Identifying the natural polyphenol catechin as a multi-targeted agent against SARS-CoV-2 for the plausible therapy of COVID-19: an integrated computational approach

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Abstract

The global pandemic crisis, coronavirus disease 2019 (COVID-19), caused by the severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) has claimed the lives of millions of people across the world. Development and testing of anti-SARS-CoV-2 drugs or vaccines have not turned to be realistic within the timeframe needed to combat this pandemic. Here, we report a comprehensive computational approach to identify the multi-targeted drug molecules against the
SARS-CoV-2 proteins, which are crucially involved in the viral–host interaction, replication of the virus inside the host, disease progression and transmission of coronavirus infection. Virtual screening of 75 FDA-approved potential antiviral drugs against the target proteins, spike (S) glycoprotein, human angiotensin-converting enzyme 2 (hACE2), 3-chymotrypsin-like cysteine protease (3CL\textsubscript{pro}), cathepsin L (CTSL), nucleocapsid protein, RNA-dependent RNA polymerase (RdRp) and non-structural protein 6 (NSP6), resulted in the selection of seven drugs which preferentially bind to the target proteins. Further, the molecular interactions determined by molecular dynamics simulation revealed that among the 75 drug molecules, catechin can effectively bind to 3CL\textsubscript{pro}, CTSL, RBD of S protein, NSP6 and nucleocapsid protein. It is more conveniently involved in key molecular interactions, showing binding free energy ($\Delta G$\textsubscript{bind}) in the range of $-5.09$ kcal/mol (CTSL) to $-26.09$ kcal/mol (NSP6). At the binding pocket, catechin is majorly stabilized by the hydrophobic interactions, displaying $\Delta E_{vdW}$ values: $-7.59$ to $-37.39$ kcal/mol. Thus, the structural insights of better binding affinity and favorable molecular interaction of catechin toward multiple target proteins signify that catechin can be potentially explored as a multi-targeted agent against COVID-19.

**Graphical Abstract**

**Key words:** SARS-CoV-2; COVID-19; multi-targeted drug; catechin; free energy landscape
Introduction

There are different members of Coronaviridae family of virus, which often cause mild, moderate to severe respiratory symptoms in humans [1]. Recently, the novel coronavirus [2019-nCoV or severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2)] appears to be the highly infectious and contagious virus of this family [1, 2]. Although SARS-CoV-2 shares a high level of genetic similarity with SARS-CoV, the infection rate of SARS-CoV-2 is much higher than suspected [3]. The molecular and structural organization of the virus includes an envelope, non-segmented, positive-sense RNA which codes for several structural proteins such as spike (S) protein, envelope (E) protein, membrane (M) protein and the nucleocapsid (N) proteins and also 16 putative non-structural proteins [NSPs, encoded by replicase complex (orf1ab)] [1, 4–6]. During the last couple of months, dozens of coronavirus vaccines have already been tested, and approximately more than hundreds are either under development or in the preclinical investigation [5, 7]. However, the success of these efforts remains elusive. Therefore, the need of the hour is to identify novel and effective measures to control the pandemic [3, 7, 8].

The major setback in identifying drugs/vaccines against COVID-19 is the lack of knowledge about the mechanism of action of the virus, molecular targets and network of associated molecular interactions. Recently, several targets have already been discovered that interact with SARS-CoV-2, such as human angiotensin-converting enzyme 2 (hACE2), transmembrane protease serine 2 (TMPRSS2), phosphatidylinositol 3-phosphate 5-kinase (PIKfyve), two pore channel subtype 2 (TPC2) and cathepsin L (CTSL) [6, 9, 10]. At the same time, various research groups have identified the effective inhibitors against some of these targets, such as main protease Mpro (3CLpro) [11, 12], envelope (E) protein [13], RNA-dependent RNA polymerase (RdRp) [14] and spike (S) protein [15, 16].

Since the genes of RNA viruses (including SARS-CoV-2) are genetically variable [17–19], they can quickly accumulate genomic mutations through an error-prone viral reverse transcriptase, which advances their adaptation inside the human host. This further adds to the difficulty in designing active antiviral therapeutics against RNA viruses [9]. Moreover, most of the antiviral drugs today are single target drugs designed against specific viral enzymes, which are essential for viral interaction, replication or invasion. Therefore, the high rate of mutations in these single viral drug targets has been main cause for reduced susceptibility of currently available antiviral drugs [20].

Nevertheless, finding the compounds having efficacy for multiple molecular targets, remained a preferable approach over combination therapy to avoid unwanted drug interactions [9, 23]. Additionally, the drugs designed for multiple protein targets are extensively preferable for the treatment of infectious, inherited and complex diseases due to low treatment cost, less drug dosage and minimal side effect and drug–drug interactions. Therefore, with the immediate requirement of multi-targeted strategies against the novel coronavirus SARS-CoV-2, with either biologically active drug-like molecules or approved drugs are in pressing priority [6, 24, 25]. Recent advancements in the computational techniques have proven their efficacy for identifying the potential drug candidates [7, 26–28]. Considering the improvement, reliability and accuracy of computational methods, it has become a suitable choice to design structure-based drugs [21, 29–31]. Keeping these facts in mind, we adopted a multi-target drug discovery approach to hit various druggable targets of SARS-CoV-2, which may appear highly beneficial to strike this highly mutated virus. The repurposing of FDA-approved drug molecules is safe and significantly free from off-targets binding which warrants severe toxicity [32, 33].

