Towards an optimal monoclonal antibody with higher binding affinity to the receptor-binding domain of SARS-CoV-2 spike proteins from different variants

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ABSTRACT

A highly efficient and robust multiple scales \textit{in silico} protocol, consisting of atomistic Molecular Dynamics (MD), coarse-grain (CG) MD, and constant-pH CG Monte Carlo (MC), has been developed and used to study the binding affinities of selected antigen-binding fragments of the monoclonal antibody (mAbs) CR3022 and several of its here optimized versions against 11 SARS-CoV-2 variants including the wild type. Totally 235,000 mAbs structures were initially generated using the RosettaAntibodyDesign software, resulting in top 10 scored CR3022-like-RBD complexes with critical mutations and compared to the native one, all having the potential to block virus-host cell interaction. Of these 10 finalists, two candidates were further identified in the CG simulations to be clearly best against all SARS-CoV-2 variants, and surprisingly, all 10 candidates and the native CR3022 did exhibit a higher affinity for the Omicron variant despite its highest number of mutations. The multiscale protocol gives us a powerful rational tool to design efficient mAbs. The electrostatic interactions play a crucial role and appear to be controlling the affinity and complex building. Studied mAbs carrying a lower total net charge show a higher affinity. Structural determinants could be identified in atomistic simulations and their roles are discussed in detail to further hint at a strategy towards designing the best RBD binder. Although the SARS-CoV-2 was specifically targeted in this work, our approach is generally suitable for many diseases and viral and bacterial pathogens, leukemia, cancer, multiple sclerosis, rheumatoid, arthritis, lupus, and more.
1 INTRODUCTION

The world has been in the grip of the “Covid-19” pandemic, caused by the severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2), for roughly two years and a half with no corner on Earth being saved. To date, more than 533 million cases and 6.3 million deaths are reported with no end in sight [https://www.who.int/] at the time of writing. The pandemic has quickly mobilized the scientific communities, pharmacological companies, and governmental health agencies. Vaccines have been developed in the shortest possible time and manufactured in large quantities on a global scale. Yet, the number of people to be vaccinated remains quite large. How the pandemics will develop largely depends on the populations becoming vaccinated worldwide in combination with non-pharmacological measures. Unfortunately, on one hand, there is strong resistance to accepting the vaccine, and on the other hand, poor countries do not have supplies for their populations. This worsens the situation as it gives time for new mutations of the virus to develop with a relatively high potential for breakthrough cases. Consequently, it requires continuously new research and constantly new treatment strategies. To be able to combat the infections, and hopefully to finally put an end to the pandemics, it is important to investigate the genomics, molecular structure and dynamics, mechanisms of binding, and the life cycle of the viruses.

Coronaviruses have four types of glycoproteins all taking part in the pathogenesis: spike (S), membrane (M), nucleocapsid (N), and envelope (E). The S-protein is a homotrimer sticking out of the viral surface. There is a specific site for the protease furin to cleavage it in the spike protein, and an open reading frame differentiating it from other RNA viruses. The furin site is particularly mutation-prone. The spike protein is highly flexible, being even able to rotate and swing, and compared to other coronaviruses, where the spike protein is rather rigid, it is much more efficient in binding. At the top of the spike protein, there is the receptor-binding domain (RBD) which binds to the angiotensin-converting enzyme 2 (ACE2) receptor of the host cell to promote an entry to infect the human cell assisted with host factors such as the cell surface serine protease TMRSS2 or cathepsin L by fusing with the cellular or endosomal membrane. Each chain of the S protein contains two subunits, S1 and S2. They are cleaved by TMRSS2 and both have important roles in the fusion to the host cell. S1 subunit leaks out from the S2 unit facilitating the membrane fusion between viral and cell membranes by hydrophobic amino acids which become buried in the cell membrane of the host cell. Some studies show that SARS-CoV-2 has a much higher affinity and exhibits stronger interaction for ACE2 than the RBD of the previous SARS-CoV-1. Other works measured similar affinities. Nevertheless, the RBD region of the S protein is obviously an ideal target for vaccines to induce host immune response and generate neutralizing antibodies. New variants tend to appear in the S1 subunit, for example, the Delta variant has several mutations there and also some specifically in RBD. The Alpha variant has ten changes in the spike protein sequence favoring its open state and making the RBD more accessible to ACE2 helping the virus to enter the cell. The trimeric spike has several conformations (open and closed) and orientations before fusion, and once bound it undergoes a structural transition forming a narrow bridge that also protects the virus from the immune defense. The probability to be in these two different conformational states (open and closed) is strongly dependent on the variants as revealed by experimental and theoretical works. There seems to be a correlation between this probability and the tendency of either asymptomatic or symptomatic clinical cases.

There are several therapy strategies that in principle can be used to combat SARS-CoV-2 infections. For example, in developing the vaccines, the main choices space from mRNA vaccines releasing a synthetic mRNA sequence encoding the virus spike protein from a nano-particle to virus vector vaccines where DNA encoding the spike protein is embedded in a viral vector to enter the cell. Despite some promising medicines including PAXLOVID, there is still no effective drug against Covid-19 and neutralizing monoclonal antibodies (mAbs) are currently the most attractive alternative. One approach that has been successfully used in the past to treat various infectious diseases is convalescent plasma therapy (CPT). This treatment uses plasma collected from COVID-19 recovering patients (those that have been cured successfully) to be administered to recipient patients that still did not develop a protective immune response. Neutralizing antibodies (Abs) from the donor's plasma helps in reducing the viral load in the recipient. Apart from neutralization, other mechanisms may also be responsible for the protective effect of CPT. Some studies revealed the presence in the plasma of COVID-19 convalescent patients of Abs capable of inducing antibody-dependent cell cytotoxicity (ADCC) and complement activation. Despite its success, one important limitation of CPT in the treatment of COVID-19 is represented by the diversity of virus.
variants found in the population which makes the selection of donors difficult. The large variation of Abs level in different plasma samples and serum incompatibility of recipients are yet another issue that has to be carefully evaluated when using CPT. On the other hand, mAbs or polyclonal antibody cocktails can be specifically engineered and biotechnologically produced to fight SARS-CoV-2. Spike protein is the primary target of these mAbs with four classes of them being described to date depending on the location of their target epitope on the S protein. Regardless of their source, either from convalescent blood or industrially produced, human mAbs are safe therapeutic tools and can be produced quickly. There are already more than 50 commercially available mAbs approved for the treatment of other inflammatory and immune disorders and other infectious pathogens. Several candidates against SARS-CoV-2 are by now in different trial phases. Patents have already been deposited (e.g., US 2021/0292393, US20210388066A1), and some mAbs are approved by regulatory agencies such as the FDA.

Among available mAbs, CR3022 is a class IV mAb that does not bind to an RBD epitope that overlaps the ACE2 binding site, but to a conserved region in RBD when the spike homotrimer is in “up” (or open) configuration exposing the RBD to interact with either ACE2 or binder molecules. Exploiting such conserved regions in RBD (despite requiring the spike protein to be at the up/open state), class IV mAbs have thus broad neutralizing activity against SARS-CoV-2, its variants, and other related coronaviruses (e.g. SARS-CoV-1). The broad sarbecovirus neutralizing activity from CR3022 has already been confirmed even for the Omicron variant. Due to the abrupt appearance of new variants of concern (VOCs) in a short time, improving a mAb that has its epitope on a conserved region of the RBD may constitute an advantage for developing biopharmaceuticals broadly effective against different variants of the virus. Moreover, attention has recently been turned again to the CR3022 antibody as a promising candidate for COVID-19 treatment and prevention as new experimental and computational evidence became available. The experimental study of Tian et al. and the computational study of Nguyen et al. showed that CR3022 binds RBD of SARS-CoV-2 with high affinity (K_D ~ 6.3 to 3 nM) in contrast with the results of Yuan et al. which gave a much lower affinity of binding (K_D ~ 115 nM). Other theoretical works showed similar binding affinities with a tendency for stronger complexes formed with RBD SARS-CoV-1. Despite these inconsistencies regarding the affinity of CR3022 for SARS-CoV-2 S-protein, this mAb is nevertheless an appealing candidate to be subjected to affinity maturation, especially due to its cross-reactivity. Therefore, the CR3022 antibody has been chosen in this study and will be the central part of our discussions.

