An In Silico Approach for Identification of Inhibitors as a Potential Therapeutics Targeting SARS-Cov-2 Protease

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Abstract

SARS-CoV-2 caused COVID-19 which is pandemic and is a global health emergency. Protease is the drug target for corona viruses and this enzyme processes the production of polyproteins from the viral RNA. Objective of this study is to find the inhibitors against SARS-Cov-2 protease. AutoDock 4.2 was used for docking calculations. To check different molecules, test ligands like lopinavir, ritonavir (retroviral drugs) and hydroxychloroquine (anti-malarial drug) were docked against our target enzyme protease retrieved from Protein Data Bank. With respect to docking free binding energy, it is revealed that, Hydroxychloroquine has the lowest binding energy followed by Ritonavir and Lopinavir binds significantly to target enzyme protease. The results of this study provide a solid base for the use of Hydroxychloroquine against COVID-19 treatment. The interactions from structural models at the protease active site of virus can offer a valuable guide for more strategies for structure-based medications and the development of more operative inhibitors of SARS-CoV-2 protease.

Keywords: COVID-19, Hydroxychloroquine, In Silico, Lopinavir, Ritonavir, SARS-Cov-2iProtease

1. Introduction

The outbreak of COVID-19 is caused by severe acute respiratory syndrome coronavirus-2 (SARS-CoV-i2) appears to have happening in Wuhan, China in December 201912. Afterward, it has extent within and outside China3, 4 and has develop an unparalleled global communal health issue5. As of 23rd March 2020, India has 471 confirmed cases of Covid-19, including nine deaths6. According to data from the Indian Council of Medical Research a total of 20,864 samples from 19,974 individuals have been tested for SARS-CoV2 as on 24th March 2020. A total of 482 individuals have been confirmed positive among suspected cases and contacts of known positive cases7. Thirty states and union territories in India have announced a complete lockdown to curb the spread of corona virus.

One of the best considered drug targets of corona viruses is the main protease8. Beside with the papain-like protease(s), this enzyme is important for producing the polyproteins that are translated from the viral RNA. Constraining the activity of this enzyme would block viral duplication. Meanwhile no human proteases with similar cleavage specificity are recognized, inhibitors are improbable to be lethal. Currently, there is no exact treatment beside the new virus. Thus, finding active antiviral mediators to combat the infection is immediately desirable. An effective attitude to drug discovery is to check whether the existing anti-viral medicines are operative in treating linked with viral infections. Computational approaches are dynamic part
of the medicine scheme procedure and they are presently used to acquire a deep considerate into the ligand-receptor/protein interfaces. In this current study, we evaluated the antiviral efficiency of FDA approved two retroviral drugs (Lopinavir, ritonavir) and one well-known broad-spectrum anti-malarial drug (Hydroxychloroquine) against a novel SARS-CoV-2 protease (COVID-19) or *In Silico*.

2. Materials and Methods

2.1 PASS Prediction

For activity spectra prediction of the considered drugs the program PASS was required. PASS, calculates the natural efficacy spectrum of a molecule built on the examination of structural performance associations of above one million of recognized molecules\(^9\). These molecules possess over 80 different biological actions. The normal exactness of prediction is nearly 95%.

2.2 Molecular Docking Methods

For molecular docking, Auto-Dock 4.2 software was used\(^{10}\). The free energy (DG) binding of SARS-CoV-2 viral protease with the selected compounds was created by means of this molecular docking package.

Docking is a computational simulation method of a ligand binding to a receptor or enzyme and expects the favored orientation of binding of one molecule to the 2nd to form a steady complex. To predict the attraction and activity of binding of the minor molecule to their enzyme targets by using scoring functions docking is used. Therefore, docking shows significant role in the rational design of medicines. The sensitivity of docking calculations concerning the geometry of the involvement ligand displays that even minor changes in the ligand structure can lead to big changes in the geometries and scores of the subsequent docked poses.

2.2.1 Selection of Ligand

Anti-viral and anti-malarial medications were recognized as potential corona virus inhibitors from diverse literature evaluations. The three-dimensional structure files of the selected compounds were downloaded in SDF format from the PubChem 3-D was used for molecular docking. Figures 1(a, b, & c) show the structures of selected different ligands.

![Ligand Structures](image)

**Figure 1.** Test ligands (a-c): (a): Lopinavir, (b): Ritonavir, (c): Hydroxychloroquine.

2.2.2 Selection of Target

The main COVID-19 protease remained used as a target to novelty repurposing candidates over computational selection amongst clinically accepted drugs. The study identified a list of FDA permitted two retroviral drugs and one anti-malarial medicine that may form hydrogen bonds to key residues of amino acids within the binding pocket of viral protease and may too have a higher tolerance to conflict mutations. The crystal 3D structure of SARS-CoV-i2 protease (PDB ID: 6LU7) remained obtained from *Protein-Data Bank*\(^{11}\) (Figure 2).
Meanwhile this protease has its crystal structure in a state that signifies the pharmacological target for the progress of new medicines to treatment diverse infectious diseases. The preparation of the target enzyme 6LU7 with the Auto-Dock tools software intricate addition of all H2 atoms to the enzyme, which is a step essential for accurate calculation of fractional atomic charges. The ligand and all water molecules were detached to make the structure for docking. Gasteiger charges are considered for each atom of the protein in AutoDock 4.2 instead of Kollman charges which were used in the earlier versions of this package.

