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Binder design for targeting SARS-CoV-2 spike protein: An in silico perspective

Ali Etemadi a, Hamid Reza Moradi a, Farideh Mohammadian b, Mohammad Hossein Karimi-Jafari c, Babak Negahdari a, Yazdan Asgari a, Mohammadali Mazloomi a,∗

a Medical Biotechnology Department, School of Advanced Technologies in Medicine, Tehran University of Medical Sciences (TUMS), Tehran, Iran
b Department of Medical Biotechnology, Faculty of Advanced Medical Sciences, Tabriz University of Medical Sciences, Tabriz, Iran
c Department of Bioinformatics, Institute of Biochemistry and Biophysics, University of Tehran, Tehran, Iran

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ABSTRACT

Introduction: The COVID-19 pandemic is now affecting all people around the world and getting worse. New antiviral medications are desperately needed other than the few approved medications that have shown no promising efficacy so far.

Methods: Here we report three blocking binders for targeting SARS-CoV-2 spike protein to block the interaction between the spike protein on the SARS-CoV-2 and the angiotensin-converting enzyme 2 (ACE2) receptors, responsible for viral homing into the alveolar epithelium type II cells (AECII).

Results: The design process is based on the collected natural scaffolds and using Rosetta interface for designing the binders.

Conclusion: Based on the structural analysis, three binders were selected, and the results showed that they might be promising as new therapeutic targets for blocking COVID-19.

1. Introduction

The number of cases worldwide diagnosed with new Coronavirus (SARS-CoV2) is increasing exponentially (Thakur et al., 2020). There are a lot of interventional clinical trials on SARS-CoV-2 (Courtemanche et al., 2020). Drug repurposing is one of the main approaches in combating the coronavirus scourge (dos Santos, 2020). Antiviral, anti-inflammatory, and supportive agents, ACE2 blockers, and convalescent plasma therapy are being tested. Other new treatments such as photobiomodulation are ongoing but still, there is a few adequate and proven medicine to its treatment (Fekrazad, 2020).

Coronaviruses are zoonotic single-stranded RNA viruses and SARS-CoV-2 belongs to the second group of this family. Their genomic sequences are highly similar (especially between SARS-CoV-1 and SARS-CoV-2) therefore their protein structures have striking resemblance. Spike glycoprotein (S protein) of the virus is the key molecule employed for cell penetration by binding to the receptor-binding domain (RBD) of angiotensin-converting enzyme 2 (ACE2). It is the only homing receptor recognized for the virus till date. 83% of the alveolar epithelium type II cells (AECII) express ACE2, and making the respiratory system the main site of injury. The virus not only uses ACE2 as the receptor but also diminishes its protective role against lung damage (Wan et al., 2020). Also, the presence of ACE2 in other organs results in multi-organ dysfunction (M.-Y. Liu et al., 2020). Reports indicate that 76.5% of the amino acid sequence of the S protein is the same between SARS-CoV-1 and SARS-CoV-2, besides their matching 3D structure of RBD in the spike protein. SARS-CoV-2 shows a higher binding affinity than SARS-CoV-1 which can be an explanatory factor for the fast spread of the virus around the world (Xu et al., 2020).

Owing to the designability and selectivity, protein therapeutics show great promise for the future. As intermediate, small proteins (2–20 kDa) fill the gap between small molecules and antibodies due to their lower molecular weight, these small proteins are highly soluble and stable compared to antibodies (140 kDa). However, they have some pitfalls such as lower serum half-life. Considering the advancement in drug delivery methods, it is possible to overcome these problems in the nearest future (Vazquez-Lombardi et al., 2015). With the advent of high-throughput techniques such as in-silico design and experimental
evaluation, it is possible to rapidly produce de novo binder proteins in a massively parallel way. This previously was thought to be impossible (Chevalier et al., 2017). Some inventions are being done on the binding of macrocyclic peptides for inhibiting the entry of the virus (Istifli et al., 2020). On the other end of the drug spectrum (W. Liu et al., 2020), in-silico prediction has been utilized to assess binding of the phytochemicals (small molecules) such as RBD and cellular proteases binding monoterpenoids (TMPRSS2, Cathepsin B, and Cathepsin L) (Wang et al., 2019).