Traditionally, mAbs production is based on immunization by hybridoma technology which uses an optimized immunogenic specific antigen to infect a host animal and then to isolate its short-lived mature B-cells from the spleen which can produce antigen-specific mAbs. These specific B-cells are then fused with immortal myeloma cells to obtain hybridomas which are able to secrete large amounts of the specific mAb. Promisingly, mAbs discovery for SARS-CoV-2 has benefited more recently also from in vitro approaches like phage and yeast display technologies. Despite their tremendous positive impact on therapeutic mAbs development, all these techniques present also challenges and limitations whose description falls beyond the scope of this paper but can be found in two excellent recent reviews. Both hybridoma and in vitro techniques sometimes lead to mAbs with lower than expected affinity and specificity. To alleviate this drawback, supplementary in vitro affinity maturation is used to increase the potency of the designed antibodies resulting in lower injected doses and side effects. The two approaches in use today in this respect include random mutagenesis and chain shuffling/site-directed mutagenesis. Both of them introduce mutations in the complementarity-determining regions (CDR) sequences (the part of the variable chains in immunoglobulins), either at random or at specific positions, followed by affinity screening of the expressed mutant using display technologies to select the best-improved antibody. Although there are successful studies that have used in vitro affinity maturation alone, several challenges related to display technologies may limit its efficient applicability. These include the limited sub-space of the possible mutations effectively accessible to display approaches, the time-consuming and laborious process of building sub-libraries, the risk of non-specific binding or loss of stability and not in the last place the long time needed and prohibitively expensive price to complete experiments.

To successfully develop potent therapeutic Abs many recent studies employ, alongside experimental in vitro techniques, computational approaches which can provide detailed structural information on the atomic-scale regarding the antibody-epitope interactions and predict possible improvements.
Attempts to computationally optimize existing Abs for SARS-CoV-2 were already reported in the literature following the work of Giron and co-authors that provided an optimized CR3022. A large number of such studies emphasize how computational tools are being routinely used for complementing experiments in the Ab design process. At present, a large diversity of algorithms for computational Ab studies are available, from those that predict the structure of an Ab based on its primary sequence only (i.e. antibody modeling), to the ones that make use of detailed structural information from X-ray diffraction data to design improved antibodies from existent ones (i.e. antibody design). Availability of experimental data regarding the complex between the Ab to be improved and its antigen (Ag) greatly improves the prediction accuracy although antibody-antigen (Ab-Ag) computationally modeled structures could be used as well. In short, the general workflow of computational design of novel paratopes includes: (i) generation of new CDRs from experimental antibody databases and their grafting/modeling on the antibody framework, (ii) CDRs sequence redesign introducing mutations, (iii) antibody-antigen docking according to the known epitope and relative orientation of the partners, (iv) binding energy evaluation using a scoring function. Eventually, these steps are performed in many rounds to encounter a better binding antibody. The generally used algorithms available today include: RosettaAntibodyDesign, AbDesign, OptCDR, and OptMAVEn. In addition to this general protocol, other molecular modeling techniques can be applied to improve the accuracy of predictions. Molecular dynamics (MD) simulations, with more than 60 years of history, represent a well-established method for analyzing the physical movement of atoms and molecules, with a long history of applicability in biological sciences. This method has the advantage of introducing the time dimension in the simulations, this way being able to capture a wide variety of biological processes like conformational changes or protein folding and how biological molecules will respond to post-translational modifications or mutations. Atomistic MD has been successfully used in the past in Ab design studies for affinity maturation of camelid nanobodies against alpha-synuclein, a weak immunogenic antigen, of bevacizumab antibody for an increased affinity to vascular endothelial growth factor A (VEGF-A), and of a toll-like receptor (TLR4) targeting Ab, to mention just a few examples. However, different steps of Ab-Ag interaction take place on different time scales, which cannot be effectively covered by conventional atomistic simulations only.

In this paper, we propose a novel multiscale approach for in silico affinity maturation of the mAb CR3022 against RBD of SARS-CoV-2 in order not only to improve binding but also to understand the molecular determinants on different time/length scales which can be exploited to further modify and modulate the binding affinity. Our approach did include an initial conventional Ab design stage followed by complementary evaluations of the best candidates using constant-pH Monte Carlo (MC) calculations, constant-charge coarse-grain (CG) MD calculations, and constant-charge atomistic MD simulations. We then selected, as the result of the affinity maturation process, the candidate that showed improved affinity in all four types of evaluations. A strong point in favor of our approach is that it uses complementary methods for binding affinity estimation, with different physical bases ranging from rigid-body long-range interaction assessment to local conformational rearrangements upon intimate Ab-Ag binding. Furthermore, our multiscale approach allowed us to test our candidates against 11 different strains of the SARS-CoV-2 virus. We identified key residues of the 6 CDRs in the light (L) and heavy (H) chains important for binding located either at the interaction interface or distant to it, together with mutations that improve binding like S35K (CDR L1), S72E (CDR L1), Y110R (CDR L3), and Y39W (CDR H1). Also, the total net charge of the mAb proved to be an important parameter that should be considered when aiming to design better binders for RBD of SARS-CoV-2.

2 THEORETICAL METHODS

Modern virology, immunology, and related fields are expanding their achievements by incorporating theoretical approaches in their daily practice. In fact, computer simulations have been playing a key role in the investigation of many biological systems including pharma and viruses. Examples are found in several study cases aimed at the understanding of a broad spectrum of phenomena, ranging from the elucidation of capsid formation to the design of antiviral molecules. Canonical simulation methods - well established in more fundamental scientific areas and material sciences - are now routinely used to enhance the understanding of the biomolecular interactions involved in infectious diseases. Following a biophysical approach, classical MD and MC methods are techniques that have a long history of successful applications in many scientific problems as standalone theoretical studies or in combination with experimental techniques. Other bioinformatics algorithms are completing theoretical resources and
helping to address challenging problems in the biomolecular world 68,87–89.

Some previous theoretical studies with CR3022

As mentioned above, CR3022 has been suggested as a promising therapeutic option to neutralize SARS-CoV-2 9,16,29,90, and is often explored in theoretical studies 33,34,91,92. In a pioneer computational study at the very beginning of the pandemic 34, using constant-pH MC simulations, Giron, Laaksonen, and Barroso da Silva showed that CR3022 - known to bind to SARS-CoV-2 RBD - could also bind to SARS-CoV-2 RBD. They also mapped the epitopes and identified the importance of electrostatic interactions for the corresponding Ab-Ag interface 34. Following different routes, Ding et al. 91 have proposed an efficient and reliable computational screening method based on “Molecular Mechanics Poisson-Boltzmann surface area” (MM/PBSA) to estimate binding free energy between SARS-CoV-2 RBD and ACE2 together with CR3022 and CB6 in good agreement with reported experimental values. Their scheme identifies the key residues that increase hydrophobicity and indicates that changing the sign of charged residues from positive to negative can increase the binding affinity. In comparison with standard MM/PBSA, their method is more accurate due to the introduction of electrostatic energy in the scheme.

