

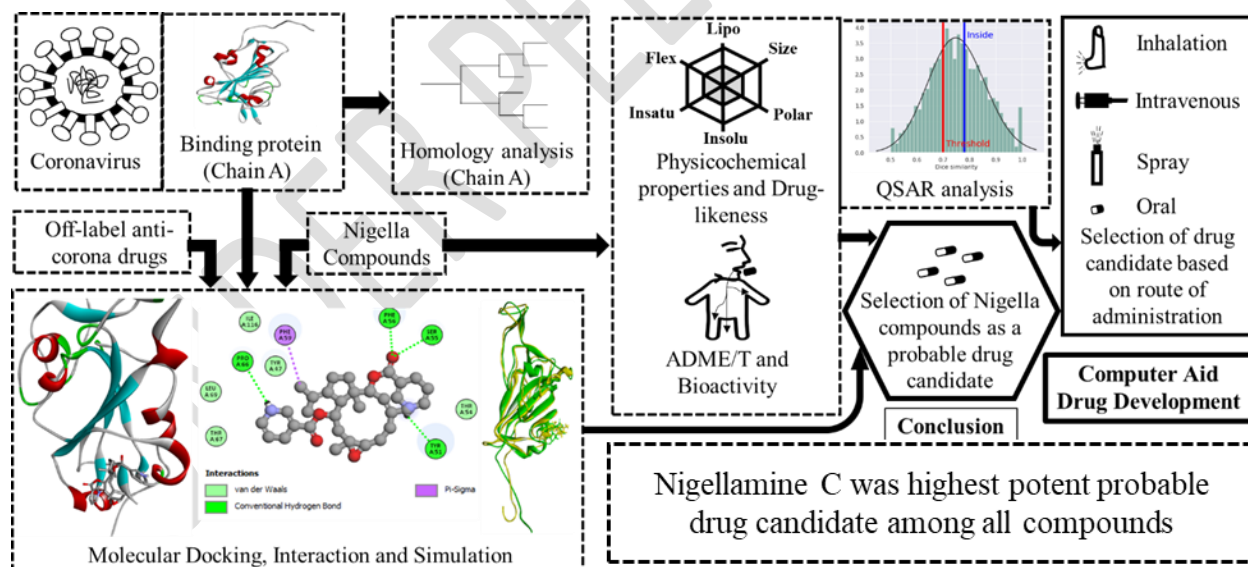
Original Research Article

In silico screening of potential drug candidate against chain A of coronavirus binding protein from major Nigella bioactive compounds

Abstract

More hazardous varieties of severe acute respiratory syndrome-related coronavirus have created major health hazards around the globe since 2019. There is no hundred percent effective drug has been developed against this virus. Bioactive compounds from plants are used as drugs or the main source of raw material for drugs against various diseases. This computational biology-dependent study aimed to identify of drug ability and inhibitor properties of major Nigella compounds against coronavirus by using molecular docking, ADME/T, bioactivity, and QSAR analysis tools. Nigellamine C showed the highest binding affinity as of -7.9 kcal/mol to chain A of subunit 1 (S1) of the spike protein of coronavirus in molecular docking. Nigellamine C showed suitable ADME properties with negligible toxic properties and good drug-likeness properties. Nigellamine C may be a potent candidate for inhalation and/or oral drug development based on QSAR analysis.

Graphical Abstract



Keywords: Black cumin; phytochemicals; COVID-19; ADME/T; QSAR

INTRODUCTION

Coronavirus is responsible for flu-like symptoms depending on the infectious disease with acute respiratory distress and an acute inflammatory state in humans (Bischof, Wolfe, & Klein, 2020). The total number of cases and deaths during the COVID-19 pandemic was 676,609,955 and 6,881,955 from January 2020 to March 2023 worldwide (JHCRC, 2023). Some therapeutic interventions such as convalescent plasma therapy, monoclonal antibodies therapy, immune response regulators, and some off-level drugs such as sotrovimab, regdanvimab, tocilizumab, infliximab, baricitinib, tofacitinib, ruxolitinib, statins, anakinra, fluvoxamine, novaferon, remdesivir, ribavirin, favipiravir, molnupiravir, sofosbuvir, daclastivir, nirmatrelvir, ritonavir, etc. are used to treat its clinical complication (Bischof et al., 2020). However, there are no COVID-19 complete curing therapeutics or agents or drugs that can cure or prevent it from spreading.

