



In Silico Screening of Traditional Herbal Medicine Derived Chemical Constituents for Possible Potential Inhibition against SARS-CoV-2

R. Srimathi¹, Muthu K. Mohan Maruga Raja² and Muthu K. Kathiravan^{3*}

¹Department of Pharmaceutical Chemistry, SRM College of Pharmacy, SRMIST, Kattankulathur, Kancheepuram – 603203, Tamil Nadu, India

²Department of Pharmacognosy and Phytochemistry, Parul Institute of Pharmacy & Research, Parul University, Waghodia – 391760, Gujarat, India

³Dr APJ Abdul Kalam Research Lab, SRM College of Pharmacy, SRMIST, Kattankulathur, Kancheepuram – 603203, Tamil Nadu, India; drmkathir@gmail.com

Abstract

The outbreak of SARS-CoV-2 has initiated an exploration to find an efficient anti-viral agent. From the previous scientific studies of traditional herbal medicines like garlic, ginger, onion, turmeric, chilli, cinchona and pepper, 131 chemical constituents were identified. The filtered search of drug-like-molecules searched using Datawarrior resulted in 13 active constituents (apoquinine, catechin, cinchonidine, cinchonine, cuprediene, epicatechin, epiprocurcumenol, epiquinine, procurcumenol, quinidine, quinine, zedoaronediol, procurcumadiol) showed no mutagenic, carcinogenic or toxic properties. *In silico* study of these 13 compounds with the best binding affinity towards SARS-CoV-2 protease was carried out. The ligands were subjected to molecular docking using Autodock Vina. Epicatechin and apoquine showed highest binding affinity of -7 and -7.5kcal/mol while catechin and epicatechin showed four hydrogen bond interactions. It is interesting and worth noticing the interaction of GLU166 residue with the ligand in most of the constituents. The effectiveness of catechin and epicatechin as an antiviral agent could be tested against COVID-19.

Keywords: COVID-19, Catechin, Epicatechin, Data Warrior, Molecular Docking, Plant Products

1. Introduction

On December 30, 2019, the epicenter of the novel coronavirus was first reported in the provinces of Hubei P. R. China. The contagious disease has seen its massive expansion in a short span of about three months by spreading to more than 190 countries and infecting 31 lakhs, causing over 2 lakh casualties worldwide as of April 28, 2020¹. In spite of the available protease inhibitors in the treatment of viral infections, WHO said there is no medicine to prevent or cure SARS-CoV-2. Analyzing the treatment strategies for

the epidemic that occurred in 2002, SARS coronavirus (SARS-CoV), it is observed that Traditional Chinese Medicine (TCM) has found to modulate T cells enhancing the host defensive mechanism². A research paper in *Lancet* indicates glycyrrhizin from liquorice root inhibited viral replication in SARS³.

Among the four classes of coronaviruses, severe acute respiratory syndrome virus (SARS- CoV-2) belongs to beta type⁴. The beta-coronavirus has glycosylated spike (S) protein invading host cells. This S protein binds to angiotensin-converting enzyme 2 (ACE2)⁵⁻⁷. This binding affinity is higher than that of

*Author for correspondence

SARS-CoV⁸⁻¹¹. The COVID-19 is a non-segmented enveloped positive sense RNA virus, β coronavirus. The genome consists of \sim 30,000 nucleotides. It is replicated by a replicase gene that codes for two proteins, pp1a and pp1ab, required for viral transcription and replication. The viral main protease (M^{Pro}) digests the polyprotein thereby causing auto-cleavage of the enzyme itself from pp1a and pp1b. Targeting M^{Pro} in the viral life cycle would prove an attractive target for the deadly virus¹².

Nature provides diversified simplest to complex structures. They have evolved over a long period of time for better interactions with biomolecules. The existing antiviral drugs from natural products are zanamivir, peramivir and lanamivir octanoate¹³. Virtual screening is one of the most powerful tool in drug discovery. The development of a novel candidate for SARS-CoV-2 by a medicinal chemist would take several months or even years till it is marketed. However, to combat the disease at the earliest, an immediate treatment is the vital requirement of the hour.

