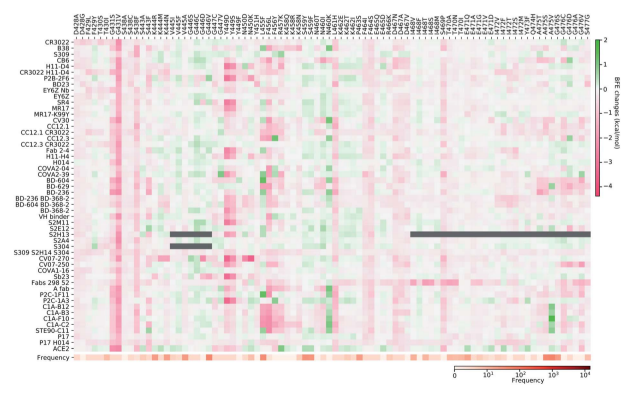


Fig. 12 Illustration of SARS-CoV-2 coil-residue mutation induced BFE changes for the complexes of S protein and 51 antibodies or ACE2 (continued from Fig. 11). Positive changes strengthen the binding while negative changes weaken the binding. Mutation frequency is presented for each mutation. Grey color indicates that PDB structures do not include residues induced by those mutations.

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P491L, Y495C, and O506P will weaken most antibody bindings to the S protein. Some residues, like A348, N460, and P521, can produce many binding-strengthening mutations for most antibodies and ACE2. For the high-frequency mutation S447N in Fig. 13, the BFE changes are mild on ACE2 and antibodies. Additionally, the N501Y mutation, one of the typical mutations in the UK B.1.1.7 variant, strengthens the infectivity but induces mixed reactions to antibodies as shown in Fig. 13.

4.3 Statistical analysis of mutation impacts on COVID-19 antibodies

First, we perform a statistical analysis of all mutation-induced BFE changes studied in the last section. Most mutations induce binding-weakening BFE changes. The total rate of negative BFE changes is 71% (i.e., 16 661 out of 23 512); for coil residues, 67% BFE changes are negative, while for helix and sheet residues, 72% and 80% BFE changes are negative, respectively. However, for ACE2, 300 out of 462 mutations (i.e., 65%) on the RBD produce positive or binding-strengthening BFE changes, showing the effect of the natural selection of mutations. In contrast, at most 200 out of 462 mutations on the RBD give rise to negative BFE changes for antibodies. More specifically, 11 antibodies have less than 100 positive BFE changes while 41 antibodies have less than 200 positive BFE changes. Interestingly, in our prediction, 4 out of the 43 single antibodies have less than 100 positive BFE changes, while 7 out of the 9 antibody cocktails have less than 100 positive BFE changes. Although antibody cocktails have mild negative BFE changes, it turns out that they have high affinities to S protein and the BFE changes are mild for positive ones as well.

Fig. 14 indicates the BFE change extreme values (maximal in cyan and minimal in pink) and average values (positive in blue

and negative in red) of the complexes of S protein and ACE2 or antibodies following mutations. The maximal BFE changes of the helix, sheet, and coil residues are 1.44 kcal mol⁻¹, 1.94 kcal mol⁻¹, and 1.00 kcal mol⁻¹, respectively, while the minimal BFE changes are -3.87 kcal mol⁻¹, -3.9 kcal mol⁻¹, and -4.38 kcal mol⁻¹, respectively. The disparity in their maximal and minimal values indicates the relatively optimal nature of the S protein and antibody binding complexes. It means that the human immune system has the ability to produce optimized antibodies for a given antigen. However, antibodies, once generated, are prone to infection by new mutants. The disparity shown in Fig. 14 also means that the SARS-CoV-2 was at an advanced stage of evolution with respect to human infection. There is not much room for SARS-CoV-2 to improve its infectivity by single-site mutations.

Many antibody cocktails, such as CR3022/H11-D4, CC12.1/CR3022, BD-236/BD368-2, BD604/BD368-2, S309/S2H14/S304, and Fabs 298/52, are relatively less sensitive to the current S protein mutations. However, some other antibodies, such as H11-D4, CV30, CC12.3, and S2H13, can be dramatically affected by SARS-CoV-2 mutations. Importantly, ACE2 is also impacted by mutations and has the largest positive BFE change on average.