Considering the global demand, finding a target, independent of any post-translational modification that can be expressed in fast-growing hosts with high scalability would be very beneficial. To date, many biopharmaceuticals have been produced in Escherichia coli (E. coli) (Rosano and Ceccarelli, 2014). E. coli is being used as a host for 30% of the approved protein therapeutics due to its easy scale-up, well-established techniques for manipulation and optimization, various genome-scale metabolic models, etc. Therefore, E. coli can be a superior host for the production of antivirals against SARS-CoV2 (Yang et al., 2020).

In this study, we aimed at designing some novel binders to block the spike glycoprotein (S protein) of the SARS-CoV-2. The design process (Fig. 1) was based on collecting a bunch of scaffolds which are natural proteins for redesigning thereby making them modified binders for targeting the S protein.

2. Methods

2.1. Scaffolds selection and relaxation

In the first step, a list of natural proteins (Supplementary T1) was collected using the advanced search in RCSB (Berman et al., 2000) based on the following criteria: 1) resolution below 2 Å, 2) X-ray diffraction method, 3) must be expressed in E. coli, 4) sequence length below 100 residues, 5) monomeric proteins, and 6) no DNAs, RNAs, ligands, and mutations in proteins. These scaffolds are natural proteins and were used as initial proteins for making binders using docking, redesign, and other steps.

To remove all clashes in scaffolds and make them more favorable for Rosetta (Nivon et al., 2013), all selected scaffolds were relaxed using the
following script (/path/to/rosetta/main/source/bin/relax.hdf5. linearxgccrelease-database/path/to/rosetta/main/database-IPDBLIST-
ignore_unrecognized_res-relax:constrain_relax_to_start_coords-relax:
coord_constrain_sidechains-relax:ramp_constraints fake-ex1-ex2-use_in-
put_sc-no_his_his_pairE-no_optHfalse-flip_HNQ).

2.2. Initial Docking and Rosetta interface design

We used SARS-CoV-2 spike protein (PDB number 6M0J) (Shang et al., 2020) as a target for docking. Patchdock (Schneidman-Duhovny et al., 2005) was used for docking of all scaffolds against the target. 2000 models per scaffold were generated. Then the FastDesign was used for the interface design of all complexes of scaffolds and the target.

177 designs were selected based on the following scores: dDG < −38, Interface SASA > 2000, Shape complementary > 0.66, Buried Unsatisfied Polar < 2 and score per res < −2.2.

2.3. Molecular Dynamics (MD)

Molecular Dynamics (MD) simulation was conducted using the GROMACS package (Berendsen et al., 1995) with an AMBER99SB-ILDN force field and TIP3P water model.

Gromacs was used to analyze the overall RMSD and RMSF after 20 ns MD simulation, and to analyze the protein conformational changes and stability under physiological conditions. 3 trajectories were analyzed for 50 ns, 2 fs per step. Also, MD simulations were done on the original target (6M0J) and promising scaffolds.

A simulation box with at least 1 nm box boundary to protein distance was used for all proteins. Periodic boundary conditions (PBC) and particle mesh Ewald (PME) methods were used for simulations to consider the long-range electrostatic interactions (Darden et al., 1993).

Energy minimization (using the steepest descent method) was done after adding ions (Cl– or Na+) to neutralize the total charge, followed by equilibration for 200 ps under the NPT (constant number of particles, pressure, and temperature) ensemble. Position restraints and the temperature were coupled to 310 K using the velocity-rescaling thermostat.

For all the designed binders and also the original scaffold and the target, root-mean-square deviation (RMSD), residue contact maps, the end-state free energy calculations (Average DELTA G binding) were recorded. In VADAR 1.8, the Shrake method (Shrake and Rupley, 1977) was utilized for the values of Van der Waals radii and the definition of polar/nonpolar accessible surface area (ASA) and charged ASA. This method did not consider a uniform radius (1.8 Å) for all side-chain atoms and any hetero-atom attached to carbon was considered polar. Also, the Voronoi procedure was chosen for volume calculation (Richards, 1977). The protein structure coordination files were examined in the main chain, stereo/packing quality, and 3D profile quality indexes (Lüthy et al., 1992). Default thresholds were considered for every IEDB measuring tool (Morris et al., 1992).

