Mutations in the SARS-CoV-2 spike RBD are responsible for stronger ACE2 binding and poor anti-SARS-CoV mAbs cross-neutralization

Masaud Shah, Bilal Ahmad, Sangdun Choi, Hyun Goo Woo

Abstract

Severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2), which causes coronavirus disease 2019 (COVID-19), is a novel beta coronavirus. SARS-CoV-2 uses spike glycoprotein to interact with host angiotensin-converting enzyme 2 (ACE2) and ensure cell recognition. High infectivity of SARS-CoV-2 raises questions on spike-ACE2 binding affinity and its neutralization by anti-SARS-CoV monoclonal antibodies (mAbs). Here, we observed Val-to-Lys417 mutation in the receptor-binding domains (RBD) of SARS-CoV-2, which established a Lys-Asp electrostatic interaction enhancing its ACE2-binding. Pro-to-Ala475 substitution and Gly482 insertion in the AGSTPCNGV-loop of RBD possibly hinders neutralization of SARS-CoV-2 by anti-SARS-CoV mAbs. In addition, we identified unique and structurally conserved conformational-epitopes on RBDs, which can be potential therapeutic targets. Collectively, we provide new insights into the mechanisms underlying the high infectivity of SARS-CoV-2 and development of effective neutralizing agents.

1. Introduction

SARS-CoV-2 causes coronavirus disease 2019 (COVID-19), which was initially reported in Wuhan city in China and disseminated around the world very quickly [1,2]. However, the origin and global spatial spread of the virus, particularly in Europe and North America, has been hotly contested [3]. SARS-CoV-2 is a novel beta coronavirus [4,5] and apparently transmits from human-to-human markedly faster than previously known coronaviruses, such as SARS-CoV and MERS-CoV [6,7]. Although the epidemiological features of SARS-CoV-2 are largely unknown, asymptomatic transmission and poor self-quarantine measurements of infected persons are thought to be the most crucial reasons for the uncontrolled spread. In the current pandemic situation of COVID-19, there is an urgent need to develop effective therapeutics and vaccines. Several pre-existing anti-viral drugs and novel vaccines including Sputnik V, Coronavac, mRNA-1273, d5-nCoV, inactivated vaccine, Covishield, and BNT162 are now on clinical trials [2,8].

Among the mode of action mechanisms of the viral infection, cell-recognition and cell-entry are the most crucial steps that determine viral infectivity and pathogenesis [9]. Like SARS-CoV, SARS-CoV-2 uses spike (S) glycoprotein to interact with the human respiratory and epithelial cells expressing angiotensin-converting enzyme 2 (ACE2) receptors [10,11]. The ectodomain of S protein is an ~1200 amino acid long class 1 fusion protein and normally exists in a trimeric metastable pre-fusion conformation of “laying down”, which can undergo conformational rearrangement and acquire an ACE2-feasible conformation, i.e., “up or standing” [11,12]. The “down” and “up” poses are differentiated due to the conformational rearrangement of the receptor binding domain (RBD, ~200 amino acid) in the S1 subunit of the S protein. The position switching of the RBD from down-to-up facilitates receptor binding while up-to-down helps the virus to evade the immune surveillance [12,13]. The availability of receptor-binding determining region (RBDR) to the ACE2 is controlled by the hinge-like conformational motion of the RBD [11]. A mutation at the 614 amino acid i.e. Asp614-Gly has been reported to enhance the up conformation of the RBD that makes the virus more infectious and susceptible to neutralizing antibodies [14,15]. Thus, S protein is indispensable for the virus survival and remains a priority target for antibodies and peptides-based biologics to curb viral entry. Although the receptor-binding mechanism of the spike protein has been widely studied and well-established now, we constructed an ectodomain model of the SARS-CoV-2 S protein in its pre-fusion form with respect to the two-conformation-states of RBD that pose

Corresponding author at: Department of Physiology, Ajou University School of Medicine, Suwon, Republic of Korea.
E-mail address: hg@ajou.ac.kr (H.G. Woo).

https://doi.org/10.1016/j.csbj.2020.11.002
2001-0370 © 2020 The Authors. Published by Elsevier B.V. on behalf of Research Network of Computational and Structural Biotechnology. This is an open access article under the CC BY license (http://creativecommons.org/licenses/by/4.0/).
a major challenge to the neutralizing antibodies and vaccines development. In addition, using the ACE2-RBD structures of SARS-CoV RBD (sRBD) and SARS-CoV-2 RBD (cRBD), we investigated their relative dynamic interaction, stability, and binding affinities. Using these structural-insights, we suggest peptides-based therapeutics that could possibly hinder the ACE2-RBD interaction. Further, we identified conformational epitopes on the cRBD that ought to be taken into account while investigating the cross-reactivity of anti-sRBD monoclonal antibodies (mAbs) or designing new mAbs and peptide therapeutics against SARS-CoV-2 S protein.

2. Methods

2.1. SARS-CoV-2 spike protein modeling

Almost all ectodomain trimeric structures of the SARS-CoV-2 spike protein that are mainly resolved through cryogenic electron microscopy (Cryo-EM) are incomplete, lacking vital structural information. This is because of poor electron density of the highly flexible looped regions that connect different motifs and subdomains in spike protein. Thus, understanding the structural insights of the trimeric spike requires extensive loop-modeling of the existing structures or new template-based modeling. As SARS-CoV and SARS-CoV-2 share significant identity in their spike proteins (>76% total identity; 92% in the S2 domain and 64% in the S1 domain) and by the time we conducted this study no ectodomain structures of the SARS-CoV-2 was reported; we used multiple SARS-CoV structures as template (PDB ID: 5X5B, 6ACG, and 5I08) to build the ectodomain (a.a 13–1122) spike protein model in Molecular Operating Environment (MOE) suit (2019.0102). The amino acid sequence of the spike protein that is among the first reported SARS-CoV-2 sequence and placed as reference sequence in NCBI was retrieved (accession #: NC_045512) and used in spike modeling. MOE modeling package uses a four-step procedure for model building i.e. initial partial geometry specification (to the templates), insertion and deletion (depending on the target and template sequence alignment), loop selection and sidechain packing (if template has missing loop structure), and final model selection and refinement (based on model scoring). Taking the RBD conformational position “up” or “down” into account, two trimeric spike models were generated. The spike-ACE2 complex was generated by structural superposition of the modelled trimeric SARS-CoV-2 “up” and SARS-CoV structure, 6ACG. The models were further refined through the structural correction package and validated through the protein geometry (this package evaluate the model through Phi-Psi angles, dihedrals, rotamers energies etc.) package distributed in MOE. For the molecular dynamics simulation analysis, the ACE2-RBD regions were truncated from the trimeric spike-ACE2 complexes of both SARS-CoV-2 and SARS-CoV (PDB: 6ACG). Protein surface and patch analyses were performed in MOE as described previously [16]. Further details about the MOE-based modeling procedure and the protein-protein interface analysis have been described in our previous studies [17–19].