More than 80% of the world's population relies on traditional medicinal plants for treating various illnesses (Akerle, 1993) because medicinal plants are a good source of secondary metabolites that are used in multidisciplinary fields such as traditional systems of medicine, food supplements, modern medicines, pharmaceutical products, folk medicines, and chemical industries (Karna, 2013). Phytochemicals from black cumin may cure COVID-19 because traditionally oil of seeds from this plant was used in curing different symptoms associated with sinusitis such as coryza, nasal congestion, headache, neck pain, earache, and toothache since the ancient era (Mahboubi, 2018). It helps to inhibit the inflammation of sinuses and respiratory airways, and microbial infections as well as shows respiratory protective activities (El Tahir, Ashour, & Al-Harbi, 1993).

The traditional uses, phytochemical analysis, and biological activities of the black cumin seeds and their applications in human and animal health including poultry have been described in many literature and reviews (Ahmad et al., 2013; Azeem, Zaib-Ur-Rehman, Asif, Arif, & Rahman, 2014; Forouzanfar, Bazzaz, & Hosseinzadeh, 2014; A.-u. H. Gilani, Jabeen, & Khan, 2004; Kooti, Hasanzadeh-Noohi, Sharafi-Ahvazi, Asadi-Samani, & Ashtary-Larky, 2016; Majdalawieh & Fayyad, 2015; Mollazadeh & Hosseinzadeh, 2014; Paarakh, 2010; Ragheb et al., 2009; Randhawa & Alghamdi, 2011; Sharma, Ahirwar, Jhade, & Gupta, 2009; Tembhurne, Feroz, More, & Sakarkar, 2014). Deferent extracts of this plant and seeds exhibited spasmolytic and bronchodilator activities (A. Gilani, Aziz, Khurram, Chaudhary, & Iqbal, 2001), preventing early and late pulmonary fibrosis and inflammation (Poursalehi et al., 2018) and showed anti-viral activity (Hanafy & Hatem, 1991; Salem & Hossain, 2000), anti-bacterial activity (Chaieb, Kouidhi, Jrah, Mahdouani, & Bakhrouf, 2011; Chowdhury, Islam, Rashid, & Ferdous, 1998; Hanafy & Hatem, 1991), anti-parasitic activity (Mahmoud, El-Abhar, & Saleh, 2002), anti-inflammatory activity (Ghannadi, Hajhashemi, & Jafarabadi, 2005; Marsik et al., 2005; Shaterzadeh-Yazdi, Noorbakhsh, Hayati, Samarghandian, & Farkhondeh, 2018), anti-allergic (Kalus et al., 2003; Nikakhlagh et al., 2011), anti-tumor activity (Kumara & Huat, 2001; Nagi & Almakki, 2009; Salomi, Nair, & Panikkar, 1991), anti-oxidant activity (Abdel-Wahhab & Aly, 2005; Houghton, Zarka, de las Heras, & Hoult, 1995), immune-modulatory activity (Haq, Lobo, Al-Tufail, Rama, & Al-Sedairy, 1999), etc. Thus, this computational biological method-dependent study will reveal the binding properties of several major metabolites to chain A of the spike protein of coronavirus with the different physiochemical and drug-related properties ability identification for helping future drug development.

MATERIALS AND METHODS

This in silico study has been designed to find out some effective metabolites that can help appropriate drug design against coronavirus-responsible disease. This study includes the following steps.

Protein chain selection and properties analysis

The S1 subunit C-domain serves as the RBD for most coronavirus. Chain A of this subunit plays a vital role in receptor binding and entry of various components of the virus into a cell. Chain A of spike protein S1 of coronavirus was searched in the NCBI protein databank (<https://www.ncbi.nlm.nih.gov/protein/2425373090>). Chain A of Spike protein S1 (GI: 2425373090 and PDB: 7XNF) which was released on 2023-01-11 in the protein bank has been downloaded in FASTA format from NCBI. Physicochemical properties include the amino acids number, molecular weight, theoretical pI, aliphatic index, grand average of hydropathy (GRAVY), Instability index, half-life, and charged residues number had been determined by using ProtParam analysis tools (Gasteiger et al., 2005). The function and family of this protein chain had been identified by the NCBI Conserved Protein Domain Family search tool (<https://www.ncbi.nlm.nih.gov/Structure/cdd/wrpsb.cgi>).

