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Exploring the inhibitory potential of *Saussurea costus* and *Saussurea involucrata* phytoconstituents against the Spike glycoprotein receptor binding domain of SARS-CoV-2 Delta (B.1.617.2) variant and the main protease ( $M_{pro}$ ) as therapeutic candidates, using Molecular docking, DFT, and ADME/Tox studies



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# ABSTRACT

The B.1.617.2 Delta variant is considered to be the most infectious of all SARS-CoV2 variants. Here, an attempt has been made through in-silico screening of 55 bioactive compounds from two selected plants, Saussurea costus and Saussurea involucrata as potential inhibitors of two viral proteases, main protease Mpro (PDB ID:6LU7) and the RBD of SGP of Sars-CoV-2 B1.617.2 Delta variant (PDB ID:7ORB) where the binding energy, molecular interactions, ADMET/Tox, chemical descriptors and Quantum-Chemical Calculations were explored. Molecular docking results demonstrated that the three top docked compounds formed relatively stable complexes within the active site and displayed remarkable binding energy in the order of Tangshenoside III, Rutin and Hesperidin (-9.35, -9.14 and -8.57 kcal/mol, respectively) with Mpro and Rutin, Tangshenoside III and Hesperidin (-9.07, -7.71 and -7.57 kcal/mol) with RBD of SGP. These compounds are non-Mutagen and non-carcinogen. Therefore, according to the Lipinski's Rule of Five they exhibited three violations concerning hydrogen acceptor, donor and molecular weight. However, based on the Quantum-Chemical Calculations results the selected ligands have effective reactivity, as they showed lower band gaps. The difference of the  $E_{LUMO}$  and  $E_{HOMO}$  was low, ranging from 0.0639 to 0.0978 a.u, implying the strong affinity of these inhibitors towards the target proteins. Among the three inhibitors, Rutin exhibited higher reactivity against two viral proteases, main protease (Mpro) and the Sars-CoV-2 B1.617.2, as the band energy gap was lowest among all the three phytochemicals, 0.0639 a.u This could indicate that Rutincan be potential anti-viral drug candidates against the existing SARS-CoV-2, the B.1.617.2 Delta variant.

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# 1. Introduction

At the end of year 2019 a novel Severe Acute Respiratory Syndrome Coronavirus (SARS-CoV-2) with human-to-human transmission, was identified [1]. COVID-19 outbreak was first identified in the city of Wuhan, China in December 2019 as an epidemic; afterward it quickly propagated in Chinese territory and outside. World Health Organization declared it, two months later, as a pandemic on March 11, 2020 [2]. SARS-CoV-2 spread rapidly, causing the disease known as COVID-19, has since infected over 198 million people, leading to at least 4.2 million deaths worldwide of August 1st, 2021 [3]. The Alpha Variant of Concern (VOC) (formerly known as the Kent VOC, B.1.1.7, or S gene negative) had been the dominant strain previously, but it has rapidly been replaced on May 19, 2021, by the Delta VOC formerly known as the Indian Delta VOC or B 1.617.2 [4].

The most promising targets so far identified have been the spike protein, RNA-dependent RNA polymerase (RdRp), and the papain-like protease 3CLpro, also known as main protease (Mpro) [5,6]. It is essential for processing the polyprotein that led to the proteolytic activation of the viral functional proteins [7]. Since the

\* Corresponding author. E-mail address: houchi.selma@univ-setif.dz (S. Houchi). main protease (Mpro) is required for viral gene expression and replication, it is a promising target for antagonists to treat novel coronavirus disease.

SARS CoV-2 utilizes the receptor-binding domain (RBD) of trimeric spike glycoprotein (SGP) to gain quick entry into bronchial epithelial cells by binding with a human receptor, Angiotensin-Converting Enzyme-II (ACE-II) on the host cells that located on many body organs [8,9]. It is well reported that the utilization of ACE-II receptor blockers exhibit adverse effect such as hyperkalemia and angioedema [10]. The efficacy of currently licensed COVID-19 vaccines against B.1.617.2 is unknown [4]. Currently, several inhibitors are proposed by researchers across the globe as potential drug targets for the treatment of this disease but they still lack optimum specificity and efficacy; and to date, no highly effective therapy for treating coronavirus infections has been made available, although it possesses 12 mutations in its spike protein relative to the wild type SARS-CoV-2 first detected in Wuhan, China. Thus, blocking of the RBD of SGP, which interacts with amino acid residues of ACE-II could be an attractive target domain for the treatment of the development of new antiviral drugs anti-B.1.617.2 infection and prevention of the risk of multi-organ failure.

It is well known that, over the past decades, medicinal plants have been attracting attention in the search for bioactive compounds to develop new drugs and healthy foods among which polyphenols, lipids, polysaccahrides, sterols, terpenes. Furthermore, various plant extracts were reported to have high antioxidant, anticancer activity and to influence anti-inflammatory responses. However, plant based medicines are widely used due to their wide acceptance and therapeutic effects with minimal side effects [11,12]. Hence, at such a time, herbal medicine have shown experimentally promising results in SARS CoV-2 inhibition. Recently, various studies have highlighted the implication of a variety of phytoconstituents which has the potential to be developed as antiviral agents for SARS-CoV-2 [13–15].

The genus *Saussurea* DC. of the flowering plant family Asteraceae comprises about 300 species in the world. A family that is rich with well-known medicinal plants of highly economic importance. Many of which have been intensively studied for their phytochemical constituents and medicinal applications, including *S. lappa.* [16] and *S. involucrata.* [17]. Traditional clinical applications were considered guide for bioactivity evaluations, and many correlations were found between the pharmacological actions in traditional records of the herb and bioactivities confirmed in modern studies [18]. *S. lappa.* and *S. involucrata* are traditionally used to treat a wide spectrum of disorders; its pharmacological evidence is grounded by long history of usage and promising experimental results.

Saussurea costus (Falc.) Lipschitz, a perennial herbaceous plant, one of the best-known species within this genus, is commonly known as costus in English. It has several synonyms, such as *Aucklandia costus* Falc., *Aplotaxis lappa* Decne., *Saussurea costus* (Falc.) Sch.-Bip., *Aucklandia lappa* Decne, *Saussurea lappa* (Decne.) C. B. Clarke., *Theodorea costus* O. Kuntz [19]. Several workers have reported on the different biological activities of *Saussurea costus* in various *in vitro* and *in vivo* test models. Pharmaceutical effects investigations of extracts from *Saussurea costus* roots have been found to exhibit anti-inflammatory [20], hepatoprotective [21], anti-ulcer [22], anticancer [23], would-healing [24], antifungal and antimicrobial [25], anticonvulsant [26], larvicidal [27] and protein tyrosine phosphatase 1B inhibitory [28] activities.

Another Saussurea species, *S. involucrata* Karel. & Kir. with large translucent cream-colored bracts concealing the huge inflorescence, are named 'glasshouse plant' [29]. According to "The Plant List", *S. involucrata* Matsum. & Koidz is the only accepted name for the herb, with asynonym "*S. involucrata* (Kar. & Kir.) Sch. Bip." [30].

More recent studies have shown that this plant have antiinflammatory, analgesic [31]. Immunomodulatory [32], antioxidative [33], antihypoxic and neuroprotective [34], cardiovascular diseases [35], allergies and asthma [36].

The research and the development of a drug molecule "drug discovery" is often an extremely long, tedious and very expensive process at high risk of failure. For each new drug, the time elapsing between the first step, during which a relevant biological target in a given disease process is identified, until a drug is placed on the market is estimated at an average of 12 to 14 years [37]. An alternative approach, such as virtual screening or rational drug design, are now routinely used to guide drug discovery. The use of these in silico techniques could improve the efficiency and save the cost of drug discovery [38].Virtual screening approaches based on the structure of the target, more specifically the molecular docking, predict possible modes of interaction between a ligand and the therapeutic target and provide a way of studying interactions at the molecular level and are, therefore, an indication of the biological activity of new molecules taking into account only structural criteria.

In this context, we conducted computational screening study on a library of 55 isolated molecules from *Saussurea costus* and *Saussurea involucrata*, investigated the ADMET/tox studies, the DFT calculations and the binding affinity, of these compounds with the key target RBD of SGP SARS-CoV-2 B1.617.2 and the main protease (M<sub>pro</sub>) through molecular docking analysis.

# 2. Materials and methods

## 2.1. Molecular modeling platform

Molecular docking and ADMET/Tox studies were performed using Molecular Operating Environment (MOE) version 2019.10 molecular modeling software and Discovery Studio (DS) 2.0 Software (Studio 2.5, Accelrys, Co. Ltd., San Diego, CA, USA), respectively. DFT calculations were performed by Gaussian 09 suite of program at 6–31 G (d,p) basis set.

