IN Silico approach of some selected honey constituents as

SARS-CoV-2 main protease (COVID-19) inhibitors

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Abstract:

The huge attack of coronavirus disease 2019 (COVID-19) over all the world forces the researcher around the world to study the crystal structure of the main protease M^{pro} (3-chymotrypsin-like cysteine enzyme) which is the essential enzyme for coronavirus processing the polyproteins and its life cycles. And by the way, the inhibition of this enzyme active site becomes the target of all scientists of drug discovery in order to overcome this disease. In this study, we have used the molecular modeling approach to evaluate the activity of different active compounds from honeybee and propolis to inhibit the presented sars-cov-2 main protease via Schrödinger Maestro v10.1. the presented study resulted in six main compounds possess high binding energy with the receptor active site of COVID-19 main protease. we hope this study being the way for honeybee constitution as an effective ligand for sars-cov-2 main protease inhibition and be in the medicinal study of anti-COVID-19 therapeutic drugs.

Keywords: *SARS-CoV-2*, honeybee, propolis, Molecular docking, Structure activity relationship (SAR), M^{pro} inhibition.

Introduction:

The crystal structure of *coronavirus COVID-19* main protease was studied by medicinal chemists around all the world to develop antiviral drugs. It was mentioned that the *COVID-19* doesn't produce many proteins and by the way, there are not many targets for coronavirus inhibition. The three-dimensional structure of coronavirus main protease is like that of *SARS-CoV M^{pro}*, its RNA genome is about 82% like *SARS-CoV* which belongs to the *betacoronavirus* genus [1,2]. The *COVID-19* has been named *SARS-CoV-2* and its drug target is M^{pro} (3-chymotrypsin-like cysteine enzyme) which is the essential enzyme for coronavirus processing the polyproteins [3-5].

 M^{pro} yields about 790 kDa polypeptide at different 11 spots of the large polyprotein which is the key to viral reproduction [6]. The inhibition of these active sites of the enzyme is the target point

for producing *anti-COVID-19* drugs. But due to there is no known cleavage specificity of similar human proteases, unlikely the inhibitors may be toxic [7].

Natural products play an attractive role in the production of drugs for several diseases without harmful side effects [8]. Bee's productions have been used in medicine for several diseases included tumor treatment and immune-related diseases [9,10]. Especially, honey and propolis produced by bees have been possessed several biological activities like anti-inflammatory, antibiotic, antifungal, antiviral, antioxidant, anti-cancer, and immunomodulatory, hepatoprotective effects [11-13].

General composition of propolis is about 50% resins, 30% waxes, 10% essential oils, 5% pollen, and 5% of various organic compounds included polyphenols and flavonoids, and more than 200 constituents have been elucidated [14-17]. Honey is the main bee's production from different plant and so the chemical composition of honey depends mainly on the source and environment of the plant which reflected on its biological activity. The dark honey, *Manuka Honey*, has attracted attention for its biological properties, especially as an anti-viral effect. Several studies have been shown that there are many flavonoids and polyphenolic compounds in Manuka honey constituent [18-21] which suggested being accountable to its anti-microbial, anti-viral, and antioxidant effects [22-25].

According to the great biological effect of honeybee and propolis, in the presented study six selected compounds which have previous attention on their biological effect [23, 26] namely, 3-phenyllactic acid, caffeic acid phenethyl ester (CAPE), lumichrome, Galangin, chrysin, and caffeic acid used to study their potent anti-COVID-19 effect in silico approach (molecular docking), hoping this study being the way for honeybee constitution as an effective anti-COVID-19 drugs.