Fragment quality application was used in Robetta to pick the best 3- and 9-mer fragments scores and used to predict 3D protein structures based on comparative modeling in Robetta (Kim et al., 2004). PyMol was used for structure visualization (DeLano, 2014).

3. Result

3.1. Docking after binder design

Patchdock and Cluspro were applied for global or blind docking without providing any binding sites. 167 binders from the outputs of RosettaDesign were selected and docked against the target. Fig. 2 shows the results for all docked designs against the target using ClusPro and Patchdock.

From the top 10 binders selected from both Patchdock and Cluspro, two (3HGL_1.78_2.1_78 or BIN78 and 3HGL_1.45_1.1_32 or BIN32) were found to be the same in both Patchdock and Cluspro outputs and were highly promising. These binders were bound to the same binding site of interest. Also, the binding sites were the same as the site targeted by RosettaDesign. Predicted PyMol RMSD based on the align method for Rosetta binder with BIN78 ClusPro and PatchDock models were 0.263 and 0.393, respectively (Fig. 3A). Both ClusPro and PatchDock predicted that the BIN78 bound to the desired pocket on the target with minimum RMSD.

A closer look at BIN78 (Fig. 3B), revealed that two main helices interacted with the main groove on the target. There were two π-π interactions between the BIN78 and the target: F28 with F202/236 and also H7 with Y251. Ddg scores for BIN32 and BIN78 were −43.998 and −41.691. Table 1 shows some predicted scores for all selected binders.

To check whether our designed binders had enough binding scores, we compared the weighted scores in ClusPro for designed binders BIN78, BIN12, BIN91, and SARS coronavirus spike receptor-binding domain complexed with its receptor (PDB number 6MOJ). The weighted scores for BIN78, BIN32, BIN91 were −747.3, −814, and −771.8, respectively. However, this score for the complex of S protein and ACE2 was −719.6.

3.2. Web logo and structure prediction

The bite signals or highly frequent residues in all sequences were 30

for each binder, 2000 outputs were checked. The top designed binders against the target. For this reason, binding sites were calculated using MMPBSA embedded in AmberTools.

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promising in predicting the 3-dimensional structure. The RMSD of binders. The results of the modeling showed that BIN91 was highly selected binders.

Rosetta was used for the comparative modeling of all selected binders. The results of the modeling showed that BIN91 was highly promising in predicting the 3-dimensional structure. The RMSD of BIN32, BIN91, and BIN78 with native scaffolds (3HGL) were 2.81, 1.97, and 2.88 Å, respectively (Fig. 5).

3.3. MD analysis

To investigate the conformational behavior of the binder’s models and native scaffold, MD simulation was used during the 50 ns simulation (Fig. 6). The minimum and maximum RMSD score for native scaffolds and the binders were close and the traces for the binders were followed by the traces for native scaffolds, although the RMSD for BIN91 was a little different from the others.

The radius of gyration (Rg) was analyzed to check protein structure compactness and stability of the binders, and also the target and native scaffolds. The Rg values for the binders in the binding form with the target were similar and larger than the target and the native scaffold alone. The contact maps for all three binders also showed the similarity between the binders. Likewise, minimum distances between the atoms in all binders and the native scaffold were analyzed and showed that the binders and native scaffolds were close to each other.

The results for distances between the specified residues (77–195, 19–244, 22–247, 32–149, 61–219, and 61–235) showed that the average distances between these residues were 2.2 nm, 0.9 nm, and 1.2 nm for BIN78, BIN32, and BIN91, respectively.

MMPBSA in AmberTools showed that the binding energy of BIN32, BIN78 and BIN91 in binding states with the SARS-coronavirus spike protein (PDB number 6M0J) were -22.43, -20.91 and -17.01, respectively (Average DELTA G binding reported in kcal/mol).