In an attempt to find the use of traditional herbs, an in-silico based screening¹⁴ was carried out by using computational methods to scientifically prove the efficacy of these herbs against SARS-CoV-2. Chemical constituents of herbs having a long history of traditional use against infectious disease were selected for the study¹⁵. We herein report the identification of natural compounds in comparison with N3 inhibitor (N-[(5-methylisoxazol-3-yl) carbonyl] alanyl-l-valyl-n~1~--((1r,2z)-4-(benzyloxy)-4-oxo-1-[(3r)-2-oxopyrrolidin-3-yl]methyl}but-2-enyl)-l-leucinamide)¹⁶. The study will be further extended to see its effect on ACE 2 and inflammation mediators. In addition, ADMET, toxicity, drug likeliness was predicted along with molecular dynamics simulations¹⁷.

2. Material and Methods

The validated computational approach was developed by collecting the 3D structures of chemical constituents of traditionally used plant products. The downloaded database was further subjected to toxicity screening using Datawarrior v.4.5.1 software¹⁸. The filtered constituents were identified for their binding affinity towards the target using Autodock Vina¹⁹. The constituents with greater binding affinity were then

docked using Autodock²⁰. Figure 1 depicts the protocol of this study.

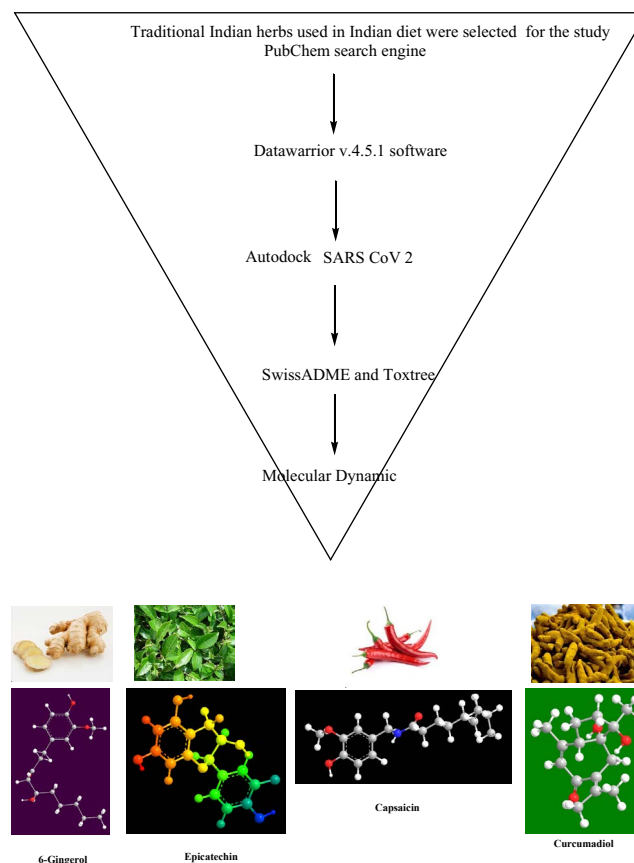


Figure 1. The screening strategy and key chemical constituents from Indian plants.

2.1 Generation of Plant Product Database

The 131 chemical constituents were identified from traditionally used plant materials²¹ such as garlic, ginger, onion, turmeric, chilli, cinchona and pepper. Their chemical structures were collected from PubChem search engine. The dataset collected consisted of major and minor active constituents, which were then screened for toxicity properties, mutagenic, carcinogenic and druglikeness score. Datawarrior v.4.5.1 software was used. Lipinski's rule of five (RO5) and Veber's rule was applied. The filtered ligands were selected for further processing.

2.2 Preparation of 6LU7 Structure

The three-dimensional structure of SARS-CoV-2 main protease in complex, with an inhibitor N3 was obtained

from the Protein Data Bank [PDB: 6LU7]²². Further, water molecules and inhibitors N3 (Michael acceptor inhibitor) complexed with the protein were removed using Swiss PDB viewer and saved as *pdb* file for further virtual screening and docking.

2.3 Virtual Screening using Autodock Vina

Virtual screening of the ligands was carried out using Autodock Vina. Vina is more efficient in calculating the binding affinity (kcal/mol), and low energy binding affinity indicates stronger binding of the ligand with the receptor. Vina was selected over AutoDock 4 for its accuracy and speed in selecting the compounds which showed better binding affinity towards the target. The ligand files were prepared as *.pdbqt* and the macromolecule was added in the receptor. The grid box was centered on the active site at $-11.062 \times 5.943 \times 70.793$ in the dimensions of x, y and z using 1.000 \AA spacing.

