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Molecular docking and ADMET study of bioactive compounds of *Glycyrrhiza glabra* against main protease of SARS-CoV2

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**ABSTRACT**

Recent pandemic situation of COVID-19 is caused due to SARS-CoV2 and almost all the countries of the world have been affected by this highly contagious virus. Main protease (Mpro) of this virus is a highly attractive drug target among various other enzymes due to its ability to process poly-protein that is the translated product of the SARS-CoV2 RNA. The present study demonstrates molecular docking study of *Glycyrrhiza glabra* (Gg) active compounds such as Glycyrrhizic acid (GA), Liquiritigenin (L) and Glabridin (G) against the Mpro. Docking studies shows that these active compounds bind strongly with some of the amino acid residues in the active site of Mpro and inhibits the enzyme strongly. GA, L, and G are proposed to be strong inhibitors of the enzyme and the amino acids: His 41, Gly 143, Gln 189, Glu 166, Cys 145, Thr 25, Asn 142, Met 49, Cys 44, Thr 45 and pro 168 present in the active site of Mpro were shown to make non-covalent interaction with these compounds. In silico ADMET properties prediction also shows that Gg active compounds had good solubility, absorption, permeation, non-toxic, and non-carcinogenic characteristics. Our finding concludes that all of the three active compounds of Gg have the potential to be strong inhibitors for Mpro of SARS-CoV2 but glycyrrhizic acid has a high binding affinity and a good ADMET properties than the other two.

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Selection and peer-review under responsibility of the scientific committee of the National Conference on Functional Materials: Emerging Technologies and Applications in Materials Science.

**1. Introduction**

Wuhan, Hubei province of China was the very first location where the severe acute respiratory syndrome coronavirus2 (SARS-CoV2) outbreak recorded [1]. From Wuhan city of China this virus had spread across more than 200 countries with America and Brazil as the epicentre as of now. COVID-19 caused by SARS-CoV2 has affected 8, 242, 999 people worldwide and killed as many as 445, 535 people, global data as of 18th June 2020, 4.57 pm CEST. (WHO coronavirus disease Dashboard), World Health Organization (WHO) has stated the COVID-19, a public health emergency of international concern on January 30, 2020 [2]. SARS-CoV-3, MERS-CoV [4] and SARS-CoV2 [5] are the three categories of coronavirus that causes the deadly pneumonia in humans. Severe acute respiratory syndrome-2 (SARS-CoV2) is an enveloped, positive-sense single-strand RNA placed in a beta coronavirus family of Coronaviridae [6]. As compared to the other two, SARS-CoV2 is highly contagious and spread the infection more quickly and severely, leading to large number of deaths and thus arising a pandemic situation [7,8].

The SARS-CoV2 has a genome ranging from 29 to 30 kb in length [9]. The recently proposed drugs targets the main protease (Mpro also known as 3CLpro) which is one of the best targets among the other enzymes in the virus [10]. This enzyme participates in the processing of polyprotein that is translated from the RNA of the same virus [11]. These polyprotein are processed and converted into mature non-structural proteins using two proteases PLpro and CLpro encoded by ORF1 (open reading frame) [12]. A non-structural protein of coronavirus plays a significant role in replication and transcription during the infection [13]. Main protease operates at more than 10 cleavage sites on the polyprotein 1ab which is nearly 790 kDa [14]. Mpro is a homodimer having two identical chains and the recognition sequence is Leu-Gln ↓ Ser, Ala, Gly and cleaved through the Cys-His dyad protease in which -SH group of cysteine serve as a nucleophile in the proteolysis [15]. Inhibiting the main protease activity would block transcription or replication. The amino acid residues of the active site
of M\textsuperscript{pro} participating in substrate binding includes Cys145 –His41 dyad, and Phe140, Thr45, Arg188, Asp187, Met49, Asn142, Met165, Gln189, His172, Glu166 \[16\]. An enormous amount of active components of medicinal plants effective against the viruses. Among these medicinal plant, \textit{Glycyrrhiza glabra} is the one plant, its active compounds may have the potential to act against various viruses \[17\]. \textit{Glycyrrhiza glabra} (Gg) commonly called ‘sweet wood’ and ‘liquorice’ belongs to the family Fabaceae. It has been effective against many RNA viruses like H5N1 virus, Hepatitis C virus, influenza A virus, Newcastle disease virus, H1N1 virus, Rotavirus, severe acute respiratory virus etc. and DNA viruses such as Varicella Zoster virus, Epstein-Barr virus and Herpes Simplex virus etc. \[18\].