3. Spatial epitope prediction of mAbs docking onto SARS-CoV-2 RBD

For the mAbs, CR3014 and CR3022, modeling, an antibody modeling package of MOE was used and only the single chain variable fragments (scFv) were constructed [20]. The complementarity-determining regions (CDR) were annotated and numbered as described previously [21,22]. Structural data of other mAbs including 80R, m396, F26G19 and s230 were obtained from PDB [23–26]. For mAbs docking, a built-in protein–protein docking procedure was used in MOE suit. In docking simulation, CDR regions of mAbs were considered as ligand-sites instead of entire scFv regions. Conformational epitopes of cRBD were predicted using Epipred implemented in SAbPred web server, utilizing the CDR information of an input mAb and predicting conformational epitopes on a target protein [27,28]. By calculating geometric fitting and knowledge-based asymmetrical antibody-antigen scoring, the epitopes of the cRBD were predicted and ranked on the basis of combined conformational matching of the antibody-antigen structures. The score of the epitope is calculated using the following formula:

\[
\text{Epitope Score} = \sum_{n \in C} \frac{1}{d(n)} \Pr(T_{ab}, T_{ag})
\]

where \(T_{ab}\) and \(T_{ag}\) are the amino acid types of the antibody and antigen residues, respectively, which belong to node \(n\).

3.1. Molecular dynamics (MD) simulations

MD simulations were performed using GROMACS 2019.3. The RBD-ACE2 complexes were solvated with TIP3P water filled cubic box of dimension boundaries extended to 10 Å from protein atoms. To neutralize the charge of the simulation system the Na⁺/Cl⁻ counter ions were added and energy minimization was performed using CHARMM37 force field [29] and steepest descent algorithm. The energy-minimized systems were simulated for 0.2 ns with an NVT ensemble followed by 0.2 ns with NPT ensemble for equilibration, under constant temperature and pressure, respectively. The temperature and pressure were coupled with V-rescale and Parrinello-Rahman barostat methods [30], respectively. The long-range electrostatic interactions were computed using the particle mesh Ewald algorithm [31], while LINCS algorithm was applied to constrain the bond lengths [32]. After temperature and pressure equilibration, MD simulations were carried out for 30 ns for each system. A detailed procedure has been described in our previous studies [16,18].

3.2. Binding free energy analysis

Since both the cRBD and sRBD bind to the same ACE2 protein, we used Molecular mechanics Poisson-Boltzmann surface area (MM-PBSA) approach [33] to calculate the relative binding free energies of both complexes. In GROMACS, the built-in tool g_mmpbsa and APBSA were called for the MM-PBSA calculations. For g_mmpbsa analysis, the dielectric constant of the aqueous solvent was set to 80, and the interior dielectric constant was set to 4; the surface tension constant g was set to 0.022 kJ/mol. As g_mmpbsa tool is compatible with older versions of GROMACS (versions 5 or lower), the “tpr” files created by GROMACS 2019.3 were recreated through GROMACS 5.1 and used for binding energies calculations, as described previously [34]. The relative binding energies of the complexes were approximated according to the following energies terms.

\[
\Delta G_{\text{bind}} = \Delta G_{\text{MM}} + \Delta G_{\text{sol}}
\]

Which is further extended as:

\[
\Delta G_{\text{MM}} = \Delta E_{\text{MM}} + \Delta G_{\text{polar}} + \Delta G_{\text{non-polar}}
\]

where as;

\[
\Delta E_{\text{MM}} = \Delta E_{\text{cov}} + \Delta E_{\text{elec}} + \Delta E_{\text{disp}}
\]

and;

\[
\Delta G_{\text{sol}} = \Delta G_{\text{polar}} + \Delta G_{\text{non-polar}}
\]

where \(\Delta G_{\text{MM}}\) is gas-phase molecular mechanical energy change and \(\Delta G_{\text{sol}}\) is the solvation free energy change upon binding. All these changes were computed via ensembles which are averaged over a
set of conformations sampled throughout the simulation trajectories at 0.01 ns time interval. The $\Delta E_{\text{traject}}$ is calculated through molecular mechanics (MM) by combining three energy terms: the van der Waals energy change ($\Delta E_{\text{vdW}}$), the electrostatic energy change ($\Delta E_{\text{ele}}$), and the covalent energy change ($\Delta E_{\text{cov}}$) which further consists of change in the bond ($\Delta E_{\text{bond}}$), angle ($\Delta E_{\text{angle}}$) and torsion terms ($\Delta E_{\text{torsion}}$). The solvation free energy ($\Delta G_{\text{sol}}$) is computed by combining both polar and non-polar energies.

4. Results and discussion

4.1. Structural modeling of the SARS-CoV-2 spike and ACE2 interaction

A full-length spike protein is composed of S1 and S2 subunits, which further contains sub-domains and motifs with distinct functions (Fig. 1A). Based on the hinge-like motion of the RBD of S1 subunit, the trimeric S protein exists in a transiently symmetric or asymmetric conformation. Recent studies revealed that the cRBD, like other coronaviruses, exhibit stochastic breathing-like movement, facilitating receptor binding to the exposed RBD and subsequent shedding of the S1 subunit [11,12]. This hinge-like motion of the RBD is considered as one of the immune evasion strategy of the SARS-CoV-2 where the virus masks its RBD from host antibodies in down conformation and makes it available for the ACE2 in up position [12,13]. Owing to the limitation in the availability of crucial structural information in the Cryo-EM SARS-CoV-2 spike protein structure (PDB ID: 6VSB, we modelled the monomeric and trimeric models of the SARS-CoV-2 S protein in different conformational states (with respect to the RBD position, and with or without ACE2) using SARS-CoV spike proteins as template. When compared with Cryo-EM structure, we could see a high degree similarity in the structural folds of both modeled protein and reported spike EM-structures in up and down conformations. To understand the structure more clearly and validate our

---

**Fig. 1.** Modeling of the SARS-CoV-2 spike protein and its ACE2 binding mechanism. A) Domain architecture of the monomeric full-length S protein and the 3D monomeric spike model corresponding to a.a. 13–1122 in the ectodomain is shown. The furin-cleavage site at the junction of S1-S2 junction is highlighted with yellow arrow. B) The RBDs are displayed in receptor accessible “up” (cyan color) and non-accessible “down” (yellow color) conformation on the trimeric spike. ACE2 makes steric clash with the adjacent RBD if bound to RBD$^{\text{down}}$. In RBD$^{\text{up}}$ conformation, single trimeric spike can accommodate up to three ACE2 molecules, depending on the number of RBDs in up conformation. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)
model, the Cryo-EM structure and our models were superimposed in standing (in the presence of ACE2) and laying conformation. Total root mean square deviation (RMSD) of the backbone atoms of S protein in standing pose was 2.93 Å while that of the RBD region was 1.45 Å (Supplementary Fig. 1A). By contrast, RMSD values in the laying pose were 2.56 Å and 1.38 Å, respectively. This validation supports our model and further suggests that the structure modelled by Cryo-EM poorly capture the looped and highly flexible regions in spike protein, which is critical for the binding of ACE2 and antibodies. To investigate that the up conformation of RBD is indispensable for ACE2 binding, we constructed models where ACE2 was bound to both up and down RBD in trimeric spike.

