Probing the competitive inhibitor efficacy of frog-skin alpha helical AMPs identified against ACE2 binding to SARS-CoV-2 S1 spike protein as therapeutic scaffold to prevent COVID-19

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Abstract
In COVID-19 infection, the SARS-CoV-2 spike protein S1 interacts to the ACE2 receptor of human host, instigating the viral infection. To examine the competitive inhibitor efficacy of broad spectrum alpha helical AMPs extracted from frog skin, a comparative study of intermolecular interactions between viral S1 and AMPs was performed relative to S1-ACE2p interactions. The ACE2 binding region with S1 was extracted as ACE2p from the complex for ease of computation. Surprisingly, the Spike-Dermaseptin-S9 complex had more intermolecular interactions than the other peptide complexes and importantly, the S1-ACE2p complex. We observed how atomic displacements in docked complexes impacted structural integrity of a receptor-binding domain in S1 through conformational sampling analysis. Notably, this geometry-based sampling approach confers the robust interactions that endure in S1-Dermaseptin-S9 complex, demonstrating its conformational transition. Additionally, QM calculations revealed that the global hardness to resist chemical perturbations was found more in Dermaseptin-S9 compared to ACE2p. Moreover, the conventional MD through PCA and the torsional angle analyses indicated that Dermaseptin-S9 altered the conformations of S1 considerably. Our analysis further revealed the high structural stability of S1-Dermaseptin-S9 complex and particularly, the trajectory analysis of the secondary structural elements established the alpha helical conformations to be retained in S1-Dermaseptin-S9 complex, as substantiated by SMD results. In conclusion, the functional dynamics proved to be significant for viral Spike S1 and Dermaseptin-S9 peptide when compared to ACE2p complex. Hence, Dermaseptin-S9 peptide inhibitor could be a strong candidate for therapeutic scaffold to prevent infection of SARS-CoV-2.

Keywords COVID-19 · SARS-CoV-2 · S1 spike protein · Antimicrobial peptide · Brevinin · Dermaseptin · Magainin · Ocellatin · Conformational sampling · Semi-empirical calculation · Conventional molecular dynamics · Steered molecular dynamics

Introduction
Owing to current global pandemic COVID-19, the entire world is facing serious consequences, since the day of coronavirus outbreak in Chinese wet markets. The causative agent of COVID-19 is found to be novel β-coronavirus SARS-CoV-2, which is a pivotal member of the sarbecovirus family that would be species-specific to humans and other related mammals [1–4]. Spike protein (S) of SARS-CoV-2 is composed of S1 and S2 subunits. S1 retains a receptor-binding domain (RBD) made up of five β-stranded sheets (β1, β2, β3, β4, and β7) from 334th to 528th positions that are arranged antiparallel to a loop area between β1 and β2 strands [3, 5, 6]. In particular, this loop region in S1-RBD
consisting of 56 amino acids from 446th to 502nd residue positions forms the most unique region that specifically binds to the peptidase domain of the angiotensin-converting enzyme 2 (ACE2) receptor [7].

Distinctly, ACE2 is a zinc-containing metalloenzyme protein found on the surface of lungs, kidneys, and a number of human cells [8]. ACE2 is the functional host cell receptor for the virus to enter into the host cell and begin infection [9]. Henceforth, S1 (RDB) prevails as an appealing therapeutic target for SARS-CoV-2, for which the chemical inhibitors [10–13] and the peptide-based drugs [14–16] are developed in the line-up of therapies for COVID-19. However, there are series of complications in existing COVID-19 therapies [17]; henceforth, the antimicrobial peptides (AMPs) might be considered as an alternative therapy for SARS-CoV-2 infection.