Thereby, we screened 75 FDA-approved antiviral drugs against known targets of SARS-CoV-2 [34, 35]. We have taken both human proteins as well as viral targets as a strategy. The targets were chosen as the recently published structures of SARS-CoV-2 proteins complexed with drug molecules. Following seven targets were identified: (1) the hACE2 interacting with the (2) transmembrane viral spike (S) glycoprotein at receptor-binding domain (RBD), which forms homotrimers protruding from the viral surface [6, 7, 15, 16]; (3) the highly immunogenic, antigenic and abundantly expressed viral nucleocapsid (N) protein, which plays essential roles in viral genome packaging by formation of helical ribonucleoproteins [36]; (4) main protease Mpro (3CLpro), an essential viral enzyme for processing the polyprotein complexes that are translated from the viral RNA [37]; (5) the human endosomal cysteine protease CTSL required for viral entry [7, 38]; (6) non-structural viral protein NSP6 which dwells in the endoplasmic reticulum (ER) and has role in the generation of autophagosomes [39] and (7) the NSP12, along with the two other cofactors NSP7-NSP8 as a complex, aiding increased RdRp template binding and processivity. Firstly, the virtual screening of 75 FDA-approved antiviral drugs was performed against these targets to select a highly potent multi-targeted agent. The best binding poses of antiviral drugs with target proteins display a wide range of binding affinities. Among them, (+)-catechin (catechin) emerged as a multi-targeted agent that can effectively bind with five target proteins: RBD, CTSL, nucleocapsid protein, 3CLpro and NSP6.

Catechin (flavan-3-ol) is a natural phenol and a major chemical component of sinecatechin, a first FDA-approved herbal drug for the treatment of external genital warts caused by human papilloma virus (HPV) infections. Topical ointment Veregen is marketed in Canada and it is a purified form of catechins, extracted from the leaves of Chinese green tea, which comprises 80% catechins. It is well recognized for the antiviral activity, anti-bacterial activity, anti-inflammatory and for the immunosuppressive actions [34]. Further, we have acquired several data contributing to the stable structural dynamics of the protein–ligand complex, including free energy landscape (FEL), which validates the binding efficacy of drug molecules using molecular dynamics (MD) simulation and molecular mechanics–Poisson–Boltzmann Surface Area (MM-PBSA). Thus, identifying the catechin as a novel multi-targeted agent may provide the structural basis for the designing strategy of potential drug molecules, targeting SARS-CoV-2 in the therapy of COVID-19.

Materials and methods

Protein structure retrieval

The three-dimensional coordinates of protein structures were taken from the Protein Data Bank (www.rcsb.org) [40]. The co-crystallized X-ray structure of SARS-CoV-2 spike receptor with ACE2 (PDB ID: 6M0J) [41], 3-chymotrypsin-like cysteine protease (3CLpro) (PDB ID: 6M2N), CTSL (PDB ID: 6F06) [41], crystal structure of nucleocapsid protein (PDB ID: 6M3M) [36], RdRp (PDB ID: 6M71) [42], non-structural protein 6 (NSP6) [39, 43] and cryo-electron microscopy structure of RdRp enzyme with
remdesivir and NSP12-NSP7-NSP8 complex (PDB ID: 7BV2) [14] were taken as the targets for the virtual screening of selected FDA-approved antiviral agents as shown in the supporting information, Supplementary Figure S1 available online at https://academic.oup.com/bib.

Virtual screening and molecular Docking
The virtual screening of FDA-approved antiviral compounds against the selected SARS-CoV-2 proteins was performed using Glide, Schrodinger, LLC [44–46]. Glide involves three-step filtering methods, standard precision, extra precision and the selection of best docked compounds by integrating coulombic and van der Waals (vdW) interaction energies and Glide scoring function. CORINA v2.64 software package [47] was utilized to add the missing hydrogen atoms and optimize the sdf format structures of the ligands. The lowest energy three-dimensional structures of ligands were generated using Ligprep [48]. The ionization/tautomeric states of the selected compounds were taken care of by Epik and a maximum of up to 32 conformations was generated per ligand, using the Schrodinger protocol [49–51]. The molecular interactions of docked complexes were analyzed using PyMol [52] and LigPlot [53].

MD simulation
MD simulations were carried out on the coordinates of protein–ligand complexes using GROMACS-2018.1 [54], with the protein interactions approximated using CHARMM36 force field [55]. The ligand parameters were generated utilizing CGenFF server [56]. Each protein–ligand complex was placed in the center of a cubic simulation box with 10 Å distance to the edges and solvated with TIP3P water molecules. The counterions (Na+ or Cl−), 0.15 M, were added to neutralize the system. The periodic boundary condition was defined in the x, y and z directions [57], and the electrostatic interactions were evaluated using particle-Ewald summation [57], and a cut-off of 10 Å was used for calculation of vdw interactions. The resulting systems were energy-minimized by steepest descent and conjugate gradient algorithms. Energy minimization was performed for 50 000 steps. Equilibration was first performed for 500 ps in an NVT ensemble and for the subsequent 500 ps in an NPT ensemble. Temperature and pressure were set at T = 300 K and 1 bar, which was controlled by a Parrinello–Rahman pressure [59], and Parrinello–Rahman pressure [59], respectively. The integration step of 2 fs was used. Each system was simulated for 200 ns and the snapshots were saved every 10 ps for further analysis. All production runs were performed on CUDA-enabled Tesla GPU machine (DELL T640 with V100 GPU) and OS Centos 7 [60, 61].