## 2.2. Library preparation

A total of 55 compounds were included in the screening (Table 1), encoded by *S. lappa.* and *S. involucrate* medicinal plants, after a profound literature review. Structure Data Format of these bioactive phytoconstituents were retrieved from the PubChem database (www.pubchem.ncbi.nlm.nih.gov) alongside the reference inhibitors (N3 and Hydroxychloroquine for M<sub>pro</sub> and Spike glycoprotein, respectively).

## 2.3. Molecular docking studies

As molecular targets, three important proteins for SARS-CoV2 infection were obtained from protein data bank of the Research Collaboration for Structural Bioinformatics (RCSB), website (http: //www.rcsb.org/pdb) in PDB formats. This including the Main Protease (Mpro) (also called 3C-like protease \_ 3CLpro), which represents a pivotal role in the propagation of SARS-CoV-2. The crystal structure of this enzyme was downloaded in complex with its inhibitor N3, (PDB ID: 6LU7; 2.16 A°). The crystalline structure of the L452R mutant receptor binding domain (RBD) of SARS-CoV-2 Spike glycoprotein (SGP) was downloaded in complex with COVOX-75 and COVOX-253 Fabs (PDB code: 7ORB; 2.5 A°). However, the crystalline structure of the Native Human Angiotensin Converting Enzyme-Related Carboxypeptidase (ACE2) was downloaded without inhibitor (PDB ID: 1R42; 2.20 A°).

Informations of the tested library of active phytoconstituents from S. lappa. and S. involucrata.

no	Compounds	Source	Refs.	PubChem CID	Formula	Type compounds
1.	8-cedren-13-ol	S.costus	[27]	519,545	C15H24O	Sesquiterpenoid
2.	(7Z,10Z,13Z)	S.costus	[16]	556,280	$C_{16}H_{26}O$	Trienyl derivative
	7,10,13-Hexadecatrienal					-
3.	$\alpha$ -Cyclocostunolide	S.costus	[39]	385,663,116	$C_{15}H_{20}O_2$	Sesquiterpenoid lactones.
4.	$\alpha$ -Curcumene	S.costus	[27]	92,139	C <sub>15</sub> H <sub>22</sub>	Sesquiterpenoids
5.	α-Selinene	S.costus	[27]	10,856,614	C <sub>15</sub> H <sub>24</sub>	Sesquiterpenoids
6.	$\beta$ -sitosterol	S.costus	[39]	348,274,860	$C_{29}H_{50}O$	Steroids
7.	Alloisoimperatorin	S. involucrata	[40]	5,317,436	$C_{16}H_{14}O_4$	Coumarins
8.	Apigenin	S.costus	[41]	5,280,443	$C_{15}H_{10}O_5$	Flavones, (flavonoïdes)
~		S. involucrata	[17]	4.64.440		
9.	Arbusculin B	S.COSTUS	[42]	161,442	$C_{15}H_{20}O_2$	Sesquiterpenoid lactones.
10.	Arctigenin	S. involucrata	[43]	64,981 100 528	$C_{21}H_{24}O_6$	Linnen skusssides
11.	Porganton	S. Involucrata	[41]	100,526	$C_{27}\Pi_{34}U_{11}$	Europocoumoring
12.	Caffeic acid	S costus	[41]	689 043		Hydroxycinnamic acid
13.	Catechin	S costus	[41]	9064	C15H14Oc	Flavones (flavonoïdes)
15.	Chlorogenic acid	S.costus	[41]	1.794.427	C <sub>16</sub> H <sub>18</sub> O <sub>9</sub>	Hydroxycinnamic acid
		S. involucrata		, . ,	10 10 5	5
16.	Chrysoeriol 7-O-glucuronide	S. involucrata	[29]	14,630,703	$C_{22}H_{20}O_{12}$	Flavonoid-7-o-glucuronides
17.	Costunolide	S.costus	[44]	5,281,437	$C_{15}H_{20}O_2$	Germacranolide
18.	Daucosterol	S.costus	[45]	5,742,590	$C_{35}H_{60}O_{6}$	Steroids-like
19.	Dehydrocostuslactone	S.costus	[39]	73,174	$C_{15}H_{18}O_2$	Guaianolide sesquiterpene
						lactone
20.	Edultin	S. involucrata	[41]	5,317,013	$C_{21}H_{22}O_7$	Furanocoumarin
21.	elemol	S.costus	[16]	92,138	$C_{15}H_{26}O$	Sesquiterpenoid
22.	Epicatechin gallate	S.costus	[39]	107,905	$C_{22}H_{18}O_{10}$	Flavanois
23.	Epicatechin Epigallogatochin gallato	S.costus	[41]	/2,2/6	$C_{15}H_{14}O_6$	Flavanois
24. 25	Epiganocalecinii ganale	S.costus	[41]	5 281 461	$C_{22}\Pi_{18}O_{11}$	Flavallois
25. 26	Callic acid	S costus	[42]	370	C_H_O_	Phenolic acids
20.	Hesperidin	S costus	[41]	10.621		Flavanone glycoside
28.	Hispidulin	S. involucrata	[46]	5.281.628	C16H12O6	Monomethoxyflavone
29.	hispidulin-7-glucoside	S. involucrata	[46]	5,318,083	C <sub>22</sub> H <sub>22</sub> O <sub>11</sub>	Flavonoid-7-o-glucusides
30.	Hydroxybenzoic acid	S.costus	[41]	135	$C_7H_6O_3$	phenolic derivative
31.	Isopimpinellin	S. involucrata	[41]	68,079	$C_{13}H_{10}O_5$	8-methoxypsoralens
32.	Isorhamnetin	S.costus	[41]	5,281,654	$C_{16}H_{12}O_7$	Flavonols
33.	Jaceosidin	S. involucrata	[47]	5,379,096	$C_{17}H_{14}O_7$	6-o-methylated flavonoids
34.	Kaempferol	S. involucrata	[48]	5,280,863	$C_{15}H_{10}O_{6}$	Flavonols
35.	Luteolin	S.costus	[41]	5,280,445	$C_{15}H_{10}O_6$	Flavone
20	Lutadin 7 shuasaida	S. involucrata	[17]	5 290 627		Flavoraid 7 a chronoidea
30. 27	Luteoini-7-giucoside	S. Involucrata	[17]	5,280,037	$C_{21}H_{20}U_{11}$	Alpha bydroxy acid
38	Nenetin	S. involucrata	[41]	5 317 284	$C_{4} G_{16} O_{5}$	Flavonoid
39.	Nepetin-7-glucoside	S. involucrata	[29]	12.314.010	C22H22O12	Flavonoid-7-o-glucusides
40.	Oroselol	S. involucrata	[41]	160.600	C14H12O4	Coumarins
41.	Osthol	S. involucrata	[41]	10,228	$C_{15}H_{16}O_3$	A derivative of coumarin
42.	Protocatechuic acid	S.costus	[41]	72	$C_7H_6O_4$	Dihydroxybenzoic acid
		S. involucrata	[50]			
43.	Quercetin	S. involucrata	[46]	5,280,343	$C_{15}H_{10}O_7$	Flavonols
44.	Quercetin-3-glucoside	S.costus	[41]	25,203,368	$C_{21}H_{19}O_{12}^{-}$	Flavonoid-3-o-glycosides
		S. involucrata	[29]	5 202 452		
45. 46	Quercitrin	S. involucrata	[29]	5,280,459	$C_{21}H_{20}U_{11}$	Flavonoid-3-o-glycosides
40.	Quillic aciu Butin	S.costus	[41]	6308 5 380 805	$C_7 H_{12} U_6$	Flavonolo
-1/.	Kuuni	S involucrata	[ <sup>+1</sup> ] [46]	3,200,003	C <sub>27</sub> H <sub>30</sub> O <sub>16</sub>	ιανυποιδ
48.	Spathulenol	S.costus	[27]	92.231	C15H24O	Sesquiterpenoids
49.	Syringin	S. involucrata	[41]	5,316,860	C <sub>17</sub> H <sub>24</sub> O <sub>9</sub>	Phenolic glycoside
50.	Tangshenoside III	S. involucrata	[51]	11,968,970	$C_{34}H_{46}O_{17}$	Phenylpropanoid
51.	Vaginidiol diacetate	S. involucrata	[41]	9,997,718	C <sub>18</sub> H <sub>18</sub> O <sub>7</sub>	Organic Compounds
52.	Valerenol	S.costus	[16]	91,699,505	$C_{15}H_{24}O$	sesquiterpenoids
53.	Vulgarol B	S.costus	[39]	91,748,781	$C_{15}H_{24}O$	sesquiterpene ketone
54.	Xanthotoxol	S. involucrata	[39]	65,090	$C_{11}H_6O_4$	furanocoumarin
55.	Zaluzanin	S.costus	[42]	12,445,012	$C_{17}H_{20}O_4$	Sesquiterpene lactone

# 2.3.1. Ligand and protein structures preparation

*Retrieval and protein structure preparation of active sites.* The three targets proteins, 6LU7, 7ORB and 1R42 used in this study, was protonated where hydrogen atoms were added with their 3D geometry, corrected for any found errors in the connection or type of different atoms, and then energy minimized at the end of the preparation steps. This step followed by site-finder to define and isolate the same binding pocket of the co-crystallized native inhibitor as dummy atoms over helix for the docking step.