Considering the possibility of ACE2 binding to the down RBD where the RBD of the adjacent spike protomer may exist in up or down conformation, we observed that in both cases ACE2 may not bind to the down RBD due to the steric hindrance of the adjacent RBDs (Fig. 1B). On the contrary, ACE2 dock well onto the RBD in up conformation. One spike trimer could possibly accommodate two ACE2 proteins simultaneously, depending on the occurrence of two standing RBD at the same time. Depending on the state of ACE2, which exist in monomeric (solution form) as well as membrane-bound dimeric form [23], the one spike trimer–two ACE2 notion could be possible in two ways. First, two soluble ACE2 bind to two up RBDs on the same spike trimer (shown in
conserved between the two viruses with few mutations including in this structure was taken from SARS-CoV, which is considerably in our model; nonetheless, the chimeric structure lacks the crucial shares structural folds with our model and recently reported Cryo-through X-ray crystallography[38]. Overall, this chimeric structure study has delineated the interface of ACE2 with a chimeric RBD supporting the reliability of our computational model. Another strated the interaction between SARS-CoV-2 RBD and ACE2 by per-

lysine 417 mutation in cRBD plays a crucial role in ACE2 recogni-

Table 1

| RBD-ACE2 interface (0 ns) | RBD-ACE2 interface (20 ns) |
|--------------------------|---------------------------|
| Virus | RBD | ACE2 | Bond | g^mol | Dist Å | Atom | RBD | ACE2 | Bond | g^mol | Dist Å | Atom |
| SARS-CoV | Arg426 | Gln329 | H | ~35.88 | 2.99 | – | Tyr481 | Lys353 | H | ~9.80 | 2.89 | b |
| | Gly482 | Lys353 | H | ~7.20 | 2.80 | b | Tyr436 | Asp38 | H | ~5.80 | 2.75 | – |
| | Tyr491 | Gln37 | H | ~4.90 | 2.56 | – | Gly488 | Gln354 | H | ~4.00 | 2.80 | bb |
| | Tyr436 | Asp38 | H | ~3.70 | 2.53 | – | Thr486 | Asn330 | H | ~3.40 | 2.83 | b |
| | Asp463 | Ser19 | I | ~1.49 | 3.61 | b-b | Asp463 | Gln24 | H | ~2.60 | 2.78 | – |
| | Asn479 | Gln35 | H | ~1.30 | 3.18 | – | Thr486 | Asp355 | H | ~1.70 | 2.60 | – |
| | Thr486 | Tyr41 | H | ~1.20 | 2.99 | – | Tyr481 | Gln34 | A | ~0.70 | 3.80 | b |
| | Asn473 | Gln24 | H | ~1.00 | 2.89 | – | Gln493 | Lys31 | H | ~1.10 | 2.85 | – |
| | SARS-CoV-2 | Lys417 | Asp30 | H | ~30.83 | 2.71 | – | Gln493 | Lys31 | H | ~1.10 | 2.85 | – |
| | | | | | | | | | | | | |
| | Tyr505 | Gln37 | H | ~4.60 | 2.57 | – | Tyr449 | Asp38 | H | ~2.80 | 2.58 | – |
| | Gly502 | Lys353 | H | ~3.60 | 3.10 | bb | Asn501 | Lys353 | H | ~2.80 | 3.05 | b |
| | Gln498 | Lys353 | H | ~2.90 | 3.00 | – | Gln474 | Gln24 | H | ~2.70 | 2.78 | – |
| | Gln506 | Gln325 | H | ~2.60 | 2.82 | – | Asn487 | Gln24 | H | ~2.40 | 2.85 | – |
| | Asn487 | Gln325 | H | ~1.60 | 2.80 | – | Gln498 | Lys353 | H | ~1.60 | 3.03 | – |
| | Gln493 | His34 | H | ~0.50 | 3.07 | – | Thr500 | Lys353 | H | ~1.40 | 2.71 | b |
| | | | | | | | | | | | | |
| | Tyr489 | Phe28 | A | ~0.60 | 3.75 | b-b | Gln498 | Tyr41 | A | ~0.60 | 4.52 | – |

VH, variable heavy chain; VL, variable light chains; H, hydrogen bond; A, aromatic bond; I, ionic (electrostatic) bond; bb, backbone atoms. | 3402–3414

Fig. 1B); second, two ACE2 in the membrane-bound dimeric form bind to two up RBD on two separate spike trimers (supplementary Fig. 1B). As a dimeric ACE2 is known to accommodate two RBD simultaneously [23], we suggest that soluble ACE2 may bind to the spike RBD more readily and stoichiometrically. This is because the binding of membrane-bound ACE2 is highly likely dependent on the spatial arrangement of the spike trimers on the virus surface, whereas soluble ACE2 can bind to any RBD available in up conformation. This is why soluble ACE2 or its derivative efficiently block the SARS-CoV-2 cell entry [35,36].

Next, we sought to analyze and compare the interface cRBD-ACE2 with that of sRBD-ACE2. Sequence and structure comparison analyses revealed that RBDR of SARS-CoV-2 was substantially vari-

cable compared to that of SARS-CoV, which harbored few conserved motifs. The average RMSD for the whole cRBD and sRBD was ~1.1 Å, whereas the average RMSD for RBDR deviated by ~2–3 Å owing to the glycine insertion and other mutations (Fig. 2A). Both cRBD and sRBD established similar contacts with ACE2 residues due to the conserved motifs, although they are vari-

able in RBDR region (Table 1). Interface analyses revealed that the electrostatic contact between Arg426 and Gln329 in sRBD-ACE2 was analogous to that of Lys417 and Asp30 contact in cRBD-ACE2 (Fig. 2B). However, this interaction was transient and break after the Asp30 of ACE2 established an intrachain contact with the nearby His34 (explained later). By performing protein patch analysis, we demonstrated that the standing cRBD exposes Lys417 that establishes strong electrostatic interaction with Asp30 of the ACE2, although this patch remains buried in the laying position of the cRBD (Fig. 2C). This finding indicates that the lysine 417 mutation in cRBD plays a crucial role in ACE2 recognition, which is otherwise substituted by hydrophobic valine in sRBD.