Material and Methods:

Preparation of honeybee's chemical compounds (Ligand):

The 2D structure of the six selected chemical compounds of honeybee and propolis were obtained in the SDF format from *PubChem* database. They were subjected to *LigPrep* module of Schrodinger Maestro v10.1, and converted to three-dimensional 3D structure by including ionization, variation, stereochemical, correction, and energy minimization and optimization of geometry. Minimization using Optimized Potentials for Liquid Simulation-2005 force field (OPLS_2005). Ionization state were carried out at PH 7.0 +/- 2.0 using *Epik* module. Generating tautomer, desalted, varying their other chiral center, and a single low energy ring confirmation per ligands. The optimized ligands

were then used for docking. The 2D structure of selected compounds and their PubChem CID are shown in Fig 1.

Figure 1: Two dimensional *2D* structures of six selected honeybee chemical compounds *COVID-19 main protease Identification and Preparation:*

The three dimensional crystal structure of *COVID-19* main protease in complex with *N-(2-phenylethyl)methanesulfonamide* (PDB ID: 5R7Y) was downloaded from the protein data bank (https://www.rcsb.org) in 1.65A° resolution. Protein structure was refined by assigning their bond orders, adding their missing hydrogen atoms and disulfide bonds and removing water molecules within 5 A° of the heteroatom. Optimizing the side-chain hydroxyl groups Asn, Gln, and His states using OPLS_2005 force field. Restrained minimization of the input protein coordinates by 0.3 A° *RMSD* (Root Mean Square Deviation).

Receptor Grid Generation:

The interaction between different prepared ligand and receptor protein was studied by setting up the receptor grid generation. The already bound ligand was excluded from the grid generation and the site of docked ligand is confined to the enclosing box, centroid of docked pose and in similar size to the workspace ligand.

Molecular Docking (Glide Docking):

The prepared ligands and protein were docked using Schrödinger maestro software v10.1. using SP flexible ligand mode, with 10 possess per ligand. Energy minimized possess were performed with final scoring using *Glide score*, the lowest glide scores (the best docked pose) for each ligand were recorded, as well as, the RMSD value (root mean square deviation) between the observed X-ray crystallographic of protein (native structure) and the predicted confirmation of input ligand geometry were computed. The 3D structures of the docked protein and the binding interaction distance of each ligands with the receptor were analyzed using *PyMOL* software.

Results and Discussion:

According to the several reported studies on some of honeybee constitutions as antiviral compounds [23-27], and a survival need of an effective ant-covid-19 drugs without harmful side effect, the presented study directed to computational drug discovery.

Docking simulation studies of the six selected honeybee constitution to the covid-19 main protease using the advanced molecular docking program (Schrödinger maestro v10.1) revealed very interesting results. The details of all docking score, glide score, potential energy, and RMSD for the ligands (1-6) to the selected protein is summarized in table 1. It was shown that all ligands have good binding affinity towards the receptor in different ways. Compounds 2,4,5 have more glide score arising from electrostatic attraction as well as more than one hydrogen bond between hydroxyl groups of ligand and different amino acid residues. The most binding modes affinity of ligands 1-6 with the receptor are shown in figure 2.

Table 1: Docking results for ligands 1-6 with COVID-19 main protease:

Number	Ligands	Potential	Docking	Glide	RMSD
		energy	Score	score	
1	3-phenyllactic acid	34.546	-5.867	-5.868	0.049
2	Caffeic acid phenylethyl	46.07	-6.383	-6.386	0.048
	ester (CAPE)				
3	Caffeic acid	14.22	-4.387	-4.387	0.035
4	Chrysin	63.126	-6.097	-6.103	0.047
5	Galangin	74.258	-6.295	-6.307	0.044
6	Lumichrome	94.141	-5.205	-5.205	0.040

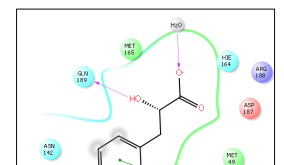
CAPE residue was stabilized within the receptor by interacting through electrostatic (bi-bi stacking) of phenyl ring with aromatic ring of (HIE-41) of the receptor with 4.2 A°, while the other closing residues of amino acid (THR-24) and (THR-26) are having strong hydrogen bond with the two hydroxyl group of *CAPE* ligand (cf. Fig.2).