MD analysis
The obtained MD trajectories were analyzed using GROMACS utilities. The structural order parameters that we measured are: root-mean-square deviation (RMSD), the radius of gyration (Rg), solvent-accessible surface area (SASA), root-mean-square fluctuation (RMSF) and hydrogen bond (H-bond) interactions. H-bonds were defined by a distance cut-off of 3.5 Å between the donor and acceptor atoms and by an angle cut-off of 30°. Similarly, a hydrophobic interaction was defined by the condition that the distance between two residues (i and j, with |i − j| > 3) is less than 4.5 Å. Principal component analysis performed using the projection of principal components (PCs), PC1 and PC2, along the native structure [61, 62] and gmx-sham utilized for the FEL [63, 64].

Binding free energy estimation
The binding free energy of the protein–ligand complexes was evaluated using MM-PBSA, which describes the structural and molecular stability of the ligands in the active site of the protein [27, 28, 65, 66]. The binding free energy of a protein–ligand complex (∆Gbinding) can be written as,

$$\Delta G_{\text{binding}} = (G_{\text{complex}}) - (G_{\text{receptor}}) - (G_{\text{ligand}}),$$

where $G_{\text{complex}}$ represents the free energy of the protein–ligand complex; $G_{\text{receptor}}$, the free energy of protein; $G_{\text{ligand}}$ is the free energy of ligand and 〈〉 represents the ensemble average.

Excluding the entropy term (T∆S), the above equation for the binding free energy can be approximately written as,

$$\Delta G_{\text{binding}} \approx \Delta E_{\text{MM}} + \Delta G_{\text{solv}},$$

where $\Delta E_{\text{MM}}$ is the change in the average molecular mechanics interaction energy (gas phase) upon ligand binding computed as the sum of the changes in the bonded and non-bonded (electrostatics and vdw) interactions upon ligand binding ($\Delta E_{\text{MM}} = \Delta E_{\text{bonded}} + \Delta E_{\text{electrostatics}} + \Delta E_{\text{vdw}}$). $\Delta G_{\text{solv}}$ is the change in solvation free energy upon ligand binding. Further, $\Delta G_{\text{solv}}$ can be written as,

$$\Delta G_{\text{solv}} = \Delta G_{\text{POL}} + \Delta G_{\text{NP}},$$

where $\Delta G_{\text{POL}}$ is the change in the polar part of the solvation free energy, and $\Delta G_{\text{NP}}$ is the change in the non-polar part of the solvation free energy as a result of ligand binding to the proteins. PB equation was used for the estimation of the polar part of the solvation free energy, and the non-polar part was estimated with a surface area-based approach. Binding free energy ($\Delta G_{\text{binding}}$) for the protein–ligand complex was estimated using the MMPBSA.py script of the AMBER Tools [67]. An ionic strength of 0.15 M and a solute dielectric constant value of 2 was used for the PBSA calculations. Considering the convergence issues associated with the MM-PBSA calculation, only last 50 ns data were used.

Results and discussion
Targeting SARS-CoV-2 proteins for identifying multi-target agents
The development of novel molecules is not thought to be realistic in the timeframe needed to impact on the pandemic of SARS-CoV-2 [32, 35]. Thus, there is a clear need for new treatments that will reduce the mortality rate while we wait for a vaccine or the novel drug molecules. The repurposing of FDA-approved drugs often provides an advantage to identify the promising drug molecules in the shorter time which can be undertaken for the medical application without delay [68, 69]. Furthermore, it is more cost-effective and less risky as compared to the de novo drug discovery [32, 34, 35, 70]. However, the ability of a chemical compound to work as a drug lies in its competency to bind efficiently to a druggable target. While there exist plenty of methods to evaluate the binding affinity of a ligand toward a target starting from very reliable and accurate alchemical free energy methods (computationally very costly) to
less accurate docking methods, molecular docking and virtual docking remain the first choice to screen chemicals as they offer reasonable accuracy with modest computational efforts [26, 28, 29]. Considering this, we have first performed molecular docking study to screen the efficacy of 75 FDA-approved drugs against various important druggable targets of SARS-CoV-2 [34]. Our docking study revealed that some of the FDA-approved drugs have excellent interactions with particular target, displaying satisfactory docking scores. Docking scores of the top 10 antiviral drug molecules corresponding to each protein are enumerated in Table 1. Results demonstrate the best binding affinity of antiviral drugs: ritonavir, dolasetravin, tenofovir, tinofovir, bocaprevir, catechin and zanamivir toward the target proteins: 3CLpro, RdRp, ACE2, CTSL, NSP6, nucleocapsid protein and RBD, respectively (Supplementary Figures S2 and S3 available online at https://academic.oup.com/bib). Interestingly, the naturally derived polyhydroxy molecule catechin showed multi-targeted action against all seven targets; however, its promising binding capability (with the cut off range -5.0 kcal/mol) was noticed against the five important targets engaged in the invasion and survival of SARS-CoV-2 in to the human cells. We found that catechin effectively binds to 3CLpro, CTSL, NSP6, nucleocapsid protein and RBD of S protein, showing the docking scores in the range of −5.79 to −8.34 kcal/mol.