Lagoumintzis et al. 93 used in silico methods in their studies of the recent hypothesis that SARS-CoV2 would interact with nicotinic acetylcholine receptors (nAChRs) and disrupt the regulation of nicotinic cholinergic system (NCS) and the cholinergic anti-inflammatory pathway. They used ROSETTA and multi-template homology modeling to study a sequence from a snake venom toxin to predict the structure of the extracellular domains of nAChRs (“toxin binding site”). Using the “High Ambiguity Driven protein-protein DOCKing” (HADDOCK) approach 94, they found a protective role of nicotine and other cholinergic agonists and observed that CR3022 and other similar mAbs show an increased affinity for SARS-CoV-2 Spike glycoprotein. To study the molecular mechanisms in SARS-CoV-2 S protein binding with several mAbs, Verkhivker and Di Paola 95 performed all-atom and CG simulations with mutational sensitivity mapping, using the BeAtMuSiC approach and perturbation response scanning profiling of SARS-CoV-2 receptor-binding domain complexed with CR3022 and CB6 antibodies, complementing it with a network modeling analysis of the residue interactions. Their results provide insight into allosteric regulatory mechanisms of SARS-CoV-2 S proteins, where the mAbs are modulating the signal communication. This provides a strategy to target specific regions of allosteric interactions therapeutically. Recently, Riahi et al. 96 presented a combined physics-based and machine learned-based computational mAb engineering platform to improve the binding affinity to SARS-CoV-2. They minimized (protonated, if needed) the Protein Data Bank (PDB) structures 97 using the “Molecular Operating Environment” (MOE) program 98, and continued with ROSETTA 52,99 and FastRelax ROSETTA 100 to later apply machine learning. MOE was used for residue scanning in combination with two machine learning models: TopNETTree (for local geometry of protein complexes) and SAAMBE3D (for a variety of chemical, physical as well as sequential, and mutation properties). CR3022 and two other mAbs (m396 and 80R) were used as templates for their diversified epitopes, complexed with SARS-CoV-2 RBD. Their results suggest combining these three mAbs for higher neutralization activity. Nguyen et al. 33 found electrostatic interactions explaining the higher binding affinity of CR3022 for SARS-CoV-2 RBD<sub>2</sub> than the 4A8 mAb in their all-atom and CG MD simulations (including steered MD). They used the Jarzynski equality to estimate the non-equilibrium binding free energy. They analyzed H-bonds and non-bonded contacts and used Debye–Hückel theory to model the electrostatic interactions for both RBD and N-terminal domain binding sites containing charged residues. Their results indicate that effective mAb candidates should contain many charged amino acids in the regions binding to spike protein. As the important residues of the spike protein involved in the binding are positively charged (Lys and Arg), the mAbs should correspondingly contain negatively charged residues (Asp and Glu) as anticipated before 34. Martí et al. 101 have applied classical MD simulations and accelerated MD (aMD) for enhanced sampling. They have included the complexes between the RBD of SARS-CoV-2 spike (S) glycoprotein and CR3022 or S309 antibodies and the ACE2 receptor. Using MD simulations, they calculated the potential of mean force to obtain the free energy profiles for the complexes with the RBD. With their protocol, they could explore a large part of the conformational space accessible to RBD-ACE2/CR3022/S309 complexes. They found the affinity in protein-protein complexes to follow the decreasing order: S/CR3022 > S/309 > S/ACE2. Shariatifar and Farasat 92 have also performed MD simulations for SARS-CoV-2 RBD complexed with CR3022 and its modifications and calculated the free binding energies. They used the FastContact software to select mutations favorable for the wild type to
produce two variants of CR3022 based on their amino acid binding conformations, showing a clear affinity enhancement compared to the wild type.

**Proposed framework for the current study**

Some of the methods mentioned above were combined in this work (see Figure 1), specifically: (a) a structural-bioinformatics-based methodology to explore macromolecules as potential candidates with higher RBD affinity using an existing experimental RBD-CR3022 complex as a template (steps 1 and 2 in Figure 1), (b) classical MD simulations combined with an enhanced sampling to precisely quantify the free energy of interactions for the RBD-binder complexation (steps 3 and 5 in Figure 1), and (c) a quicker MC sampling of a more simplified protein-protein model to investigate a larger number of complexes and search towards an optimal binder (steps 4, 7 and 8 in Figure 1). Additional atomistic simulations were also performed to explore more details of the Ab-Ag interface (step 6 in Figure 1). Such a combination of tools allows us to explore different aspects of the complex physical interactions involved in the Ab-Ag complexation process. The pros and cons of these approaches, together with the procedure in which they were combined here, are presented below. Being the Ab-Ag complexation at least a three-step process [(i) approximation of the two macromolecules (long-range interaction), (ii) structural rearrangements at short-range separation, and (iii) the “lock” phase at the close-contact level], the combination of these tools is useful to investigate all the involved phases. All the presently studied systems do not incorporate the conformation of the transition state between the open and closed states of the spike protein. It is assumed that the RBD is already available to interact with the other biomolecules.

![Figure 1](image)

**Figure 1:** Scheme for the multiple scales in silico protocol, consisting of an initial structural-bioinformatics-based methodology to explore macromolecules as potential candidates (steps 1 and 2), constant protonation state CG MD (steps 3, and 5), constant-pH (variable protonation state) CG MC simulations (steps 4, 7, and 8), and an atomistic constant charge MD simulation. At the end of this cycle, an optimized mAb with a higher binding affinity is obtained. See the text for more details.

**Structural-bioinformatics-based approach**

RAbD approach

The RosettaAntibodyDesign (RAbD) tool was employed as the first step to suggest potential binders.
candidates for the next phases (see Figure 1). The RAbD protocol consists of alternating outer and inner Monte Carlo design cycles. Each outer cycle consists of randomly choosing CDRs (L1, L2, L3, H1, H2, H3) from clusters in the RAbD database and then grafting that CDR’s structure onto the antibody framework in place of the existing CDRs (GraftDesign). The program then performs N rounds of the inner cycle, consisting of sequence design (SeqDesign) followed by energy minimization. Each inner cycle introduces mutations and structurally optimizes the backbone and repacks side chains of the CDR chosen in the outer cycle to optimize interactions of the CDR with the antigen and other CDRs. Through all the steps the Ab-Ag complex is modeled at the atomistic level. Considering that any antibody has 6 CDR’s (i.e. L1, L2, L3 on the light chain and H1, H2, H3 on the heavy chain) one has to decide which of these CDR’s should be modified with respect to the original structure (CR3022 as given by PDB id 6w41). We ran 3 sets of calculations: (A) all CDR’s were considered for full design (GraftDesign and SeqDesign) 55; (B) all CDR’s except L1 were considered for full design (GraftDesign and SeqDesign), L1 in the original fragment of CR3022 mAb (for the sake of simplicity, we shall refer to the fragment of CR3022 simply as CR3022 from now on) is an extended loop that makes a large surface area contact with the RBD of antigen (wildtype/Wuhan sequence) which stabilizes the Ab-Ag interaction. So, L1 was subjected only to SeqDesign, not to GraftDesign as the latter one also modifies the length of the CDR; (C) all CDR’s except L1 and H3 were considered for full design (GraftDesign and SeqDesign). L1 was modeled as in step (B). Among the 6 CDR’s, H3 - located in between the H and L chains - is the only one not canonical, i.e. it does not adopt classifiable conformations (clusters of conformations). When visually analyzing the RAbD generated conformations, it became obvious that some candidates with both very high scores and Ab-Ag interface surface areas were not realistic (Figure S1). For example, large Ab-Ag interface surface areas encountered in some cases were due to very long H3 CDRs. However, care should be taken when predicting H3 CDR, especially for long ones. Thus, H3 was not subjected to GraftDesign or SeqDesign in this scenario.

Poses from all three sets of calculations were considered for further evaluations, manually excluding the unrealistic ones. In total, we generated a number of 91,800 candidates for (A), 72,000 candidates for (B), and 72,000 candidates for (C) scenarios.

From the entire pool of candidates, we have selected the best improved mAbs using two criteria: (I) the complex must have an Ab-Ag interface score below -150 REU (ROSETTA Energy Units) (the native interface score is -65 REU), and (II) an Ab-Ag interface surface area larger than 1900 Å² (the native complex interface surface area is 2060 Å²). The interface surface area is important as a larger total surface area means a higher specificity due to shape complementarity. Similarly, a larger hydrophobic contact area means a higher affinity while a larger polar contact area means a higher specificity. There were 41 Ab-Ag complexes that fulfilled the imposed criteria (Table S1), which were considered for further evaluations, manually excluding the unrealistic ones. According to the ROSETTA scoring function, all these antibody-antigen complexes present better affinities for SARS-CoV-2 RBD (wild type sequence) than the original CR3022 antibody. The first 10 best candidates (P01 to P10) from the final RAbD list plus the native complex were considered for the next analyses.