*Ligands preparation.* Structures of the phytoconstituents of *Saussurea* genus were surveyed and downloaded from the PubChem database in structure data format (SDF). Then, Energy minimizing of these ligands was done under the following conditions: temperature = 300 K, pH = 7. Furthermore, the geometry was performed using the field strengths in the MMFF94x implanted in MOE and Austin model 1 (AM 1) with gradient value of 0.0001 kcal/mol. Then, the compounds were saved in mdb format as a new database.

#### 2.3.2. Docking

The dock tool of Molecular Operating Environment (MOE) 2019.0102 software was used for fitting of the isolated compounds' database into the active site of SARS-CoV-2 and ACE-2 enzymes. Docking site was selected as dummy atoms, alpha triangle as the placement methodology, and London dG as the scoring methodology. The docking process was run followed by evaluation of poses. Poses with the highest energy scores and best ligand-enzyme interactions were selected and recorded.

# 2.4. Chemical descriptors calculation

Lipinski's Rule of Five [52] was used to investigate phytoconstituents that were selected for this study. Filters like Molecular weight of the ligand (<500 Da), high lipophilicity (LogP<5), number of hydrogen bonds donors (<5), number of hydrogen bond acceptors (<10) and lip druglike were used to carry out the further selection of this compounds. Violation of more than 2 of the above-stated parameters debarred further analysis of a particular molecule. Parameter details were calculated using Molecular Operating Environment (MOE) 2019.0102 software.

## 2.5. ADMET analysis

A computational study of the top ten docking inhibitor compounds of each protein was carried out for the prediction of pharmacokinetics ADMET (Absorption, Distribution, Metabolism, Excretion and Hepatotoxicity) properties. In this module, eight mathematical models, such as solubility and its level, HIA (human intestinal absorption), ADMET AlogP98, ADMET PSA-2D, BBB penetration level PPB (plasma protein binding), ADMET CYP2D6 and hepatotoxicity were predicted via Discovery Studio (DS) 2.0 Software (Studio 2.5, Accelrys, Co. Ltd., San Diego, CA, USA)

# 2.6. Toxicity risks assessment

Another method used to determine the therapeutic compatibility of the drug is toxicity prediction by using the USFDA (US FDA, United States Food and Drug Administration) standard toxicity risk predictor software TOPKAT in the Discovery Studio. Toxicity by TOPKAT and prediction, TOPKAT rat female NTP probability and prediction, developmental toxicity potential, aerobic biodegradability, ocular irritancy, skin irritancy and Daphnia  $EC_{50}$ .

# 2.7. Quantum chemical calculations

All Density Functional Theory (DFT) calculations were performed by using the Gaussian 09 program package [53] and the output files were visualized by means of Gauss View 5.0.8 program [54]. Quantum chemical calculations such as optimized geometries and corresponding molecule orbitals (MOs) energies, Mulliken atomic charge and molecular electrostatic potential (MEP) were obtained at B3LYP (Becke's three parameter hybrid model using the Lee-Yang Parr correlation functional) [55,56] level of density functional theory (DFT) with 6–31G(d,p) basis set [57].

## 3. Results and discussion

#### 3.1. Phytochemical constituents of S. lappa. and S. involucrata

The tested library of active constituents from *S. lappa*. and *S. involucrata* comprised phytochemicals that cover major classes of natural products. Phytochemical studies of the two *Saussurea* plants showed the presence of terpenes (sesquiterpenes

and steroids); polyphenols (flavonoids and lignans); aromatic acids (phenolic and hydroxycinnamic acids) and others.

Closely-related plants species often share similar secondary metabolites and bioactivities. The two *Saussurea* species shows resemblance in bioactivities as well as phytochemical constituents (References against each phytochemical constituents candidate plants have been shown in Table 1).

Saussurea is a medicinal plant of immense medicinal importance having a variety of compounds and diversity in the pharmacological spectrum. It is used to treat fever, headache, cough, bronchial asthma [58], diarrhea [59], cholera [60], tuberculosis, vomiting, dyspepsia [58,59], gastric ulcer [59], abdominal pain [58,59]. Different studies reported that Saussuria active compounds have antiviral activity including SARS-CoV-2. Prawiro et al. searched to make evidence-based medicine that Honey, Saussuria coctus and Nigella may cure the Covid-19. They divided mice into two groups, the first group as control received PBS as a placebo. Then the second group, received Honey, Saussuria coctus and Nigella sativa. After administration regiments a long three weeks, they sacrificed the mice and evaluated the immune responses markers that are Th2, Th17 for cellular, and NK cells, TGF- $\beta$ , IL-17A, sIgA, IL-4, IL4, B-def, and IgG for humoral. As results, they reported a statistically deference in the cellular immune responses markers, Th2 and Th17, between control with treatment group (p = 0.05) and a deference in the markers of humoral immunity (p => 0.05). However, the markers of humoral immunity (IL-17A) have no statistical difference [61]. Another study evaluated the crude extract prepared from Saussuria coctus for its antiviral activity against Hepatitis B Virus (HBV). Two compounds Costunolide and Dehdrocostuslactone suppressed the expression of Hepatitis B surface antigen (HBsAg) in a dose dependent manner with an  $IC_{50}$ value of 1.0 and 2.0 mM in Human hepatoma Hep3B cells. Significant suppression was also observed in human hepatoma cell line HepA2 derived from the HepG2 cells [62]. These findings suggest that Saussuria coctus may have the potential to develop as specific anti-virus drugs such as anti- SARS-CoV 2.

## 3.2. Docking studies

Plants are sources of phytoconstituents which has the potential to be developed as antiviral agents for SARS-CoV-2, as has been reported by previous studies [13–64]. Since ligand binding to a protein of interest is the first step in drug discovery, molecular docking is widely used to predict and identify ligands that fit into the binding pocket of a protein of interest [65]. Molecular docking is not only cost effective but also time effective approach to identify a promising therapeutics agent of COVID-19.

# 3.2.1. Evaluation test of the program used

Several methods are used to evaluate the performance of the different programs of docking for each application. The ability of an algorithm to find the correct location of the ligand in relation to its receptor is usually determined by means of the deviation root mean-square-deviation (RMSD) of the model designed by the software against the crystal structure. The RMSD test was performed on complexes formed between the SARS-CoV-2 receptors and different ligands. The obtained molecular docking results were considered valid since redocking the orginal ligands (N3). The accepted value is a maximum difference of 2Å beyond which the prediction is considered as inadequate [66]. The obtained molecular docking results were considered valid since redocking the crystalized ligands (N3 and Hydroxychloroquine) furnished great ligand superposition with root-mean standard deviation (RMSD) below 2 Å. Scores, RMSD\_refine values, and diverse interactions of the best top scoring ligands with the amino acids of the Mpro, RBD of SGP and ACE-2 pockets are described in Table 2.

Molecular docking results and interactions of the top scoring compounds of Saussurea species with SARS-CoV-2 proteins and ACE-2.