Molecular interaction of catechin as multi-targeted agent

The structure-based virtual screening protocol bestowed catechin as the effective multi-targeted agent which effectively hits 3CLpro, CTSL, NSP6, nucleocapsid protein and RBD (Figure 1 and supporting information, Supplementary Figure S3 available online at https://academic.oup.com/bib). The SARS-CoV-2 protein, 3CLpro plays a critical role in the replication of the virus particles and is a potential target for anti-coronaviruses inhibitors screening. The active site of 3CLpro consists of Cys-His catalytic dyad (Cys145 and His41), which is highly conserved in the CoVs family, also referred as the main protease, Mpro [71]. The molecular docking result shows that catechin nicely fits in the active site of 3CLpro with the highest docking score, −8.34 kcal/mol, among all selected five target proteins. It displays H-bonds with Thr26, Met49, Arg188 and Gln189, whereas Leu27, His41 and Leu58 are involved in hydrophobic interactions. Phenolic hydroxy participate in H-bonding with Ser216, and hydroxy at benzopyran moiety showed H-bonding with Met161. Benzopyran ring was stabilized by the hydrophobic interaction with Trp26, His163 and Ala214.

In the CoVs family, nucleocapsid acts as a multifunctional RNA-binding protein and plays an indispensable role in regulating viral RNA transcription/replication and the modulation of host cell metabolism [73]. The recently solved N-terminal domain of SARS-CoV-2 nucleocapsid consists of antiparallel β-sheets at the core, protruding β-hairpin and short 3₁₀-helix [36]. The molecular interaction of catechin with nucleocapsid shows that phenolic hydroxy formed H-bond interaction with

Table 1. Molecular docking scores (kcal/mol) of the FDA-approved potential antiviral drug molecules (best 10 hit molecules) with SARS-CoV-2 proteins

| S. No. | 3CLpro | RdRp | ACE2 | CTSL | NSP6 | Nucleocapsid protein | RBD (S protein) |
|-------|--------|------|------|------|------|----------------------|-----------------|
| 01    | Ritonavir | -8.39 | Dohazevir | -12.43 | Tenofovir | -10.43 | Didanosine | -6.98 |
| 02    | Catechin | -8.29 | Bocaprevir | -10.73 | Tenofovir | -7.83 | Tenofovir | -6.42 |
| 03    | Nelfinavir | -8.28 | Didanosine | -10.63 | Tenofovir | -7.83 | Tenofovir | -4.54 |
| 04    | Nelfinavir | -8.28 | Didanosine | -10.43 | Tenofovir | -7.83 | Tenofovir | -4.54 |
| 05    | Indinavir | -8.24 | Zalcitabine | -10.01 | Tenofovir | -7.50 | Tenofovir | -4.58 |
| 06    | Beclabuvir | -8.16 | Zalcitabine | -10.01 | Tenofovir | -7.50 | Tenofovir | -4.58 |
| 07    | Tenofovir | -8.16 | Zalcitabine | -9.05 | Tenofovir | -7.42 | Tenofovir | -4.58 |
| 08    | Abacavir | -8.16 | Zalcitabine | -9.05 | Tenofovir | -7.42 | Tenofovir | -4.58 |
| 09    | Trifluridine | -8.16 | Zalcitabine | -9.05 | Tenofovir | -7.42 | Tenofovir | -4.58 |
| 10    | Telbivudine | -8.16 | Zalcitabine | -9.05 | Tenofovir | -7.42 | Tenofovir | -4.58 |
Figure 1. Molecular interactions of FDA-approved antiviral drug, catechin at the binding pocket of potential target proteins, RBD of S protein, 3CL\textsuperscript{pro}, CTSL, nucleocapsid and NSP6, using LigPlot. The bar chart showing the target proteins specific antiviral drugs binding affinity (kcal/mol) obtained through virtual screening (represented by green bars) and the comparative binding affinity (kcal/mol) with catechin shown with red bars.

Ala\textsubscript{108}, and phenyl moiety imparted hydrophobic interaction with Trp\textsubscript{5}, Ala\textsubscript{109} and Ile\textsubscript{110}. Hydroxy group at benzopyran moiety was stabilized by the H-bond interaction with Asn\textsubscript{28} and Ser\textsubscript{31}. Additionally, this benzopyran moiety was stabilized through hydrophobic interactions with His\textsubscript{98} and Ile\textsubscript{99}. Another important CoVs protein, non-structural viral protein 6 (NSP6), plays an essential role in viral RNA synthesis by sequestering the membrane of ER of the host cell [74]. A strong affinity of catechin with NSP6 was also discerned. Indeed, catechin contoured well in the active site of NSP6, displaying a satisfactory docking score (~6.68 kcal/mol). It was noticed that catechin forms tight interactions in the active site through three H-bond interactions and several hydrophobic interactions. Hydroxy groups of catechin are found to be involved in H-bonding with
is seen up to 200 ns, suggesting stable interaction of catechin
RMSD of equilibrium can be seen up to 90 ns. We find a slight drop in the
∼ rise in RMSD the simulation time. The RMSD plot of 3CLpro shows an initial
attained equilibrium in 0–5 ns and remained stable throughout
active site of the protein very quickly (Figure 2A). These systems
complex, we observed that catechin achieved stability in the
like RMSD, complexes by measuring various structural order parameters
We assessed the conformational stability of the protein–ligand