During the preparation of our manuscript, Yan R et al. demonstrated the interaction between SARS-CoV-2 RBD and ACE2 by per-

forming Cryo-EM analysis [37], which is in line with our study supporting the reliability of our computational model. Another study has delineated the interface of ACE2 with a chimeric RBD through X-ray crystallography [38]. Overall, this chimeric structure shares structural folds with our model and recently reported Cryo-

EM structure [37] and validates the cRBD-ACE2 interface residues in our model; nonetheless, the chimeric structure lacks the crucial mutation of Val-to-Lys417. This is because the core (scaffold) RBD in this structure was taken from SARS-CoV, which is considerably conserved between the two viruses with few mutations including Val-to-Lys417 (Supplementary Fig. 1C). As the high resolution crystal structure of the SARS-CoV-2 RBD with ACE2 has been reported during the course of this study, we validated our ACE2-RBD model by comparing the structure and interface with recently reported structure (6LZG) resolved through X-ray diffraction [39]. As expected, there was negligible differences in the overall RBDs (backbone atoms RMSD = 1.26 Å) structures and the ACE2-RBD interface residues were same in both complexes, including Lys417 (Fig. 2D). Therefore, we suggest that our computational model provides detailed understanding about the structural varia-

tion in cRBD and its interface with ACE2.

4.2. Mutation of Lys417 in cRBD may facilitate stronger interaction with ACE2

Differing from SARS-CoV and SARS-related CoVs, the S protein of SARS-CoV-2 has furin cleavage site at the S1/S2 boundary as well as exhibits similar or more binding affinity towards ACE2, which might be responsible for the efficient spread of SARS-CoV-2 [40]. In addition to these two points, we next sought to identify mutations in cRBD that play critical roles in the stronger binding-tendency towards ACE2 as compared to sRBD. As the information is limited about the static conformation of a protein complex considering the changes of the binding interface in physiological con-
dition, we simulated the complex structure of cRBD-ACE2 and compared this with the sRBD-ACE2 complex. The distances between interface residues were monitored as a function of time to trace the shifting, breaking, or formation of new bonds. Surface Plasmon resonance (SPR) and bio-layer interferometry (BLI) analy-

ses have shown that cRBD-ACE2 interaction is stronger than sRBD-ACE2 interaction [11,40,41]. Supporting this, we also observed that the total number of hydrogen bonds remained similar throughout the simulation time in both sRBD-ACE2 and cRBD-ACE2 models (Fig. 3A). This result may imply that the stronger binding affinity of cRBD toward ACE2 might be attributed to the stronger interaction of Lys417-Asp30 compared to Arg426-Glu329. Interestingly, when the minimum interaction distances with respect to the simu-
lation time was monitored, we observed that Lys417-Asp30 pair was more compact as compared to Arg426-Glu329 pair. Initially the residues in both pairs were ~1.4 Å apart; however, the Arg426-Glu329 pair separated by 2.6 Å, but the Lys417-Asp30 pair remained intact until the midpoint of the simulation. The bonds between both pairs broke at the same time point and remained separated by ~5 Å till the end of simulation (Fig. 3A). The relative strength and variation in the bond distance between these two pairs have been recently evaluated computationally, which were
substantially in line with our findings [42]. These strong yet transient electrostatic contacts can partly explain the phenomena of receptor recognition and S1 shedding upon enzymatic cleavage of the S1-S2 junction. S protein transiently utilizes the RBD of S1 subunit for receptor recognition and sheds them during cell internalization. Thus, faster SARS-CoV-2 transmission as compared to SARS-CoV is, at least in part, might be facilitated by the robust Lys417-Asp30 interaction.

In addition, we observed that Tyr449, Tyr489, Gln493, and Asn501 in cRBD established strong hydrogen bonds with the interface residues of ACE2 and remained intact throughout the simulation (Fig. 3A). These results indicate that these residues are equally responsible for the relatively stronger interaction of cRBD with ACE2. To demonstrate our results more clearly, we captured the motions of these interface residues in animations (Supplementary movies 1 and 2), and calculated binding free energies for each conformation. The binding affinity and interface analysis of the RBD-ACE2 complexes (Fig. 3A) show the RMSD of the backbone atoms of the RBD-ACE2 complexes with reference to the 0 ns conformation are shown. Total number of hydrogen bonds between RBD-ACE2 interfaces is shown in the second to top panel. The minimum contact distances, as a function of time, between the oppositely charged and other important residues at the RBD-ACE2 interface are shown in the third and bottom panels, respectively. The interfaces (PDB frames extracted at 20 ns of simulation) of ACE2 with cRBD (top) and sRBD (bottom) are depicted. B) The binding free energies of the sRBD-ACE2 and cRBD-ACE2 are estimated through MM-PBSA and tabulated here. All energies were computed via ensembles, which are averaged over a set of conformations sampled throughout the simulation trajectory at 0.01 ns time interval. Statistical significance of the differential energies was calculated through t-test. The p-values (<0.05) indicate the significance of calculated energies and their differences between sRBD-ACE2 and cRBD-ACE2 complexes. C) Structure based cRBD-ACE2 blocking peptides designing. HW1 and HW2 peptides are suggested in this study and SBP1 and LCB1 are reported RBD-binding peptides.
plex along the simulation time. The vdW, electrostatic, and SASA energies of the cRBD-ACE2 were relatively stronger than that of sRBD-ACE2. Besides, the polar solvation energy of cRBD-ACE2 was relatively higher than sRBD-ACE2, which may compensate the differences in the other energies of these complexes, resulting in overall slightly higher total binding free energies for the cRBD-ACE2 (see Fig. 3B). These data are in agreement with recently reported study who estimated the binding-free energies of sRBD and cRBD-ACE2 complexes through AMBER tool [43]. Collectively, our structural modeling analyses could demonstrate stronger

Fig. 4. Epitope mapping of the cRBD and complementarity-determining region (CDR) annotation of the mAbs. A) Anti-sRBD mAbs (single-chain variable fragments (scFv)) and their CDRs are shown. B) The variable light (VL) and variable heavy (VH) chains of the scFv regions of the reported anti-SARS-CoV-2 RBD mAbs are superimposed and the CDR regions are annotated according to Chothia and Lesk numbering scheme. C) The epitope prediction was validated through sRBD-F26G19 complex (PDB ID: 3BGF). The tabular interface is reported in the crystal structure while red boxes in the aligned sequences show the EpiPred predicted epitope. D) Conformational epitopes predicted with reference to six known anti-sRBD mAbs are highlighted and encircled. Residues participating in epitopes are indicated with arrows in the aligned cRBD and sRBD a.a. sequences (the arrow colors correspond to their respective epitopes). (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)
cRBD-ACE2 interaction compared to sRBD-ACE2 interaction. In addition, we demonstrated that Lys417 mutation may allow cRBD-ACE2 contact more readily, which may facilitate the rapid transmission of SARS-CoV-2 compared to SARS-CoV.