Homology analysis and phylogenetic tree

Homology searching was carried out in NCBI blastp suite where the database was standard database of non-redundant protein sequences (nr) and the algorithm was blastp (protein-protein BLAST)

(https://blast.ncbi.nlm.nih.gov/Blast.cgi?PROGRAM=blastp&PAGE_TYPE=BlastSearch&LINK_LOC=blasthome). Multiple sequence alignment and phylogenetic trees were made by using Molecular Evolutionary Genetics Analysis version 11 (Tamura, Stecher, & Kumar, 2021).

Structure prediction and validation of chain A of Spike protein S1

Various chemical and structural properties of secondary structure were predicted by using PSIPRED (http://bioinf.cs.ucl.ac.uk/psipred/&psipred_uid=53bf5940-baf9-11ed-948f-00163e100d53)(Buchan & Jones, 2019) and *SOPMA* (https://npsa-prabi.ibcp.fr/cgi-bin/npsa_automat.pl?page=/NPSA/npsa_sopma.html) (Combet, Blanchet, Geourjon, & Deleage, 2000) online tools.

Preparing and validation of macromolecule

The 3D structure of the target protein was determined using SWISS-MODEL web-based server (<https://swissmodel.expasy.org/interactive>) on homology modeling where the server automatically performs BLASTp search to identify templates for each protein sequence (Waterhouse et al., 2018). The Cryo-EM structure of PCoV_GX spike glycoprotein A was selected for homology modeling from the searched result. The identity of the sequence was 100%, the GMQE value was 0.85, the QSQE value was 0.60, the method was EM and the oligo-state was homo-trimer. This modeled homo-trimer peptide was analyzed by PROCHECK(Laskowski, MacArthur, Moss, & Thornton, 1993) (Job ID: 1279619) and ProSA(Sippl, 1993; Wiederstein & Sippl, 2007). The 3D model structure was visualized by BIOVIA Discovery Studio Visualizer (Studio, 2008) and deleted chain B, Chain C, and attached ligands. No water molecule was found. Polarized the remained chain A monomer by BIOVIA Discovery Studio Visualizer and analyzed again by PROCHECK (Job ID: 1279624).

Ligand selection and preparation

Six major metabolites of *N. sativa* were selected and the 3D structures of these compounds were downloaded as SDF files (.sdf) from PubChem (<https://pubchem.ncbi.nlm.nih.gov/>). All six compounds were nigealanine (PubChem CID 12116700), nigellamine C (PubChem CID 101341399), nigellamine D (PubChem CID 101727452), nigellicine (PubChem CID 11402337), nigellidine (PubChem CID 136828302) and nigellone (PubChem CID 398941). Four different antiviral therapeutic agents that are commonly used in COVID-19 off-level treatment were also selected for comparison after the deep literature review (Bischof et al., 2020). The 3D structures of these therapeutic drugs were also downloaded as SDF files (.sdf) from PubChem named ribavirin (PubChem CID 37542), favipiravir (PubChem CID 492405), remdesivir (PubChem CID 121304016) and nirmatrelvir (PubChem CID 155903259). All compounds were converted from SDF files (.sdf) file to the PDB format (.pdb) using the Open Babel graphical user interface (GUI) tool (O'Boyle et al., 2011). The energy minimization of all ligands and control drugs was performed by using PyRx tool (Dallakyan & Olson, 2015).

Molecular docking and simulation

The conversion from the PDB files (.pdb) to the PDBQT files (.pdbqt) format and the molecular docking of individual ligands into the target chain A of spike protein S1 was performed by using the PyRx tool (Dallakyan & Olson, 2015). Vina search space was maximized for macromolecule in Pyrx where the center was X: 164.3046, Y: 177.8894, and Z: 202.6309, and the dimension (angstrom) was X: 69.4436, Y: 41.6859, and Z: 67.7703. Molecular dynamics simulation was done by using myPresto version 5 (Kasahara et al., 2016). Crystal water solvate was used, the system size was a sphere, and the margins (Å) were 8 margins from molecular size. Center coordination was the center of gravity of the molecule and the ion was automatically neutralized. Hydrogen atoms were added to the ligands and the partial charge was also added by the Gasteiger method.