**CG molecular dynamics (MD) with an enhanced sampling approach for free energy calculations**

Umbrella sampling (US) constant-protonation state MD simulations were employed to evaluate the free energy of binding of the WT and the top ten Ab-Ag complexes selected from the above described RAbD calculations (Step 3 in Figure 1). In these simulations, also known as constant-charge simulations, the protonation states of titratable groups are set at the beginning of the simulation and remain constant. For the efficiency of calculations, a reduced representation of the interacting partners and solvent was adopted, using the SIRAH 2.2 coarse grain (CG) force field (ff). This approach was successfully used in the past for estimations of free energy of binding in the case of Ab-Ag complexes. During the US procedure, the geometry of the mAb was restrained by applying weak harmonic position restraints (20 kJ mol⁻¹nm⁻² force constant) on the CG beads corresponding to the protein backbone. The distance between the center of mass (COM) of mAb and the RBD was considered as the reaction coordinate. Cylindrical positional restraints (as defined in the GROMACS 2019 suite) were applied on the RBD to allow for its movement relative to mAb along the reaction coordinate. A total number of 36 windows were used for US simulations which ensured a sufficient histogram overlapping for accurate results. Within each window, a US potential with a force constant of 1500 kJ mol⁻¹nm⁻² was applied to the COM of the RBD. Ab-Ag complexes were simulated for 16 ns in each US window, the first 2 ns being excluded from the analysis. For each Ab-
A fast constant-pH coarse-grained (CG) simulation approach for free energy calculations on a large scale

The ten selected fragments of mAbs candidates obtained from the RAbD analysis were also submitted to exhaustive investigations utilizing a fast constant-pH CG biophysical model specially designed for protein-protein complexation \[^{106-112}\] – see step 4 in Figure 1. With the term “constant-pH” we refer here to a class of simulation methods where the amino acid charge is allowed to change during the simulation run as a function of the variable surrounding at a given pH (given as input parameter) \[^{61,113}\]. These cost-effective CG simulations are less expensive than the US calculations with the CG SIRAH ff described above and have the benefit to allow the description of amino-acid charge variations (due to protonation state variations) occurring at constant pH. This implies that important electrostatic interactions as the charge regulation mechanism are properly included in the model \[^{114-118}\]. This approach is aimed to capture only the main features of the complexation phenomena with a clear emphasis on the electrostatic interactions. Different studies have highlighted the importance of these interactions for the host-pathogen and antigen-specific antibody interfaces \[^{33,34,78,119-122}\]. Successful applications of this simplified model to several biomolecular systems have been previously reported in the literature including viruses proteins and the SARS-CoV-2 RBD-mAb interactions \[^{34,109-112,123,124}\]. We shall refer to this electrostatic model as FORTE (Fast Coarse-grained Protein-protein Electrostatic Interaction). The core of such a model is the fast proton titration scheme (FPTS) \[^{125,126}\] combined with the possibility to translate and rotate the macromolecules using the Metropolis MC method \[^{127}\]. Charged and neutral spherical beads of different radii mimicking titratable and non-titratable amino acids, respectively, interact via Coulombic and van der Waals terms \[^{109,110}\]. Protein coordinates given by the structural-bioinformatics-based approach were directly converted into this amino acid model. Atoms not belonging to the protein and the solvent were removed from the input structures.

Early ideas of such a model are rooted in the works of Marcus \[^{128}\] and Jönsson and co-authors \[^{113,129,130}\]. The reduction in the degrees of freedom together with the description of the proteins at the mesoscopic level in a continuum solvent model is a clear advantage from a computational point of view. For instance, the smaller amino acid (glycine) is modeled by four sites in the SIRAH ff while a single bead is used in FORTE. The reduction is even more significant for larger residues such as aspartic acid where the drop is from 11 to 1. This can result in a decrease of the computing time by a factor of 11\(^2\) making it possible to apply it in a large-scale scenario. On the other hand, the main drawbacks are the assumption of a rigid body description to model the macromolecules and the ambiguity involved in the choices of the van der Waals contributions \[^{34,109}\]. Yet, to form the RBD-mAb complex, the two molecules have to come close before conformational changes happening at short-range separation distances can be important to add additional attraction and/or stabilize the formed complex \[^{102}\]. These limitations have not been critical in previously studied cases. The present outcome also contributes to this direction as discussed in the results section below. More details of this electrostatic protein-protein model are given elsewhere \[^{34,109,110}\]. All calculations with FORTE were performed at pH 7, 150mM of NaCl, and 298K. After equilibration, at least 3\times10^9 MC steps were run during the production phases. Three replicate runs were carried out for each simulated system. They were also used to estimate the corresponding standard deviations.

The main quantity of interest extracted from these biophysical simulations was the free energies of interactions (binding free energies), or PMF \([\beta w(r)]\), where \(\beta=1/K_BT\), \(K_B=1.380\times10^{-23}\) m\(^2\) kg s\(^{-2}\) K\(^{-1}\) is the Boltzmann constant, and \(T\) is the temperature, in Kelvin, as a function of the macromolecules separation distances \((r)\). They were directly calculated from their center-center pair radial distribution functions \([\beta w(r)]=-\ln g(r)]\) and sampled using histograms during the production phase of the MC runs. For long enough simulations, \(\beta w(r)\) is typically obtained with good accuracy and is able to reproduce experimental trends \[^{34}\]. The relatively lower computation costs in comparison with other theoretical approaches allow the repetition of the calculations at different physical-chemical conditions (e.g. different solution pHs) and macromolecular systems (e.g. RBDs with several different mutations) using high-performance computers. This is a key
aspect to investigate the binding affinities for a set of several RBDs (with all possible mutations of interest), and different binder candidates and performing further optimization of the best ones. Many runs are needed which are prohibitive with more elaborated molecular models.

Different sets of calculations were carried out with FORTE. Initially (see step 4 in Figure 1), the binding properties of the 10 best binder candidates (protein P01 to P10) that were obtained from the structural-bioinformatics-based analysis (using RAbD) were tested to form complexes with the RBD from the wild type (wt) SARS-CoV-2 (Wuhan sequence): RBD<sub>wt</sub> + Px → RBD<sub>α</sub> + Px, where x ranges from 1 to 10 (i.e., the ten top candidates). β<sub>wt</sub>(r) was calculated for all systems. Simulations with the fragment of the original mAbs CR3022 (WT) were also performed for comparison using the coordinates from the PDB id 6w41 (X-ray data with a resolution of 3.08Å, pH 4.6) 16. All these simulations were repeated for RBDs built up with sequences with different mutations (see step 7 in Figure 1) present in some relevant VOCs: (a) N501Y (Alpha/B.1.1.7), (b) K417N, E484K and N501Y (Beta/B.1.351), (c) K417T, E484K and N501Y (Gamma/P.1), (d) L452R, T478K and E484Q (Delta/B.1.617.2), (e) G339D, S371L, S373P, S375F, K417N, N440K, G446S, S477N, T478K, E484A, Q493R, G496S, Q498R, N501Y, Y505H (Omicron/B.1.1.529), and variants of interests (VOI) 32: (a) L452R (Epsilon/B.1.427/B.1.429), (b) E484K (Eta/B.1.525), (c) E484K and N501Y (Iota/B.1.526 NY), (d) L452R and E484Q (Kappa/B.1.617.1), and (e) Y453F (mink). A three-cycle process was followed in this optimization pipeline for mAb engineering with higher affinity. After all possible ALA substitutions were tested one per time, the best single mutation obtained from this first cycle was incorporated into the binder resulting in a binder’’. This new protein binder’’ contains a single ALA mutation in a specific position that resulted in the higher RBD<sub>α</sub> affinity among all tested possibilities. The process was restarted with this new macromolecule (i.e. binder’’) being subjected to another TAS loop. After this second loop, the new binder (binder’’’’) has a new amino acid substitution in its sequence together with the previous one already incorporated by the ALA single mutation in the first cycle (i.e. at this point, the binder’’’’) has two ALA replacements [A-A]. A third cycle of TAS was also done using the binder’’’’ with the double ALA mutation to explore the effect of an additional ALA replacement in its sequence (A-A-A). Besides TAS, equivalent tests were also done with the substitution of any residue by GLU in a “theoretical glutamic acid scanning”. This acid residue was chosen based on a previous work where we could observe some dependency on an increase in the binding affinities with a decrease in the total net charges of the binders 34. This process with three cycles was repeated for GLU as done with ALA. All these replacements by either ALA or GLU were combined in different ways (A-A-A, A-A-E, A-E-E, and E-E-E). Each possible combination was evaluated to guide the selection of the systems that improved the binding affinity for the RBD<sub>α</sub>. Only up to three simultaneous substitutions in the wildtype mAb were tested to avoid a complete mischaracterization of the original template protein (for both mAbs candidates classified as P<sub>best</sub> or P<sub>best-1</sub>). This could result in a complete unfolding of the macromolecule and/or an increase in the chances for