2.4 Molecular Docking

The screened ligands using AutoDock Vina were further subjected to docking simulation using AutoDock 4.0 suite as molecular-docking tool. The protein was downloaded from the protein data bank and prepared using the protein preparation wizard of the Graphical User Interface program "AutoDock Tools" by removing the polar hydrogens, by addition of Kollman charges and was saved in *.pdbqt* format. The ligands were defined with their torsional roots and were allowed to rotate freely. The grid was centered in the active site region and the grid box size was set at 60, 60 and 60 \AA and the x,y and z dimensions were $-11.062 \times 5.943 \times 70.793$ respectively. AutoDock 4.0 program using the Lamarckian Genetic Algorithm (LGA) was chosen to identify the best conformers and the top 10 conformers were generated for each compound and studied.

2.5 Bioavailability and Pharmacokinetic Prediction

The physicochemical properties of the top ligand were subjected to SwissADME²³ and Toxtree v2.6.13 software²⁴.

3. Results and Discussion

3.1 Initial Screening of Active Chemical Constituents from Plant Products

The natural plant products with antiviral activity was identified by their major and minor active chemical constituents. About 131 structures from plants such as ginger, garlic, onion, turmeric, cinchona, neem, chilli and pepper were downloaded from PubChem database in *.sdf* format file. From ancient times, it is believed that natural plant products used as traditional foods have a great healing power against microbes. Initial screening of the constituents was performed using Datawarrior v.4.5.1. Software to eliminate the undesirable compounds by following Lipinski's RO5, which states that molecular weight <500 Daltons, LogP should be lower than -1.5 and higher than 6.5, hydrogen bond acceptor <10 and hydrogen bond donor <5. The chemical constituents that defy this criterion were eliminated. The Total Polar Surface Area (TPSA) greater than 180 \AA^2 and having rotatable bonds higher than 14 were also removed.

About 30 compounds did not satisfy Lipinski's RO5, 24 were found to have toxicity properties and 64 compounds were eliminated as they failed to have a druglikeness score of 0. It was identified that only 13 chemical constituents passed all the parameters and were used in the study. They are apoquinine, cinchonidine, cinchonine, cupreidine, epi-quinine, quinine, guanidine, epiprocurcumenol, procurcumiadiol, procurcumenol, zedoronediol, catechin and epicatechin. These compounds were pre-processed and used in docking simulations. Further, the drug likeness of the compounds was considered using the drug likeness score and the compounds were selected that had a score of above 0 (Tables 1, 2).

3.2 Binding Affinity of the Screened Chemical Constituents

With the identification of 13 chemical constituents that showed drug-like-properties, the calculation for the compound with the best binding affinity towards

Table 1. The Pharmacokinetic properties, oral bioavailability and toxicity prediction

S.No	PUBCHEM ID	Pharmacokinetics				Med Chem PAINS	Med Chem PAINS	Oral Bioavailability	
		GI Absorption	Bioavailability Score	CYP inhibitor	Solubility Index			Veber	Egan
1.	101600159	High	0.55	CYP2D6	Good	0	0	Good	Good
2.	9064	High	0.55	None	Good	0	0	Good	Good
3.	101744	High	0.55	CYP2D6	Moderately soluble	0	0	Good	Good
4.	90454	High	0.55	CYP2D6	Moderately soluble	0	0	Good	Good
5.	54991	High	0.55	CYP2D6 & CYP3A4	Good	0	0	Good	Good
6.	72276	High	0.55	None	Good	1	1	Good	Good
7.	10263440	High	0.55	None	Good	0	0	Good	Good
8.	10448938	High	0.55	CYP2D6	Moderately soluble	0	0	Good	Good
9.	189061	High	0.55	None	Good	0	0	Good	Good
10.	441074	High	0.55	CYP2D6	Moderately soluble	0	0	Good	Good
11.	3034034	High	0.55	CYP2D6	Moderately soluble	0	0	Good	Good
12.	101792719	High	0.55	None	Good	0	0	Good	Good
13.	14633011	High	0.55	None	Good	0	0	Good	Good