4.3. Druggability of the ACE2-RBD interface and decay-based peptide inhibitors

With the help of structural information provided by the binding analysis of SARS-CoV-2 spike protein and ACE2, many groups have been actively involved in designing peptide antidotes that are able to block the receptor binding of the virus. A peptide, S471–503, has been actively involved in designing peptide antidotes that are able to hinder ACE2-sRBD interaction and thus SARS-CoV entry into the cell, as confirmed in vitro [44]. By comparing the cRBD region corresponding to the S471–503 (ALNCYWPPLNDGYFYYTTGTIGYQ-PYRVVVLSFEL) peptide, we found that the N-terminal (bold letters) of S471–503 and the corresponding cRBD region were considerably different; however, the C-terminus portion (non-bold letters) was 100% identical to cRBD (see Fig. 2A, aligned sequences). Owing to the difference in the N-terminal half, S471–503 may not hinder SARS-CoV-2 cell entry as it exhibited in the SARS-CoV related study. We suggest that a peptide HW1, FNCYFPPLLQSYGFQPITNGVCYGQ-PYRVVVLSFEL may abolish the interaction between cRBD and ACE2. However, the c-terminal region, PYRVVVLSFEL, of HW1 is predominantly hydrophobic and belongs to the core region of the RBD (Fig. 3C). Overall, this peptide is physiologically unstable and may aggregate, losing its’ target specificity in solution.

Alternatively, we observed that the helical region of the ACE2 remains stable during simulation that establish strong electrostatic and hydrophobic contacts with the RBDR of both SARS-CoV and SARS-CoV-2 (Table 1, Supplementary movies 1 & 2), probably holding the potency to bind RBD. In fact, the interface between sRBD and ACE2 has been exploited in the past and ACE2-derived peptides have been used to block the SARS-CoV cell entry. A peptide, P6 (EEQAKFLKFHNHAEFLYQSS–GLKGDFR), constructed by the glycine linkage of two separate segments of ACE2 has been able to exhibit efficient antiviral activity (IC50 = 0.1 μM) [45]. Others and we found that cRBD and sRBD interact with the overall same helical peptide (α1 helix) of the ACE2 with some differing interface residues from the RBDs (see Table 1). Particularly, the two glutamic acids, EE, at the N-terminus of ACE2 did not make stable contacts with both s- and cRBDs and were exposed to the solvent. Lysine 353 (bold letter in P6) is crucial for RBDR binding, establishing multiple stable hydrogen bonds with both s- and cRBDs (Fig. 3A, Table 1). Thus, we suggest that a peptide HW2, QAKFLKFHNHAEFLYQSS-linker-GLKGDFR (Fig. 3C) may hold the optimum capacity to engage RBD and halt its binding to ACE2. More recently, a short peptide SBP1 derived from the α1 helix of ACE2 was shown to bind SARS-CoV-2 RBD that was expressed in insect; however, the results were not reproduced with human and other insect-derived RBDs [46]. This finding suggests that either the α1 helix of ACE2 is not sufficient to bind RBD or it loses helicity and RBDR-binding ability in solution state. A new peptide, P6 (EEQAKFLKFHNHAEFLYQSS–GLKGDFR), constructed by the glycine linkage of two separate segments of ACE2 has been able to exhibit efficient antiviral activity (IC50 = 0.1 μM) [45]. Others and we found that cRBD and sRBD interact with the overall same helical peptide (α1 helix) of the ACE2 with some differing interface residues from the RBDs (see Table 1). Particularly, the two glutamic acids, EE, at the N-terminus of ACE2 did not make stable contacts with both s- and cRBDs and were exposed to the solvent. Lysine 353 (bold letter in P6) is crucial for RBDR binding, establishing multiple stable hydrogen bonds with both s- and cRBDs (Fig. 3A, Table 1). Thus, we suggest that a peptide HW2, QAKFLKFHNHAEFLYQSS-linker-GLKGDFR (Fig. 3C) may hold the optimum capacity to engage RBD and halt its binding to ACE2. More recently, a short peptide SBP1 derived from the α1 helix of ACE2 was shown to bind SARS-CoV-2 RBD that was expressed in insect; however, the results were not reproduced with human and other insect-derived RBDs [46]. This finding suggests that either the α1 helix of ACE2 is not sufficient to bind RBD or it loses helicity and RBDR-binding ability in solution state. A new study has shown that stabilizing the helical fold of α1 helix (LCB1 & AHB1-2, Fig. 3C) retains its RBDR-binding capacity and effectively inhibit the SARS-CoV-2 cell entry [47]. Conclusively, we suggest that merely α1 helix derived peptides, in the absence of structural constraints or LGKGDFR motif, may poorly bind RBD and hinder its binding to ACE2.

4.4. Identification of epitopes on cRBD that bind to SARS-CoV-2 mAbs

SARS-CoV-2 and SARS-CoV belong to the genus betacoronavirus of the family coronaviridae and share considerable sequences in the RBD region of S protein [6,40], which allow researchers to delineate the cross-reactivity of anti-sRBD mAbs with cRBD. The binding parameters of all epitopes are listed in Table 2. We should note that merely aligning the α1 helix derived peptides, in the absence of structural constraints or LGKGDFR motif, may poorly bind RBD and hinder its binding to ACE2.
To investigate whether the previously known SARS-CoV mAbs can bind cRBD, we explored the binding potential of the cRBD with the SARS-CoV mAbs including 80R [23], m396 [25], F26G19 [24], s230 [26], CR3014, and CR3022 [48]. The structures of mAbs were obtained from PDB [for 80R [23], m396 [25], F26G19 [24], and s230 [26]] or modeled them [for CR3014 and CR3022 [30]] (for details see Methods). The variable heavy (VH) and variable light (VL) chains of scFv regions in these mAbs were aligned and their CDRs were annotated. These models revealed that the VL-CDR1 of CR3022 and s230 were relatively longer and more similar as compared to the VL-CDR1 of the other mAbs; in addition, the VH-CDR3 of s230 was more expanded than those of the other mAbs (Fig. 4A). Differences in the sequence and length of the CDRs indicate that these mAbs recognize distinct epitopes on the RBD and may not overlap thoroughly.

Over a short period, more than two dozen of SARS-CoV-2 S protein neutralizing mAbs have been identified and structurally elucidated. We compared the sequence and structures of CDR regions of these mAbs with that of CR3022 and F26G19. We found that the immunoglobulin G heavy-chain variable region 3 (in other words the VH-CDR3) and to some extent the VL-CDR1 in these mAbs are diversified and utilized to target the RBD of spike protein (Fig. 4B, Supplementary Tables 1 & 2).

Next we sought to predict conformational epitopes of cRBD using structural information of the mAbs. To ensure the authenticity of the epitope prediction, the co-crystal structure of sRBD-cRBD

Fig. 5. RBD-mAbs interface and their protein ligand interaction fingerprints (PLIF) analysis. A) The binding interface of SARS-CoV mAbs are displayed with respect to the predicted epitopes on cRBD (Green = Epitope 1, Cyan = Epitope 2, Red = Epitope 3; explained in Fig. 4). B) PLIF analysis of the CR3014 and CR3022 mAbs with cRBD. C) CR3014 cluster around epitope 2, which is highly variable between sRBD (left) and cRBD (right). CR3022 cluster near epitope 3, which is conserved between cRBD and sRBD. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)
Interface residues of cRBD with docked CR3014 and CR3022 mAbs.