Several active compounds from Gg have been reported that inhibit the virus’s growth. Some studies showed that Glycyrrhizin (glycyrrhizinate; glycyrrhizic acid) inhibit the binding of the virus to the target cells, and also beneficial in controlling the viral replication and found to have an important antiviral activity. Studies on clinically isolates of severe acute respiratory syndrome Virus [FFM-1 and FFM-2] gave important information about the anti-viral property of glycyrrhizic acid \[19,20\]. Another active compound, Glabridin (G) isolated from Gg roots. It has been used as a traditional medicine in Asia and has anti-inflammatory, anti-osteoporotic, anti-nephritic, anti-atherogenic, estrogenic, and anti-oxidant, regulation of metabolism, neuro-protective and skin-whitening properties. This is also reported to have vasodilatation property and used for the treatment of many diseases like bacterial, bronchitis infection \[27,28\]. Liquiritigenin (L) has anti-cancer, anti-oxidative, antimicrobial activities, hepatoprotective, immune regulatory, and cardio-protective effects and also has anti-platelet aggregation, anti-tumorigenic, anti-angiogenic, anti-allergic properties \[29\].

For the virtual screening and structure-based drug designing, molecular docking is commonly used in bioinformatics based research and is accustomed to predict non-covalent interaction (mainly hydrogen bonding) of macromolecule (receptor) with the ligand (drug molecule). AutoDock is a suited program for docking and virtual screening which calculates the grid internally (automatically), for the atom types that are required. A more improved version, AutoDock Vina achieves within the average accuracy of the binding pose prediction and bring order of magnitude faster than previously developed AutoDock4 \[30\].

OSIRIS Property Explorer is employed to determine drug likeness. It provides drug relevant information whenever a structure is valid and underlines various properties with a high risk of inadmissible effects such as poor intestinal absorption, mutagenicity or drug conform behaviour through different customization characteristics. This also determines the clog P (Logarithm of compounds partition coefficient between water and n-Octanol) which determines the hydrophilicity of the compounds. Larger the clogP value, lower the hydrophilicity and thus poor the permeation and absorption. Log S suggest solubility; lower the log S value, higher the solubility which enhances the sorption. The Topological polar surface area (TPSA) suggest the surfaces that belong to the polar atoms and molecules in the compound. Larger TPSA value is related with the least permeability of the membrane. The compounds that have larger TPSA value will be a better substrate for p-glycoprotein which is liable for efflux of drug from the cell and thus reduced TPSA was beneficial for drug-like property. Some studies also anticipated that a molecule having a better penetration property through CNS should have lesser value of TPSA \[26\].

A good drug candidate can be consumed in the desired time as well as distributed through the entire body for its efficient action and metabolism. Another factor is toxicity which is often very vital and dominates the Absorption, Distribution, Metabolism, Excretion behaviour of the drugs. Some studies have shown that drug failed at clinical trials due to its adverse side effect and their toxicity which has been proven to be very detrimental and expensive in the drug development. In silico, ADMET and drug-likeness prediction help in the discovery of new targets and compounds with anticipated biological activities \[31\]. Human intestinal absorption [HIA] is an important ADMET properties. The exploitation of drugs in the body is such a complex process that can hardly be analysed by statistical models. HIA is one of the important steps during compounds (drug) transporting to their desired target \[31\]. Blood-brain barrier (BBB) is used to predict various features of the CNS vasculature. CNS vessel does not have pores in the endothelial cell and have various properties that regulate the transport of ions, molecules, and cells which make CNS vessel very prohibitive in nature and provides an interference for the delivery of compounds into the central nervous system \[32\]. Carcinogenicity means the ability of the substance to cause cancer. P-glycoprotein substrate is a compound that utilizes the P-glycoprotein transporter for several exercises like excretion and absorption of drugs and other vital exercises \[33\]. These properties are often used for the study of ADMET behaviour of the drugs.

The main objective of the present study is to carry out molecular docking analysis of \textit{Glycyrrhiza glabra} active compounds, Glycyrrhizic acid, Liquiritigenin and Glabridin against the main protease (M\textsuperscript{pro}) one by one followed by molecular interaction study (hydrogen bond prediction between target and drugs), drug-likeness behaviour and ADMET prediction to confirm the efficiency and efficacy of these active compound against SARS-CoV2.