Table 2. The key features of active constituents

S. No	Compound	Michael acceptor	Leadlikeness	Synthetic Accessibility
1.	Apoquinine	No	Yes	4.25
2.	Catechin	No	Yes	3.50
3.	Cinchonidine	No	Yes	4.18
4.	Cinchonine	No	Yes	4.18
5.	Cuprediene	No	No (1 violation) MW>350	4.47
6.	Epicatechin	No	Yes	3.50
7.	Epiprocurcumenol	Yes	No (1 violation) MW<250	4.04
8.	Epiquinine	No	Yes	4.34
9.	Procurcumenol	Yes	No (1 violation)MW<250	4.04
10.	Quinidine	No	Yes	4.34
11.	Quinine	No	Yes	4.34
12.	Zedoaronediol	Yes	Yes	4.13
13.	Procurcumadiol	Yes	Yes	4.30

SARS-CoV-2 protease was carried out. In comparison to N3, the five best scoring chemical constituents were apoquinine from cinchona with highest binding affinity of -7.5 kcal/mol, epicatechin from neem with -7.0 kcal/mol, catechin from neem with -6.8 kcal/mol, procurcumenol from turmeric and quinidine from cinchona both having similar score of -6.6 kcal/mol (Table 3).

3.3 Molecular Docking

The constituents that showed higher binding affinity towards the protein were docked with SARS-CoV-2 main protease to estimate the binding energies. Molecular docking is a tool that predicts the conformation of a ligand in the active site of the receptor. SARS-CoV-2 M^{Pro} is a dimer which consists of protomer A and protomer B. Each protomer is made up of three domains, Domain I (8-102), Domain II (102-184) which are antiparallel β -barrel structures and Domain III (201-303) which is a five α helix structure arranged in antiparallel structure. The substrate binding site of COVID-19 virus M^{Pro} is located in a cleft between Domain I (residues 8-101) and II (residues 102-184) and it has a Cys-His catalytic dyad. The binding pocket identified within the residues 189-191 of the long strand and was selected as the active site of the protein.

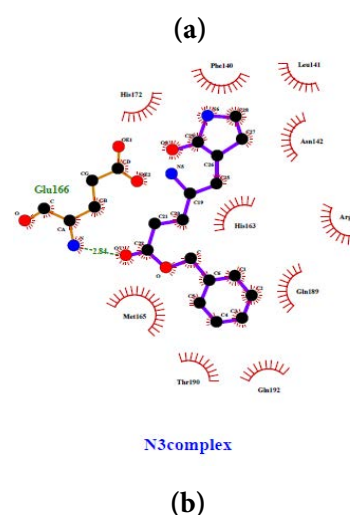
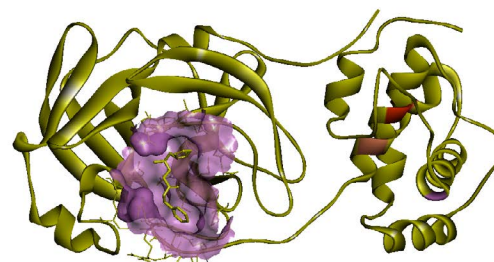


Figure 2. (a) SARS-CoV-2 main protease is represented in yellow. The N3 ligand docked in 6LU7 is shown in the hydrophobic pocket (b) 2D interaction maps of N3 inhibitor complex with pocket amino acids.

Table 3. Binding energy, affinity and key amino acid interactions calculations using AutoDock Vina

S.No	Compound	Binding Affinity (kcal/mol)	Binding Energy (kcal/mol)	Hydrogen Bonds Formed
1.	Apoquinine	-7.5	-7.7	GLU166
2.	Catechin	-6.8	-8.38	GLU166,THR190,GLN192,ASP187
3.	Cinchonidine	-6.5	-7.59	GLN189
4.	Cinchonine	-6.6	-7.49	GLU166
5.	Cuprediene	-6.5	-7.31	No Hydrogen Bonds are formed
6.	Epicatechin	-7	-7.29	ASP187,GLN189,GLN192
7.	Epiprocurcumenol	-6.2	-6.83	GLU166
8.	Epiquinine	-6.4	-6.83	GLU166
9.	Procurcumenol	-6.6	-7.11	GLU166,HIS163
10.	Quinidine	-6.6	-7.26	GLN189
11.	Quinine	-6.1	-7.48	GLN189
12.	Zedoaronediol	-6.1	-6.56	GLU166,GLN189
13.	Procurcumadiol	-5.8	-6.79	GLU166,GLU189
14.	N3	-7.04	-7.04	PHE140,GLU166,HIS163