5 Mutation impacts on COVID-19 vaccines

The increasing number of infection cases and deaths, the global spread situation, and the lack of prophylactics and therapeutics give rise to an urgent need for the prevention of COVID-19. Vaccination is the most effective and economical means to control pandemics.³⁵ Currently, 248 vaccines are in various clinical trial stages, as reported in an online COVID-19

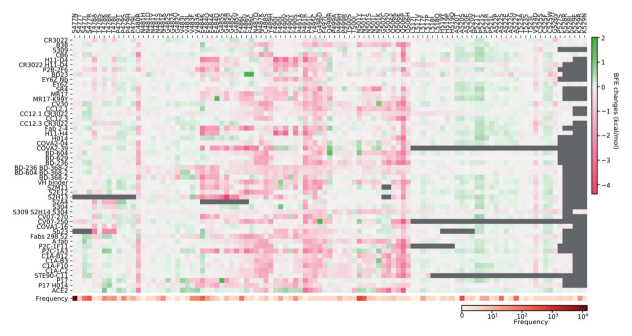


Fig. 13 Illustration of SARS-CoV-2 coil-residue mutation induced BFE changes for the complexes of S protein and 51 antibodies or ACE2 (continued from Fig. 12). Positive changes strengthen the binding while negative changes weaken the binding. Mutation frequency is presented for each mutation. Grey color indicates that PDB structures do not include residues induced by those mutations.

Treatment Vaccine Tracker And (https://covid-19tracker.milkeninstitute.org/#vaccines_intro). Broadly speaking, there are four types of coronavirus vaccine in development: virus vaccines, viral-vector vaccines, nucleic-acid vaccines, and protein-based vaccines, as shown in Fig. 1. The first type of vaccine is the virus vaccine, which injects weakened or inactivate viruses into the human body. A virus is conventionally weakened by altering its genetic code to reduce its virulence and elicit a stronger immune response. A biotechnology company, Codagenix, is currently working on a "codon optimization" technology to weaken viruses, and its weakened virus vaccine is in development.85 Unlike a weakened virus, an inactivated virus cannot replicate in the host cell. A virus is inactivated by heating or using chemicals, which induces neutralizing antibody titers and has been proven to be safe. 86 At this stage, both Sinopharm, which works with the Beijing Institute of Biological Products and Wuhan Institute of Biological Products, and Sinovac, which works with Institute Butantan and Bio Farma, are developing inactive SARS-CoV-2 vaccines that are in phase III clinical trials.

The second type of vaccine is the viral-vector vaccine, which is genetically engineered so that it can produce coronavirus surface proteins in the human body without causing diseases. There are two subtypes of viral-vector vaccine: the nonreplicating viral vector and the replicating viral vector. On February 25, 2021, the World Health Organization (WHO) granted an emergency use listing (EUL) for a vaccine developed by AstraZeneca and the University of Oxford, which is a nonreplicating viral vector vaccine. Moreover, there are 3 nonreplicating viral vector vaccines in phase III trials as well. They work by taking a chimpanzee virus and coating it with the S proteins of SARS-CoV-2. The chimp virus causes a harmless infection in humans, but the spike proteins will activate the immune system to recognize signs of a future SARS-CoV-2 invasion. Notably, booster shots may be needed to retain longlasting immunity. Furthermore, at this stage, only one replicating viral-vector vaccine is in phase II. The University of Hong Kong, in cooperation with Xiamen University and Wantai Biological Pharmacy, is developing such a replicating viral vaccine, which tends to be safe and provoke a strong immune response.

The third type of vaccine is nucleic acid vaccines, which include two subtypes: DNA-based vaccines and RNA-based vaccines. At least 40 teams are currently working on nucleicacid vaccines since they are safe and easy to develop. The DNA-based vaccine works by inserting genetically engineered blueprints of the viral gene into small DNA molecules such as plasmids for injection. Moreover, the electroporation technique is employed to create pores in membranes to increase DNA uptake into cells. The injected DNA will produce mRNA by transcription with the help of the nucleus in human cells. Such an mRNA will translate viral proteins (mostly spike proteins), which are dutifully produced by cells in response to the genes, alarm the immune system, and should produce immunity. Currently, there is one DNA-based vaccine in phase III. Similar to DNA-based vaccines, RNA-based vaccines provide immunity through the introduction of RNA, which is encased in a lipid coat to ensure that it enters into cells. Two RNA-based vaccines have been granted authorization for emergency use in many countries. One is designed by BioNTech, which cooperates with Pfizer, and the other one is from Moderna.