2. Methods and materials

2.1. Selection of ligand

Glycyrrhizic acid (PubChem CID: 14982), Liquiritigenin (PubChem CID: 114829), and Glabridin (PubChem CID: 124052) were recognized as potential SARS-CoV inhibitors from literature survey and maybe also effective against SARS-CoV2. The structure of Liquiritigenin and Glabridin, and Glycyrrhizic acid were downloaded from PubChem (http://pubchem.ncbi.nlm.nih.gov) two control drug that is lopinavir (PubChem CID: 92727) and rotinavir (PubChem CID: 39622) also downloaded and converted their file format from SDF file to PDB file [Supplementary Fig. 1 (a) – (c)].

2.2. Selection of target molecule (Receptor/M\textsuperscript{pro}): Active site and sequence analysis

The crystal structure of the main protease (PDB ID- 6Y2E) downloaded from PDB database https://www.rcsb.org/structure/6Y2E, DOI: https://doi.org//10.2210/pdb6Y2E/pdb [25] and saved as PDB format. Its functional domain and the number and name of amino acid residues present in the pocket of the active site of 6Y2E were verified using ScanProsite (ExPaSy) (http://prosite.expas.org) (supplementary figure s4) \[21\] and CDD webserver was used to determine the conserved sequence of 6Y2E \[22\]. Sequence analysis of the main protease of SARS-CoV2 was retrieved using ProtParam, ProtScale (http://web.expasy.org) \[23\].

2.3. Preparation of ligands (\textit{Glycyrrhiza glabra} active compounds) and main protease (M\textsuperscript{pro})

The Ligand was prepared by adjusting ionization, torsion, degree of freedom and stereo-chemical variation using AutoDock MG Tool. Energy of the main protease was minimized based on GROMAS6 4381 algorithm using Swiss Model Viewer, water was removed and Polar hydrogens were added to the main protease
of SARS-CoV2 followed by calculation of gasteiger Charges using the AutoDock MG Tool and saved in PDBQT file format. The receptor grid was developed using AutoDock Grid tool with grid dimensions of 30Å × 30Å × 30Å with 0.375 Å spacing and the grid box was set at -15.692, -32.457, 2.498 in X, Y and Z dimension [24].

2.4. Molecular docking of Glycyrrhiza glabra (Gg) active compounds with Mpro

The crystal structure main protease (Mpro) downloaded from RCSB Protein Data Bank (PDB ID: 6Y2E) with a resolution of 1.7 Å and R-value free of 0.222 and R-value work of 0.171. Molecular docking was performed using in silico docking software, AutoDock Vina [30] with the three active compounds of Gg, glycyrrhizic acid, liquiritigenin and glabridin one by one against 6Y2E. The docking was done using command Prompt and the other prerequisite conditions before the docking like the ligand and enzyme preparation has been established. PyMol and BIOVIA Discovery Studio Visualizer was used to visualize the binding interaction between the active compounds of Glycyrrhiza glabra with 3D structure of Mpro (6Y2E) of SARS-CoV2 [25] and Simulation was done with CHARMM36 force field in Discovery Studio visualisation tool.

2.5. Drug likeness prediction

OSIRIS Property Explorer (http://www.organic-chemistry.org/ prog/peo/) [26] is employed to determine likeness of drugs. Properties like Log S calculation, TPSA, Clog P calculation, molecular mass, and drug-likeness based on fragment and drug score of all the three active compounds have been established.

2.6. ADMET properties prediction

ADMET refers to Absorption, Distribution, Metabolism, Excretion, and Toxicity. It contains the pharmacokinetic profile of a compound (drug molecule) and plays a significant role in determining its pharmacodynamics activities. Taking all the three active compounds into account, properties like bioavailability, brain penetration, oral absorption, carcinogenicity, and other human intestinal absorption properties of the active compounds have been deter-

Fig 1. Sequence alignment using CDD webserver represent His41 and Cys145 as highly conserved amino acids (denoted by #) in all coronaviruses and actively participated in cleavage of polyprotein of SARS-CoV2 into structural proteins.
minded using admetSAR webserver (http://lmmde.ecust.edu.cn: 8000/) [31–34].