The binding energies estimated for the filtered chemical constituents were in the range of -6.56 to -8.38 kcal/mol. The specific interactions of chemical constituent with M^{Pro} is depicted in Figure 2 in comparison to the standard. It was identified from the literature sources that Glutamate at P1 position is a major requirement and seven active constituents showed interactions with GLU166 except cinchonidine, epicatechin, quinine and quinidine. The other major interactive amino acids surrounding the docked complex include HIS163, HIS164, LYS145, PRO168, ARG188, GLN189, GLN192, CYS145, LEU141, and THR190.

Catechin showed the highest binding energy of -8.38kcal/mol. There were significant interactions with the receptor binding site including GLN166, LEU167, PRO168, HIS164, MET165, and CYS145 (Figure 3). It was found to form four prominent hydrogen bonds with THR190, GLN192, GLU166, and ASP187. The five hydroxyl groups in catechin are freely available to form multiple hydrogen bonds. It is further evident from the previous studies that the co-crystallised ligand N3 was found to show similar interactions. The effectiveness of catechin as an antiviral agent could be tested against COVID-19. But it should also be noted that many polyphenolic compounds are metabolized into their conjugated forms and accumulation of metabolized compounds are different among various tissues.

Epicatechin is an isomer of catechin with *cis*-configuration and forms five hydrogen bond with GLN192, THR190, GLN 189, HIS 164 and ASP187.

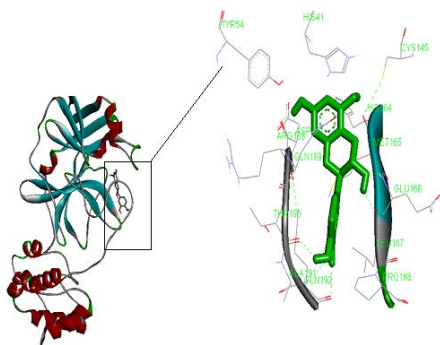


Figure 3. Docking pose of the constituent with highest docking energy Catechin and its interactions.

It had shown promising effects in the treatment of Mayaro fever caused due to the outbreak of Mayaro virus in Brazil. Epicatechin, a folklore medicine showed promising antiviral activity against hepatitis C virus, Mayaro virus and it was also observed that it had the ability to inhibit the viral replication in lower doses than the cytotoxic dose. A lower binding energy of -7.29 kcal/mol further confirms a better binding towards the substrate and this could be considered as an potential constituent in the treatment of COVID-19.

Analyzing the interactions of procurcumenol from turmeric, showed three major hydrogen bonds with the amino acids HIS163 and GLU166. The previous research studies on curcumin has shown better antiviral activities against hepatitis C virus and used in treatment of jaundice and other liver diseases. Liver impairment is seen as an arising concern of patients with COVID-19 due to the administration of high doses of potentially hepatotoxic antivirals, antibiotics and steroids. Zedoaronediol and procurcumadiol from curcumin showed three hydrogen bonding, of which two are with GLU166 and one with GLU189 with a binding energy of -6.56kcal/mol and -6.79kcal/mol respectively. The physicochemical properties of the thirteen active constituent are provided in Table 4. The toxicity of the active constituents along with its HOMO, LUMO calculation are reported in Tables 5 and 6 respectively.