ADME/T and bioactivity analysis in human

The ADME was predicted using SwissADME (DeLano, 2002) and pKCSM (Pires, Blundell, & Ascher, 2015) web-based molecular tools. Lipinski, ≤ 10 , NH or OH ≤ 5 (Lipinski, Lombardo, Dominy, & Feeney, 2012); Ghose, molecular weight was between 160 and 480 Da, WlogP is between -0.4 to 5.6, molar refractivity (MR) was between 40 to 130 for the total number of atoms; the qualifying range was between 20 and 70 atoms in a small molecule (Ghose, Viswanadhan, & Wendoloski, 1998, 1999); Veber, ≤ 10 rotatable bonds and a TPSA equal to or less than 140 \AA^2 with 12 or fewer H-bond donors and acceptor (Veber et al., 2002); Egan, WLOGP ≤ 5.88 and TPSA ≤ 131.6 , respectively (Egan, Merz, & Baldwin, 2000); Muegge, molecular weight between 200 to 600 Da, XLOGP between -2 and 5, TPSA ≤ 150 , number of rings ≤ 7 , number of carbon atoms > 4 , number of heteroatoms > 1 , number of rotatable bonds ≤ 15 , H-bond acceptor ≤ 10 , H-bond donor ≤ 5 (Ranjith & Ravikumar, 2019); and bioavailability score, $F > 10\%$ (Martin, 2005). Medicinal chemistry friendliness was predicted by PAINS (Baell & Holloway, 2010), Brenk (Brenk et al., 2008), lead-likeness (Teague, Davis, Leeson, & Oprea, 1999), and synthetic accessibility (from 1 is very easy to 10 is very difficult) methods. The Molinspiration server was used to evaluate the biological activities of the compounds where the bioactivity score < -5.0 is regarded as a biologically inactive compound, -5.0 to < 0 as a moderately active compound, and > 0 is regarded as a biologically active compound (Grob, 2021). ProTox-II server was used to evaluate the toxicity of the compounds (Banerjee, Eckert, Schrey, & Preissner, 2018).

QSAR model using powerful machine learning algorithm

STopTox web portal was used to identify hazardous potentiality in QSAR (Quantitative Structure-Activity Relationships) model (Borba et al., 2022). The acute toxicity tests included acute oral toxicity (OECD TG 401, 420, 423, and 425 and compounds 8495) in the rat model, acute dermal toxicity (OECD TG 402 and compounds 1884) in rabbit and rat models, acute inhalation toxicity (OECD TG 403 and 436 and compounds 681), skin irritation and corrosion (OECD TG 404 and compounds 1012) by Draize test in the rabbit model, eye irritation and corrosion (OECD TG 405 and compounds 3547) via Draize test in the rabbit model and skin sensitization (OECD TG 429 and 442 and compounds 1000) via LLNA test in mouse and guinea pig. Random Forest was the machine-learning algorithm for all tests. The predicted fragment contribution of toxic effects was accompanied by the map of the atomic contributions to toxicity where the red regions with continuous lines indicated the fragment was predicted to increase toxicity and green regions with dashed lines indicated the fragment was predicted to decrease toxicity.

RESULTS AND DISCUSSION

Protein chain selection and properties determination

Chain A (GI: 2425373090) is a receptor-binding domain (RBD) of the S1 subunit of severe acute respiratory syndrome coronavirus 2 spike (S) proteins (superfamily member cd21480). According to NCBI Conserved Protein Domain Family, this part plays the most important role in viral attachment, fusion, and entry into host cells, and serves as a major target for the development of neutralizing antibodies, inhibitors of viral entry, and vaccines. The summary of this analysis included pssm-Id.: 424109, Cd length: 223, bit score: 394.08, and e-value: 1.91e-141. The physicochemical properties of Chain A from S1 subunit C-domain included the amino acids: 215, molecular weight: 24210.41, theoretical pI: 8.73, aliphatic index: 73.81, grand average of hydropathicity (GRAVY): -0.230, instability index: 19.43 and stable, estimated half-life: 1 hours (mammalian, reticulocytes, in vitro), total number of negatively charged residues (Asp + Glu): 15 and total number of positively charged residues (Arg + Lys): 20.

Homology analysis and phylogenetic tree

Homology analysis and phylogenetic tree from NCBI blastp proved that the segment of Chain A from S1 subunit one was a part of the coronavirus glycoprotein family (Figure 1) where the homology method was MUSCLE, the statistical method was neighbor-joining and the bootstrap replications was 1000.