AMPs are typically cationic and amphipathic molecules that have evolved over millions of years, providing almost all multicellular animals with the first line of defense [18]. In particular, anurans (frogs and toads) exposed to both aquatic and terrestrial environments have evolved to secrete AMPs in their skin to merely survive against many pathogens [19]. According to the Antimicrobial Peptide Database-3 (APD3), out of 3000 AMPs reported, over 189 AMPs have demonstrated antiviral activity. Specifically, the potential alpha-helical AMPs obtained from frog skin retain a good antiviral activity [20, 21]. In the protein–peptide interaction analyses of AMPs focusing on S1 Middle East Respiratory Syndrome (MERS) infection, the frog skin AMPs such as Magainin, Brevinin, and Dermaseptin have been documented to be the best docked peptides with high binding affinities [22]. Moreover, we previously reported the antimicrobial properties of broad spectrum ocellatin peptides [23], which prompts us to include it in this investigation and to assess its antiviral activity. Based on it, we intended to evaluate the efficacy of potential frog skin alpha-helical AMPs with its closest analogues, such as Brevinin-1BYa (1BYa), Brevinin-1BYc (1BYc), Dermaseptin-S4 (DS4), Dermaseptin-S9 (DS9), Magainin-1 (M1), Magainin-2 (M2), Ocellatin-1 (O1), and Ocellatin-F1 (OF1) that might inhibit spike protein S1 (RDB) of SARS-CoV-2 preserving the low hemolytic activity and high antimicrobial activity.

Brevinin (1BYa and 1BYc) peptides obtained from the skin secretion of Californian foothill-yellow legged frog Rana boylii [24] have been reported to have a remarkable antiviral potency against human immunodeficiency virus-1 (HIV-1), herpes simplex virus (HSV), and Ebola virus [25, 26]. Also, Dermaseptin (DS4 and DS9) peptides which were extracted from the South American tree frogs’ dermal secretions [27, 28] hold an antiviral action for HIV-1, HSV-1, HIV-2, HSV-2, human papilloma virus (HPV), rabies virus, and SARS-CoV-2 [16, 29–33]. Magainin peptides (M1 and M2) isolated from the skin of African clawed frogs, Xenopus laevis [34, 35] showed an antiviral activity against HSV-1, HSV-2, HIV-1, and SARS-CoV-2 [36–39]. Ocellatin O1 and Ocellatin OF1 peptides were isolated from the skin secretion of Brazilian pepper frogs such as Leptodactylus ocellatus and Leptodactylus labyrinthicus respectively [24] wherein, both the ocellatin peptides have shown to possess a wide variety of antibacterial, antiviral, and antifungal activities in experimental studies [23, 40–42]. Synergic antiviral effects between OF1 and alkaloid bufotenine have shown to limit BHK-21 cell lines advanced to rabies viral infection [43]. The present study identifies a potential therapeutic scaffold among these 8 alpha helical peptides that could serve as a potential competitive ACE2 inhibitor to impede S1 (RDB) from binding to ACE2 and thus regulates the SARS-CoV-2 entry.

Materials and methods

Data set

First, the sequences of frog skin alpha-helical AMPs with its closest analogues sequences, viz., 1BYa (P84111), 1BYc (P84113), DS4 (P80280), DS9 (Q1EN15), M1 (P11006), M2 (P11006), O1 (P83951), and OF1 (C0HKF0) constituting the range of 23–27 residue length were retrieved from the UniProt database [44]. Besides, the three-dimensional (3D) structures of 1BYa (6G4U), DS4 (2DD6), M2 (2MAG), and OF1 (5UA8) available in the Protein Data Bank (PDB) were retrieved [45]. With these available 3D structural templates, the tertiary structures of their respective analogues, viz., 1BYc, DS9, M1, and O1 were modeled, through the PEP-FOLD3 server. This program follows a de novo method for estimating peptide structural orientations based on their amino acid sequences, in which the conformations of consecutive residues are calculated with the help of a hidden Markov model (HMM). Using the YASARA package, all models were energy minimized [46]. A YASARA2 force field with a 10.5 cutoff was used to perform steepest descent energy minimization on peptides, and the peptides were optimized geometrically through explicit solvent. Using the PROCHECK tool, the stereo chemical quality of energy-minimized models was confirmed by a Ramachandran plot [47]. Furthermore, the receptor 3D structure of viral S1 spike glycoprotein (ID: 6M0J: E) with a resolution of 2.45 Å was obtained from the PDB.