4. Conclusion

All the thirteen chemical constituents showed better binding affinity than the standard. From the docking study conducted, it could be understood that in most of the constituents the interaction of the ligand with the receptor through formation of hydrogen bond with aminoacid GLU166 marks its step in substrate recognition thereby necessitating the inhibitory mechanism. Epicatechin and apoquinine showed highest binding affinity of -7 and -7.5kcal/mol while catechin and epicatechin showed four hydrogen bond interactions leading to locking of the inhibitor in the binding pocket in a better way. It is interesting and worth noticing the interaction of GLU166 residue with the ligand in most of the constituents. Furthermore, studies on the dynamics of the best scoring constituent

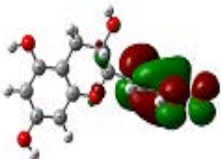
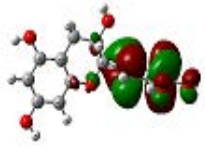
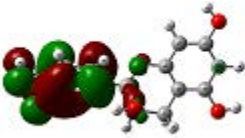
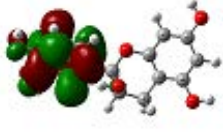
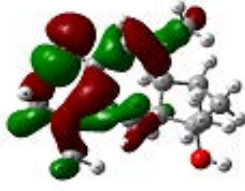
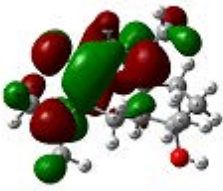


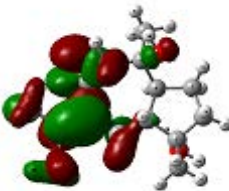
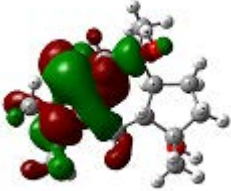
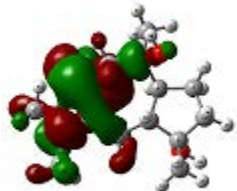
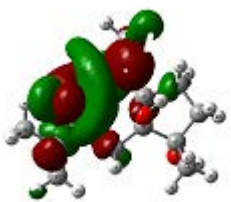
Table 4. The physicochemical properties of the best thirteen ligands

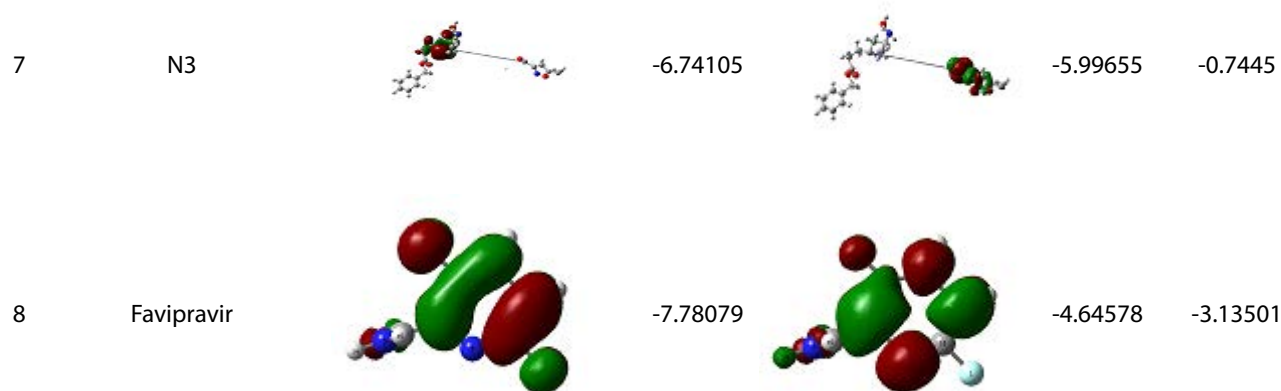
S.No	PUBCHEM ID	Physicochemical Properties MW (g/mol)	LogP (o/w)	H-Acc	H-Bond	TPSA
1.	101600159	310.39	2.34	4	2	56.59
2.	9064	290.27	0.85	6	5	110.38
3.	101744	294.39	2.78	3	1	36.36
4.	90454	294.39	2.78	3	1	36.36
5.	54991	352.43	2.77	5	1	62.66
6.	72276	290.27	0.85	6	5	110.38
7.	10263440	234.33	2.71	2	1	37.30
8.	10448938	324.42	2.81	4	1	45.59
9.	189061	234.33	2.71	2	1	37.30
10.	441074	324.42	2.81	4	1	45.59
11.	3034034	324.42	2.81	4	1	45.59
12.	101792719	252.35	2.09	3	2	57.53
13.	14633011	250.33	1.98	3	2	57.53

Table 5. The mutagenicity and carcinogenicity prediction of the best thirteen ligands