Thus, our molecular docking study evidently indicates that catechin strongly interacts with these five crucial targets associated with SARS-CoV-2, which clearly designates the multi-targeted action against SARS-CoV-2 [75]. However, considering the approximations made in molecular docking (lack of receptor flexibility and conformational entropy, lack of information about the number and free energy of water molecules in the binding site of the protein, etc.) to allow fast screening of chemicals, the dynamics of the protein–ligand interactions are overlooked, and therefore, might not explain the stability of the ligands in the active site of the protein. In the cellular system, biomolecular interactions are dynamic in nature, and the conformational flexibility is an intriguing property of proteins which triggers the biological functions and molecular recognitions [27, 30, 76]. A better understanding of protein–ligand interactions requires an accurate description of the spatial orientation of ligands at the active site of the protein, conformational dynamics which modulates the drug binding, interaction energy and molecular stability [77–79]. To understand the biomolecular interactions at atomic resolution, MD simulation is an efficient and well-established method which mimics the flexible nature of biomolecules, protein conformational changes, protein–ligand interactions, structural perturbation and provides a more realistic picture with atomic details in reference to time [61, 80, 81]. Thereby, to gain a deeper insight into the structural dynamics and stability of catechin binding with SARS-CoV-2 proteins, multiple MD simulations were performed for the period of 200 ns [82, 83]. Additionally, the energetic contribution of binding pocket residues to accommodate the drug molecule, catechin, is estimated using MM-PBSA [28, 84].

Conformational stability of protein–ligand complexes
We assessed the conformational stability of the protein–ligand complexes by measuring various structural order parameters like RMSD, Rg, SASA and RMSF, as shown in Figure 2. On comparing the Co-RMSD of RBD, CTSL and nucleocapsid protein complex, we observed that catechin achieved stability in the active site of the protein very quickly (Figure 2A). These systems attained equilibrium in 0–5 ns and remained stable throughout the simulation time. The RMSD plot of 3CLpro shows an initial rise in RMSD ∼0.2 nm, which settles gradually, and a stable equilibrium can be seen up to 90 ns. We find a slight drop in the RMSD of ∼0.1 nm around ∼100 ns and the undisrupted trajectory is seen up to 200 ns, suggesting stable interaction of catechin in the binding pocket of CTSL. The trajectory of NSP6 with catechin shows slightly large deviations in RMSD during 0–80 ns; thereafter, a gradual drop in RMSD can be seen, which attains equilibrium around ∼120 ns. Notably, the stable conformational dynamics of NSP6–catechin is observed up to 200 ns. Thus, the shorter equilibration time taken by RBD, CTSL, nucleocapsid and 3CLpro to achieve a steady equilibrium suggests a better equilibrated and stabilized protein–ligand complex structure compared to NSP6. However, the stable trajectory of the NSP6–catechin complex during 120–200 ns signifies that the ligand is spatially well occupied and stabilized with the molecular interactions at the binding pocket of NSP6.

To further understand the structural stability of the protein–ligand complexes, we determined the compactness of the protein structure by computing the radius of gyration (Rg). The Rg plots represented in Figure 2B show that the structural dynamics of RBD, CTSL and nucleocapsid protein and 3CLpro remain quite stable throughout the simulation time. The structural integrity of these four proteins was observed to be intact with the average Rg values, 1.83 ± 0.01 nm, 1.54 ± 0.01 nm, 1.50 ± 0.01 nm and 2.19 ± 0.01 nm, respectively. The slight deviations in the Rg plot of NSP6 can be seen during 0–80 ns; after that, the steady equilibrium is noted till the end of simulation at 200 ns, which signifies the stable structural dynamics of the NSP6–catechin complex with an average Rg value, 2.23 ± 0.03 nm. The initial perturbation in the Rg trajectory may indicate the spatial adjustment of the ligand in the binding site of NSP6.

Another important quantity that we measure and analyze to probe the conformational stability of the protein–ligand complex is the SASA. The solvent environment around the protein plays a key role in maintaining the protein fold and governs the protein–ligand interaction processes, orientation and stability. Interestingly, we find that the SASA plots of all five protein–ligand complexes (RBD, CTSL, nucleocapsid protein, 3CLpro and NSP6) remain fairly equilibrated during the entire simulation period (0–200 ns) which provides clear evidence of the stable conformational dynamics of protein–ligand interactions (Figure 2C). The average values of the structural order parameters, RMSD, Rg and SASA are shown in the supporting information, Supplementary Table S2 available online at https://academic.oup.com/bib.

Next, we investigated the binding stability of the catechin at the active site of the respected proteins by monitoring the time evolution plots of the average distance from the center of binding pocket to the ligand, as shown in the supporting information, Supplementary Figure S4 available online at https://academic.oup.com/bib. During the period of 200 ns simulation, the average distance of catechin from the binding site of all five proteins ranges between 0.33 nm and 0.40 nm. Although, the peaks of sharp drifts appeared transiently at ∼80 ns and 140 ns for RBD and nucleocapsid protein, respectively, the overall distance of catechin to the active site remains favorable for the stable molecular interaction. It is worth noting that the average distance plots of CTSL, 3CLpro and NSP6 remain stable throughout simulation time. Thus, this analysis provides an elegant evidence of the spatially well-fitted catechin orientation in the binding sites of proteins.