Protein		Binding energy		Bonds between atoms of	of compounds and residue	s of active site	
	ligand	Score (Kcal/mol)	RMSD refine		Involved Receptor		Bond length
				Atom of ligand	residues	Bond type	(A°)
	Tangshenoside III	_9 3479	1 6416	06	Thr 24	H-donor	2.90
	lungshenoside in	5.5-175	1.0410	011	Asn 142	H-donor	3.06
				C49	Met 49	H-donor	3.94
				C39	His 41	H-Pi	4.35
	Rutin	-9.1420	1.2988	04	Cys 145	H-donor	4.08
				08	Cys 145	H-donor	3.55
				09	Thr 26	H-donor	2.89
				012	Cys 145	H-donor	3.71
				04	Gly 143	H-acceptor	2.77
				07	His 163	H-acceptor	3.23
				010	Thr 26	H-acceptor	3.02
	Hesperidin	9 5670	1 6920	6-ring	GIII 189 Asp 142	PI-H U donor	3.07
	Hesperiali	-8.3072	1.6829	07	ASII 142 Mot 165	H-donor	3.04
				010	The 190	H-donor	2.86
				010	Cvs 145	H-donor	3.97
				012	Cys 145	H-acceptor	3.05
				6-ring	Thr 25	Pi-H	4.13
				6-ring	Asn 142	Pi-H	3.87
	Arctiin	-8.5658	1.5173	06	Met 165	H-donor	3.31
	Quercitrin	-8.4104	1.2280	05	Cys 145	H-donor	3.57
	-			011	Thr 26	H-donor	3.28
				03	His 163	H-acceptor	3.01
				6-ring	Gln 189	Pi-H	4.09
	Luteolin-7-glucoside	-8.0322	1.8745	03	Leu 141	H-donor	2.67
				05	Cys 145	H-donor	3.13
				028	Met 165	H-donor	3.89
				03	Ser 144	H-acceptor	2.91
				04	His 163	H-acceptor	3.14
				08	His 41	H-pi	3.28
				6-ring	Gln 189	pi-H	3.48
	Quercetin-3-glucoside	-7.9070	1.5491	04	Cys 145	H-donor	3.54
				08	Cys 145	H-donor	3.63
				013	Cys 145	H-donor	4.10
				03	Gly 143	H-acceptor	3.37
				04	Ser 144	H-acceptor	3.08
				06	His 163	H-acceptor	3.12
				6-ring	Gin 189	рі-Н	4.79
				6-ring	Gin 189	pi-H	3./1
	Deventerel	7.0.411	1 2721	6-ring	GIII 189	рі-н	4.24
	Daucosteroi Uicpidulin 7 glucosido	-7.8411	1.3/21	01	Giu 166	H-acceptor	3.21
	Thispiduliii-7-glucoside	-7.7850	1.5550	04 C12	Cys 145	H-donor	3.55
				03	Cly 143	H-acceptor	3.00
				06	His 163	H-acceptor	3.05
	$\beta$ -sitosterol	-75755	19469	01	Thr 26	H-donor	2.81
70RB	Rutin	-9.0665	1.7144	04	Phe 347	H-donor	2.91
				015	Ser 399	H-donor	2.82
				04	Ser 349	H-acceptor	2.83
	Tangshenoside III	-7.7140	1.9781	011	Lys 356	H-acceptor	3.11
				012	Tyr 351	H-acceptor	3.27
					Ala 348	pi-H	3.95
	Hesperidin	-7.5719	1.7773	06	Arg 346	H-donor	2.76
				08	Glu 340	H-donor	2.92
				09	Lys 356	H-acceptor	3.25
				010	Lys 356	H-acceptor	3.14
	Arctiin	-7.2114	1.4840	04	Ser 399	H-donor	2.83
				06	Arg 346	H-donor	3.10
				07	Arg 346	H-acceptor	2.94
	Daucosterol	-7.1701	1.9301	05	Glu 340	H-donor	2.81
	Epicatochin gallate	6 0922	1 7670	03	ASII 354 Sor 240	H-acceptor	3.11 2.77
	Epicateciiii gallate	-0.3023	1.7079	04	Δla 352	H-donor	2.77
				09	Ser 399	H-dopor	2.07
				011	Ser 399	H-donor	3.04
	Quercitrin	-6.8851	18457	08	Ser 399	H-donor	2.86
	Quercetin-3-glucoside	-6.694	1 5691	010	Ser 399	H-donor	2.00
	2			01	Arg 346	H-acceptor	2.93
				06	Arg 346	H-acceptor	3.20
	Hispidulin-7-glucoside	-6.6385	1.6345	03	Ser 399	H-donor	2.85
				07	Arg 346	H-donor	3.09
				6-ring	Arg 346	Pi-cation	2.99
	Edultin	-6.4001	1.9467	02	Asn 354	H-acceptor	2.64
				07	Arg 346	H-acceptor	3.18
1R42	Rutin	-8.9636	1.239	05	Glu 406	H-donor	2.80
				010	Asp 367	H-donor	3.15
				C20	Glu 406	H-donor	3.31
				07	Gln 442	H-acceptor	3.27
				09	Lys 441	H-acceptor	2.94
	Hesperidin	-8.4456	1.421	04	Asp 367	H-donor	2.84
				08	Asp 367	H-donor	2.87
				(2)	ASP 367	H-donor	3.56
				012	LYS 441	H-acceptor	2.94
	Tangchono-: 4- W	0 1205	1 176	010	GIN 522 Chi 275	H-acceptor	3.15
	rangsnenoside III	-0.1300	1.170	010	GIU 373 Asp 200	H-001101	2.84
				05	Asii 230 Ivs 441	H-acceptor	3.07
				00	Lys TTI	i i-acceptoi	J.12

## 3.2.2. Binding energy and molecular interaction studies

The molecular docking has been performed using the Molecular Operating Environment (MOE) 2019.0102 software to evaluate the binding mode of ligand and interactions in the active site. Our screening was performed against three major key target RBD of spike glycoprotein SARS-CoV-2 B1.617.2, the main protease ( $M_{pro}$ ) and ACE-2.

Docking analysis of SARS-CoV-2 Mpro. According to the 3D structure of SARS-CoV-2 Mpro, the active Mpro homodimer comprises two protomers, constituting three domains. Amino acid residues 8-101 constitute Domain I, amino acid residues 102-184 constitute Domain II, and amino acid residues 201-306 constitute Domain III. Before analyzing ligand-receptor interactions, we have defined binding site residues of Mpro enzyme. Binding site of Mpro enzyme contains Thr24, Thr25, Thr26, Leu27, His41, Met49, Tyr54, Phe140, Leu141, Asn142, Gly143, Ser144, Cys145, His163, His164, Met165, Glu166, Leu167, Pro168, His172, Asp187, Arg188, Gln189, Thr190, Ala191, and Gln192 residues. The S1 binding site is formed by Phe-140, Asn-142, Ser-144, Cys145, His-163, His-172, and Glu-166 side chains, Leu-141, Gly-143, His-164, and Met-165 backbones. The side chains of His-41, Val 42, Asn-119, Thr-25, Cys-145, Gly-143 together with the backbone of Thr-26 define S1' site. His41 and Cys145 form the catalytic dyad in the active site [67] and His164 is essential for enzyme activity. His163. His172 and Glu166 are believed to provide the opening gate for the substrate in the active state of the protomer [68], and Thr24, Thr26 and Asn119 are predicted to play roles in drug interactions [69,70].

The formation of hydrogen bonds defines the integrity and stable nature of each protein–ligand complex. Table 2 demonstrates the binding energies and different bonds of the best top ten complexes. Most of the docked compounds (1–55) formed relatively stable complexes within the active site of  $M_{pro}$  with Hydrogen bonds; showing moderate to promising energy scores in the range of -4.61 to -9.34 kcal/mol as compared to the inhibitor N3 (-7.44 kcal/mol). Most phytochemicals exhibited higher binding score than the standard. On the competition of docking, total of seven metabolites, namely Tangshenoside III, Rutin, Hesperidin, Quercitrin, Luteolin-7-glucoside, Quercetin-3-glucoside and hispidulin-7-glucoside showed effective binding with HIS41 or CYS145 of catalytic dyad along with multiple interactions with other amino acid residues in active site of SARS CoV2-Mpro.

The binding interactions of the top two docked complexes demonstrated that Tangshenoside III had three hydrogen bonds with Mpro, at Thr 24, Asn 142, Met 49 and one H-pi bond were observed for His 41. Rutin exhibited a higher number of hydrogen bonds, forming seven hydrogen bonds three at Cys 145, and four at Thr 26, Gly 143, His 163, Thr 26. This complex also formed a pi-H bond at Gln 189. These amino acids are predicted to play a major role during chemical interactions with these compounds and the inhibition of Mpro. The detail of interaction and visualization of the docking results of Tangshenoside III and Rutin with Mpro are provided in Figs. 1 and 2. Ghosh et al., [71] reported that polyphenols like broussochalcone A, papyriflavonol A, 3'-(3methylbut-2-enyl)-3',4',7-trihydroxyflavane, broussoflavan A, kazinol F, and kazinol J had good interaction with the catalytic residues of His41 and Cys145 of Mpro. These amino acids interactions with the SARS-CoV-2 main protease Mpro are similar to our findings. Previous studies reported that the inhibitor N3 docks in the active binding site of 6LU7 and forms hydrogen bonds with PHE 140, GLY 143, THR190, HIS 164, GLU166 and GLN189 [72] and these results similar to our finding. A recent study performed by Cherrak et al. showed that Rutin possesses high docking scores of -9.2 kcal/mol and interacts with Mpro via conventional H-bonds at Thr26, Tyr54, Leu141, and Glu166,  $\pi$ -cation interactions at His41,  $\pi$ -alkyl interactions at Met49,  $\pi$ -sulfur interactions at Cys145, and C-H bond interactions at Met165 [73]. Another investigation reported that Rutin was found to have, with Mpro, a docking score of -9.16 kcal/mol, it was considered to be a potential Mpro inhibitor. Rutin was predicted to form hydrogen bonds involving Cys145, Asn142, Gly143, and Thr190, with additionally the possible formation of  $\sigma$ - $\pi$  stacking interaction with Gln189. Notably, the major binding affinity was based on the presence of a hydroxyl group, which presented the key to anchoring and blocking the substrate into the active pocket of the catalytic center [74].