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Ab-Ag aggregation. Moreover, at this point, the number of different simulated systems was too large, and the computational costs started to be prohibitive even for a relatively cheaper CG model. Finally, the most promising mAb (identified as $P_{\text{best}}$ in our classification) designed with these three engineering cycles, i.e. the one with the higher $\text{RBD}_{\text{wt}}$ binding affinity, was tested with the other RBDs from the main VOCs and VOIs. Additionally, titration simulations employing the FPTS were used to provide the main physical-chemical properties of the individual macromolecules (all from the wild-type to P10 and each new macromolecule produced by the binding optimization technique with a single, double and triple mutation). In these runs, the total net charge numbers of these binders were computed in the absence of the RBD, i.e. the FPTS was employed for a single binder in an electrolyte solution. Such a large set of net charges data was used to further investigate the possible correlations between the binding affinities and the mAb’s net charge. This was useful to complement the previous initial analysis performed with a quite smaller number of pairs of RBD-binders in comparison with this larger data set.

The process is systematically repeated for all residues. Only one residue is replaced by ALA per time. This procedure corresponds to step 8 in Figure 1.

**Atomistic MD simulations**

In order to enhance our understanding of the intimate interactions of our best candidate with $\text{RBD}_{\text{wt}}$, and to allow for local conformational re-arrangements at the interaction interface, we used high-resolution models, based on all-atom classical force fields, to simulate Ab-$\text{RBD}_{\text{wt}}$ complexes. Constraint-free MD simulations (with the constant-protonation state) of the wild-type CR3022-RBD (PDB id 6w41) and $P_{\text{best}}$-$\text{RBD}_{\text{wt}}$ ($P_{\text{best}}$ is generically used here to refer to the best mAb identified with Rosetta score and in the previous CG simulations) complexes have been performed using the Amber99SB-ILDN ff 136 for protein description and the TIP3P 137 model for water (step 6 in Figure 1). The Ab-$\text{RBD}_{\text{wt}}$ models have been solvated with a sufficient number of solvent molecules to ensure a minimum distance between periodic images of 2.2 nm. Ions have been added to neutralize the solvated systems. The length of the simulations was 50ns at constant pressure (1 bar) and temperature (298K). The GROMACS 2019.3 suite 107 has been used for all atomistic MD simulations.

3 RESULTS AND DISCUSSION

**ROSETTA-designed antibody structure: identifying the top best binder candidates**

The general workflow of the antibody design procedure consisted of generating a large number of Ab-Ag structures using RosettaAntibodyDesign software 55, followed by ranking of the generated models according to the ROSETTA scoring function. In short, the RAbD algorithm is meant to sample the diverse sequence, structure, and binding space of an antibody-antigen complex. We have generated 235,000 candidates who were ranked according to the ROSETTA scoring function. From the total number of designed candidates, only 383 showed better affinities compared with the native CR3022-$\text{RBD}_{\text{wt}}$ complex (Figure 3). From this list, we choose 10 complexes (according to the criteria described in the “Theoretical Methods” section) that were further evaluated by free energy ($\Delta G$) calculations using both umbrella sampling (US) and FORTE (CDRs primary sequences are presented in Figure 4). We then considered, as the most promising mAbs candidates, the structures that give consensus results in all three types of evaluations.

![Diagram](image-url)
Figure 3. 2D plot of RAbD score against antibody-antigen interface SASA (solvent accessible surface area). The horizontal line corresponds to the CR3022/RBD_{wt} complex RAbD score given in Rosetta Energy Units (REU). The green square indicates the area in the two-dimensional plot that corresponds to the mAb candidates' selection criteria. The P01, i.e. the best candidate according to the Rosetta score, is indicated in this plot with an arrow.

Considering that any computational method has its limitations and accuracy, evaluating the Ab-Ag complexes with complementary computational approaches, and selecting the ones that are consistent in all evaluations, strongly increase the reliability of predictions. Thus, the observed improved affinity of the models could be considered to result from the real physics/chemistry of the complex interactions in the systems and not from any particular algorithm-related issues.

Figure 4. Sequences of the CDRs (A) and the structures (B) of the 10 selected mAbs for evaluations using US/SIRAH and FORTE methods. Dots represent identities. Residue letters in (A) were colored according to similarity with their counterparts in the reference native CR3022 sequence. In (B) RBD_{wt} (up) was colored in red and each candidate mAb (down) in a different color.

RBD_{wt}-mAb free energy of interactions explored with umbrella sampling CG simulations

Free energy calculations using US MD calculations revealed that from the list of the top 10 best RAbD
candidates only P01, P05, P06, P09, and P10 showed better binding affinities to RBD<sub>wt</sub> compared with the native CR3022 (see Figure 5 for their PMF profiles). Although all these candidates show statistically significant improved affinities compared with CR3022, the errors in estimating the binding free energies for P05, P06, P09, and P10 did not allow us to distinguish between them in terms of affinity. Among the candidates P01 seems the best, giving statistically significant better affinity of binding compared with both the native CR3022 and the P05, P06, P09, and P10 sets.

However, caution must be taken when considering absolute values obtained by US-CG SIRAH to compare with more rigorous atomistic or experimental data. As the results of Patel et al. (2017) showed, only the ΔΔG = ΔG<sub>mutant</sub> - ΔG<sub>wild-type</sub> should be regarded quantitatively, as the granularity of the SIRAH-CG model may affect the absolute values of the calculated ΔG from the PMF profiles. On the other hand, ΔDG computations based on SIRAH-CG did fit well with experimental data and are much more efficient than atomistic US approaches, which of course give more precise absolute values for ΔG. Therefore, it is meaningless to use the CG-US results for a direct comparison with experimental data like K<sub>B</sub>.

The free energy evaluations based on CG-US showed ΔΔG values of -18.3 kcal/mol for P01 and -7.1 kcal/mol to -10.8 kcal/mol for P01-P05-P06-P09-P10 group relative to the wt CR3022 antibody. These values are comparable to experimental kinetic data found in the literature for stabilizing mutations induced in Ab-Ag complexes.

![Figure 5](image)

Figure 5. Averaged RBD<sub>wt</sub>-mAb PMF profiles for the candidates that give statistically significantly better binding affinity than CR3022 in the constant-protonation state MD simulations with the SIRAH CG force field using the umbrella sampling method. The vertical bars represent the estimated standard deviations of the minimum of the PMF profiles. See the text for more details.

**RBD-mAb free energy of interactions explored with FORTE**

Despite its approximations, the simplified CpH CG protein-protein model can reproduce the main trends obtained from using the more elaborate models (see above). Figure 6 shows the minimum values of the free energy of interactions [βw<sub>min</sub>] for the complexation between the SARS-CoV-2 RBD<sub>wt</sub> with the fragment of CR3022 and with all engineered mAbs given by RabD from P01 to P10. These new mAbs candidates all have a higher RBD<sub>wt</sub> affinity as observed for the simulations with SIRAH ff too (see Figure 5). They can be ranked as P01/P06 > P02/P05/P08/P09/P10 > P04 > P03 > P05 in terms of their binding affinities. It is virtually impossible to distinguish the proteins within the groups P01/P06 and P02/P05/P08/P09/P10 due to their estimated standard deviations. Conversely, there is no difficulty in determining that the two best candidates are P01 and P06. Both proteins have virtually identical free energy profiles for their complexation with the RBD<sub>wt</sub> [-0.788(6)K<sub>T</sub>T and -0.790(6)K<sub>T</sub>T for P01 and P06, respectively]. Yet, the mAb P01 was selected as P<sub>best</sub> to follow the consensus with the best binder found in the previous biophysical simulations with the SIRAH ff (see Figure 5). Since the physical description is not the same between the two models, this reinforces the conclusion that P01 has special features for a stronger binding with the RBD<sub>wt</sub>.