We further performed RMSF analysis to evaluate the positional fluctuation of each amino acid around its average mean position (Figure 2D). This analysis provides a clue about the mobility of atomic fluctuations related to the structural stability of molecular interaction during the simulation. Usually, the higher values of RMSF are often associated with loops or may be the terminal residues, whereas the lower RMSF values indicate the rigid conformation of stable secondary structures
of α-helices and β-sheets. The result indicates that all the complexes show equilibrium fluctuations, except NSP6. The plot shows that the values of RMSF significantly vary for all residues of NSP6 in comparison to the other proteins, RBD, CTSL, nucleocapsid protein and 3CLpro, respectively. The RMSF plots of these proteins show an average atomic fluctuation <0.15 Å for amino acid residues, which belong to the stable secondary structure, and the regions which displayed high fluctuations represented the atomic flexibility of loops. The structure of NSP6 consists of a helical structure at the core, capped by antiparallel β-sheets and two small helices. The binding pocket is characterized by α-helix-2-4 and α-helix-6-7 and antiparallel β-sheets (β1 and β2) which are enclosed by a hydrophobic loop (Phe235-Tyr242) connecting helices and β-sheets. The RMSF plot shows on average, high fluctuations for the residues belonging to longest loop (Val84-Leu110) connecting α-helix-3 and α-helix-4, loops (Gln257-Ser262 and Leu275-Pro282) connecting to small two helices (α8 and α9) with β-sheets, respectively.

The average residual fluctuations observed were reasonably lower for terminal residues of α-helix-2 (Phe42-Phe59), α-helix-3 (Lys63-Met86), α-helix-4 (Lys109-Arg129), α-helix-6 (Ala157-Thr196), α-helix-7 (Tyr175-Tyr196) and N-terminal of connecting loop (Phe235-Tyr242) and β1 (Asp243-Val246) which are actively involved in molecular interactions with drug molecules. Further, we also established the conformational stability by analyzing the secondary structural contents, which were observed intact during the simulation (supporting information, Supplementary Figure S5 available online at https://academic.oup.com/bib). The structural snapshots of protein–ligand interactions captured at the time interval of 20 ns described in the supporting information Supplementary Figure S6 available online at https://academic.oup.com/bib. Thus, the overall results clearly indicate the stable conformational dynamics of the target proteins complexed with antiviral drug, catechin.

Another parameter, H-bonds interaction is one of the major players in governing the ligand stability at the active site of the

Figure 2. Time evolution plot of the structural order parameters of the antiviral drug, catechin-docked complex with target proteins: RBD, 3CLpro, CTSL, nucleocapsid and NSP6. (A) The RMSD of backbone Cα-atoms, (B) radius of gyration (Rg), (C) SASA plots, (D) RMSF showing the average fluctuation of amino acid residues and (E) the propensity of H-bonds interaction between the proteins and catechin during the period of simulation (200 ns) at 300 K.
with ligand (H7). 3CLpro showed six H-bond interactions with residues, Tyr451 (OH) with ligand (H7 and H8) and Asp442 (O).

result shows that catechin formed H-bond interaction with observed to be stable during the last 50 ns of simulation. The ∼were regained at

were formed between nucleocapsid–catechin, which ∼catechin which could be seen up to 60 ns, which were regained at 100 ns. Nevertheless, three H-bonds were observed to be stable during the last 50 ns of simulation. The result shows that catechin formed H-bond interaction with residues, Tyr451 (OH) with ligand (H7) and H8) and Asp442 (O) with ligand (H7). 3CLpro showed six H-bond interactions with catechin which could be seen up to 150 ns, but only three remained stable during the last 50 ns, which were formed between the residues, Asp187 (O)—ligand (H7), His164 (HE2)—ligand (O2) and Thr26 (O)—ligand (H14). CTSL formed three H-bonds, which were observed consistent up to 170 ns. It showed maximum occupancy with Asp162 (OD2)—ligand (H7), Asp160 (OD1)—ligand (H8) and Asp160 (OD2)—ligand (H8). Three H-bonds were formed between nucleocapsid–catechin, which were observed with Arg46 (HN)—ligand (O1), Asp56 (OD1)—ligand (H7) and Thr44 (HG1)—ligand (O2). NSP6 showed the possibility of three-four H-bonds, however, there were two H-bonds between Thr238 (O)—ligand (H14) and Thr130 (HG1)—ligand (O1), having maximum occupancy during the last 50 ns of simulation. These results indicate that catechin is stabilized by an average of two-three H-bonds at the binding pocket of proteins.

Essential dynamics

Protein function is regulated by switching between various conformations. The modular nature of proteins to switch between various states is governed by the collective motion of protein, which is intrinsic to many biological processes and plays a crucial role in the transmission of biological signals. For a protein to be functional, a reasonable amount of flexibility, as well as rigidity is required, specifically for the residues in the binding site. Essentially, a tighter interaction would restrict the motion of the protein, thereby not allowing it to sample some conformations required for its activity. Therefore, in order to understand the collective motion of protein occupied in the conformational space during the simulation, we applied the dimension reduction method, essential dynamics (ED) analysis by the projection of the first two PCs, PC1 and PC2. The PC1 and PC2 were calculated by diagonalizing the covariance matrix of eigenvectors to define the essential subspace in which most of the protein dynamics occur. The dynamic motion of proteins obtained through the projection of PC1 and PC2 are shown in Figure 3. It is apparent from these plots that the collective motion of proteins, RBD, CTSL and nucleocapsid protein is localized in a small conformational space in comparison to 3CLpro and NSP6, which revealed consistent results corresponding to the structural order analyses, RMSD, Rg and SASA, as described in Figure 2A–C. The well-defined small clusters of RBD, CTSL and nucleocapsid protein clearly indicate the reliability and stability of the complex structure with catechin. The ED plot of 3CLpro displays a slight increase in the conformational phase space, which can be seen along the PC2, which suggests that protein navigated the broad conformational space before achieving the ensemble of the dynamically equilibrated state. Contrary to this, NSP6 experiences a wide region of phase space. In fact, it explored a large conformational space in comparison to the other four proteins, which represent the overall higher flexibility of the protein. Thus, we observed a significantly compact structure of RBD, CTSL and nucleocapsid protein with 3CLpro as compared to NSP6, which may facilitate the vital interactions with catechin.