Protein-peptide docking and interaction analysis

To determine binding interactions of SARS-CoV-2 spike glycoprotein (S1) with all eight peptides, viz., S1-1BYa, S1-1BYc, S1-DS4, S1-DS9, S1-M1, S1-M2, S1-O1, and S1-OF1, the site-specific docking was performed using
the HADDOCK (High Ambiguity Driven protein–protein DOCKing) program [48]. The HADDOCK program implements docking based on the data-driven approach that supports a wide range of experimental data, and the best binding solution of complexes was categorized based on desolvation, Vander Waals, restraint violation, and electrostatic energies with buried surface area [48]. Furthermore, total number of intermolecular non-covalent interactions among the docked complex, such as hydrogen bonds, hydrophobic contacts, cation-π interactions, and aromatic-aromatic, were predicted, using Protein Interaction Calculator (PIC) [49]. Furthermore, the hydrogen bond interactions were visualized via the PyMol visualization tool [50].

**Conformational sampling**

Subsequently, the conformational ensembles of four preferred protein-peptide complexes were generated, using the tCONCOORD program [51, 52]. Wherein, tCONCOORD depicts position constraints of complexes, using Gromacs index files [53–55]. Using the Vega ZZ environment, geometrical observable measurements such as root mean square deviation (RMSD) and polar surface area (PSA) of peptide trajectories were predicted and visualized with the Xmgrace tool [56, 57]. The conformational free energies of peptide conformers were estimated, using the distance-scaled finite ideal-gas reference (DFIRE) program, based on free-energy score and knowledge-based potential [58]. Furthermore, the secondary structural profiles of conformers were computed via a Define Secondary Protein Structure (DSSP) algorithm of GROMACS package [59] and their corresponding radar map representation was made through Microsoft Excel.

**Semi-empirical QM/MM calculations**

AMPAC-11 package was used to perform semi-empirical QM/MM calculations of peptide structures in which the structures were optimized using Austin Model 1 (AM1) parameters [60]. Wherein, the frontier molecular orbital energies, including the highest occupied molecular orbital (HOMO) and the lowest unoccupied molecular orbital (LUMO), were estimated for peptides in the HOMO-5 to LUMO+4 range applying Koopmans’ theorem [61].

**Conventional MD calculations**

Conventional molecular dynamics (cMD) based on Newtonian equations of motion was performed in GROMACS for S1-ACE2p and S1-DS9 complexes. In order to perform conventional MD simulations, the CHARMM36 force field was used [62]. The cubic box was generated, followed by the solvation of TIP3P water molecules. Furthermore, by embedding protein with appropriate ions, the system was neutralized. However, proceeding on to the actual dynamics, the system’s energy was reduced via steepest descent minimization, followed by appropriate NPT and NVT equilibrations. Finally, using an NVIDIA DGX 1 GPU accelerator, the production MD simulations of S1-ACE2p and S1-DS9 protein complexes were conducted for 100 ns. Furthermore, the torsional angle analyses and the principal component analysis (PCA) were performed for the generated S1-ACE2p and S1-DS9 complex trajectories. Furthermore, the contact atoms among S1-ACE2p and S1-DS9 docked complexes with respect to their key binding residues were computed through a Ligand Protein Complex-Contacts of Structural Units (LPC-CSU) program [63].

**Steered MD calculations**

The steered molecular dynamics (sMD) of the S1-ACE2p and S1-DS9 complexes were performed, using YASARA [64]. The dynamics were carried out in a solvent box containing water molecules using the AMBER03 force field at a constant temperature of 298 K. While executing the SMD, a particle-mesh Ewald long-range interaction and periodic boundary condition were used and the physiological pH of 7.0 was retained. The peptides from the S1 spike protein were pulled in the preferred direction using a steering potential, which kept the S1 center mass constant. An extrinsic steering force with a constant pulling acceleration of 1500 pm/picoseconds² was used to extract the peptides from S1, and forces were calculated for every 10 ps.