S.No	PUBCHEM ID	Negative for genotoxic carcinogenicity	Negative for nongenotoxic carcinogenicity	Potential <i>Salmonella typhimurium</i> TA100 mutagen based on QSAR	Potential carcinogen based on QSAR
1.	101600159	Yes	Yes	No	No
2.	9064	Yes	Yes	No	No
3.	101744	Yes	Yes	No	No
4.	90454	Yes	Yes	No	No
5.	54991	Yes	Yes	No	No
6.	72276	Yes	Yes	No	No
7.	10263440	No	Yes	No	No
8.	10448938	Yes	Yes	No	No
9.	189061	No	Yes	No	No
10.	441074	Yes	Yes	No	No
11.	3034034	Yes	Yes	No	No
12.	101792719	No	Yes	No	No
13.	14633011	No	Yes	No	No

Table 6. E_{HOMO} and E_{LUMO} of best six chemical constituents and two standard ligands

S.No	Molecule	HOMO	E HOMO	LUMO	E LUMO	Energy gap (eV)
1	Catechin		-8.26652		-4.26286	-4.00366
2	Epicatechin		-8.23468		-4.23489	-3.99979
3	Epiprocurcumenol		-9.42708		-5.52443	-3.90265
4	Procurcumenol		-9.35143		-5.50457	-3.84686
5	Zedoaronediol		-9.22898		-4.77068	-4.4583
6	Procurcurmadiol		-9.16177		-5.32416	-3.83761



would help to understand the stability of the constituent in the binding site. The effectiveness of catechin and epicatechin as an antiviral agent could be tested against COVID-19.

5. References

- WHO. Naming the coronavirus disease (COVID-19) and the virus that causes it [Internet]. [cited 2020 Apr 8]. Available from: [https://www.who.int/emergencies/diseases/novel-coronavirus-2019/technical-guidance/naming-the-coronavirus-disease-\(covid-2019\)-and-the-virus-that-causes-it](https://www.who.int/emergencies/diseases/novel-coronavirus-2019/technical-guidance/naming-the-coronavirus-disease-(covid-2019)-and-the-virus-that-causes-it).
- Liu RP, Ge J De, Zhong Y, Zheng Q, Sun R. Traditional Chinese medicine for treatment of COVID-19 based on literature mining of targeting cytokine storm. *Chinese Traditional and Herbal Drugs*. 2020; 51(5):1096–105.
- Li Y, Liu X, Guo L, Li J, Zhong D, Zhang Y, Clarke M, Jin R. Traditional Chinese herbal medicine for treating novel coronavirus (COVID-19) pneumonia: Protocol for a systematic review and meta-Analysis. *Systematic Reviews*. 2020; 9(1). <https://doi.org/10.1186/s13643-020-01343-4>. PMID:32268923 PMCid:PMC7138957
- Lerner K. SARS, MERS, and the Emergence of coronaviruses. *Worldmark Global Health and Medicine Issues* [Internet]. 2016. Available from:
- Yan R, Zhang Y, Li Y, Xia L, Guo Y, Zhou Q. Structural basis for the recognition of SARS-CoV-2 by full-length human ACE2. *Science*. 2020; 367(6485):1444–8. <https://doi.org/10.1126/science.abb2762>. PMID:32132184. PMCid:PMC7164635.
- Ludwig S, Zarbock A. Coronaviruses and SARS-CoV-2: A Brief Overview. *Anesthesia and Analgesia*. 2020. <https://doi.org/10.1213/ANE.0000000000004845>. PMID:32243297. PMCid:PMC7173023
- Wang X, Xu W, Hu G, Xia S, Sun Z, Liu Z, et al. SARS-CoV-2 infects T lymphocytes through its spike protein-mediated membrane fusion. *Cellular and Molecular Immunology*. 2020. <https://doi.org/10.1038/s41423-020-0424-9>
- Wang Q, Zhang Y, Wu L, Niu S, Song C, Zhang Z, et al. Structural and functional basis of SARS-CoV-2 entry by using human ACE2. *Cell*. 2020. <https://doi.org/10.1016/j.cell.2020.03.045>. PMID:32275855 PMCid:PMC7144619
- Kaladhar SVGKD. Effects of drugs on spike glycoprotein of sars-cov 2 in control of covid-2019. *International Journal of Advanced Research*. 2020; 8(3):918–24. <https://doi.org/10.21474/IJAR01/10706>
- Law S, Leung AW, Xu C. Severe Acute Respiratory Syndrome (SARS) and Coronavirus disease-2019 (COVID-19): From causes to preventions in Hong Kong. *International Journal of Infectious Diseases*. 2020; 94:156–63. <https://doi.org/10.1016/j.ijid.2020.03.059>. PMID:32251790 PMCid:PMC7195109
- Giron CC, Laaksonen A, Silva FLB da. *pbioRxiv*. 2020;2020.04.05.026377.
- Rut W, Groborz K, Zhang L, Sun X, Zmudzinski M, Hilgenfeld R, et al. Substrate specificity profiling of SARS-CoV-2 Mpro protease provides basis for anti-COVID-19 drug design. *bioRxiv*. 2020. <https://doi.org/10.1101/2020.03.07.981928>. PMID:32014497
- Rabaan AA, Al-ahmed SH, Sah R, Tiwari R, Iqbal M, Patel SK, et al. SARS-CoV-2 / COVID-19 and advances in developing potential therapeutics and vaccines to counter this emerging pandemic virus - A Review.