Hesperidin inhibit the cleavage activity of the Mpro in a dosedependent manner in cell-free and cell-based assays, with an IC<sub>50</sub> of 8.3  $\mu$ M [75]. A previous study also showed that the docked hesperidin compound against Mpro was -13.51 [76], and according to a previous investigation, the best hesperidin position against SARS-CoV-2 Mpro had a score of -10.1 [77,78]. Furthermore, Rameshkumar et al., docked 36 flavonoid to the active site of Mpro and it was observed that the compound agathisflavone had shown the highest binding energy value of -8.4 kcal/mol and interacts with Mpro by t hydrogen bonds at Lys102, Thr111, Ser158, His246 followed by Dracorubin and Cupressuflavone that showed dock score of -8.2 kcal/mol [79]. As presented in a previous study reported that several potentially active compounds have best docking scores which are cannabinoids, rhoifolin, pectolinarin, morin, kaempferol, epigallocatechin gallate, herbacetin, and hesperidin against 3CLpro were -8, -8.2, -8.2, -7.8, -7.8, -7.2, and -8.3, respectively [80].

Docking analysis of SARS-CoV-2 B1.617.2 spike glycoprotein. The B.1.617.2 Delta variant is considered to be the most infectious of all variants and as of June 2021 has become one of the most transmissible variants with the highest number of reported cases, followed by the B.1.617.1 [81,82]. Variants are characterized by multiple mutations in the surface spike glycoprotein. Mutations in the RBD can change the ability of the virus spike protein to bind to and enter the host cell. The high rate of mutation and recombination in SARS-CoV2 makes it difficult for scientist to develop specific anti-CoV2 drugs and vaccines.

According to global reports and Health agencies, in comparison with SARS-CoV-2 from Wuhan-2019, the new variants Delta B.1.617.2 present 5 mutations in S protein T19R; L452R; T478K; P681R; D950N [83]. L452R/T478K in B.1.617.2 are assumed to play roles in infectivity, transmissibility of the virus and leads to an increase in virulence by reducing the antibody binding affinity, as well as immune evasion [84].

All the 55 active constituents from *S. lappa.* and *S. involucrata* were individually docked to the active site of 70RB. As compared to the inhibitor, Hydroxychloroquine (binding energy = -7.409), Rutin had shown the highest binding energy value of -9.0665 kcal/mol. it was observed that the key amino acids include Phe 347, Ser 399 and Ser 349 were involved in the hydrogen bond interactions (donor and acceptor). It followed by Tangshenoside III and Hesperidin, that showed binding energy of -7.71 and -7.57 kcal/mol, respectively.

The binding interactions of the top docked complexes demonstrated that Rutin forming three hydrogen bonds at Phe 347, Ser 399 and Ser 349. Tangshenoside III had two hydrogen bonds at Lys 356 and Tyr 351and one pi-H bond were observed for Ala 348. Hesperidin exhibited four hydrogen bonds, two at Arg 346, Glu 340 and two at Lys 356. Arctiin had three hydrogen bonds at Ser 399 and two at Arg 346 (Table 2). These amino acids are predicted to play a major role during chemical interactions with these compounds and the inhibition of RBD of spike glycoprotein. The detail of interaction and visualization of the docking results of Rutin and Tangshenoside III with SARS-CoV-2 B1.617.2 spike glycoprotein is provided in Figs. 3 and 4. Solo and Doss [85] tries



Fig. 1. 2D and 3D structure of the Tangshenoside III docking results with the target Main protease Mpro of SARS-CoV-2.



Fig. 2. 2D and 3D structure of the Rutin docking results with the target Main protease Mpro of SARS-CoV-2.



Fig. 3. 2D and 3D structure of the Rutin docking result with the target spike protein (B.1.617.2) of SARS-CoV-2.

to explore the inhibitory activity of 50 phytochemicals against the spike protein of the Delta SARS-CoV-2. They used Hydroxychloroquine, as a standard in the docking analysis and they reported that this standard had a binding affinity of -6.5 kcal/mol with the delta variant. Solo and Doss identified the 3,5,3'Trimethoxy-6,7:4',5'-bis(methylenedioxy) flavone as having the highest binding affinity of -8.7 kcal/mol for both the target proteins, in comparison to the other 49 phytochemicals. The major Hydrogen bond interactions were observed with Glu340, Val341, Ala344, Phe347, Ala348, Ser349, Tyr351, Ala352, Lys356, Ser 399 and Tyr451. Except Val341, Ala344 and Ala348 that exhibited a pi-H bond, Amino acids involved in the Hydrogen bond interactions of Delta SARS-CoV-2 spike found in our study were similar with the report of Solo and Doss.



Fig. 4. 2D and 3D structure of the Tangshenoside III docking result with the target spike protein (B.1.617.2) of SARS-CoV-2.

Docking analysis of ACE-2. The S-protein is made up of two subunits, the S1 subunit is involved in host cell receptor recognition and is the receptor-binding domain (RBD) and S2 subunit is responsible for fusion of the viral membrane and the host cellular membrane [86]. Cell entry of SARS CoV-2 depends on two consecutive steps, firstly binding of the viral spike (S-protein) to host cellular receptors followed by priming of S-protein by cell proteases. Recently, researchers showed that SARS CoV-2 uses the ACE-2 receptor for entry [87] and the serine protease TMPRSS2 for priming of S-protein. It adheres to the host cell's surface receptor, human angiotensin-converting enzyme 2 (hACE2), allowing viral cellular entry via endosome formation and/or plasma-membrane fusion [88,89]; ACE-2 is an integral membrane protein type I [90]. Due to its key role in the process of SARS-CoV-2 infection, this structural component has been confirmed as the entry point for the SARS-CoV-2 virus to infect human cells [91]. The SARS CoV-2 virus can colonize any organ that has ACE-2 receptors such as lung endothelial and alveolar type II cells, epithelial cells in the upper esophagus, enterocytes in the ileum and colon, epithelial cells in the bile, heart muscle cells, proximal tubules in the kidneys, and urothelial cells in the bladder [92].

SARS-CoV-2 binds to ACE2 ten times more strongly than other coronaviruses, making SARS-CoV-2 more infectious than others [93]. So the spike protein of SARS-CoV-2 was predicted to have a strong binding affinity to human ACE-2. Mutations in the RBD can help to enable strong affinity and binding capacity to ACE2, leading to higher transmissibility [94–96]. Moreover, The Indian (B.1.617.2 or Delta) variant has different mutations in the RBD of SARS-CoV-2: L452R and T478K. Thus, targeting the ACE2-RBD interaction site may be a viable solution that may restrict the coronavirus from entering into the cells by inhibiting the interaction between ACE-2 and the RBD [97]. During the COVID-19 pandemic, various molecular docking studies have targeted the interaction regions between the RBD and ACE2, performing docking for drug repositioning or using chemotherapeutic libraries (or using both methods) to search for drugs or compounds that can inhibit the interaction between RBD with ACE-2 [98-101]. The majority of studies have reported that bioactive molecules, have a higher binding potential than antiviral drugs (hydroxychloroquine and remdesivir) used in SARS-CoV-2 spike protein and ACE-2 [102,103]. In this context, molecular docking was performed to tested the interaction of bioactive compounds from two selected plants, Saussurea costus and Saussurea involucrata with ACE2 to determine their potential to inhibit its binding with RBD of the S-protein.