2.2 Docking Procedure

For ligand conformational incisive, we take the 'Lamarckian-Genetic Algorithm (LGA)', which is a mixture of a genetic algorithm and a native search algorithm. This algorithm initially builds a population of entities, being a diverse casual conformation of the docked enzyme. Each distinct protein is then mutated to attain a slightly diverse translation and alternation and the local search algorithm then achieves energy minimizations on a user-specified amount of the population of individuals. The entities with the low subsequent energy are moved to the succeeding generation and the procedure is then repetitive. This algorithm is called Lamarckian while every novel group of entities is allowable to receive the local search variations of their parents.

To get many docked structures, Auto-Dock was run numerous times, and used to examine the expected docking energy. Rapid energy assessment was attained by pre-calculating nuclear affinity capacities for every atom in the compound molecule. The binding sites of the target enzyme for these molecules in the Auto Grid process were designated on the patterns of founded ligand-binding pockets. Auto-Dock Tools deliver various approaches to examine the outcomes of docking-simulations such as, structural resemblance, and other limitations like inter-molecular energy, visualizing the binding site and its energy and inhibition constant. The energy of interaction of every atom in the ligand was met. For each ligand, 10 best postures were made and scored using Auto-Dock 4.2 scoring purposes.

3. Results and Discussion

3.1 Computational Prediction of Anti-Viral Activity

For three selected FDA accepted drug molecules PASS calculation of anti-viral activity was achieved. Calculation was conceded out by means of PASS online version. The anti-viral activity was calculated for all particular compounds, with Pa values in range of 0.323–0.602 (Table 1). The predicted Pa values for the most of selected composites were less than 0.5, representing their virtual novelty compared to the structures of the drugs from the PASS training set. All the three compounds Pa values are higher (lopinavir 0.538; ritonavir 0.602; Hydroxychloroquine 0.623). However, the computational prediction of anti-viral activity results states that, the compounds exhibiting anti-viral effect.

3.2 Molecular Docking Prediction

Binding energy is the preliminary parameter which is generated as a consequence of molecular docking. It gives us the knowledge of strength and attraction of the interaction between the ligand and the target enzyme. The higher the binding energy is, the frailer the interaction is and vice versa. Therefore during any docking study, we propose to look for the ligand which shows the minimum binding energy, thus the best attraction amongst the test molecules. Among the test compounds in this study, hydroxychloroquine displayed the lowermost binding energy of $-8.30$ Kcal/Mol. The binding energy of the lopinavir and ritonavir was higher ($-6.11$ & $-8.25$) than the hydroxychloroquine. The binding energies of the tested compounds have been showed in Figure 3 and Table 1.
Using the same parameters, FDA approved two retroviral drugs and one anti-malarial drug were docked to the ‘SARS-CoV-2 protease’ (PDB ID: 6LU7) enzyme (Table 1). All tested compounds showed very worthy binding scores and taking into explanation moreover the calculation results by PASS. Anti-malarial drug, hydroxychloroquine shows higher binding energy values than the other compounds.

The amino acids of SARS-CoV-2 virus involved in the interaction with test compound lopinavir are GLU:166, ASN:142, SER:144, HIS:163,164 PHE:140, CYS:145, LEU:141,167 ARG:188, MET:165, GLN:189,192, ALA:191, THR:190 (Figure 4). The interaction of these amino acids states that, the inhibition of SARS-CoV-2 protease with lopinavir with one hydrogen bond formation.

GLY:143,170, GLU:166, LEU:141,27,167, PHE:140, CYS:145, ASN:142, HIS:41, ASP:187, MET:49,163 ARG:188, GLN:189, ARG:188, MET:165, TYR:54, ASN:142, HIS:41,172,163,164, PHE:140, LEU:141, CYS:145, ASN:142, GLN:189, ALA:191, GLY:143, GLU:166. These hydrophobic interactions donate expressively to the maintenance of the compound-SARS-CoV-2 protease enzyme complex.
As shown in Figure 8 and 9, the hydroxychloroquine ring is positioned exclusive the cavity bounded by the amino acid residues Met165, Cys145, His172, and His163. It is significant to reference that the occurrence of the $H_2$ bond with Gln189 is typical of various anti-viral inhibitors.

Outcomes from the *In Silico* molecular docking study maintained the great inhibitory efficacy of the two retroviral drugs and one anti-malarial compound since they could launch $H_2$ bonds with Glu166, Gly143, and Gln189, the residues of amino acids contributing in the 30-processing response. They also binds with the amino acid residues Lys159, Glu152, Gln148, and Asn155 involved in the strand transmission reaction that caused in an inhibition of SARS-CoV-2 protease activity. Thus, the projected binding interactions of the dynamic molecules with the protease by the docking study evidently established their inhibitory strength towards catalytic response of the protease.
4. Conclusion

The present study concludes that, two retroviral drugs and one anti-malarial drug showed efficient docking score. These outcomes afford a strong foundation for the use of hydroxychloroquine, lopinavir and ritonavir for CORONA management. Moreover, the dynamic ligands inhibited the catalytic response of protease by blocking the residues of amino acids intricate in the processing and strand transmission reactions. The interactions by the structural model at the protease active site can afford a valuable guide for additional strategies for structure-based medicines and development of new operative inhibitors of SARS-CoV-2 protease.

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