3.4. Structural analysis

Structural analysis was done on all 177 designed proteins. Since there was no perfection, each protein had some pros and cons. Therefore, by comparing their characteristics, with each other and with the main 3HGL protein, the 10 top proteins with enhanced or comparable parameters were selected (Table 2). Among them, Bin91 had the highest score in the docking experiments. Additionally, two modified proteins, Bin78 and Bin32, had a good performance in both Cluspro and Patchdock sections. For positive control, a Human Immunodeficiency Virus (HIV) entry inhibitor, Enfuvirtide (T20) was used. It has 36 amino acids with a helical secondary structure and a molecular weight of around 4.5 kDa. With the Trademark name of Fuzeon™, Enfuvirtide is considered the first small protein antiviral drug in the market (De Clercq and Li, 2016).

All binders comprised of 78 amino acids. The analysis based on ProtParam indicated that the designed proteins’ molecular weight was around 8500 Da. The results from 2Struc indicated 71.8% of the 3HGL is composed of alpha-helix, with 9% as helix-turn, and 19.2% as loops. All three top binders showed an increment in the proportion of helix-turn secondary structure. BIN91 changed to 14.1% and BIN32, 78 to 15.4%. To compensate for this, the alpha-helices decreased to 66.7% for BIN91 and 65.4% for BIN32, and BIN78. Enfuvirtide is composed of 90.9% alpha-helix and 3% helix-turn. Coils constitute the remainder of the structure.

One of the factors employed in classifying these candidates was the MolProbity score which is based on the clash score and geometrical parameters. It was normalized to mirror a crystallographic resolution. The main 3HGL protein had a score of 0.76, and a mild clash of <40 Å, with 6 bad bonds detected. Moreover, this model had a Ramachandran and Rotamer value in the allowed region. The chosen candidate proteins all had lower MolProbity scores which indicated a high quality from a structural standpoint of the model (Fig. 7A). The three top candidates had a MolProbity score of 0.5 which demonstrated high quality in their structure. The MolProbity results suggested that there were no preliminary geometrical problems with the binders, but this does not imply that the overall structure was flawless. To augment the scrutiny on assessing the quality of the structures, VADAR was implemented. It calculated more than 30 structural parameters and presented graphs and tables. In most of the structures, VADAR algorithms detected at least 1 residue that had 1 possible problem in the main chain and Stereo/Packing quality index. This might have resulted from a coordination defect or a systemic problem such as the wrong choice of the standard
None of the structures in the candidate pool had a 3D profile quality above 0. Inferentially, this implied there was no problem in the local environment, packing, and hydrophobic energy of the structures. Among the three final candidates, BIN78 had a better stereo/packing quality index and overall better performance in this section. Enfuvirtide PDB structure as positive control showed a low MolProbity score and many problems were detected with its 3D profile quality index. Ramachandran plots produced by VADAR in Fig. 8 indicate that, for BIN32 and BIN91, 92% of $\phi$ and $\psi$ angles are in the core region, and 7% are in the allowed region. Additionally, 93% of BIN78 $\phi$ and $\psi$ angles are in the core region, and 5% are located in the allowed region. For Enfuvirtide, 90% of residue angles are in the core region, and only 9% belong to the allowed region. No $\phi$ and $\psi$ angle outliers were detected in any structure.

Table 1
Rosetta predicted scores for Bin78, Bin32 and Bin91.

| Design name | $ddG$ | Interface SASA | Shape complementarity | Buried Unsatisfied Polar atoms | score_per_res | MMPBSA $^a$ | Van der Waals energy | Electrostatic energy |
|-------------|-------|----------------|-----------------------|--------------------------------|---------------|-----------------|--------------------|--------------------|
| Bin78       | -41.69 | 2004           | 0.666                 | 0                              | -2.381        | -20.91          | -209.93 kJ/mol     | -24.54 kJ/mol       |
| Bin32       | -43.998| 1966           | 0.663                 | 1                              | -2.219        | -22.43          | -154.74 kJ/mol     | -42.71 kJ/mol       |
| Bin91       | -37.206| 1701           | 0.673                 | 2                              | -2.292        | -17.01          | -164.15 kJ/mol     | -93.58 kJ/mol       |

$^a$ MMPBSA was predicted by AmberTools.