3. Results and discussion

3.1. Sequence analysis and active site identification

Sequence analysis of SARS-CoV2 M\textsuperscript{pro} revealed a theoretical PI value of 5.95 with an extinction coefficient of 33640 M\textsuperscript{-1} cm\textsuperscript{-1} and absorbance (1 g/L) computed 0.995. The Aliphatic index of the M\textsuperscript{pro} was 82.12 and shows a stability index of 27.65. All these result shows that SARS-CoV2 M\textsuperscript{pro} is a very stable macromolecule. The secondary structure of SARS-CoV2 main protease revealed that the alpha helix and the beta strand were maximally present (supplementary table 1). The ExPASy Prosite and CDD webserver result shows that His\textsuperscript{41} and Cys\textsuperscript{145} has a vital role in the catalytic activity of the main protease of SARS-CoV2 and also has been conserved in all the different types of coronaviruses as depicted in Fig. 1.

3.2. Molecular docking analysis of control drugs with the M\textsuperscript{pro} of the SARS-CoV2

The molecular docked pose of the minimum binding affinity conformer of the Lopinavir and Ritonavir (control drugs) demonstrated that they bind to the active site of the M\textsuperscript{pro} protein of the SARS-CoV2. Fig. 2 showed that Lopinavir binds through two conventional H-bond with Asn\textsuperscript{142}, two carbon H-bond with Thr\textsuperscript{25} and Gln\textsuperscript{189}, and two alkyl bond with His\textsuperscript{41} respectively. However, Ritonavir also bind firmly to the active site of the main protease and it stabilized the M\textsuperscript{pro} active site through four hydrogen Bond in which two are conventional hydrogen bond and rest two are carbon hydrogen bond. Asn\textsuperscript{142} and Thr\textsuperscript{25} amino acid residue formed conventional and carbon hydrogen bond respectively. Whereas Cys\textsuperscript{44} and Cys\textsuperscript{145} formed alkyl bond and His\textsuperscript{41} and Pro\textsuperscript{168} shows Pi-alkyl interaction, besides other residues also found to be bound to control drugs through Vander Waals interaction depicted in 2D (two dimensional) plot (Fig. 3). The binding affinity energy of the control drugs are mentioned in table 1. If we compare the binding affinity energy of control drugs, Lopinavir and Ritonavir, we observed that Lopinavir have highest negative binding affinity with −7.6 Kcal/mol followed by Ritonavir with −7.3 Kcal/mol [35].

3.3. Molecular docking analysis of Gg active compounds with the M\textsuperscript{pro} of the SARS-CoV2

The molecular docked pose of the minimum binding affinity conformer of the Gg active compounds that is glycyrrhizic acid, Glabridin and Liquiritigenin demonstrated that they also firmly goes and bind to the active site of the M\textsuperscript{pro} protein of the SARS-CoV-2 (Fig. 4 and table 2). Glycyrizic acid (GA) binds within the M\textsuperscript{pro} active site through several non-covalent interactions. GA binds active site through one conventional H-bond with Glu\textsuperscript{166} and three carbon H-bond with Ser\textsuperscript{46}, Pro\textsuperscript{168} and Asn\textsuperscript{142}. GA also found to be bound to active site through alkyl and Pi-alkyl bond with Cys\textsuperscript{145}, Leu\textsuperscript{27} and His\textsuperscript{41} respectively (Fig. 7). Glabridin stabilizes the M\textsuperscript{pro} active site through hydrogen bond with His\textsuperscript{41} and Glu\textsuperscript{166} and through alkyl and Pi-alkyl with Cys\textsuperscript{145} and Pro\textsuperscript{168} respectively and play an important role in the stabilisation of the active site of the main protease (Fig. 6). Likewise Liquiritigenin also binds to the active site and formed several non-covalent interaction in which Thr\textsuperscript{25} and Thr\textsuperscript{45} are found to bind with the Liquiritigenin through hydrogen bond and, Met\textsuperscript{49}, and Cys\textsuperscript{145} formed Pi-sulfur interaction to Liquiritigenin L also firmly binds to the His\textsuperscript{41} through Pi-Alkyl that stabilize the active site (Fig. 5). On comparing the binding affinity of the control drugs with the binding affinity of Gg active compounds, we found that GA have the highest negative binding affinity energy than Lopinavir and ritonavir with −8.0 Kcal/mol, G have lower binding affinity than Lopinavir but have similar binding affinity with ritonavir with −7.3 Kcal/mol whereas L have lowest binding affinity among all ligands [36].