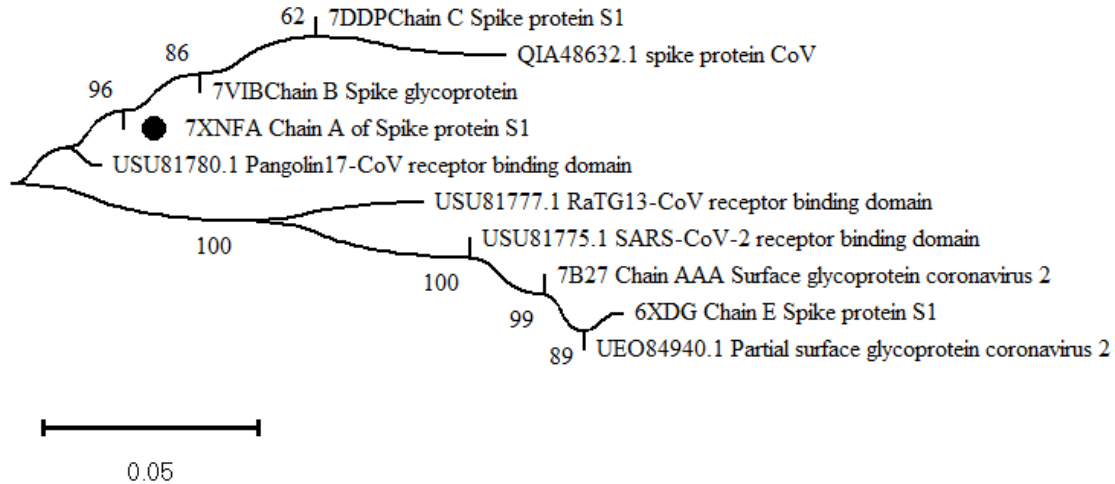


Figure 1. Phylogenetic tree of chain A from S1 subunit and other related protein parts.

Structure prediction of chain A of spike protein S1

The secondary structure of chain A of spike protein S1 of coronavirus was analyzed by PSIPRED and SOPMA (Id.: 9254fa70e19f, started: 20230313-075340 and name: chainx0 A of Spike protein S1). The analysis results of PSIPRED are visualized in Figure 2. In SOPMA, the random coil was found to be the most predominant at 56.28% followed by extended strand, alpha helix, and beta-turn were 27.44, 9.30, and 6.98 %, respectively.