The fourth type of vaccine is the protein-based vaccine, which aims to inject viral proteins directly to human bodies to trigger immune readiness. The protein subunit vaccine is one of the subtypes of the protein-based vaccine. More than 80 teams are working on vaccines with viral protein subunits, such as spike proteins and membrane (M) proteins. Another subtype of the protein-based vaccine is the virus-like particle (VLP) vaccine. VLP vaccines closely resemble viruses. However, they are not infectious since they do not contain viral genetic material. Their non-replicating properties provide a safer alternative to weakened virus vaccines; the HPV vaccine or newer flu vaccines are

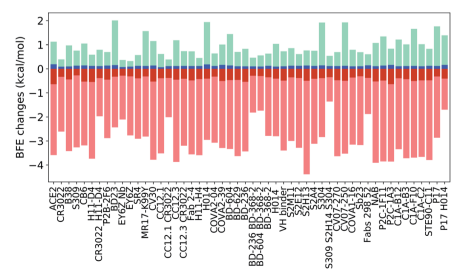


Fig. 14 Illustration of SARS-CoV-2 mutation-induced maximal and minimal BFE changes in cyan and pink for the complexes of S protein and 51 antibodies or ACE2, and average of positive and negative BFE changes in blue and red. Here, the maximal change strengthens the binding while the minimal change weakens the binding for each complex.

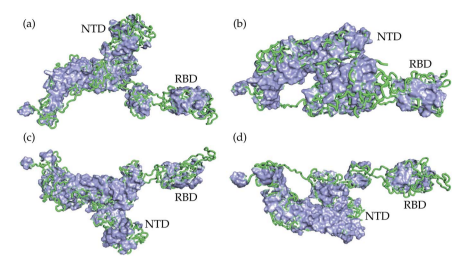


Fig. 15 The 3D rotational structure of SARS-CoV-2 S protein. The random coils of S protein are drawn with green strings and the other secondary structure is described with a purple surface. (a) 3D structure of S protein. (b) 3D structure of S protein that is rotated 90° based on (a). (c) 3D structure of S protein that is rotated 180° based on (a). (d) 3D structure of S protein that is rotated 270° based on (a).

VLP vaccines. Currently, 22 teams are working on VLP vaccines for future prevention of COVID-19.

Secondary structures of antigenic determinants

Since the structural basis of antibody CDRs, or paratopes, is random coils, we hypothesize that CDRs favor antigenic random coils as complementary epitopes, i.e., antigenic determinants. 87,88 Fig. 15 depicts the 3D structure of S protein, where the random coils are drawn with green strings, and the other secondary structure is described with the purple surface. It shows that the RBD and the NTD mostly consist of random coils. The RBD is the antigenic determinant of 43 structurally known SARS-CoV-2 antibodies; meanwhile, the NTD is the binding domain of antibodies 4A8 and FC05 and antibody 2G12 also binds to the S2 domain with random coils, which confirms

our hypothesis. More detailed analysis considered the random coil percentages of antibodies' paratopes which are summarized in Table S1 of the ESI.† It reveals that antibodies predominantly contact residues in random coils of S protein. Most of the antibody paratopes had greater than 90% random coil content.

Fig. 16 shows the secondary structure of the S protein. The red, blue, and green colors represent helix, sheet, and random coils of S protein. It can be seen that the S protein mostly consists of random coils, which means that there are many other potential antigenic epitopes on the S protein for antibody CDRs. We believe that the emphasis on direct binding competition with ACE2 in the past^{66,67,80} has led to the neglecting of many important antibodies that do not bind to the RBD. Therefore, we suggest that researchers pay more attention to antibodies that do not bind to the RBD.

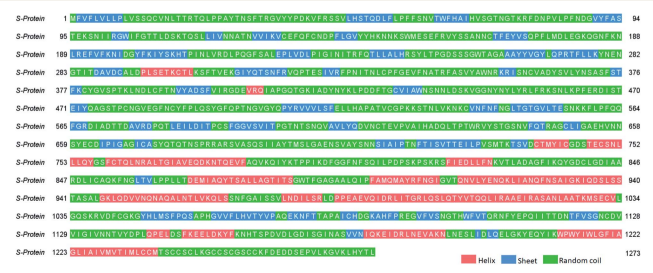


Fig. 16 The secondary structure of S protein. The red, green, and blue colors represent helix, sheet, and random coils of S protein.