Molecular docking studies with ACE-2 have shown that Rutin, Hesperidin, and Tangshenoside III have the highest binding energy values of -8.9636, -8.4456 and -8.1305 kcal/mol, respectively. Sebsequently, these compounds bind strongly to ACE-2 and may hinder the substrate accessibility and its subsequent inhibition. Rutin involves five hydrogen bonds with Glu 406, one with Asp 367, one with Gln 442 and one with Lys 441. Hesperidin also shows favorable interactions with ACE-2 through five hydrogen bonds three with Asp 367, one with Gln 522 and one with Lys 441 (Table 2). The detail of interaction and visualization of the docking results of these complexes with ACE-2 is provided in Figs. 5 and 6. Furthermore, another study conducted by Chen and Du (2020) demonstrated that the molecular docking studies of hesperidin with the ACE-2 enzyme showed that hesperidin can bind to ACE-2 with a predicted binding energy of -8.3 kcal/mol and binding sites at Glu-479, Arg-482, Ser-611 and Tyr-613. These results suggest that hesperidin may bind to ACE-2 and thus block 2019-nCoV infection [77]. Rutin has been identified as one of the three compounds (glycyrrhizin, rutin, and pheophorbide A) with a high binding affinity -6.9 kcal/mol to the ACE2 receptor.166 It created an interaction with the following residues: Phe4. Leu29, Asp30, Asn33, Val93, Ala99, Leu100, Ala387, Asp350, Ala387, Gln388, Pro389, Leu391, and Arg393 [104]. The molecular docking studies also showed that baicalin might bind strongly to the ACE-2 enzyme with a predicted binding energy of -8.46 kcal/mol and binding at Asn-149, Arg-273, and His-505. Based on the potential binding to ACE-2, it can be suggested that baicalin is a promising candidate for 2019-nCoV treatment. In addition, Scutellarin has the potential to bind to ACE-2 with a projected binding energy of -14.9 kcal/mol and binding sites at GLU-495, UNK-957, and ARG-482 [77].

#### 3.3. Screening through pharmacokinetic properties

Drug-like properties and pharmacokinetic properties are intrinsic characteristics of drugs that may need to be optimized independently from pharmacodynamics properties during drug development. It is a balance among molecular properties affecting pharmacodynamics and pharmacokinetics of small molecules.

#### 3.3.1. Chemical descriptors calculation

The drug-likeliness of top ten drug ligands for both SARS-CoV-2 Mpro and spike proteins was evaluated through Lipinski's rule of five parameters and the values are summarised in Table 3 (values of all the molecules are mentioned in supplementary Data (Table S1). This rule is based on physicochemical parameters of the tested ligands, including: Molecular weight (MW) not greater than 500 g/mol; A partition coefficient log P less than or equal five;



Fig. 5. 2D and 3D structure of Rutin docking results with the target ACE-2.



Fig. 6. 2D and 3D structure of Hesperidin docking results with the target ACE-2.

number of hydrogen bond donors (HBD) (NH and OH groups) not greater than five; and number of hydrogen bond acceptors (HBA) (O and N atoms) not exceeds ten. Drugs having log P ranging from 0 to 5, have high possibility of oral absorption. Not more than one violation of these parameters will reduce the druggability of the molecule [105].

All the screened compounds were found to have little violation and it was believed to effective despite of few violations, except edultin that was fully in agreement to Lipinski's rule of five (without violation).  $\beta$ -sitosterol and Epicatechin gallate also qualified Lipinski's rule of five with one violation concerning Log P and hydrogen donor, respectively. Except for  $\beta$ -sitosterol Daucosterol all other drug molecules showed the value of Log P less than 3.53, which indicates that the molecules are more likely to be in the hydrophilic environment and are favorable for their drug-likeness. Daucosterol has another violation regarding the molecular weight. His molecular weights is slightly more than the recommended values but within the permissible limit. Quercitrin, Luteolin-7glucoside and Quercetin-3-glucoside exhibited two violation regarding hydrogen acceptor and donor as well as Arctiin showed two violations regarding hydrogen acceptor and molecular weight. Tangshenoside III, Rutin and Hesperidin exhibited three violations concerning hydrogen acceptor, donor and molecular weight.

## 3.3.2. ADMET parameters screening for drug likeness

The determination of pharmacokinetic properties of drugs at the preliminary stages of drug discovery is essential to bring the drug up to clinical trial [106]. These properties such as absorption, distribution, metabolism, excretion and hepatotoxicity (ADMET) are important in order to determine the role in drug discovery and success of the development of compound for human therapeutic use. To determine the drug-likeliness of our compounds, we calculated their ADMET properties. ADMET prediction was done using Discovery Studio 2.5 (Accelrys, San Diego, USA) software. Prediction is based on two 2D molecular descriptors, namely AlogP\_98 and PSA. As shown in Fig. 7, Compounds with more than 90% absorbability would lie within the 99% confidence ellipse as the green ellipse Compounds falling out of the ellipse are those with less than 30% absorbability. The results of ADMET were displayed in Table 4, revealing eight descriptors, such as the absorption of

Detailed analyses of parameters of Lipinski's filter for the drug-likeness. The elaborated comparative analysis of all the seven drugs concerning parameters of Lipinski's filter.

Ligand	MW	Lip acc	lip-Don	Log p (o/w)	Lip drug- like	Violation
Tangshenoside III	726.72	17	9	-1.49	0	3
Rutin	610.52	16	10	-1.11	0	3
Hesperidin	610.56	15	8	-0.77	0	3
Arctiin	534.6	11	4	0.56	0	2
Quercitrin	448.38	11	7	0.80	0	2
Luteolin-7-glucoside	448.38	11	7	0	0	2
Quercetin-3-glucoside	463.37	12	7	-0.11	0	2
Daucosterol	576.8	6	4	6.15	0	2
hispidulin-7-glucoside	462.4	11	6	-0.06	0	2
$\beta$ -sitosterol	414.71	1	1	8.07	1	1
Edultin	386.4	7	0	3.53	1	0
Epicatechin gallate	442.38	10	7	3.38	1	1



Fig. 7. Prediction of drug absorption for active phytoconstituents from S. lappa. and S. involucrate considered for anti-SARS-CoV-2.

Та	ble	4

	Solubility	Absorption			Distribution		Metabolism	Toxicity
Ligand	ADMET Aqueous Solubility (Level)	Intestinal Absorption level	ADMET AlogP98	ADMET PSA-2D	BBB penetration level	Plasma Protein Binding	ADMET CYP2D6 binding	Hepato-toxicity
Tangshenoside III	-4.583 (2)	3	-0.274	258.78	4	False	False	False
Rutin	-6.182 (1)	3	-1.158	270.1	4	False	False	True
Arctiin	-3.057 (3)	3	1.814	154.1	4	False	False	False
Hesperidin	-4.489 (2)	3	-0.431	237.4	4	False	False	True
Quercitrin	-3.888 (3)	3	0.589	189.8	4	False	False	True
Luteolin-7-glucoside	-3.328 (3)	3	0.238	189.8	4	False	False	True
Quercetin-3-glucoside	-2.98 (3)	3	-1.003	207.1	4	False	False	False
Daucosterol	-5.55 (2)	2	6.337	101.12	4	False	False	False
hispidulin-7-glucoside	-3.426 (3)	3	0.464	177.91	4	False	False	True
$\beta$ -sitosterol	-8.256 (0)	3	8.084	20.81	4	True	False	False
Edultin	-4.856 (2)	0	3.502	87.62	2	False	False	True
Epicatechin gallate	-5.538 (2)	3	3.339	180.87	4	True	True	True

ligand	Ames mutagenicity	TOPKAT Mouse female NTP probability	TOPKAT Mouse male NTP probability	TOPKAT Rat female NTP probability	TOPKAT Rat male NTP probability	Aerobic bio-degradability	Ocular irritancy	Skin irritancy	TOPKAT Daphnia EC <sub>50</sub> (mg/L)
Tangshenoside III Rutin	Non- Mutagen Non- Mutagen	0,597,672 0,208,873	0,708,292 0,609,889	0,310,728 0,0,827,222	0,578,401 0,651,442	Degradable Degradable	Mild	None Weak	0.05599 1.41631
Hesperidin Arctiin	Non- Mutagen Non- Mutagen	0,368,231 0,621,887	0,278,522 0,614,135	0,246,047 0,2685	0,576,179 0,667,288	Degradable Degradable	Mild	None None	1.71086 0.27347
Quercitrin Luteolin-7-alucoside	Non- Mutagen Non- Mutagen	0,129,232 0 203 474	0,604,399 0.613.408	0,0,644,857 0 111 993	0,643,589 0.586.446	Degradable Degradable	Mild Moderate_severe	None Week	2.53329 3.08780
Quercetin-3-glucoside	Non- Mutagen	0,382,166	0,548,789	0,16,454	0,668,277	Degradable	Moderate-severe	None	6.39441
Daucosterol	Non- Mutagen	0,987,038 (Carcinogen)	0,99,373	0,183,303	0,293,199	Degradable	Moderate-severe	None	0.152622
hispidulin-7-glucoside	Non- Mutagen	0,371,303	0,578,767	0,272,004	0,692,262	Degradable	Moderate-severe	None	4.27525
$\beta$ -sitosterol	Non- Mutagen	0,979,079	0,995,694	0,280,137	0,355,373	Degradable	None	Weak	0.217316
Edultin	Non- Mutagen	0,584,079	0,706,606	0,364,043	686,881	Degradable	Mild	None	0.811667
Epicatechin gallate	Non- Mutagen	0,384,233	0,612,907	0,281,324	0,580,833	Degradable	None	Strong	1.61585