(Richards, 1974). None of the structures in the candidate pool had a 3D profile quality above 0. Inferentially, this implied there was no problem in the local environment, packing, and hydrophobic energy of the structures. Among the three final candidates, BIN78 had a better stereo/packing quality index and overall better performance in this section. Enfuvirtide PDB structure as positive control showed a low MolProbity score and many problems were detected with its 3D profile quality index.
A calculated parameter in ProtParam was the GRAVY score. Kyte & Doolittle calculated the GRAVY score for different soluble and membrane proteins. For this purpose, they summed up all the amino acid hydropathy scores in a sequence and divided the result by the number of residues. It can be inferred from their data that there is a high probability for proteins with a negative value of GRAVY score to be soluble, and proteins with a positive value to be membranous (Kyte and Doolittle, 1982). As shown in Fig. 7C, the GRAVY index value of a large number of proteins is at the boundary and above 0. The GRAVY score of top binders and controls are represented in Table 2. BIN91 with a GRAVY score of −0.168 is likely the most soluble protein among the top 3 candidates. The other parameter that was taken from ProtParam was the instability index. Grusprad et al. proposed that the stability and instability of a protein depend on the arrangement of amino acids in a specific order. It was shown that there are some dipeptides (the smallest unit of order) in the protein's sequence which defines the stability (Feng, 2020). The instability index of all designed proteins is plotted in Fig. 7D. Although it has been stated that proteins with an instability index below
are considered to be stable, the main 3HGL protein had an instability index of 59.03.

Another important aspect of choosing these candidates is the aggregation potential of the molecules. For this purpose, the normalized parameters such as global aggregation propensity (Na4vSS), Total Hotspot Area per residue (THASAr), and the number of Hotspot areas were calculated for each candidate.

Table 2

| Design name | MolProbity | VADAR | ProtParam | Aggrescan |
|-------------|------------|-------|-----------|-----------|
|              | Ramachandran favored (Goal > 98%) | Favored rotamer (Goal > 98%) | MolProbity score | Stereo/packing quality index (α) | Main chain possible problem (β) | Instability index (γ) | GRAVY index (δ) | Na4vSS | THASAr | NnHS |
| 3HGL        | 98.68%     | 98.41% | 0.76      | 2         | 2V,1A    | 59.03   | –0.321     | –16.2 | 0.01   | 1.282 |
| 3HGL091_37  | 98.68%     | 98.41% | 0.5       | 2         | 2V,1O    | 49.19   | –0.423     | –16.9 | 0.048  | 2.564 |
| 3HGL091_114 | 100%       | 100%   | 0.5       | 2         | 3V,1O    | 57.88   | –0.274     | –12.2 | 0.058  | 2.564 |
| 3HGL178_126 | 100%       | 100%   | 0.5       | 2         | 3V       | 32      | –0.141     | –11.4 | 0.071  | 2.564 |
| 3HGL178_164 | 100%       | 100%   | 0.5       | 2         | 4V,1O    | 36.82   | –0.205     | –13.9 | 0.062  | 2.564 |
| 3HGL178_2,61| 100%       | 100%   | 0.5       | 2         | 2V       | 29.58   | –0.146     | –10.1 | 0.087  | 2.564 |
| 3HGL178_2,88| 100%       | 100%   | 0.5       | 2         | 2V       | 35.68   | –0.163     | –11.1 | 0.073  | 2.564 |
| 3HGL178_59  | 100%       | 98.36% | 0.5       | 1         | None     | 48.93   | –0.153     | –12.3 | 0.091  | 2.564 |
| 3HGL192_59  | 100%       | 100%   | 0.5       | 1         | 3V       | 52.53   | –0.201     | –11.2 | 0.025  | 2.564 |
| 3HGL209_2,88| 100%       | 100%   | 0.5       | 2         | 3V,1O    | 47.85   | –0.431     | –18.5 | 0.035  | 2.564 |
| 3HGL209_4,11| 100%       | 100%   | 0.5       | 2         | 3V,1O,1A| 31.22   | –0.168     | –13.2 | 0.095  | 2.564 |
| 3HGL178_1,91| 100%       | 100%   | 0.5       | 3         | 4V,1O    | 30.36   | 0.021      | –4    | 0.116  | 3.846 |
| 3HGL045_1,39| 100%       | 100%   | 0.5       | 1         | 2V       | 58.28   | 0.119      | 1.7   | 0.13   | 2.564 |
| 3HGL078_2,78| 100%       | 100%   | 0.5       | 3         | 3V       | 62.65   | –0.875     | –20.5 | 0.174  | 5.556 |

(o) Indicates Possible Problems in the Protein by assigning a score to these three criteria: Torsion angle, Omega angle and Van der Waals radii.