Free energy landscape

FEL provides an accurate description of the minimum energy conformational ensembles of biomolecules, which is undoubtedly essential to understand the conformational transition underlying protein–ligand interactions [85]. Thus, FEL plot is constructed using Boltzmann inversion (F = −RT ln P), where P is the two-dimensional probability distribution of the first two PCs, PC1 and PC2, as reaction coordinates. Figure 4 shows that the binding of catechin with proteins occurs through the minimum free energy pathway. The structural ensemble derived from FEL shows that the catechin-bound complex with RBD navigated the broad conformational space, clustered in the different energy basins, distributed along the PC1. However, these energy minima separated through the low transition barrier (<2.0 kcal/mol) indicates that with small excursion, the ensemble states of RBD can easily move out from one energy basin to another. This may be the reason we observed a small and consolidated cluster of stable populations in the ED analysis. The FEL plot of 3CLpro shows the appearance of two distinct populations confined to two different energy basins, separated with high transition barrier (>4.0 kcal/mol, which signifies the population of loosely and tightly ligand bound conformations of the protein. The conformational ensemble occupying the small energy basin represented the population of the equilibrium phase, which readily achieved a stable equilibrium. These (ed)ible ensembles of stable complex transverse to a broad and deep energy basin. The complex with CTSL shows single but elongated energy minima, which depicts the heterogeneous population of different sub-states, but the very less transition barrier (<1.0 kcal/mol between the ensemble states suggested the stable conformation of the protein–ligand complex confined to energy basin interplays between the
subspace. Whereas, the rugged FEL with segmented small energy minima of nucleocapsid protein suggested the population of loosely bound complex [86]. The low transition barriers (∼1.5 kcal/mol) between the small energy basins indicate a more prolonged equilibration phase of complex structure. During the progression of simulation, the protein underwent structural modifications to accommodate the ligand and adopted a stable conformation; thus, the equilibrated ensemble smoothly shifted to broad and deep energy minima. Remarkably, The FEL of NSP6 shows that the stably bound conformation of the protein-ligand complex is widely populated to a single consolidated energy minimum, which provides the elegant evidence of interactions inducing the stable conformational transition of NSP6-catechin complex. Thus, the comparison of FEL results indicates the different binding stabilities of the catechin-bound protein-ligand complexes.
Figure 6. The residue decomposition plot (MM-PBSA) representing the binding energy contribution of the active site residues of five target proteins energetically stabilizing the catechin at binding pockets. (A) RBD, (B) 3CLpro, (C) CysL, (D) nucleocapsid and (E) NSP3.
Binding free energy and ligand–residue interaction decomposition

In order to understand the molecular interaction and stability associated with the binding of catechin to five different proteins of SARS-CoV-2, a detailed analysis of the binding free energy is executed through the MM-PBSA. MM-PBSA provides the best prediction accuracy in terms of energy components of bonded, polar and non-polar solvation free energy, electrostatic and vdW interactions. In addition, it supplies the residue decomposition plot, which helps to probe the contribution of amino acid residues involved in the spatial interaction to stabilize ligands at the binding pocket of the protein. This analysis was performed on the fully converged trajectory of the last 50 ns with a solute dielectric constant value of 2 and an ionic strength of 0.15 M (supporting information, Supplementary Figure S7 available online at https://academic.oup.com/bib). Results show that catechin favorably binds to all five proteins, however, it shows a wide range of the total binding free energies ($\Delta G_{\text{bind}}$) as enumerated in the supporting information, Supplementary Table S4 available online at https://academic.oup.com/bib. As illustrated in Figure 5, catechin possesses the highest binding affinity toward NSP6, with a maximum value of $\Delta G_{\text{bind}}$ = −26.09 kcal/mol, whereas the lowest was toward CTSL ($\Delta G_{\text{bind}}$ = −5.09 kcal/mol). Results show relatively more favorable contribution of vdW energies ($\Delta E_{\text{vdW}}$) = −37.39 kcal/mol and −24.76 kcal/mol for NSP6 and 3CLpro as compared to nucleocapsid protein (−16.34 kcal/mol) and RBD (−15.8 kcal/mol), respectively. Whereas the least contribution of vdW interaction (−7.59 kcal/mol) is observed for CTSL. Another binding energy component, electrostatic ($\Delta E_{\text{ele}}$) energy, which describes ligand–protein interactions, is a critical factor in determining the binding stability of ligand. The interaction of catechin with CTSL shows the major contribution of electrostatic energy −29.61 kcal/mol; however, the lowest values of $\Delta E_{\text{ele}}$ = −7.59 kcal/mol and $\Delta G_{\text{bind}}$ = −5.09 kcal/mol signify that the electrostatic energy contributes relatively less as compared to the other energies in the binding stability of catechin. Thus, the binding free energy analysis revealed that the binding of catechin to the active site of proteins are predominantly stabilized by hydrophobic interactions.