virtual toxicity studies of the best ten scoring ligands for both SARS-CoV-2 Mpro and spike glycoprotein

drugs depends on intestinal absorption, polar surface area (PSA) and AlogP98. The distribution depends on factors that include the blood-brain barrier (BBB) and plasma protein binding (PPB). Metabolism is predicted based on the Cytochrome P450 2D6 inhibition and hepatotoxicity. ADMET aqueous solubility predicts the solubility of each compound in water at 25°C. ADMET solubility descriptors predicts molar solubility of drugs within the ranges: < -8.0 (level 0 = extremely low solubility), -8.0 to -6.0 (level 1 = very low, but possible) -6.0 to -4.0 (level 2 = low), -4.0 to -2.0 (level 3 = good solubility), and -2.0 to 0.0 (level 4 = optimal solubility). Except  $\beta$ -sitosterol and Rutin ADMET Aqueous solubility logarithmic level of most of the compounds was found to be 2, 3 which indicates low or good aqueous solubility, respectively. AD-MET absorption predicts human intestinal absorption (HIA) after oral administration. Intestinal absorption of drug was determined by obtained levels: 0 (good), 1 (moderate), 2 (low), 3 (very low). Daucosterol and Edultin were found to have moderate to good HIA whereas, all other ligands were found to have very low absorption level, therefore are poor in gastro intestinal absorption. High hydrogen-bonding capacity of molecules has ebeen reported as one of the most important factors affecting intestinal absorption of new chemical entities. As the generally respected "rule of 5" suggests, any molecule having more than 5 hydrogen bond donors and 10 hydrogen bond acceptors would normally face poor absorption consequences. Easy distribution of the drug through blood brain barrier (BBB) measured by its AlogP98 value which must be less than 5. The obtained absorption levels determine drug absorption and absorption decreases inversely with the level, i.e., level 0 denotes proper absorption, level 1 denotes moderate absorption and so on. All the compounds were fallen outside the 99% ellipse (undefined). Hence, the compounds may not be able to penetrate the blood brain barrier. ADMET plasma protein binding model predicts whether a compound is likely to be highly bound to carrier proteins in the blood. Predictions are based on AlogP98 and 1D similarities to two sets of "marker" molecules. True symbolizes binding and false symbolizes non-binding. The ADMET plasma protein binding property prediction clearly suggesting that all of 12 compounds with an exception of  $\beta$ -sitosterol and Epicatechin gallate, are not likely to be highly bound to carrier proteins in the blood. ADMET CYP2D6 binding predicts cytochrome P450 2D6 enzyme inhibition using 2D chemical structure as input as well as a probability estimate for the prediction. Predictions are based on a training set of 100 compounds with known CYP2D6 inhibitions. This descriptor determines inhibitory effect by predicted classes: noninhibitor (false) and inhibitor (true). Except Epicatechin gallate all compounds are predicted as non-inhibitors of CYP2D6 suggesting that these compounds are well metabolized in Phase-I metabolism. Hence, the side effects (i.e., liver dysfunction) are not expected upon administration of these compounds which indicated they did not have hepatotoxicity.

The common top three ligands, according to the docking results, namely Tangshenoside III, Rutin and Hesperidin pass through the Lipinski's filter with 3 violations regarding molecular weights, hydrogen bond acceptors and donors. They were reported to show low intestinal absorption because of their physicochemical properties. Considering that hydrogen bond acceptor and donor groups influence permeability. In this context, the low oral bioavailability of compounds could be explained in terms of their calculated parameters according to Lipinski's rule, including molecular weights, hydrogen bond acceptors and donors. Thus, these compounds are predicted to be hardly transported, diffused, and absorbed than compared with the small molecules. On the other hand, the most popular and authentic rule for confirming the drug-likeness of the Ligand is Lipinski's rule. However, many drugs that do not pass through the Lipinski's filter but have immense pharmacological properties have been approved by the FDA as potential drug for



Fig. 8. Optimized molecular structures of (a) Tangshenoside III, (b) Rutin and (c) Hesperidin calculated with the methodology B3LYP/6-31 G (d,p).

clinical purposes [107]. Despite violation of some rules, approved anticancer and anti-infective drugs from natural products or their semisynthetic derivatives such as taxol and amphotericin B have also some violations but are biologically effective as drugs. Therefore, these results do not interfere with the development of these compounds as potential SARS-CoV-2 therapeutic agents [108].

# 3.3.3. Toxicity risks assessment

The toxicity predictions of the compounds were also investigated with Discovery Studio using the TOPKAT protocol. TOPKAT is a computational tool for in-silico quantitatively prediction of toxicity developed by Accelrys [109]. And is used by universities, private companies and government agencies including the US EPA, US FDA, Environment Canada, Health Canada and the Danish EPA for toxicity assessments. It computes and validates assessments of the toxic and environmental effects of chemicals solely from their molecular structure. TOPKAT employs robust and cross-validated Quantitative Structure Toxicity Relationship (QSTR) models to predict a probable value of toxicity . Moreover, with the help of these QSTR models, the product of a structure descriptors and its correspond-







Fig. 9. Presentation of the energy levels, energy gaps, and frontier molecular orbitals of (a) Tangshenoside III, (b) Rutin and (c) Hesperidin.

ing coefficient, it calculates probability values and evaluates toxicity through this descriptors contribution. It follows the criterion of checking the components in the optimal predictive space (OPS), and when they lie outside then the results were considered as unreliable.The Optimum Prediction Space (OPS) technology is implemented in TOPKAT as the methodology used to identify model applicability domain, providing a means of cheking whether the compounds under investigation are well represented in the models. The toxicity profiles calculated for all the compounds are tabulated in Table 5. Our results indicate that all ligands are non-



Fig. 10. Molecular electrostatic potential maps of (a) Tangshenoside III, (b) Rutin and (c) Hesperidin at the B3LYP/6-31G(d,p) level of theory.

Mutagen, non-carcinogen with Rat female NTP. Therefore, the toxicity of the ligands was found to be suitable for the development into a medical drug. Thus, they experience significant first-pass effect.

## 3.4. Quantum-Chemical Calculations

After sorting the common top three drug ligands (Tangshenoside, Rutin and Hesperidin) based on their highest negative binding energies value. These drug ligands are quite similar with different binding energies towards the three targets. The above-mentioned drug ligands were subjected to Quantum-Chemical Calculations studies.

#### 3.4.1. Geometry optimization

Optimized geometries of the investigated compounds obtained from DFT calculations using B3LYP/6–31G(d,p) method are given in Fig. 8. The absence of imaginary frequencies confirmed the global minima on the potential energy surface. Calculated geometric parameters of the investigated compounds are given in Supplementary Data (Table S2).

# 3.4.2. Frontier molecular orbital analysis

The distributions and energy levels of the HOMO and LUMO orbitals, computed with the B3LYP/6–31G(d,p) level for Tangshenoside III, Rutin and Hesperidin are shown in Fig. 2. To understand the relationship between electron delocalization and reactivity of

Tangshenosio	le III, Rutir	and Hesperidin	geometrical	parameters	calculated	in a.u at	the B3LYI	P/6-31G(d, <b>j</b>	p) level of theory.
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Compound	НОМО	LUMO	Egap	IP	EA	μ	χ	η
Tangshenoside III	-0.1440	-0.0462	0.0978	0.1440	0.0462	-0.0951	0.0951	0.0489
Rutin	-0.1703	-0.0809	0.0639	0.1703	0.0809	-0.1256	0.1256	0.0395
Hesperidin	-0.15238	-0.07922	0.07316	0.15238	0.07922	-0.1158	0.1158	0.0365

 $\Delta E = E_{HOMO} - E_{LUMO}, IP = -E_{HOMO}, EA = -E_{LUMO}, \mu = 1/2(E_{HOMO} + E_{LUMO}), \chi = -(E_{LUMO} + E_{HOMO})/2, \eta = 1/2(E_{LUMO} - E_{HOMO}).$ 

studied compounds, density plots of the HOMO and LUMO composition of studied compounds can be used (Fig. 9). As shown in Fig. 9, the positive region is presented in red and the negative one in green to further determine the relationship between delocalization and reactivity, global descriptors were applied. Important factors used to describe inhibition activity of selected phytochemicals plant against two viral proteases, main protease (Mpro) and the Sars-CoV-2 B1.617.2 and ACE-2 are HOMO and LUMO energies and HOMO-LUMO energy gaps (Egap) [110]. The structure with the lower LUMO energy is a weak electron acceptor while the structure with the higher HOMO energy is a good electron donor [111]. The HOMO and LUMO energy values as well as Egap values of the studied compounds are shown in Table 6.