**Statistical analysis**

A nonparametric statistical method of Wilcoxon matched pair testing was done using StatPlus software version-7 (AnalystSoft, www.analystsoft.com/en) for the conformational sampling data of RMSD and PSA. In evaluating the statistical significance of complexes, this specific test is deemed similar to Student’s t-test for matched pairs and the probability value (P value) less than 0.05 was determined to be statistically significant [65].

**Results and discussion**

**Binding effect of protein-peptide interactions**

First, the structural orientations of antiviral peptides were modeled, using proficient structure prediction platforms. Wherein, the modeled structures exhibited the alpha helical content, confirmed by visualization of their 3D structures (Supplementary Fig. 1). The modeled 3D structures were also corroborated by Ramachandran plot analysis, where all the modeled peptides were found to be sterically
stable and structurally robust. Subsequently, the site-specific docking of S1 spike protein with alpha helical peptides was performed and the representations of those eight docked complexes such as S1-1BYa, S1-1BYc, S1-DS4, S1-DS9, S1-M1, S1-M2, S1-O1, and S1-OF1 were observed (Fig. 1). Consequently, the interaction between peptides and RBD of S1 were quantified and assessed with the help of molecular docking (Table 1).

Furthermore, the inhibitory efficacy of eight docked complexes was compared to that of the S1-ACE2 complex (ID: 6M0J) by quantifying intermolecular interactions among all docked complexes (Table 3). Regarding the inter-molecular interactions, about 43 interactions were quantified between S1 and ACE2, which serves as a reference to evaluate the corresponding interactions between antiviral peptides and S1 receptor. Similar levels of intermolecular interactions between the antiviral peptides and S1 receptor could insinuate that they could produce similar interactions that were observed between S1 and ACE2. Therefore, the peptides from each category of frog antiviral peptides which produce similar levels of interactions with S1, just as ACE2 does, were utilized for subsequent analysis. About 51 intermolecular interactions were found at the interface of S1 receptor and DS9 peptide while, about 41 and 32 inter-molecular interactions are represented as cartoon illustrated with various colors, viz., black, orange, brown, pink, red, blue, cyan, light green, and dark green, respectively. Similarly, S1 is represented as surface model.

**Table 1** Binding score estimation among S1-peptide docked complexes obtained through HADDOCK tool

| S. no | Peptide complexes | HADDOCK score |
|-------|-------------------|---------------|
| 1     | S1-1BYa           | −64.0         |
| 2     | S1-1BYc           | −42.0         |
| 3     | S1-DS4            | −35.5         |
| 4     | S1-DS9            | −82.9         |
| 5     | S1-M1             | −46.2         |
| 6     | S1-M2             | −69.4         |
| 7     | S1-O1             | −49.2         |
| 8     | S1-OF1            | −49.0         |

**Fig. 1** a) S1 docked peptide complexes, b) S1-ACE2p peptide complex derived from S1-ACE2 experimental complex structure (6M0J). The peptides-ACE2p, 1BYa, 1BYc, DS4, DS9, M1, M2, O1, and OF1 are represented as cartoon illustrated with various colors, viz., black, orange, brown, pink, red, blue, cyan, light green, and dark green, respectively. Similarly, S1 is represented as surface model.
interactions were found at the juncture of peptide-receptor for S1-M1 and S1-OF1 complexes respectively (Table 2); their corresponding interface representations in terms of hydrogen bonds and hydrophobic interactions are observed (Fig. 2).