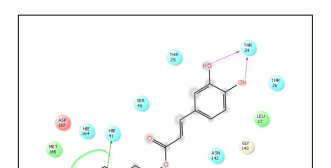
Chrysin is binding with (SER-46), (THR-24), and (THR-26) through hydrogen bond with 2.4, 2.6, 2.1 A°, respectively, beside strong electrostatic (bi-bi stacking) interaction of phenyl ring with (HIE-41) with 3.8 A°.

Galangin interacted with the receptor with glide score -6.307 Kcal/mol through hydrogen bond with two amino acid residues (SER-46), (THR-24), as well as other bi-bi interaction with (HIE-41). In the other way, *caffeic acid* interacted with high affinity with COVID-19 main protease by binding its hydroxyl groups with two amino acid residues ((GLN-189), (HIE-164) of the receptor by hydrogen bond with 2.0, 2.8 A°, respectively, beside other bi-bi interaction with (HIE-41) with 4.1 A° (cf.Fig.2).

The other two ligands *Lumichrome* and *3-phenyllactic acid* posses moderate binding affinity with the receptor with glide score -5.205 and -5.867 Kcal/mol, respectively. *Lumichrome* and *3-phenyllactic acid* binding within the pocket through bi-bi stacking with (HIE-41), while *3-phenyllactic acid* possess also strong hydrogen bond with only one amino acid residue (GLN-189) (cf. Fig. 2).

The three-dimensional crystal graphical structure of the docked six ligand 1-6 with the COVID-19 main protease are presented in Figure 3.





3-phenyllactic acid

Caffeic acid phenylethyl ester CAPE

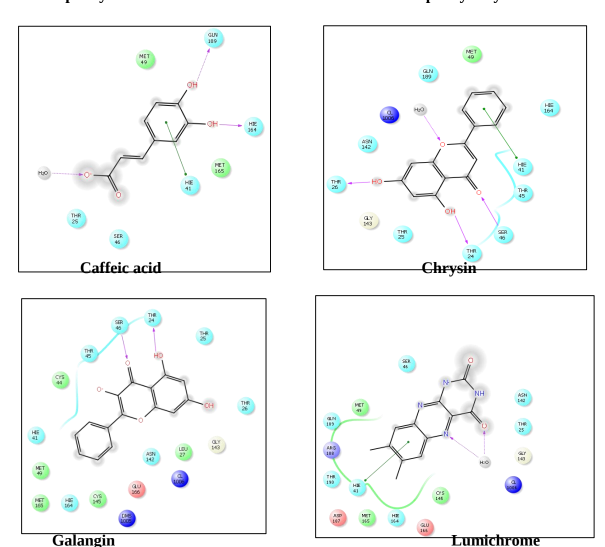
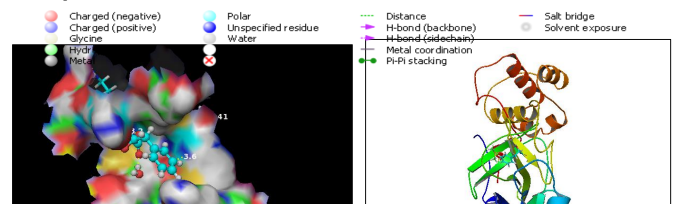
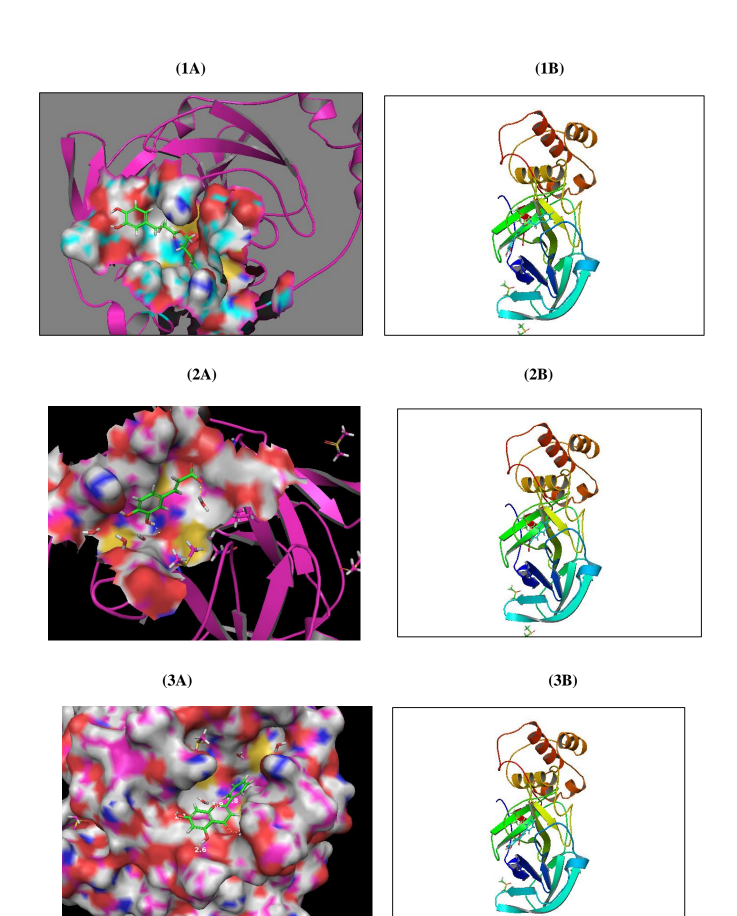