Although P06 was not found equivalent to P01 according to the PMF obtained with the US SIRAH MD (see Figures 5 and 6), FORTE calculations indicate that it could still be an interesting candidate since it scores as the best together with P01. Therefore, P06 was assumed as P<sub>best</sub>, i.e. the second-best binder. The fact that FORTE provided these two binders (P01 and P06) with equivalent binding affinities instead of a single one as seen in the previous approach could be due to a couple of reasons: a) a smother energetic landscape in this CG model eventually is not able to describe the same roughness of the phase space, b) the presence of the charge regulation mechanism in the FORTE approach (absent in the constant-protonation state MD simulations) that give rise to an extra attraction, c) the lack of structural rearrangements in the FORTE approach. Nevertheless, even in the US/SIRAH ff approach, P06 is a better binder than the native CR3022 (see Figure 5).
An interesting comment here is that both CG simulations revealed some subgroups of binders with similar binding free energies among the member of each subgroup (within the error estimation). This suggests that Abs produced from previous infections or by the seroconversion of a given vaccine could have a chance to neutralize more than one strain in agreement with clinical reports. It also illustrates that a weaker response can be obtained depending on the specific mutations as also expected. Equivalent reasoning can be done for therapeutic molecules designed to prevent the attachment of the spike protein via the RBD in the human cell. For instance, Wang and co-authors described a potent mAb from covalent patients that works well against 23 variants.

It is worth reminding that a limitation of this CG model is the precise quantification of the absolute numbers in the reported values for the PMF. It is especially difficult to quantitatively (not qualitatively) reproduce experimental $K_d$ values and/or typically $\beta w(r)$ values obtained by more detailed force field descriptions. The same problem described above for SIRAH ff might be more severe for FORTE due to its higher granularity. Yet, any ambiguity in the interpretation of the obtained data can be solved when results are interpreted in relative terms. For instance, comparing the measurements between a set of similar macromolecules under the same experimental conditions. This is exactly the case for this comparison between different pairs of RBDs and a putative binder. Both CG approaches are useful for distinguishing between stabilizing and destabilizing CDR modifications, for ranking different candidates relative to the wild-type and in-between them. Qualitative comparison can still be made as shown in Figure 6.

After validating the theoretical predictions obtained with FORTE by comparison with the results from US on CG simulations with the finer-grained SIRAH force field, we expanded its analysis and tested the binding affinities of these Rosetta-designed fragments of mAbs to the selected mutated form of SARS-CoV-2 RBDs. It is particularly interesting to assess if the engineered mAbs are good binders also for the present VOCs (Alpha, Beta, Gamma, Delta, and Omicron) and VOIs (Epsilon, Eta, Iota, Kappa, mink). Such information would be vital for any practical therapeutic applications of these mAbs in the future. The free energy data at the same experimental conditions as in Figure 6 for several pairs of RBDs and mAbs was compiled and displayed as a heatmap-style plot in Figure 7. In this Figure, the darkest blue is the case with the highest binding affinity. Conversely, the darkest red is given to the system with the lowest binding affinity. This is observed for the native CR3022 (referred to here as the WT mAb) interacting with any studied RBD. All putative Rosetta-designed mAbs improved the binding features in comparison with the native CR3022. As it can also be seen, the general behavior for all RBDs is similar to what was observed for the interaction with the SARS-CoV-2 RBDwt (see Figure 6). All molecules P01 to P10 have the potential to block the interaction between virus and host cell preventing the RBD to be available to anchor ACE2. The two best candidates for all these RBDs identified by FORTE were P01 and P06. This reinforces the conclusions concerning their potential ability to neutralize both the wild-type SARS-CoV-2 virus and the main present variants. Considering the estimated error bars, all Rosetta-designed proteins follow the same trend seen in Figure 6 for RBDwt. They also tend to respond with similar affinities to all studied RBDs, i.e., a good binder for one specific RBD is a good binder for another RBD too. However, there are interesting exceptions. For example, P03 has a relatively lower affinity for RBDDelta, RBDIota, RBDKappa, and RBDmink in comparison with the RBDwt. Surprisingly, both the native CR3022 and all

Figure 6: Minima free energy of interactions values $[\beta w_{\min}]$ measured for the SARS-CoV-2 RBDwt-mAbs complexation at pH 7 and 150mM of NaCl by the CpH MC simulations (FORTE), CG-SIRAH Umbrella sampling (CG-SIRAH/US), and the RAkB in REU scores, for the wildtype RBDwt and different fragments of mAbs candidates (P01 to P10). The reference (WT) is the data for the complexation with the native fragment of CR3022 (from the PDB id 6w41). The error bars were calculated using the three replicates for each simulation system. The minima values obtained by classical MD with SIRAH ff are included in the figure for the sake of comparison. Values from the RAbD in REU are also reported for comparison.
Rosetta-designed binders show a higher affinity for RBD<sub>Omicron</sub>. This suggests that both CR3022 and its derived-mAbs have a good chance to neutralize the Omicron variant.

Figure 7: Heatmap with the minima free energy of interactions values measured for the SARS-CoV-2 RBD-mAb complexation at pH 7 and 150mM of NaCl by the CpH MC simulations (FORTE) for the RBD of the main critical variants and different Rosetta-designer binder candidates (P01 to P10). The reference (WT) is the native fragment of CR3022. All values of β<sub>W</sub> are given in K<sub>B</sub>T units. The maximum estimate error is 0.01. See the text for a description of mutations considered in each case.

**RBD-Ab local interface interactions explored with atomistic MD simulations**

We will discuss in the following the structural determinants at close-contact distances, identified by atomistic simulations, responsible for the observed better affinity of P01 compared to CR3022 against RBD<sub>wt</sub>. Mutations designed in the P01 candidate were located in all six CDRs on both the light (L) and heavy (H) chains. These mutations improved specificity and affinity by favoring the formation of supplementary salt bridges and hydrogen bonds as well as hydrophobic interactions. Protein-protein interactions are complex processes that, at short distances, are dependent on the interface properties consisting of shape complementarity, size, short-range electrostatic interactions, polar and hydrophobic interactions. Specificity in protein-protein binding is mainly given by electrostatic interactions while non-specific and van der Waals interactions are the driving force for increased affinity. We have used atomistic MD simulations to comparatively evaluate P01 with the wild-type CR3022 in terms of different interaction contributions to RBD<sub>wt</sub>-Ab binding. The simulations were analyzed for obtaining the probability distribution functions (PDF) of three interface interaction descriptors (see Figure 8), namely the number of i) Ab-Ag salt bridges, ii) non-polar contacts and iii) hydrogen bonds (between polar-polar and polar-charged residues). Thus, for each saved MD trajectory frame the number of salt-bridges formed between charged residues of the mAb and RBD<sub>wt</sub> was calculated. A salt bridge was considered to exist between acidic/basic residues if the distance between the heavy atoms of the charged moieties (i.e. side-chain carboxyl and amino) was shorter than 3.5 Å. Using the entire pool (full trajectory) of calculated salt-bridge numbers, a histogram was extracted with respect to the number of salt bridges (i.e. the distribution of a number of frames with a particular number of salt bridges). The histogram was then normalized considering the area under the curve to be 1 to obtain the PDF of mAb-RDB<sub>wt</sub> salt-bridges number. The number of non-polar contacts was obtained by counting, for each frame, the hydrophobic residue atom pairs (mAb-to-RBD<sub>wt</sub>) within a distance shorter than 3.5 Å. A similar procedure as above was applied to calculate the histogram and PDF of non-polar contacts. For polar interaction histogram and PDF, we counted the number of hydrogen bonds between polar residues and polar to charged residues in the mAb-RBD<sub>wt</sub> pair. It can be seen from the plots in Figure 8 that the main contributions to the increased affinity of P01 to RBD<sub>wt</sub>, compared with the wild-type CR3022, come from an increased number of salt-bridges and a higher hydrophobic contact area to RBD<sub>wt</sub>. The average number of salt-bridges at the P01 interface to RBD<sub>wt</sub> is ~4 while for the wild-type CR3022 is ~3. The Lys35 residue, which replaced a Ser35 in the CR3022, is an important player in short-range electrostatic interactions at the P01-RBD<sub>wt</sub> interface as it is strategically placed on the tip of the L1 loop and has the optimum side-chain length to form a strong salt-bridge with the acidic Glu in position 516 on the RBD<sub>wt</sub>. The average number of hydrophobic contacts between P01 and RBD<sub>wt</sub> (~95) is significantly bigger than for the CR3022-RBD<sub>wt</sub> pair (~125), which will result in a higher binding affinity. Four mutations in P01, that replace original residues in CR3022 with more hydrophobic ones, contribute to this increased hydrophobic contact area: W40F, S110A, G111A, I112L. They form a hydrophobic patch in the middle of the mAb–RBD<sub>wt</sub> interface that interacts with Lys378(4 methylene groups)-Tyr380-Pro-384-Tyr369-Val382 on the RBD<sub>wt</sub> counterpart.
To get a detailed picture of the RBD<sub>wt</sub>-P01 interaction interface, we have analyzed the impact of the introduced mutations on the local molecular environment of the complex, which is discussed below for each CDR separately.