To ease the intense burden levied upon computational resources, we have chosen ACE2p (S1 interacting peptide derived from ACE2) instead of the entire ACE2 protein. Besides, visualizing the peptide protein complex interaction revealed that ACE2p interaction with S1 is almost on par with ACE2 complete protein interactions with S1 (Table 3). Hence, it would be a prudent choice to choose ACE2p over ACE2. Hydrogen bonds are deemed to be one of the crucial factors in ascertaining intermolecular interactions between protein and peptides [66]. Notably, the interacting regions of S1 with the peptides are similar to that of ACE2 binding site. The binding site predominantly comprised of loop region with 56 amino acids spanning between β1 and β2 strands of RBD. Strikingly, all 4 peptides interact with loop region of S1-RBD, such as 1BYc peptide interacts with amino acids of S1 in the range of 448–498, while peptide DS9 forms hydrogen bond with S1 residues ranging 479–500. In the case of S1-M1 and S1-OF1 complexes, similar results were found (Fig. 2). Based on these findings, it was apparent that peptide DS9 manifests considerable interaction with S1 compared to other peptides (Fig. 2 and Table 2).

Loop region of S1 makes for a preferable drug target, since there binds the ACE2p. In recent past, several studies endorsed the targeting of loop region in S1 to competitively inhibit the binding of ACE2p, thereby proficiently arresting SARS-CoV-2 entry and averting infection [6, 7, 67, 68]. The fact that both the ACE2p and peptide DS9 associates with S1 in the similar binding vicinity coupled with an observation that the interaction of DS9 (Table 2 and Fig. 2) with S1 is a bit more adhesive than the interaction of ACE2p with S1, indicating a strong plausibility of DS9 to be ACE2p’s competitive inhibitor for binding with S1, magnificently. Although DS9 was found to be more effective in interacting with S1, the efficacy of other peptides were not negated, up till now. Accordingly, the top four peptides, viz., 1BYc, DS9, M1, and OF1, which exhibited considerable interactions with S1, were utilized for subsequent evaluation.

Exploring the effects of conformational ensembles

Most proteins perform many functions based on conformational changes in their structure without losing stability under varying circumstances [69]. The root mean square deviation (RMSD) and root mean square fluctuation (RMSF) were computed to illustrate the structural stability parameters of peptide complex ensembles (Table 3). In which, S1-DS9 showed the least RMSD and RMSF values, when compared to other peptide complexes. Moreover,
The RMSD measures the mean distance moved by atoms from their average positions. It is also a traditional metric for assessing conformational stability. Higher the RMSD, lower the conformational stability [70]. Our findings indicated that, among four peptide complexes, DS9 showed the least RMSD, even lower than S1-ACE2p complex which serves as positive control. This shows that the complex formed as a result of DS9’s interaction with S1 is notably stabilized. The RMSF is a distance measure between clusters of atoms in reference to a coordinate set with a well-defined average position [70]. Table 3 shows that S1-DS9 had the least fluctuation, with a mean value of 2.38 Å, indicating that it has a stronger structural stability than other peptides (Supplementary Fig. 2), further confirmed by its replicates (Supplementary Fig. 3).

Besides, its residual flexibility is the closest to positive control, compared to other peptide complexes. To further investigate the stability of S1-AMPs complexes, the conformational free energy of their trajectories was computed (Fig. 3). The data plotted in the form of kernel density estimation (KDE), illustrated that among four peptide complexes, S1-DS9 showed a least energy. The lower the...
free energy, the higher the thermodynamic stability [71]. Therefore, S1-DS9 is deemed to be the most stable complex. Also, it was found that S1-DS9 complex energy was quite closer to the positive control, S1-ACE2p complex energy compared to others (Table 3). Furthermore, the secondary structural profile predicted for peptide conformations from trajectory analysis showed a considerable variation in dynamic pattern of alpha helix represented as radar chart (Fig. 4). When interacting with a lipid membrane, an alpha helical peptide regulates its helicity by the distribution of intramolecular hydrogen bonds; conversely, when interacting with receptor residues, it tends to non-covalently interact with the receptor residues, resulting in perpetuation of peptide helicity.

In comparison to DS9, other three docked complexes improperly interacted with S1 receptor during protein-peptide simulations, resulting in turn or random coil. However, a notable conservation of secondary structural feature was observed in DS9, that its overall alpha helical content of a peptide retained throughout the simulation, which influenced its interaction with S1 receptor. This further corroborated aforementioned findings which endorses DS9 to be a suitable candidate against S1.