Corresponded to Pro462 in sRBD (see Fig. 1A). A previous study that binds longitudinally to RBD (Table 3).

Anti-sRBD mAbs have CDR that are very specific and recognize by the structural differences of binding regions between them.

Can bind to ACE2 with high affinity. This could be partly explained interestingly, cRBD escapes from the anti-sRBD mAbs even though cRBD non-conserved or non-epitope regions of cRBD, and therefore suggest that these mAbs, 80R, s230, and F26G19 mAbs recognize epitope regions (Fig. 5A), suggesting that these mAbs may not bind and F26G19 were partly overlapped onto the residues at non-binding in the previous SPR and BLI experiments[11,41]. m396 (epitope 2) of cRBD; this could possibly explain their negative a part of the overlapping residues at the hypervariable RBDR region

In addition, we observed a mutation at Ala475 in cRBD, which corresponded to Pro462 in sRBD (see Fig. 1A). A previous study has shown that CR3014 mAb was not effective on the mutant Pro462Leu viruses, although it could prevent lung damage and SARS-CoV shedding in ferrets [48]. We found a glycine insertion mutation in the same loop (475-AGSTPCNVG-483), lengthening the loop RMSD to 2–3 Å (see Fig. 1C). This might be the reason why the previous BLI study could not demonstrate the binding of cRBD with CR3014 [41].

With this concern, we performed epitope mapping and protein ligand interaction fingerprints (PLIF) analyses to further evaluate the binding of CR3014. CR3014 was clustered around the same AGSTPCNVG-loop, a part of epitope 2, implying that the epitope 2-targeting sRBD mAbs may not be able to bind cRBD (Fig. 5B). Likewise, the sRBD mAbs recognizing the epitopes in variable RBDR regions may also not be able to bind cRBD. Thus, we suggest that mAbs or therapeutic peptides that bind to a conserved epitope on RBD may hinder the interaction of both SARS-CoV-2 and SARS-CoV-2 spike with ACE2.

CR3022 has been reported to completely neutralize the CR3014 escape SARS-CoV mutants (i.e., Pro462Leu) and synergize the neutralizing effect of CR3014 without competing with its epitopes [48]. A recent study has also demonstrated that CR3022 does not compete with ACE2-binding site of cRBD and exhibit binding to cRBD in the BLI analysis; in contrast, other mAbs, such as CR3014, m396, and MERS-CoV neutralizing mAb m336 were not able to bind cRBD (Fig. 5B). CR3022 was clustered over Arg24 and Arg26 and interacted with Glu19, which we designated as epitope 3 in the cRBD (Fig. 5B, C). Recently, Park et al. have performed computational analyses to demonstrate whether the previously known anti-MERS-CoV and anti-SARS-CoV mAbs can bind and neutralize cRBD [49]. However, this study did not consider the fact that the spike protein, particularly the RBD of SARS-CoV and MERS-CoV, have significant variations, which may preclude neutralization of cRBD by anti-SARS-CoV and anti-MERS-CoV mAbs [50]. In their docking analysis, the binding of CR3022 and s230 overlapped on the same interface of cRBD. In contrast, we and Tian et al. [41] have demonstrated that there is no overlap between CR3022 and the ACE2-binding region of cRBD (see Supplementary Fig. 2A). A crystal structure analysis also revealed that s230 bind to the ACE2-binding region of sRBD [26].

To validate these results, we superimposed the structures of s230-sRBD and ACE2-sRBD complexes, which revealed that ACE2 and s230 were overlapped with the same interface of sRBD.

F26G19 was used as control. We observed that epitope 1 completely overlapped with the experimental result, supporting the reliability of our analysis (Fig. 4C). Among the predicted cRBD epitopes, the residues in epitope 2 were mainly composed with highly variable regions between sRBD and cRBD (cyan color arrows in the aligned sequences). In contrast, the residues of the epitope 1 and 3 were significantly conserved between sRBD and cRBD (epitope 1, 93%; epitope 3, 100%, Fig. 4D). This result implies that the anti-SARS-CoV sRBD mAbs recognizing epitope 1 or epitope 3 could bind the cRBD and may hinder its receptor binding. However, the epitope 2 region was highly variable between cRBD and sRBD, therefore the anti-sRBD mAbs recognizing epitope 2 may not be able to bind or neutralize cRBD.

4.5. Highly conserved epitopes of cRBD are promising target for anti-SARS-CoV-2 agents

Recent studies comprising SPR and BLI analyses have demonstrated that the sRBD mAbs including m396, 80R, s230, and CR3014 are not able to recognize cRBD [11,41], although the reason for failure was not understood. To evaluate the reason, we placed or docked the scFv regions of these sRBD mAbs onto cRBD revealing their interface residues (Table 2), s230 and 80R interacted with a part of the overlapping residues at the hypervariable RBDR region (epitope 2) of cRBD; this could possibly explain their negative binding in the previous SPR and BLI experiments [11,41]. This result implies that the anti-SARS-CoV sRBD mAbs recognizing epitope 1 or epitope 3 could bind the cRBD and may hinder its receptor binding. However, the epitope 2 region was highly variable between cRBD and sRBD, therefore the anti-sRBD mAbs recognizing epitope 2 may not be able to bind or neutralize cRBD.

Table 3

| mAB Type | cRBD | scFv | Bond | ΔGcal/mol | Dist Å | Atom |
|----------|------|------|------|----------|------|------|
| CR3014   | VL   | Asp96| Gln27| H        | -2.00| 3.21 | b- |
|          |      | Asp96| Tyr92| H        | -2.40| 2.62 | –  |
|          |      | Asp97| Asp1 | H        | -13.963| 2.80 | -b |
|          |      | Ser128| Tyr32| H        | -2.00| 3.74 | –  |
|          |      | Lys131| Ser91| H        | -0.600| 3.28 | -b |
|          |      | Pro132| Thr94| H        | -1.60| 3.01 | -b |
|          |      | Glu134| Arg54| H        | -26.192| 2.78 | –  |
|          |      | Asp136| Asn58| H        | -0.50| 3.48 | b- |
|          |      | His188| Asp71| H        | 0.60| 3.27 | b- |
| CR3022   | VL   | Arg26| Gly61 | H        | -14.70| 3.63 | b- |
|          |      | Arg135| Tyr56| H        | -8.00| 2.75 | –  |
|          |      | Arg135| Ser58| H        | -3.40| 2.96 | –  |
|          |      | Phe133| Lys36| H        | -2.90| 2.87 | -b |
|          |      | Arg24| Tyr56 | H        | -2.60| 3.24 | -b |
|          |      | Glu9 | Gly30 | H        | -6.80| 2.80 | b- |
|          |      | Arg26| Ser102| H        | -3.80| 2.88 | –  |
|          |      | Arg26| Tyr14 | H        | -2.50| 2.66 | b- |
|          |      | Leu4 | Thr33 | H        | -2.30| 2.78 | -b |
|          |      | Asn29| Thr54 | H        | -2.90| 2.61 | -b |

VH, variable heavy chain; VL, variable light chains; H, hydrogen bond; A, aromatic bond; I, Ionic (electrostatic) bond; bb, backbone atoms. Residue numbers of the cRBD should be tallied with the top numbers in Fig. 2A.
We also found that CR3022 did not compete with the ACE2 and CR3014 interfaces of cRBD, which was consistent with the results of previous studies for SARS-CoV [48] and the recent SARS-CoV-2 [41] (Supplementary Fig. 2A). Notably, we found that CR3022 recognized a highly conserved region partly overlapping with epitope 3 (Fig. 5C).