**L1 CDR design.** The tyrosine residue in position 31 in L1 CDR was replaced by Asn in P01, which easily participates in hydrogen bonding with both Thr430 and Asp428 on RBD<sub>wt</sub>. The interaction of the native Tyr31 with Thr430 is impeded due to the longer side chain of tyrosine compared with asparagine. One key mutation present in L1 is S35<sup>K</sup> which replaces a neutral serine residue with a positively charged lysine. The original Ser35 residue is located on the tip of the L1 loop that penetrates deep into the Ab-Ag interface stabilizing the complex. Replacing the serine with lysine in this position provides a strong salt bridge formation between the long basic side chain of Lys35 and the acidic Glu in position 516 on the RBD<sub>wt</sub>, an interaction not present in the native CR3022. Replacement of Asn with Gln in position 37 provided further stabilization as the longer side chain of glutamine compared with asparagine allowed for a new interaction of Gln37 on the antibody with His519 on the RBD<sub>wt</sub>.

**L2 CDR design.** The replacement of the neutral hydrophilic Ser72 with an acidic Glu residue in P01 introduces a strong salt bridge formation between the long basic side chain of Lys35 and the acidic Glu in position 516 on the RBD<sub>wt</sub>, an interaction not present in the native CR3022. Replacement of Asn with Gln in position 37 provided further stabilization as the longer side chain of glutamine compared with asparagine allowed for a new interaction of Gln37 on the antibody with His519 on the RBD<sub>wt</sub>.

**L3 CDR design.** Another mutation that increases binding specificity and affinity was introduced in the L3 loop replacing Tyr110 with Arg in P01. The substitution of neutral tyrosine with the positively charged arginine allows for a salt bridge formation with the neighboring Asp428 on RBD<sub>wt</sub>. The P01 designed L3 CDR has two supplementary residues Pro135 and His136 compared with the native CR3022. This makes the L3 loop more extended towards RBD<sub>wt</sub>, increasing the Ab-Ag contact surface area. The L3 loop also contains the mutation of Thr112 to Tyr112 which, due to the new conformation of L3, allows for hydrogen bond formation between Tyr112 and three partners on RBD<sub>wt</sub>: the side chain of Glu414, the backbone carbonyl oxygen atom of Pro412, and the side chain of the acidic Asp427, interactions not present in the native CR3022.

**H1 CDR design.** The Tyr residue in position 28 was replaced by Gly. This mutation seems advantageous because the smaller side chain of glycine and the particular conformation of the designed H1 loop allowed the side chain of Tyr369 from the RBD<sub>wt</sub> to become buried on the mAb surface. The hydroxyl moiety of Tyr369 can also participate in hydrogen bonding with the side chain -NH group of Trp39, a residue that represents another mutation in the Rosetta-designed H1 loop: Y39W. Trp40 was replaced by a more hydrophobic Phe residue, whose side-chain inserts between the cycle of Tyr380 and the long hydrocarbon chain of Lys387, thus hydrophobically stabilizing the Ab-Ag interaction.

**H3 CDR design.** Serine in position 133 was mutated to a His residue which inserts in between G381 and Asp427 on the RBD<sub>wt</sub>, forming a hydrogen bond-mediated bridge between the -NH group of G381 and the backbone -C=O group of Asp427.
Physical insights to design a more efficient monoclonal antibody for the RBD

Individual residue electrostatic contributions to the complexation RBD<sub>wt</sub>-P01 were investigated through the TAS procedure described in Section 2. Each new protein produced with the ALA single mutation (i.e. binder') is referred to here as P01'. All P01 ionizable residues were replaced by ALA one per time to give P01', and the corresponding $\beta_{w(r)}$ functions for the complexation RBD<sub>wt</sub>-P01' were estimated by the FORTE approach. The differences in the $\beta_{w_{\text{min}}}$ values observed for P01 and P01' [$\Delta\Delta G=\beta_{w_{\text{min}}}(P01')-\beta_{w_{\text{min}}}(P01)$] are portrayed in Figure 9. The purples lines indicate the maximum estimated errors (0.01 K<sub>B</sub>T). Free energy shifts larger than 0.01 K<sub>B</sub>T are considered significant. Positive shifts indicate mutations that negatively impact the complexation process. These amino acids (ASP-9.LC, GLU-17.LC, ASP-72.LC, ASP-88.LC, ASP-100.LC, GLU-11.HC, GLU-23.HC, GLU-67.HC, ASP-83.HC, and ASP-137.HC) are critical for this molecular mechanism, i.e., replacing one of them with a neutral residue reduces the RBD<sub>wt</sub> affinity which is not good to block the virus-cell interaction. For instance, the substitution of the ASP residue in position 88 at the light chain (“ASP-88.LC” at L2) by ALA decreases the binding affinity by 0.03 K<sub>B</sub>T units. Conversely, negative values of $\Delta\Delta G$ correspond to mutations that can improve the capability of P01 to bind the RBD<sub>wt</sub> which is potentially useful to prevent the viral entry into the host cell. Note that most of the key amino acids that favor the RBD<sub>wt</sub>-P01 complexation are acidic residues (ASP and GLU). There are just a couple of exceptions (e.g. ARG-77 at the light chain) but always with smaller differences than the estimated standard deviations ($\Delta\Delta G<0.01$ K<sub>B</sub>T). The neutralization of a basic ionizable group (e.g. LYS-13 at the heavy chain) implies a reduction of the total net charge of the binder and an improvement in its binding features ($\Delta\Delta G_{\text{LYS-13.HC}}=-0.03$ K<sub>B</sub>T). Indeed, it was noticed before that a decrease of the binder net charge gives a higher RBD<sub>wt</sub> affinity. This information is a useful insight to design a more efficient therapeutic binder that can avoid the attachment of the RBD to the human cell.

The most important cases where the single ALA mutation in P01 has a stronger influence on the SARS-CoV-2 RBD<sub>wt</sub>-P01 complexation were mapped in the molecular structure. This can be seen in Figure 10. Interestingly not only amino acids at the antigen-antibody interface are critical for the complexation. In fact, amino acids buried inside the protein structure can also have an important influence on the complexation due to the long-range nature of the electrostatic interactions and the electrostatic coupling between the titratable groups. This is in line with a broader view of epitopes (and paratopes) defined as “electrostatic epitopes”, where any ionizable group can affect the interactions and drive the complexation.