3.4. Drug likeness prediction

The result of the OSIRIS Property Explorer (http://www.organicchemistry.org/prog/peo/) shown in table 3. Out of the three active compounds, GA shows a very low Clog P than G and L and thus GA
Table 1: Molecular Docking analysis: Binding energy and the type of molecular interaction of control drugs against M\textsuperscript{pro}. UNK: Glycyrrhiza glabra active compounds.

| Control Drugs | Binding Affinity (Kcal/mol) | Donor Atom      | Distance | Acceptor Atom   | Category          | Types of interaction |
|---------------|-----------------------------|-----------------|----------|-----------------|--------------------|----------------------|
| Lopinavir     | 7.6                         | ASN142:HD22     | 2.39     | UNK: O          | Hydrogen Bond      | Conventional H-bond  |
|               |                             | UNK:H           | 1.75     | ASN142:OD1      | Hydrogen Bond      | Conventional H-bond  |
|               |                             | UNK:H           | 2.63     | CYS44           | Hydrogen Bond      | Conventional H-bond  |
|               |                             | UNK              | 1.72     | THR25:OG1       | Hydrogen Bond      | Conventional H-bond  |
|               |                             | UNK              | 3.98     | PRO168          | Hydrophobic Pi-Pi  | Stacked              |
|               |                             | UNK              | 4.98     | CYS145          | Hydrophobic Pi-Alkyl |                      |
| Ritonavir     | 7.3                         | ASN142:HD22     | 2.63     | UNK: O          | Hydrogen Bond      | Conventional H-bond  |
|               |                             | UNK:H1          | 2.43     | ASN142:OD1      | Hydrogen Bond      | Conventional H-bond  |
|               |                             | UNK:H13         | 2.61     | THR25:OG1       | Hydrogen Bond      | Conventional H-bond  |
|               |                             | UNK:H34         | 2.23     | GLN189:OE1      | Hydrogen Bond      | Carbon H-bond        |
|               |                             | UNK:C12         | 5.14     | LEU27           | Hydrophobic Alkyl  |                      |
|               |                             | UNK:C12         | 3.87     | CYS145          | Hydrophobic Alkyl  |                      |
|               |                             | UNK              | 4.55     | UNK:C12         | Hydrophobic Pi-Alkyl |                  |
|               |                             | UNK              | 5.33     | PRO168          | Hydrophobic Pi-Alkyl |                  |

Fig. 3. 3-D (three dimensional) diagram based on hydrogen bond donor and acceptor characteristics of amino acid residues (A) and 2-D (two dimensional) (B) diagram showing several interaction between M\textsuperscript{pro} and Ritonavir. PRO: Proline, THR: Threonine, LEU: Leucine, HIS: Histidine, ASN: Asparagine.

Fig 4. Three dimensional structure of main Protease based on the hydrophobicity of amino acid residues and docked compounds are shown in black.

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has high hydrophilicity and can be absorbed easily by the cell of the SARS-CoV2 thus showing drug efficacy but in terms of solubility, GA is least soluble in blood than the other two. L shows maximum solubility in the blood. TPSA of GA is also higher as compared with the other two compounds which tell the efficiency of the compounds in terms of membrane permeability. Here again GA has more potential to cross the lipid bilayer of the virus and thus can strongly bind and inhibits the enzyme (M<sub>pro</sub>). Our predicted values lie well within the drug-likeness.

3.5. ADMET prediction

ADMET (Absorption, Distribution, Metabolism, Excretion and Toxicity) properties were retrieved using admetSAR server, which shows that all Glycyrrhiza glabra active compounds i.e. Glycyrrhizic acid (GA) (Accession Number: DB13751), Liquiritigenin (L) (PubChem CID: 114829) and Glabridin (G) (PubChem CID: 124052) are having better human intestinal absorption score (table 4).

Table 2
Molecular Docking analysis: Binding energy and the type of hydrogen bond interaction of all the three active compounds of Gg against M<sub>pro</sub>, UNK: Glycyrrhiza glabra active compounds.

| Gg active compounds | Binding Affinity (Kcal/mol) | Donor Atom | Distance | Acceptor Atom | Category | Types of interaction |
|---------------------|-----------------------------|------------|----------|---------------|----------|---------------------|
| G                   | −7.3                        | HIS41:HE2  | 2.37     | UNK:O1        | Hydrogen Bond | Conventional H-bond |
|                     |                             | UNK:H     | 1.84     | GLU166:O      | Hydrogen Bond | Conventional H-bond |
|                     |                             | MET49:SD  | 5.46     | UNK:O2        | Hydrophobic  | Alkyl               |
|                     |                             | CYS145    | 4.43     | CYS145:O      | Hydrophobic  | Alkyl               |
|                     |                             | UNK:C12   | 3.99     | CYS145:O      | Hydrophobic  | Alkyl               |
|                     |                             | HIS41     | 5.34     | UNK:O2        | Hydrophobic  | Alkyl               |
|                     |                             | HIS41     | 5.32     | UNK:C12       | Hydrophobic  | Alkyl               |
|                     |                             | UNK       | 5.2      | PRO168:O      | Hydrophobic  | Alkyl               |