(β) O - indicates possible problem with Omega angle, V - indicates possible problem with fractional volume, A - indicates possible problem with fractional ASA.

(γ) Instability index below 40 is considered as stable.

(δ) GRAVY index below 0 are more likely globular above 0 are more likely membranous.

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Another important aspect of choosing these candidates is the aggregation potential of the molecules. For this purpose, the normalized parameters such as global aggregation propensity (Na4vSS), Total Hotspot Area per residue (THASAr), and the number of Hotspot areas were calculated for each candidate.
(NnHS) were chosen. It permitted direct comparison between binders and the positive control. The main 3HGL had −16 Na4vSS, which was one of the lowest among the proteins indicating its low aggregating potential (Fig. 7B). Considering the data evaluated from the analysis of datasets by the original article, proteins with less than 5.18 for Na4vSS, 0.9 THSAr, and 3.35 NnHS parameters remained soluble when they are over-expressed in E. coli. Hence, the designed protein pool was filtered by these criteria (Kolaskar and Tongaonkar, 1990). The top ten selected protein designs all had a value below 10 for Na4vSS, except BIN32 and BIN78 for which a different selection procedure was performed (Table 2). BIN97 and BIN78 had one, while BIN32 had two additional numbers of hotspots in the same range of sequence compared to the main 3HGL protein (Fig. 8). Compared to Enfuvirtide, all three top designs had an advantage in THSAr and NnHS, but when Na4vSS was considered, Enfuvirtide had a low value (−20), indicating a low aggregation propensity of this small protein.

The final major selection parameter was the antigenicity of the proteins. The average, minimum, and maximum scores based on Kolaskar and Tongaonkar and Emini methods are presented in Table 3. Kolaskar and Tongaonkar with 75% accuracy, is a semi-empirical method for locating the antigenic determinants in the sequence. It uses the physicochemical properties of residues and their frequencies in
experimentally determined epitopes (Emini et al., 1985). As Fig. 8 shows, after the modifications, all three top binders retained the first position of antigenic determinant which starts from around the 7th to 15th amino acid position. They also had another position adjacent to the first one, which starts from around the 21st to the 30th residue. All three top binders also had a third spot which was above the threshold, but it was not predicted to be antigenic in BIN78 by the webservers’ algorithm. The calculation was based on the surface accessibility scale on a product instead of addition within the window. The Emini method used the formula $S_n = (n + 4 + i) (0.37)^i$ which gave the probability for a hexapeptide sequence to be on the surface. $S_n$ greater than 1.0 shows a higher probability for a sequence to reside on the surface of the protein (Shrake and Rupley, 1973). Results from Emini demonstrated that most parts of the antigenic sequences were not exposed at the surface. BIN91 had the least average (1.028) score between the three top binders, while the other two both had an average score of 1.046. The two ends of the Enfuvirtide sequence were the most antigenic parts but the Emini method did not consider them as exposed to the surface (Fig. 8).

4. Discussion

To date, no drugs have been approved for the treatment of the
CoVid-19. Remdesivir is the sole antiviral drug that is recommended for patients with mild disease. However, it is on limited supply (Dang and Dang, 2020), and the attainment of herd immunity by vaccination may take at least a year while the infection rate of SARS-CoV-2 keeps increasing rapidly. Despite this, the effective range of the vaccines remains inconclusive based on some reports (Feng, 2020).