![Figure 9: Individual residue electrostatic contributions to the complexation RBD<sub>wt</sub>-P01 obtained with the theoretical ALA scanning procedure applied to P01. Acid and basic residues are represented by red and blue, respectively. Data for the amino acids belonging to the light (LC) and heavy (HC) chains are given in the left and right panels, respectively. Calculations were done with the FORTE approach. $\Delta G$ (in K<sub>B</sub>T units) is defined as the difference between the minimum value measured in $\beta_{w(r)}$ for the complexation RBD<sub>wt</sub>-P01' [$\beta_{w_{\text{min}}}(P01')$] and the corresponding quantity for the original P01 [$\beta_{w_{\text{min}}}(P01)$] for each new mutation. See the text for more details.](image.png)
Figure 10: Molecular structure of the RBD<sub>wt</sub> complexed with P01. The RBD<sub>wt</sub> is represented in green. All ionizable residues from the binder P01 whose single ALA mutation resulted in representative ΔΔGs (values larger than the estimated errors) are highlighted by blue (basic residues) and red (acid residues). The most important mutations (see Figure 9) are labeled and represented with vdW spheres. The letters L and H are used to refer to the light and heavy chain, respectively.

The total net charge of the mAbs and their binding affinities

As mentioned above and previously indicated, lower charged mAbs tend to have a higher RBD affinity. In Figure 11a, this correlation is explored for the WT and Px (x: 01 to 10). Although there is no perfect linear behavior, it can be seen that there is indeed this tendency. This is more clearly seen in Figure 11b, where the data from all single, double and triple mutations investigated by the theoretical ALA/GLU scanning is shown together. This data suggests that Coulombic forces are mainly responsible for driving the association of this antigen with the studied mAbs. As a matter of fact, there are now accumulative data on the importance of electrostatic interactions for this system. Certainly, other physical interactions can participate in this associative mechanism that is not solely but largely controlled by Coulombic forces. Therefore, the total net charge of the mAb is an important physical-chemical parameter to design better binders with higher affinity. This is what was next explored towards a more specific and efficient macromolecule capable of preventing viral infection.

Additional optimization of the best RBD binder

The two best candidates P01''' (LC: R18E and HC: K20E and K84A) and P06''' (HC: K20A, G61E, and K84A) obtained through the three cycles electrostatic optimization pipeline (based on the TAS procedure) were tested against the most common SARS-CoV-2 VOCs and VOIs. This was done to assess if these macromolecules could have the potential to stop the virus from working efficiently targeting RBDs from common VOCs and VOIs. Figures 12a and b show the β<sub>Wmin</sub> values for the complexes RBD-P01''' and RBD-P06''', respectively. The new optimized binders P01' and P06' are able to form stable molecular complexes with all studied RBDs. Due to the additional electrostatic optimization, they have improved RBD binding affinities in comparison with the initial template (P01 or P06) provided by the RAbD approach. Consequently, they have more chances for a successful neutralization in vitro/in vivo of the virus. Moreover, this result indicated that the integrated biophysical modeling multi-approach used here offers an efficient route to designing better macromolecular ligands. Intriguingly both putative mAbs P01''' and P06''' have a stronger affinity for some RBDs from variants that are threatening the immune response. For instance, P01''' has the highest affinity observed for both RBD<sub>Delta</sub> and RBD<sub>Omicron</sub>. This is a promising result to deal with the present crisis offering two ideal candidates to treat the pathogenic threats from different SARS-CoV-2 variants. Quite recently, a pre-print work suggested cross-reactivity specifically between antibodies for Delta and Omicron variants. P01''' is slightly better than P06''' due to its higher RBD affinity for all mutants in general (e.g. β<sub>Wmin</sub>=-0.90(1)K<sub>B</sub>T for RBD<sub>Delta</sub>-P01''', and β<sub>Wmin</sub>=-0.92(1)K<sub>B</sub>T for RBD<sub>Delta</sub>-P06'''). From a bioinformatics point of view, it is interesting to point out that an optimized mAb is solely obtained by the three cycles electrostatic optimization pipeline (CR3022) is not as good as the one achieved by the present strategy.
Figure 11: Correlation between the total net charge number of each fragment of mAb and the corresponding $\beta_{\text{min}}$ values for their complexations with SARS-CoV-2 RBD$_{\text{wt}}$. (a) Left panel: Data for each fragment of Rosetta-designed mAb (P01 to P10). (a) Right panel: Data for each mutated fragment of P01 and P06 was obtained during the theoretical ALA scanning procedure. The data for the total net charge numbers were obtained from titration simulations with a single protein in the absence of the RBD$_{\text{wt}}$ by the FPTS at pH 7. All $\beta_{\text{min}}$ values were computed from RBD$_{\text{wt}}$-complexation studies using FORTE at pH 7 and 150mM of NaCl.

Figure 12: Binding RBD affinities for optimized mAbs obtained through the three cycles electrostatic optimization pipeline. (a) Left panel: mAb P01''' based on the Rosetta-designed P01. (b) Right panel: mAb P06''' based on the Rosetta-designed P06. Data from the estimated $\beta_{\text{min}}$ values for the molecular complexation between RBDs from different variants with P01''' (P01 with three mutations LC: R18E and HC: K20E and K84A) and P06''' (P06 with three mutations HC: K20A, G61E, and K84A), respectively, are shown in blue bars. Data for the complexes with the original Rosetta-designed mAbs (RBD$_{\text{wt}}$-P01 and RBD$_{\text{wt}}$-P06) before the electrostatic optimization process and CR3022' (RBD$_{\text{wt}}$-CR3022') are given in the purple and red bars, respectively, for comparison. The purple lines are drawn to guide the eyes for the comparison with the outcomes for the RBD$_{\text{wt}}$. See the text for more details.

4 CONCLUSIONS
RosettaAntibodyDesign protocol was applied to sample the sequence, structure, and binding space of many anti-SARS-CoV-2 mAb-RBD complexes. A total of 235,000 candidates were obtained, and among these, those that showed a higher binding affinity compared to the native CR3022-RBD$_{\text{wt}}$ complex were identified, and finally, 10 complexes were chosen to be evaluated further in binding free energy calculations. Both Umbrella sampling with
SIRAH 2.0 coarse-grained force field and the in-house Fast cOarse-grained pRotein-proTein modEl (FORTE) based on fast proton titration scheme (FPTS) were utilized both giving mutually consistent results. After compiling the results at similar conditions to experiments in a heat map we can observe that all the selected RAbD produced CR3022-RBD complexes show a stronger binding affinity compared to native CR3022 and a higher potential to block the interaction between the virus and the host cell. Further, two candidates, P01 and P06, show a clearly strongest interaction.

In a closer analysis, we can find that amino acids both at the surface and also deeper inside the protein structure are critical for the complexation through electrostatic coupling between titratable groups and are characterized as “electrostatic epitopes”.

Furthermore, we can observe that the net charge of the mAbs is one important determinant in binding affinity and there is a clear nearly linear tendency for lower charged mAbs to exhibit higher affinity to RBD. The Coulombic forces are driving the antigen-antibody association, as also observed in previous studies, while other short-range interactions are important in the close-range association as well.

We have performed atomistic simulations for the best candidates to study the structural determinants behind the strong affinity of P01 to RBD. The mutations in P01 were in all six complementarity-determining regions (CDR) sequences on both light and heavy chains, improving both the specificity and affinity in favoring the formation of supplementary salt-bridges and H-bonds, as well as through hydrophobic contacts. The two best candidates P01 and P06 were tested against to most common SARS-CoV-2 mutations to hinder the virus targeting RBDs. We could see they both can form stable complexes with all RBDs included in our study thereby having the potential to neutralize the virus both in vitro and in vivo.

Our combined multi-scale approach is found to be a fast, robust, and reliable tool to design better macromolecular ligands allowing us to identify the best candidates for the different variants of SARS-CoV-2 including Omicron. It is a pragmatic approach for a short development cycle for SARS-CoV-2 diagnosis, treatment, and prevention.

This multi-approach is a general theoretical framework towards high specific antibodies for SARS-CoV-2 and can be extended to other diseases (e.g. other viral and bacterial pathogens, leukemia, cancer, multiple sclerosis, rheumatoid arthritis, lupus) both for diagnosis and therapeutic purposes.

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