During the preparation of this manuscript, a new study has shown the binding of CR3022 with cRBD through X-ray diffraction analysis, suggesting that the CR3022 Fab binds to a cryptic epitope (epitope 3 in our study) on the cRBD with substantially lower affinity as compared to sRBD [51]. Full-length CR3022 IgG also exhibited similar binding affinity towards both cRBD and sRBD, but could not neutralize SARS-CoV-2. The difference in the binding affinities of CR3022 Fab and IgG with cRBD and its inability to neutralize SARS-CoV-2 needs further investigation. We further speculated the binding mechanism of CR3022 IgG with SARS-CoV-2 trimeric spike. Yuan et al. suggested that CR3022 could bind to a trimeric spike in two or three RBD up states, but not single up conformation due to steric hindrance. We observed that the light chain constant region of CR3022 in their suggested model clash with CTD of adjacent or same spike protomer, regardless of the RBD up position (Fig. 6A). Conversely, we suggest that epitope 3 can...
accommodate the Fab and CR3022 IgG differently without any clash with the surrounding protomers. We further suggest that a single CR3022 can bind to two RBDSs of two different protomers in nearby trimers and an RBDD above can be assessed by the Fab regardless of the up or down conformation of the adjacent RBDD (Fig. 6B). Further investigations including site directed mutagenesis and binding analysis to the trimeric spike are required to confirm the authenticity of these models.

Since dozens of SARS-CoV-2 spike-neutralizing mAbs have been reported now, we wished to investigate their epitopes on the cRBD. Nineteen mAbs bound to the cRBD were obtained from PDB database and their CDRs were compared with those of sRBD binding mAbs. Most of the SARS-CoV-2 spike neutralizing antibodies, with some exceptions, were bound to the RBDR region of cRBD (Fig. 6C). Surprisingly, all three CDRs in VH region of these mAbs make contacts with the epitope 2. Whereas, the CDR3 in VL was docked into the epitope 1. This suggests that the RBDR region in cRBD is highly immunogenic due to the differences in the RBDR regions of SARS-CoV and SARS-CoV-2, producing highly specific and distinct antibodies in the body to neutralize their spike proteins. This could possibly explain the reason why anti-SARS-CoV mAbs are not effective against SARS-CoV-2. The cRBD binding mAbs can hinder its interaction with ACE2 even if they minimally bind or do not bind to the RBDR region. In fact, some mAbs can simultaneously bind to cRBD at distinct epitopes and abrogate its binding to ACE2 [52]. Hansen et al. generated nine cRBD-binding mAbs from genetically humanized mice and COVID-19 convalescent patients and identified their epitopes on cRBD through hydrogen/deuterium exchange mass spectrometry (HDX-MS). We found that the epitopes suggested by HDX-MS were non-linear and partly or fully overlapped with the three epitopes suggested in our study [52]. One mAb, REGN10987, which does not compete with ACE2, could simultaneously bind cRBD in the presence of REGN10933, which recognize epitope 2 (Fig. 6C). Similar antibody cocktail strategy has also been reported by other group, where two RBDD-neutralizing antibody bind two non-overlapping epitopes on cRBD [53]. This study suggests that BD-629 binds to the RBDR (epitope 2), and BD-368-2 binds to a non-overlapping epitope (epitope not suggested in our study) on the opposite side of epitope 2 (Fig. 6C). Overall, these findings suggest that the RBDD of SARS-CoV-2 is highly immunogenic and harbor multiple but overlapping epitopes.

In summary, we suggest that Lys417 mutation in cRBD acquires stronger electrostatic interaction with ACE2, which may facilitate faster receptor-recognition of cells. This interaction is further strengthened by electrostatic and hydrophobic contacts at the cRBD-ACE2 interface. In addition, we identified a conserved epitope on RBDD, which might be the target for developing new peptide therapeutics or mAb for neutralizing SARS-CoV-2. Undoubtedly, our findings provide new insights into the underlying mechanisms of the high infectivity of SARS-CoV-2, which may be helpful for developing new effective neutralizing agents against SARS-CoV-2.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Acknowledgements

This research was supported by grants from the National Research Foundation of Korea (NRF) funded by the Ministry of Science and ICT (MSIT) (NRF-2017M3C9A6047620, NRF-2019R1A5A2026045, NRF-2017M3A9B6061509), Republic of Korea. This study was also supported by KREONET (Korea Research Environment Open NETwork) which is man-aged and operated by KISTI (Korea Institute of Science and Technology Information). A preliminary version of this manuscript has been released as a pre-print at Research Square by Shah et al. [54].

Author contributions

M.S. S.C. and H.W. contributed toward the conceptualization of methodology; M.S. and B.A. performed the investigation and formal analysis; M.S. and H.W. wrote the original manuscript draft; H.W and S.C. supervised the study; H.W. was responsible for funding acquisition and project administration; all the authors contributed to editing and reviewing the manuscript.

Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.csbj.2020.11.002.