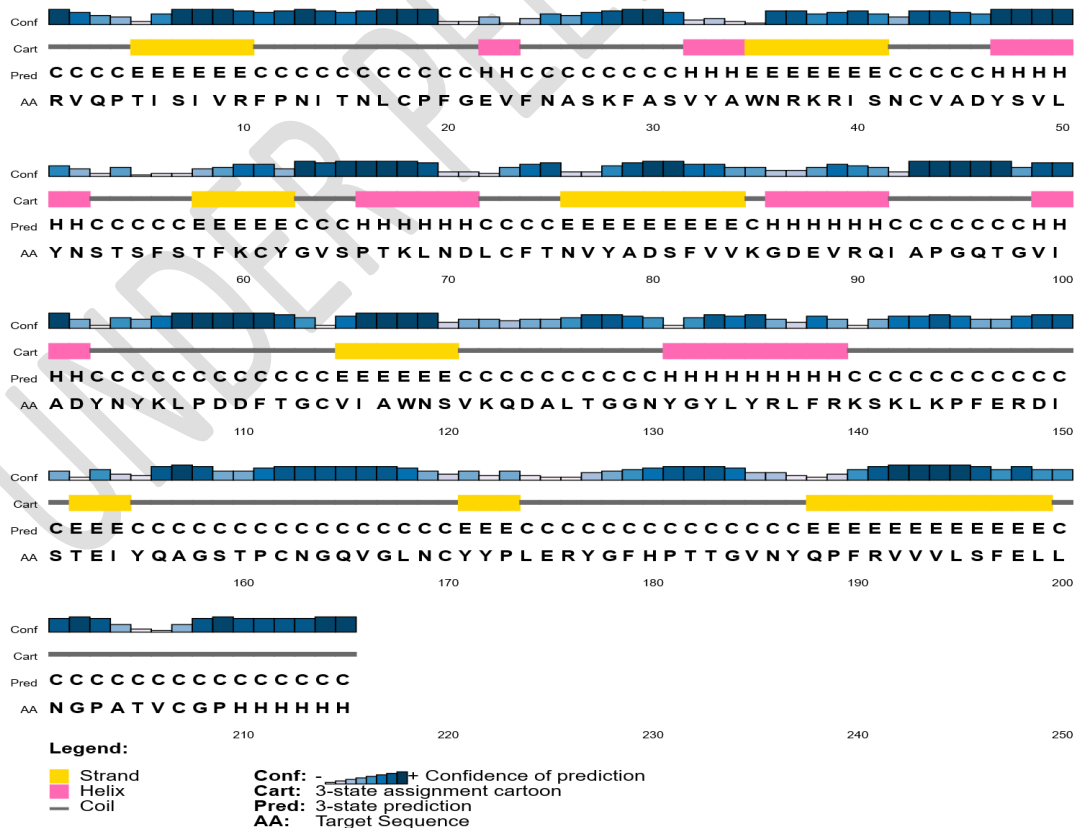


Figure 2. The structural properties of amino acids of chain A from the S1 subunit.

Preparing and validation of macromolecule

The 3D model of the Cryo-EM structure of PCoV_GX spike glycoprotein A and the cleaned polarized chain A of spike protein S1 is given in Figure 3. The summary of Ramachandran plots of the Cryo-EM structure of PCoV_GX spike glycoprotein A and the cleaned polarized chain A of spike protein S1 of coronavirus are given in Table 1. The percent of amino acid residues of the Cryo-EM structure of PCoV_GX spike glycoprotein A and the chain A of spike protein S1 fell within the most favored region in the “Ramachandran plot” based on PROCHECK result. In ProSA, the amino acid number was 209 and the Z score was 5.97.

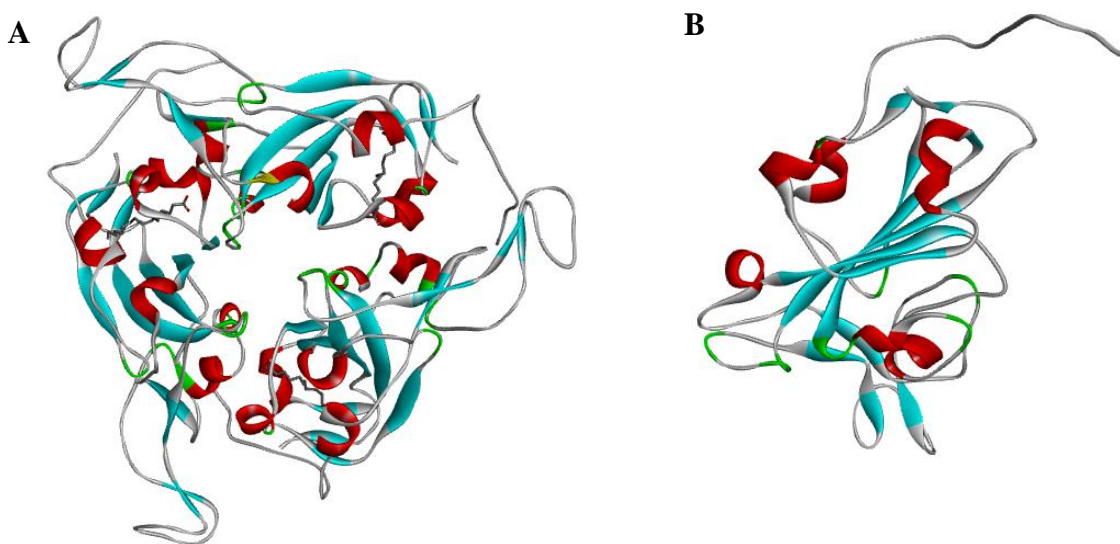


Figure 3. Visualization of glycoprotein and Chain A: (A) 3D model structure of Cryo-EM structure of PCoV_GX spike glycoprotein A with 3 chain and 3 ligands and (B) 3D model structure of only the chain A of Spike protein S1.