## 3.4.3. Global chemical reactivity

The optimized geometries obtained with B3LYP/6-31 G (d,p), were used for global chemical reactivity calculations. The reactivity parameters were calculated according to the approximations of energies  $E_{HOMO}$  and  $E_{LUMO}$  states, based on the optimization of the fundamental state geometry. The calculated parameters were: Energy Gap (Egap), ionization potential (I), electronic affinity (EA), chemical potential ( $\mu$ ), electronegativity ( $\chi$ ), and chemical hardness ( $\eta$ ). The obtained results are shown in Table 6, in which is observed that Rutin presents a higher capacity to form anions due to the electronegativity value ( $\chi$ ), which represents a greater tendency to attract electrons. Regarding chemical hardness  $(\eta)$ , Rutin shows a lower value compared to Tangshenoside III and Hesperidin; this parameteris indicative of the good interaction of this molecule with the surrounding environment. As the energy gap increases, the molecule becomes harder and more stable/less reactive [112,113]. Thus, there is short gap between HOMO and LUMO in the Rutin as compared to Tangshenoside III and Hesperidin. The values of electronic chemical potential of Tangshenoside III, Rutin and Hesperidin are presented in Table 6, ligands that have the greater electronic chemical potential (absolute values), was the less stable and more reactive. These results showed that the selected phytochemicals have effective reactivity, as they showed lower band gaps. The difference of the  $E_{IIIMO}$  and  $E_{HOMO}$ was low, ranging from 0.0639 to 0.0978 a.u, implying the strong affinity of these inhibitors towards the target proteins. Among the three phytochemicals, Rutin exhibited higher reactivity against two viral proteases, main protease (Mpro) and the Sars-CoV-2 B1.617.2, and ACE-2 as the band energy gap was lowest among all the three phytochemicals, 0.0639 a.u (Table 6).

# 3.4.4. Molecular electrostatic potential (MEP)

The molecular electrostatic potential (MEP) can predict the reactive sites of electrophilic and nucleophilic attack in a molecule. These MEPs were realized to show the electron density within the Tangshenoside III, Rutin and Hesperidin on surfaces that surround the molecules, and the locations of sites for nucleophilic and electrophilic attacks [114,115]. Red indicates negative regions promoting electrophilic attack. Here, the negative potentials are generated over the electronegative oxygen atoms whereas the H-atoms have a positive potential region in the structures and the positive regions represented by the blue color are favorable sites for nucleophilic attack. The yellow regions indicate the slightly rich electron and the green regions indicate neutral. Molecular electrostatic potentials of Tangshenoside III, Rutin and Hesperidin were calculated using B3LYP/6–31 G (d,p) methodology. The surface maps shown in Fig. 10.

## 3.4.5. Mulliken atomic charges

The Mulliken atomic charges of Tangshenoside III, Rutin and Hesperidin were calculated by DFT using B3LYP/ 6-31 G (d,p) basis set, the data were tabulated in Table S3. It showed that the C5 is the most positive charge and O26 have the most negative charge for Tangshenoside III. On the other hand, it is observed that the most nucleophilic center of Rutin is O29 which is the most electrophilic susceptibility positions, it is obvious that the nucleophilic susceptibility of the Rutin is recognized on C21 and C27 sites. However, O42 is the most negative charge of Hesperidin and their respective positively charged atom is C22. The positively charged centers are the most susceptible sites for nucleophilic attacks, electron donation. However, the most negatively charged centers are the most susceptible sites for electrophilic one. The graphical representation for Mulliken atomic charges of the Tangshenoside III, Rutin and Hesperidin is shown in Fig. 11. As it can be observed from this figure, the calculated atomic charges for the more negative atoms are the oxygen atoms while, the opposite case represented by the attached protons, are the most positive atoms. Again, this charge distribution supports the idea that the molecule donates its electrons through these groups.

#### 4. Previous studies

To the best of our knowledge, Tangshenoside III was not tested for SARS-CoV-2 inhibition or any other enzyme activity. However, previous studies provide a strong rationale for testing rutin and hesperidin in SARS-CoV-2.

Various pharmaceutical properties associated with rutin are anti-inflammatory [116], antitumor [117], antiulcer [118], antimalarial [119], antimicrobial [120], anticoagulant [121], nephroprotective [122], hepatoprotective [123]. Rutin was also found to decrease the infectivity of enteroviruses 19. It also showed effects on hepatic monooxygenase activities in experimental influenza virus infection (EIVI) [124]. Molecular docking studies suggested that rutin can act as a novel hepatitis B virus inhibitor [125]. This flavonoid has also been reported to inhibit a broad spectrum of viral proteases such as human norovirus protease 23 and NS3 serine protease, *in vitro*. A recent analysis, through *in silico* methods, suggested that the binding and therapeutic property makes the rutin a prominent lead to develop a potential inhibitor of SARS-CoV-2 Mpro, RdRp, PLpro, and S-protein to combat against COVID-19 pandemic [126].

Hesperidin is claimed to possess anti-oxidant as well as antiinflammatory properties and has the ability to hinder SARS-CoV-2 entry and replication. It exhibited anti-viral activity against the influenza virus through a significant reduction of virus replication [127]. In addition, various studies on the inhibition effects of this compounds have reported. Sakata et al. (2003) indicated that hesperidin inhibits cyclooxygenase-2 (COX-2), inducible nitric ox-



-0.654 a.u.

Fig. 11. Structures with color range Mulliken atomic charges of Tangshenoside III, Rutin and Hesperidin.

ide synthase (iNOS) proteins, which might be related to the antiinflammatory and anti-tumorigenic efficacies [128]. It comes out as a multi-potent phytochemical agent in Alzheimer's disease therapeutics exhibiting strong proteases ßsecretase-1 binding ability (BACE1), high amyloid-ß peptide (Aß) aggregation inhibition, from a phytochemical library of 200 phytochemicals through the multitarget screening to identify multi-potent [129]. Furthermore, can block coronavirus from entering host cells through ACE2 receptors which can prevent the infection. Anti-viral activity of hesperidin might constitute a treatment option for COVID-19 through improving host cellular immunity against infection and its good antiinflammatory activity may help in controlling cytokine storm [130].

# 5. Conclusion

SARS-CoV-2 spread rapidly, causing the disease known as COVID-19. Variants are characterized by multiple mutations in the surface spike (S) glycoprotein. These mutations in the RBD can change the ability of the virus spike protein to bind to and enter the host cell. The high rate of mutation and recombination in SARS-CoV2 makes it difficult for scientist to develop specific anti-CoV2 drugs and vaccines. In this workflow, we first short-listed 55 Saussurea costus and Saussurea involucrata -based phytochemicals through literature mining. The compounds were screened against the RBD of the SGP of SARS-CoV-2 Delta (B.1.617.2) Variant and the main protease  $(M_{pro})$  to identify potent inhibitors that might be able to interfere with their catalytic function. The molecular docking studies identified the three ligands namely Tangshenoside III, Rutin and Hesperidin as potent inhibitors. Rutin performed very well when docked with the Sars-CoV-2 proteins producing the highest score with the spike B.1.617.2 and the second highest score with the Mpro Sars-CoV-2.

With these encouraging results, Tangshenoside III, Rutin and Hesperidin can be further explored for structural modification and detailed investigations to arrive at possibly newer potent agents with better anti SARS-CoV-2 Delta (B.1.617.2) Variant therapeutic activity. Accordingly, we recommend further fast *in-vivo* and *in-vitro* evaluations, preclinical and clinical studies at least for the aforementioned promising compound.

## **Declaration of Competing Interest**

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

## **CRediT** authorship contribution statement

**Selma Houchi:** Methodology, Software, Investigation, Supervision, Formal analysis, Writing – review & editing. **Zakia Messasma:** Conceptualization, Formal analysis, Writing – original draft.

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## Supplementary materials

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