Table 3 Antibody disrupting mutants

Location	Mutants
Helix	E406G, I418N, Y421D, N422K, D442H, Y505S
Sheet	R355W, F400I, F400C, I402F, C432G, I434K, A435P, Q493P, V510E, V512G, L513P
Coils	V350F, W353R, I410N, G416V, G431V, Y449D, Y449S, L461H, S469P, C480R, P491R, P491L, Y495C, Q506P

5.2 Statistical estimation of mutation impacts on COVID-19 vaccines

Vaccine efficacy is an essential issue for the control of the COVID-19 pandemic. The S protein is one of the most popular surface proteins for vaccine development. However, mutations have accumulated on the S protein of SARS-CoV-2, which may reduce the vaccine efficacy. As we found in section 2, mutations are more likely to happen on the random coils of S protein, which may have a devastating effect on vaccines in development.

As shown in Fig. 14, mutations could considerably weaken the binding between the S protein and antibodies and thus pose a direct threat to reduce the efficacy of vaccines. However, there are a few obstacles in determining the exact impacts of mutations on COVID-19 vaccines. Firstly, the four types of vaccine platform can produce very different virus peptides, resulting in different immune responses, as well as antibodies. Secondly, even for a given vaccine platform, different peptides may be produced due to different immune responses caused by gender difference, age difference, race difference, *etc.* Therefore, in this work, we proposed to understand the impact of SARS-CoV-2 mutations on COVID-19 vaccines by statistical analysis. By evaluating the binding affinity changes induced by 51 existing SARS-CoV-2 antibodies, as shown in Fig. 9 to 13, we can identify

vaccine escape mutants that will strengthen the binding between the S protein and ACE2 while disrupting the binding between the S protein and antibodies. Table 3 lists a collection of the most disruptive mutations. However, this list is not complete. There are many other antibody disrupting mutations as shown in Fig. 9 to 13. For example, the infectivity-strengthening South Africa mutant E484K can cause dramatically disruptive effects on many antibodies such as H11-D4, Fab 2-4, H11-H4, COVA2-39, BD368-2, *etc.* but it also enhances the binding of other antibodies, such as B38, CV30, CC21.1, Sb23, Fabs 298 52, *etc.* The infectivity-strengthening mutation N501Y in UK B.1.1.7 variants has a disruptive effect only on a few known antibodies, including B38, CC12.3, S2M11, NAB, S309, S2H12, S304, C1A-B12, STE90-C11, *etc.*

In a nutshell, by setting up a SARS-CoV-2 antibody library with the statistical analysis based on the mutation-induced binding free energy changes, we can estimate the impacts of SARS-CoV-2 mutations on COVID-19 vaccines, which will provide a way to infer how a specific mutation will pose a threat to vaccines. This approach works better when more antibody structures become available.

Another important factor in prioritization is mutation frequency. Fig. 9–13 have provided frequency information from our SNP calling. Once a mutation is identified as a potential

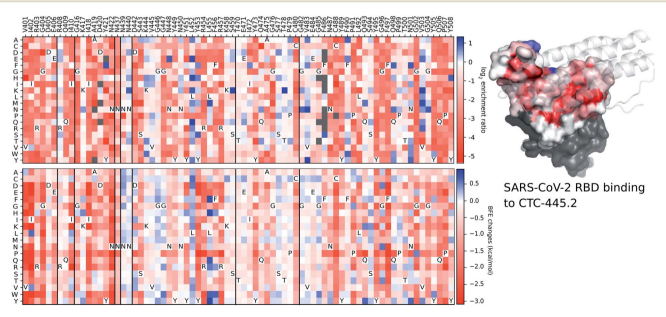


Fig. 17 A comparison between experimental deep mutation enrichment data and TopNetTree predictions for the SARS-CoV-2 S protein RBD and CTC-445.2 complex (7KL9 (ref. 89)). Top left: deep mutational scanning heatmap showing the average effect on the enrichment for single site mutants of the RBD when assayed by yeast display for binding to CTC-445.2.89 Top right: the RBD colored by average enrichment at each residue position bound to CTC-445.2. Bottom: machine learning predicted BFE changes for the CTC-445.2 and S protein complex induced by single site mutations on the RBD.

threat, it can be incorporated into the next generation of vaccines in a cocktail approach. In principle, all four types of vaccine platform allow the accommodation of new viral strains.