To further quantify the contribution of binding pocket residues to the molecular interaction of catechin with five different proteins, the free energy decomposition per residue was employed (Figure 6). The plot of free energy decomposition analysis shows that the active site residues, Agr346, Phe347, Leu441, Asp442, Lys444, Tyr449, Asn450 and Try451 energetically favor the binding stability of catechin to RBD. Remarkably, it is noted that Agr346 contributed the highest binding free energy, $\Delta E_{\text{vdW}}$ (~2.10 kcal/mol), $\Delta E_{\text{ele}}$ (~1.73 kcal/mol) and $\Delta G_{\text{bind}}$ (~2.10 kcal/mol), which indicated the favorable electrostatic and vdW interactions with catechin (Figure 6A). The protonated (−NH$_3^+$) Agr346 shows the electrostatic interactions, whereas the side chain guanidinium [−C(NH$_2$)$_2$] facilitated the hydrogen binding interaction with ligand. The binding interaction with 3CLpro shows that the amino acid residues, Leu27, His47, Ser46, Met49, His164, Met165, Asp187 and Gln189, contributed the most to the total $\Delta G_{\text{bind}}$ (~16.98 kcal/mol). Although the vdW interaction primarily stabilizes the catechin at the binding pocket of 3CLpro, the electrostatic interaction also contributes toward the observed stability by His164 (~3.80 kcal/mol) and Asp187 (~3.10 kcal/mol), respectively (Figure 6B). Indeed, catechin is predominantly stabilized in the binding pocket of CTSL through the electrostatic interaction, which is mostly contributed by the residues Asp71 (~11.31 kcal/mol), Asp114 (~11.36 kcal/mol), Asp160 (~7.61 kcal/mol) and Asp162 (~7.89 kcal/mol), respectively (Figure 6C). Figure 6D showing the free energy decomposition plot of nucleocapsid protein indicates the substantial contribution of amino acids, Thr44, Arg45, Arg46, Asp56, Arg60 and Tyr62 to energetically hold catechin at the binding pocket. Surprisingly, it is noted that Arg45 contributed to both electrostatic (−3.07 kcal/mol) and vdW interaction (−1.88 kcal/mol), but Arg46 contributed only vdW interaction (−2.83 kcal/mol). This may be the reason we observed a moderate range of total binding energy ($\Delta G_{\text{bind}}$) of value: −14.15 kcal/mol. The favorable binding of catechin with NSP6 shows the significant contribution of residues, Lys61, His62, His64, Ala65, Asp133, Asp134, Phe228, Leu237, Thr238, Leu239 and Gln290 (Figure 6E). Interestingly, it is noted that Thr238 contributed the higher electrostatic energy (~5.0 kcal/mol), whereas the maximum vdW energy (~2.22 kcal/mol) was contributed by Phe228. However, the binding pocket of NSP6 mostly consists of hydrophobic residues; thus, we observed the major collective contribution of vdW energy for stabilizing the ligand interaction.

**Conclusion**

In summary, using molecular docking and classical MD simulation, we have explored the possibility of 75 FDA-approved antiviral drugs for their potential of being used as an effective therapeutic strategy to control SARS-CoV-2 infections. The virtual screening results showed that seven therapeutic agents, ritonavir, dolutegravir, tenofovir, tinofovirafenamide, boceprevir, catechin and zanamivir, could efficiently bind to the SARS-CoV-2: 3CLpro, RdRp, ACE2, CTSL, NSP6, nucleocapsid protein and RBD of spike (S) protein, respectively. Of these, catechin has the potential to act as a multi-targeted agent, as it has the highest binding affinity toward the five crucial proteins of the virus, RBD, CTSL, nucleocapsid protein, 3CLpro and NSP6, which are essential for the invasion and infection of the host cell. Further, MD simulation, FELs and binding free estimation of catechin with the five target proteins explained the stable interactions of catechin with the critical residues in terms of occupancy of H-bonds and residue contributions to the binding free energy. Thus, our investigation bestowed promising multi-targeted agent catechin, which can be explored as an effective therapeutic agent against the SARS-CoV-2 virus to control the COVID-19 pandemic.

**Key Points**

- Virtual screening-based repurposing of FDA-approved antiviral drugs for identifying the multi-targeted agent against SARS-CoV-2.
- Catechin is identified as an effective multi-targeted agent.
- Out of seven target proteins, it shows higher binding affinity with 3CLpro, CTSL, RBD of S protein, NSP6 and nucleocapsid protein.
- Molecular interactions were evaluated through MD simulations, FELs and binding free energy estimations.

**Supplementary Data**

Supplementary data are available online at Briefings in Bioinformatics.
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Conflict of interest
Authors declare there is no conflict of interest.

Author's Contributions
Conceptualization was by C.B.M., A.P. and R.J.; methodology and analysis were taken care of by A.P., C.B.M., P.P., R.D.S., R.K.M. and R.J.; funding acquisition was by R.J.; supervision was done by A.P., R.J. and C.B.M. and writing, review and editing were by A.P., C.B.M., R.J., P.P., R.D.S., R.F. and A.M.L.

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