**Rationalizing the impact of DS9’s competitive binding efficacy**

Based upon the aforementioned results from docking and dynamics, it was apparent that DS9 could be a viable candidate to effectively bind with S1 and potentially produce the intended effect in mitigating the virulence. We further desired to substantiate the notion that DS9 binding is on par with ACE2p binding to S1 via quantum and molecular mechanical computations. Semi-empirical QM/MM global reactivity descriptors such as HOMO (highest occupied molecular orbital) denote the ability to donate electrons, whereas LUMO (lowest unoccupied molecular orbital) denotes the ability to accept electrons [72], insinuating the chemical reactivity of the compound upon interacting with S1. The lower the energy gap, the higher the reactivity [73, 74]. Findings indicate that reactivity of DS9 with S1 is almost on par with ACE2p’s reactivity with the same. Moreover, conferring

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**Fig. 4** Radar chart of secondary structural conformers of S1 spike protein and peptide complexes a) S1-1BYc, b) S1-DS9, c) S1-M1, and d) S1-OF1 where the radial values correspond to the number of residues.
to Koopmans’ theorem, HOMO–LUMO electron density energy gap (Fig. 5) determines the global hardness of DS9 and ACE2p molecules, which is known to be an estimation of their resistance to charge transfer during small perturbations [75]. Accordingly, DS9 has a greater value (0.532 eV) than ACE2p (0.454 eV), indicating that AMP has better resistive stability in chemical reactions.

Numerous proteins, in response to conformational changes in their 3D structure, dictate distinct cellular activity in varying cellular environments [69]. However, these conformational changes can be effectively investigated via conventional molecular dynamics (cMD) simulations with atomic precision [76]. Firstly, the torsional angle transition was investigated using cMd trajectory analysis, to substantiate the relevance of the side chains of key residues Y449, N487, and T500 highlighted in docking analyses of protein-peptide complexes (Fig. 6), which is essential for regulating the interaction between S1 and ACE2. Secondly, the comparative analysis was performed by overlapping the backbone angles ($\phi$; $\psi$) of all three interacting residues Y449, N487, and T500 in S1 [5], since torsional angle among amino acids is essential for maintaining the structural integrity of

![Fig. 5 HOMOL-LUMO energy gap computed via a) ACE2p and b) DS9 peptides after docking with S1](Image)
Comparing the change in torsional backbone angle of S1 upon interacting with ACE2p and DS9 peptide could effectively validate the ability of DS9 to alter the structural integrity of S1’s pathogenic conformation that can be observed from S1-ACE2p’s interaction. Accordingly, the backbone angles $\phi$ and $\psi$ of Y449 were located in range of $(-50^\circ, -100^\circ; 0^\circ, 50^\circ)$. Similarly, the backbone angles $\phi$ and $\psi$ of N487 and T500 were distributed in range of $(50^\circ, 100^\circ; -20^\circ, 40^\circ)$ and $(120^\circ, -50^\circ; -50^\circ, 50^\circ)$ respectively (Fig. 6). The preceding result implicated that the angles $\phi$ and $\psi$ of these three residues near the S1 binding site contribute significantly to the conformational modulation of DS9, which makes it a promising AMP analeptic. Table 4 reveals the range of nearest distance between atoms among S1-ACE2p and S1-DS9 docked complexes with respect to key binding residues, viz., Y449, Y487, and T500 computed.
through LPC-CSU program [63, 78]. The nearest distance between atoms of two residues in S1-ACE2p is wider (range of 2.7 to 5.4 Å) when compared to a narrow distance of S1-DS9 comparatively (range of 2.8 to 3.5 Å). This strongly suggested that DS9 has more favorable contact than ACE2p [79, 80]. Besides, the hydrophobic interaction and aromatic-aromatic interactions around key residues of DS9 and ACE2p were comparatively studied. Moreover, the destabilizing interactions (hydrophobic-hydrophilic contact) are lower in DS9 when compared to ACE2p around key residues [81]. This suggested that DS9 could possess more stable contacts towards spike protein. Therefore from Table 4, it is evident that DS9 could have more favorable binding towards spike protein when compared to ACE2p.