- Preprints. 2020; 4:1–46. <https://doi.org/10.20944/preprints202004.0075.v1>
14. Bharath EN, Manjula SN, Vijaychand A. In silico drug design-tool for overcoming the innovation deficit in the drug discovery process. *International Journal of Pharmacy and Pharmaceutical Sciences*. 2011; 3(2):8–12.
 15. Ang L, Lee HW, Choi JY, Zhang J, Soo Lee M. Herbal medicine and pattern identification for treating COVID-19: A rapid review of guidelines. *Integrative Medicine Research*. 2020; 9(2):100407. <https://doi.org/10.1016/j.imr.2020.100407>. PMID:32289016 PMCid:PMC7104236
 16. Ferner RE, Aronson JK. Chloroquine and hydroxychloroquine in covid-19. *The BMJ*. 2020; 369. <https://doi.org/10.1136/bmj.m1432>. PMID:32269046
 17. Atallah P, Wagener KB, Schulz MD. ADMET: The future revealed. *Macromolecules*. 2013. <https://doi.org/10.1002/chin.201336195>
 18. López-López E, Naveja JJ, Medina-Franco JL. DataWarrior: An evaluation of the open-source drug discovery tool. *Expert Opinion on Drug Discovery*. 2019. <https://doi.org/10.1080/17460441.2019.1581170>. PMID:30806519
 19. Trott O, Olson AJ. Autodock vina: Improving the speed and accuracy of docking. *Journal of Computational Chemistry*. 2019; 31(2):455–61.
 20. Lindstrom W, Morris GM, Weber C, Huey R. Using AutoDock for virtual screening. The Scripps Research Institute [Internet]. 2006. Available from:
 21. Lin LT, Hsu WC, Lin CC. Antiviral natural products and herbal medicines. *Journal of Traditional and Complementary Medicine*. 2014; 4(1):24–35. <https://doi.org/10.4103/2225-4110.124335>. PMID:24872930 PMCid:PMC4032839
 22. Jin Z, Du X, Xu Y, Deng Y, Liu M, Zhao Y, et al. Structure of Mpro from COVID-19 virus and discovery of its inhibitors. *Nature*. 2020. <https://doi.org/10.1101/2020.02.26.964882>
 23. Lohidashan K, Rajan M, Ganesh A, Paul M, Jerin J. Pass and Swiss ADME collaborated in silico docking approach to the synthesis of certain pyrazoline spacer compounds for dihydrofolate reductase inhibition and antimalarial activity. *Bangladesh Journal of Pharmacology*. 2018; 13(1):23–9. <https://doi.org/10.3329/bjp.v13i1.33625>
 24. Contrera JF. Validation of Toxtree and SciQSAR in silico predictive software using a publicly available benchmark mutagenicity database and their applicability for the qualification of impurities in pharmaceuticals. *Regulatory Toxicology and Pharmacology*. 2013; 67(2):285–93. <https://doi.org/10.1016/j.yrtph.2013.08.008>. PMID:23969001