AvrPtoB a *Pseudomonas syringae* pv tomato multi-domain effector protein, which plays a role in interrupting the immune response of tomato plants by inhibiting the programmed cell death and promotes disease (Feng and Cheng, 2020). To elucidate the crystal structure, Dong expressed a portion of this protein (AvrPtoB121–205 Or 3HGL) that was stable and sufficient for the interaction with its target in *E. coli* (Oh and Martin, 2011). In this study, 3HGL was used as a scaffold with the interface of its surfaces redesigned using RosettaDesign (Dong et al., 2009). This method was applied by different groups for making diagnostic and therapeutic proteins (Liu and Chen, 2016). Targeting the S protein with inhibitors is one of the promising ways of developing therapeutics (Willis et al., 2015). Not only do they inhibit the attachment of RBD to ACE2 and halt the entry of the virus, but they also preserve the ACE2. Consequently, they can function as regulators of the renin-angiotensin-aldosterone system, and protect the lung from injury by their protease function against angiotensin (Ang) 1. By attaching to the products of the ACE2, Ang 1–7, and Mas pathway they exert their anti-inflammatory, and anti-proliferative effects. Furthermore, they increase oxygenation and reduce the hypersensitivity of the airway and downregulate the apoptosis of alveolar epithelial cells. However, ACE2 is not the only enzyme that participates in the production of angiotensin. A week later, the level of this peptide gets back to normal, but in the short run, it could be quite damaging (Samavati and Uhal, 2020).

The results of ClusPro docking scored for selected binders and also spike protein against the receptor ACE2, showed that our binders are promising and might be useful for targeting the virus as it bound to the target stronger than the spike protein.

As can be seen from Fig. 5B, the substitution of caline (residue 76 in 3HGL) with aspartate (in BIN91) or threonine (in BIN32) increased the stability of the protein which may partly be due to the arginine in the 77th position of the sequence. In BIN78 substitution of valine with leucine (a methyl added as the bridge in side chain) did not change the instability index (58.28) considerably. Indicating the limited half-life of this binder. Guruprasad's method, however, does not consider the higher-order factors that may affect the results (Guruprasad et al., 1990). Buried charges may impact the folding of proteins and their stability because it is not so energetically favorable to bury a charge inside the protein structure.

There are some regions of the binders that are considered antigenic based on their sequence. Apart from the structural standpoint, other factors may contribute to the immunogenicity of the proteins like administration dose, excipient formulation, host proteins, patients’ immune/genetic background, and the aggregation of the proteins (van Beers and Bardor, 2012). The Aggroscan webserver predicted that BIN91 had the least aggregating propensity, but these predictions were based on the sequence and the folded protein may take a structure that increases or decreases the aggregation potential. Hiding some of the regions during the folding may vary the characteristics of the protein. That notwithstanding, it is possible to address such challenges with drug delivery methods (Bodier-Montagutelli et al., 2018).

5. Conclusion

Since SARS-CoV-2 inflicts a respiratory disease, the strategy of inhaled protein therapeutics could be of help. However, there are several questions to consider for this purpose such as formulation, stability, and the characterization of proteins in the droplets.

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Abbreviations

| Abbreviation | Description |
|--------------|-------------|
| ACE2         | Angiotensin-converting enzyme 2 |
| RBD          | Receptor Binding Domain |
| S protein    | spike glycoprotein |
| AECII        | alveolar epithelium type II cells |
| RMSD         | root-mean-square deviation |
| RMSF         | root-mean-square fluctuation |
| NPT          | constant number of particles, pressure, and temperature |
| PBC          | periodic boundary conditions |
| PME          | particle mesh Ewald |
| MD           | Molecular Dynamics |
| GRAVY        | grand average of hydropathy index |
| VADAR        | Volume Area Dihedral Angle Reporter |
| IEDB         | Immune-Epitope Database |
| ASA          | accessible surface area |
| DSSP         | Dictionary of Secondary Structure of Proteins |
| Na4vSS       | Normalized amino acid aggregation-propensity value, window average Sequence Sum |
| THSAr        | Total Hot Spot Area per residue |
| NnHS         | Normalized number of Hotspots |

CRediT authorship contribution statement

Ali Etemadi: Conceptualization, Data curation, Formal analysis. Hamid Reza Moradi: Resources, Software, Writing – review & editing. Farideh Mohammadian: Resources, Software. Mohammad Hossein Karimi-Jafari: Resources, Software. Babak Negahdari: Funding acquisition, Writing – original draft, Writing – review & editing. Yazdan Asgari: Writing – original draft, Writing – review & editing. Mohammadali Mazloomi: Supervision, Validation, Visualization, Writing – original draft, Writing – review & editing.

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