Furthermore, the Gibbs free energy landscape (ΔG) for ACE2p and DS9 was generated using the first two major principal components (PC1, PC2) as reaction coordinates from molecular dynamics trajectories. The 2D surface projections of the global energy minimum elucidated the cluster of stable states that occur in both S1-ACE2p and S1-DS9 complexes (Fig. 7). The Eigen vectors of collective motions in DS9 were clearly confined to a basin lesser than that of ACE2p. As a result, these differences in conformational structures between S1-DS9 and S1-ACE2p indicated that DS9 might regulate the conformational orientations of S1 in comparison with ACE2p’s influence over the same. Finally, steered molecular dynamics (sMD), a converse analysis for docking, was carried out on S1-ACE2p and DS9 complexes wherein the ligand is pulled away from the protein to evaluate its protein ligand interaction (Fig. 8). Specifically, the duration for dissociation is proportional to the interaction

| S. no | S1-peptide residues | Peptide residues | Specific contacts |
|-------|---------------------|-----------------|-------------------|
|       | Y449               | D38             | NDA (Å)          |
| S1-ACE2p |                      |                 | HB               |
|        | Y449               | Q42             | +                 |
|        | Y487               | Q24             | +                 |
|        | Y449               | F28             | +                 |
|        | T500               | Y41             | +                 |
|        | T500               | L45             | +                 |
| S1-DS9 | Y449               | W7              | -                 |
|        | Y449               | V10             | -                 |
|        | Y449               | L11             | +                 |
|        | Y449               | I14             | +                 |
|        | T500               | R3              | -                 |

NDA, nearest distance between atoms of two residues
CSA, contact surface area between two residues
HB, hydrogen bond
AAC, aromatic-aromatic contact
HHC, hydrophobic-hydrophobic contact
DC, destabilizing contact ± indicates presence/absence of specific contacts
between protein and ligand; the longer the time taken to dissociate the ligand/peptide from the protein, the higher is the binding affinity between them [64]. Accordingly, it took 35.57 ps to completely dissociate ACE2p from S1, while 37.90 ps has elapsed, to completely separate DS9 from S1. In addition, free energy versus displacement result based upon Jarynski’s theory [82, 83] has been illustrated graphically (Supplementary Fig. 4). This correlates well with docking studies and SMD analysis. Besides, it reveals more favorable binding of DS9’s interaction with S1 when compared to ACE2p.

Conclusions

Spike viral protein S1-DS9 peptide complex demonstrated the strongest intermolecular interactions and higher thermodynamic stability, thereby endorsing the potential competitive inhibitor role of DS9 in binding to S1. The complications with current COVID-19 therapy have urged the scientific community to find new treatments to tackle SARS-CoV-2. As an outcome of this study, we anticipate that identifying DS9 as a potential therapeutic peptide scaffold will lead to improvements in the design of peptidomimetics for the treatment of COVID-19 viral infection. As a future perspective, the incredible antiviral synergistic impact between DS9 peptide and drugs in preclinical trials might provide a creative lead to the development of promising antiviral drugs.

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Author contribution P. Chandra Sekar optimized the entire study and wrote the manuscript. E. Srinivasan standardized the computational method used in this study. G. Chandrasekhar made significant contributions to MD simulations and semi-empirical QM/MM calculations. D. Meshach Paul discussed the results and helped to draft the manuscript. G. Sanjay performed the conformational sampling and data analysis. S. Surya and NS. Arunraj Kumar collected the initial data set and performed docking and interaction analyses. R. Rajasekaran designed and meticulously evaluated the entire study and contributed significant intellectual material. The final manuscript was read and approved by all authors.

Data availability Not applicable.

Code availability Not applicable.

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![Fig. 8](https://example.com/fig8.png) Steered molecular dynamics data illustrating the preferred pull of peptides from S1 spike protein while upholding the S1 center mass constant.
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