| L                   | −7.0                        | UNK:H     | 2.17     | THR25:O       | Hydrogen Bond | Conventional H-bond |
|                     |                             | THR45:H   | 2.84     | UNK:O2        | Hydrogen Bond | Carbon H-bond       |
|                     |                             | MET49:S   | 5.6      | UNK:O2        | Other         | Pi-Sulfur           |
|                     |                             | CYS145:S  | 3.64     | UNK:O2        | Other         | Pi-Sulfur           |
|                     |                             | HIS41     | 5.82     | UNK:O2        | Hydrophobic   | Pi-Pi Stacked       |

| GA                  | −8.0                        | UNK:H    | 2.51     | GLU166:O      | Hydrogen Bond | Conventional H-Bond |
|                     |                             | SER46:HB2 | 2.47     | UNK:O2        | Hydrogen Bond | Carbon H-Bond       |
|                     |                             | PRO168:HD2|2.8      | UNK:O14       | Hydrogen Bond | Carbon H-Bond       |
|                     |                             | UNK:H54  | 2.46     | ASN142:OD1    | Hydrogen Bond | Carbon H-Bond       |
|                     |                             | CYS145    | 5.21     | UNK:O2        | Hydrophobic   | Alkyl               |
|                     |                             | UNK:C28   | 5.19     | LEU27:O       | Hydrophobic   | Alkyl               |
|                     |                             | HIS41     | 4.58     | UNK:C28       | Hydrophobic   | Alkyl               |

Fig. 6. 3-D (three dimensional) diagram based on hydrogen bond donor and acceptor characteristics of amino acid residues (A) and 2-D (two dimensional) (B) diagram showing several interaction between M<sub>pro</sub> and Glabridin (G). PRO: Proline, MET: Methionine, GLU: Glutamic acid, CYS: Cysteine, HIS: Histidine.
The Carcinogenic profile also shows that GA, G, and L are non-carcinogenic so it can be applied as drugs for treating COVID-19 as there would not be any bioaccumulation of compounds in the human body and these compounds would not lead to cancer in future if the patient is treated for a long duration. Acute oral toxicity was found to be higher in G followed by GA and L. It shows that L has less oral toxicity than GA which further suggests that the drugs L are not metabolized in the gastrointestinal tract before reaching the desired target. All these results have also been depicted in the form of a graph (Fig. 8 A, B and Table 4).

4. Conclusion

The present studies conclude the presence of His and Cys as the conserved amino acid of Mpro of SARS-CoV2. Molecular docking indicated that the three active compounds of Glycyrrhiza glabra...
namely glycyrrhizic acid, Liquiritigenin, and Glabridin successfully docked with the amino acid molecule at the catalytic site of the M\textsuperscript{pro} with a high negative binding affinity and formed several molecular interactions with the main protease of SARS-CoV2. They may form a complex with the M\textsuperscript{pro} of SARS-CoV2 causing inhibition of the catalytic activity of the main protease. His\textsuperscript{41} and Cys\textsuperscript{145} are conserved amino acids found in different types of coronavirus family and also plays a crucial role in enzyme catalysis and our docking result also showed that His\textsuperscript{41} and Cys\textsuperscript{145} are participated in the non-covalent interaction with all the three active compounds of Glycyrrhiza glabra. So from this study it is concluded that these active compounds exactly occupy the same location and binds these amino acid residues of the enzyme which otherwise would be occupied by the natural substrates for the enzyme (M\textsuperscript{pro}), so these compounds can be replaced as a proposed drug for the main protease of SARS-COV2. admetSAR results show that glycyrrhizic acid would be more suitable drug candidate among the other two compounds as it is more selective, potent, non-carcinogenic, and non-tumorigenic. OSIRIS property explorer had shown Glabridin to have toxicity to the reproductive system and carcinogenic, and non-tumorigenic. OSIRIS property explorer had shown Glabridin to have toxicity to the reproductive system and carcinogenic, and non-tumorigenic. OSIRIS property explorer had shown Glabridin to have toxicity to the reproductive system and carcinogenic, and non-tumorigenic. 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