6 Validation

Although the details of the methods used in this work are presented in the ESI,† we provide a validation of our deep learning prediction model, TopNetTree,⁴³ which is crucial to the credibility of this work. Specifically, we demonstrate the prediction performance of S protein mutation induced BEF changes on CTC-445.2 compared to the experimental deep mutation enrichment data.⁸⁹ More detailed descriptions of methods and datasets are provided in the ESI.†

Fig. 17 presents a comparison between experimental deep mutation enrichment data on the RBD and machine learning predicted RBD-mutation-induced BFE changes for the SARS-CoV-2 S protein and CTC-445.2 complex. In the heatmaps of Fig. 17, one can see that the predicted BFE changes have a very high correlation with the experimental enrichment ratio data. Both enrichment ratios and BFE changes describe the affinity strength of the protein–protein interaction induced by mutations. The high similarity between these heatmaps demonstrates the reliability of our machine learning predictions of BFE changes following mutations on the S protein RBD.

7 Conclusion

The coronavirus disease 2019 (COVID-19) pandemic has gone out of control globally. There is no specific medicine or effective treatment for this viral infection at this point. Vaccination is widely anticipated to be the endgame for taming the viral rampage. Another promising treatment that is relatively easy to develop is antibody therapies. However, both vaccines and antibody therapies are prone to more than 26 000 unique mutations recorded in the Mutation Tracker.

We present the most comprehensive analysis and prediction of mutation threats to vaccines and antibody therapies. First, we identify existing mutations on the severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) spike (S) protein, which is the main target for both vaccines and antibody therapies. We analyze the mechanism, frequency, and ratio of mutations along with the secondary structures of the S protein. Additionally, we build a library of 55 antibodies with structures available from the Protein Data Bank (PDB) and analyze their two-dimensional (2D) and three-dimensional (3D) characteristics by employing computational biophysics. We further predict the mutation-induced binding free energy (BFE) changes of S protein and antibody complexes using a model called TopNetTree based on deep learning and algebraic topology. The performance of our model has been extensively validated by its prediction of experimental deep mutation data. Our significant findings are as follows. First, we reveal that none of the known mutations are safe to all antibodies. On average, most mutations (i.e., 71%) will weaken the binding between the S protein and antibodies, which implies that vaccines will also be compromised by existing mutations. Additionally, we identify 31 antibody disrupting mutants that dramatically weaken

the binding between the S protein and most known antibodies. Moreover, we find that most RBD mutations (*i.e.*, 64.9%) will enhance the binding strength between the S protein and angiotensin-converting enzyme 2 (ACE2), which implies that most existing mutations will strengthen the SARS-CoV-2 infectivity. This result is consistent with the natural selection of mutations and our earlier findings. ⁸⁴ Finally, we discover that the maximal BFE change magnitudes of binding-strengthening mutations are much smaller than those of binding-weakening mutations for all antibodies, which shows that current human antibodies were optimized with respect to the original S protein and are prone to the S protein mutations. Our findings indicate the pressing need to keep developing mutation-resistant vaccines and antibody drugs and to be ready for seasonal vaccinations.

8 Data availability

Detailed mutation information is available for download at Mutation Tracker.

Author contributions

Conceptualization: Guo-Wei Wei. Data curation: Jiahui Chen, Kaifu Gao, Rui Wang. Formal analysis: Jiahui Chen, Guo-Wei Wei. Funding acquisition: Guo-Wei Wei. Investigation: Jiahui Chen, Guo-Wei, Wei. Methodology: Jiahui Chen, Rui Wang. Project administration: Guo-Wei Wei. Resources: Jiahui Chen, Kaifu Gao, Rui Wang. Software: Jiahui Chen, Rui Wang. Supervision: Guo-Wei Wei. Validation: Jiahui Chen, Guo-Wei Wei. Visualization: Jiahui Chen, Kaifu Gao, Rui Wang, Guo-Wei Wei. Writing – original draft: Jiahui Chen, Kaifu Gao, Rui Wang, Guo-Wei Wei. Writing–review & editing: Jiahui Chen, Guo-Wei Wei.

Conflicts of interest

There are no conflicts to declare.

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