Figure 2: Ligand interaction diagram of six selected honeybee constitutions (1-6) with COVID-19 main protease.





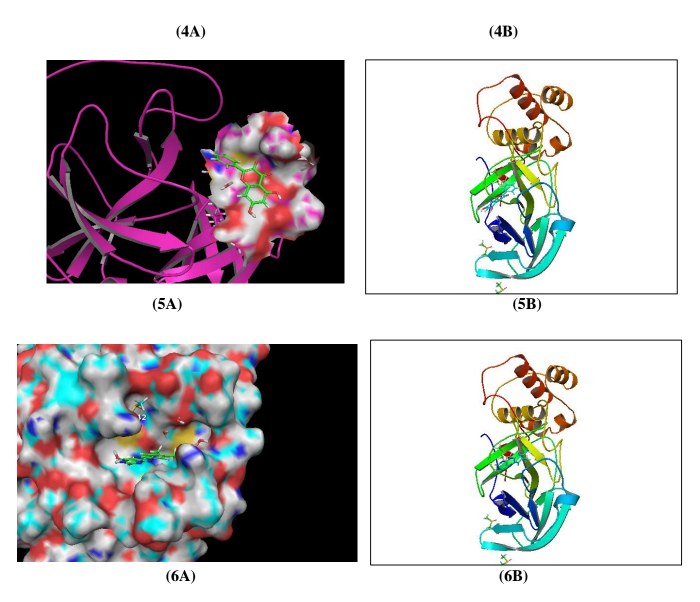


Figure 3: (A) Three-dimensional *3D* interaction diagrams of ligands 1-6 docked in the active site of COVID-19 protease using *PyMOL* software; red color reflects the high polar area, blue color reflects the mild polar area and grey color reflects the hydrophobic area. (B) Crystal Structure of COVID-19 main protease in complex with ligands 1-6.

Conclusion:

Honeybee and propolis include a wide range of flavonoid compounds with several biological activities. The presented study screened *in silico* the biological activity of six compounds present in

honeybee and propolis as antiviral components against the COVID-19 main protease. The study revealed that four compounds have strong binding affinity with good glide score and may inhibit the COVID-19 main proteas and virus replication. The study aimed for further *in-vivo* study in order to produce novel natural anti-COVID-19 drugs without any harmful side effect.

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