References

[1] WHO. Coronavirus disease 2019 (COVID-19) Situation Report – 46 6 March 2020, 2020.
[2] Belzuzi MA, Leyva-Grado VH. Overview of current therapeutics and novel candidates against influenza, respiratory syncytial virus, and middle east respiratory syndrome coronavirus infections. Front Microbiol. 2019;10:1327.
[3] Worobey M, Pekar J, Larsen BB, Nelson MI, Hill V, Joy JB, et al. The emergence of SARS-CoV-2 in Europe and North America. Science 2020.
[4] Wu F, Zhao S, Yu B, Chen YM, Wang W, Song ZG, et al. A new coronavirus associated with human respiratory disease in China. Nature 2020.
[5] Zhou P, Yang XL, Wang XG, Hu B, Zhang L, Zhang W, et al. A pneumonia outbreak associated with a new coronavirus of probable bat origin. Nature 2020.
[6] Zhu N, Zhang D, Wang W, Li X, Yang B, Song J, et al. A Novel Coronavirus from Patients with Pneumonia in China, 2019. N Engl J Med. 2020;382(8):727–33.
[7] Paraskevis D, Kostaki EG, Magiorkinis G, Panayiotakopoulos G, Sourvinos G, Tsidos S. Full-genome evolutionary analysis of the novel coronavirus (2019-nCoV) rejects the hypothesis of emergence as a result of a recent recombination event. Infect Genet Evol. 2020;79:104212.
[8] Kandimalla R, John A, Abburi C, Vallamkondu J, Reddy PH. Current Status of Multiple Drug Molecules, and Vaccines: An Update in SARS-CoV-2 Therapeutics. Mol Neurobiol. 2020;57(10):4106–16.
[9] Gallagher TM, Buchmeier MJ. Coronavirus spike proteins in viral entry and pathogenesis. Virology 2001;279(2):371–4.
[10] Letko M, Marzi A, Munster V. Functional assessment of cell entry and receptor usage for SARS-CoV-2 and other lineage B betacoronaviruses. Nat Microbiol. 2020.
[11] Wrapp D, Wang N, Corbett KS, Goldsmith JA, Hsieh CL, Abiona O, et al. Cryo-EM structure of the 2019-nCoV spike in the prefusion conformation. Science 2020.
[12] Yuan Y, Cao D, Zhang Y, Ma J, Qi J, Wang Q, et al. Cryo-EM structure of MERS-CoV and SARS-CoV spike glycoproteins reveal the dynamic receptor binding domains. Nat Commun. 2017;8:15092.
[13] Wrapp D, Wang N, Corbett KS, Goldsmith JA, Hsieh CL, Abiona O, et al. Cryo-EM structure of the 2019-nCoV spike in the prefusion conformation. Science 2020;367(6483):1269–3.
[14] Mansbach RA, Chakraborty S, Nguyen K, Montefiori D, Korber B, Gnanakaran S. Pandemic potential of SARS-CoV-2 Spike Variant D614G: Implications for a vaccine candidate. bioRxiv. 2020.
[15] Zhang L, Jackson CB, Hou H, Ojha A, Rangarajan ES, Izzard T, et al. The D614G mutation in the SARS-CoV-2 spike protein reduces S1 shedding and increases infectivity. bioRxiv. 2020.
[16] Shah M, Anwar MA, Park S, Sajri SS, Choi S. In silico mechanistic analysis of IRF3 inactivation and high-risk HPV E6 species-dependent drug response. Sci Rep. 2015;5:13446.
[17] Park S, Shin HJ, Shah M, Cho HY, Anwar MA, Achek A, et al.TLR4/MD2 specific peptides stabled in vivo LPS-induced immune exacerbation. Biomaterials 2017;126;49–60.
[18] Junaid M, Li CD, Shah M, Khan A, Guo H, Wei DQ. Extraction of molecular features for the drug discovery targeting protein-protein interaction of Helicobacter pylori CagA and tumor suppressor protein ASSP2. Proteins. 2019;87(10):837–49.
[19] Achek A, Shah M, Seo JY, Kwon HK, Gui X, Shin HJ, et al. Linear and rationally designed stapled peptides abrogate TLR4 pathway and relieve inflammatory symptoms in rheumatoid arthritis rat model. J Med Chem. 2019;62(11):5085–511.
[20] Narayanana A, Sellers BD, Jacobson MP. Energy-based analysis and prediction of the orientation between light- and heavy-chain antibody variable domains. J Mol Biol. 2009;388(5):941–53.
[21] Chothia C, Lesk AM. Canonical structures for the hypervariable regions of immunoglobulins. J Mol Biol. 1987;196(4):901–17.

[22] Murzin AG, Brenner SE, Hubbard T, Chothia C. SCOP: a structural classification of proteins database for the investigation of sequences and structures. J Mol Biol. 1995;247(3):536–40.

[23] Hwang WC, Lin Y, Santelli E, Su J, Jaroszewski L, Stee B, et al. Structural basis of neutralization by a human anti-severe acute respiratory syndrome spike protein antibody. Biochemistry. 2006;45(35):10260–6.

[24] Pak JE, Sharon C, Satkunarajah M, Auperin TC, Cameron CM, Kelvin DJ, et al. Structural insights into immune recognition of the severe acute respiratory syndrome coronavirus S protein receptor binding domain. J Mol Biol. 2009;388(4):815–23.

[25] Prabakaran P, Gan J, Feng Y, Zhu Z, Choudhry V, Xiao A, et al. Structure of severe acute respiratory syndrome coronavirus receptor-binding domain complexed with neutralizing antibody. J Biol Chem. 2006;281(23):15829–36.

[26] Walls AC, Xiong X, Park YJ, Tortorici MA, Snijder J, Quispe J, et al. Unexpected Receptor Functional Mimicry Elucidates Activation of Coronavirus Fusion. Cell 2019;178(5):1026–39 e15.

[27] Dunbar J, Krawczyk K, Leem J, Marks C, Nowak J, Regep C, et al. SAbPred: a structure-based antibody prediction server. Nucleic Acids Res. 2016;44(W1):W474–8.

[28] Krawczyk K, Liu X, Baker T, Shi J, Deane CM. Improving B-cell epitope prediction and its application to global antibody-antigen docking. Bioinformatics 2014;30(16):2288–94.

[29] Huang J, Rauscher S, Nawrocki G, Ran T, Feig M, de Groot BL, et al. CHARMM36m: an improved force field for folded and intrinsically disordered proteins. Nat Methods. 2017;14(1):71–3.

[30] Pronek S, Pašl S, Schulz R, Larsson P, Bjelkmar P, Apostolov R, et al. GROMACS 4.5: a high-throughput and highly parallel open source molecular simulation toolkit. Bioinformatics 2013;28(7):845–54.

[31] Wang H, Dommerf F, Holm C. Optimizing working parameters of the smooth mesh Ewald algorithm in terms of accuracy and efficiency. J Chem Phys 2010;133(3):034117.

[32] Hess B, Bekker H, Berendsen HH, Fraaije JG. LINCS: a linear constraint solver for molecular simulations. J Comput Chem 1997;18(12):1463–72.

[33] Kumari R, Kumar R. Open Source Drug Discovery C, Lynn A, g_mmpbsa–a GROMACS tool for high-throughput MM-PBSA calculations. J Chem Inf Model. 2014;54(7):1951–62.

[34] Anwar MA, Panneerselvam S, Shah M, Choi S. Insights into the species-specific TLRI signaling mechanism in response to Rhodobacter sphaeroides lipid A detection. Sci Rep. 2015;5:7657.

[35] Monteil V, Kwon H, Prado P, Hagelkruys A, Wimmer RA, Stahl M, et al. Inhibition of SARS-CoV-2 Infections in Engineered Human Tissues Using Human monoclonal antibody, 80R. J Biol Chem. 2006;281(45):34610–6.

[36] Andre Watson, Leonardo Ferreira, Peter Hwang, Jinbo Xu, Stroud R. Peptide Antidotes to SARS-CoV-2 (COVID-19). bioRxiv. 2020.

[37] Yan R, Zhang Y, Li Y, Xia L, Guo Y, Zhou Q. Structural basis for the recognition of the SARS-CoV-2 by full-length human ACE2. Science 2020.