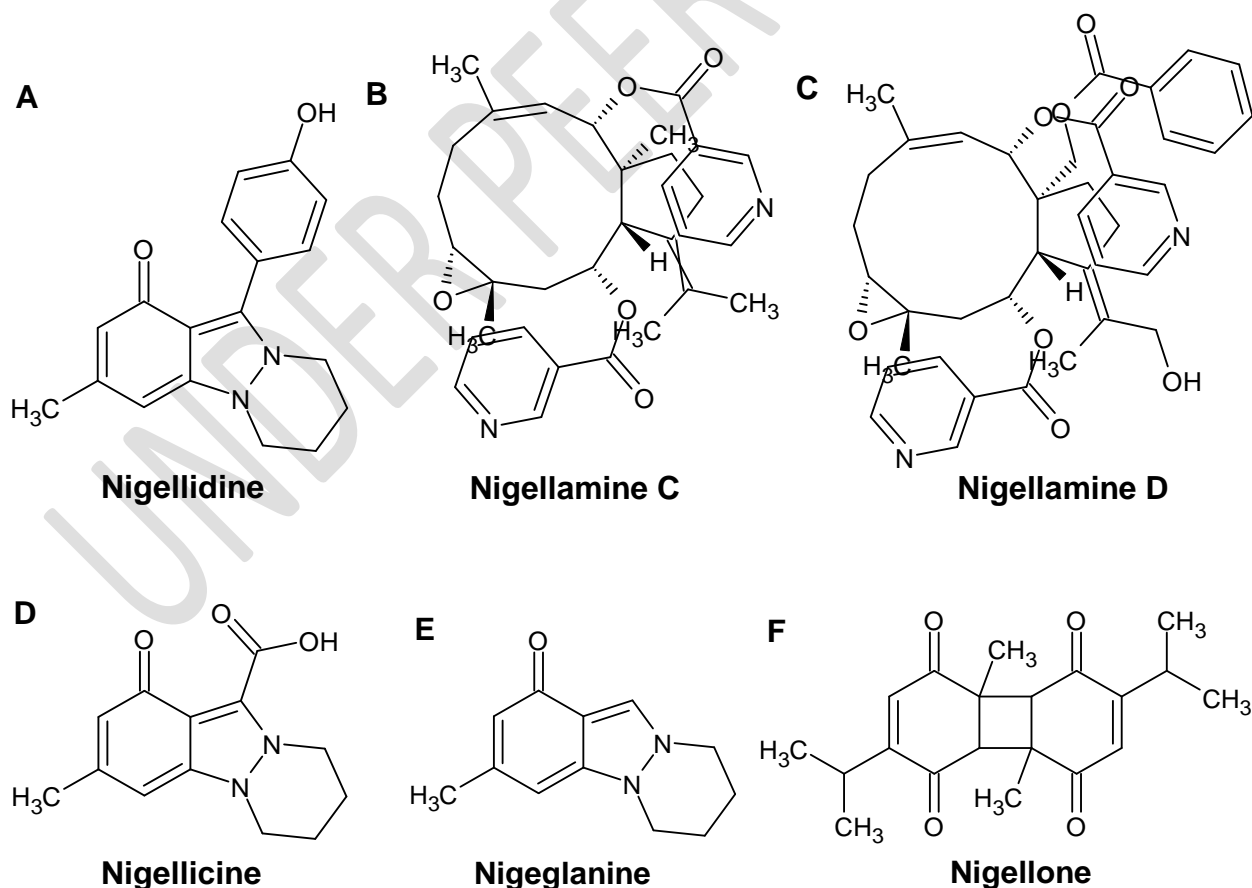
Table 1 The summary of Ramachandran plots

Cryo-EM structure of PCoV_GX spike glycoprotein A			Cleaned polarized chain A of Spike protein S1		
Plot statistics	AA residues Number	%	Plot statistics	AA residues Number	%
Residues in the most favored regions [A, B, l]	483	89.9	Residues in the most favored regions [A, B, l]	162	90.5
Residues in the additional allowed regions [a, b, l, p]	53	9.9	Residues in the additional allowed regions [a, b, l, p]	16	8.9
Residues in the generously allowed regions [~a, ~b, ~l, ~p]	1	0.2	Residues in the generously allowed regions [~a, ~b, ~l, ~p]	1	0.6

Residues in disallowed regions	0	00 (Total 100)	Residues in disallowed regions	0	00 (Total 100)
Number of nonglycine and nonproline residues	537	-	Number of nonglycine and nonproline residues	179	-
Number of end-residues (excl. Gly and Pro)	6	-	Number of end-residues (excl. Gly and Pro)	1	-
Number of glycine residues (shown as triangles)	48	-	Number of glycine residues (shown as triangles)	16	-
Number of proline residues	39	-	Number of proline residues	13	-
Total number of residues	630	-	Total number of residues	209	-

3.5 Ligand and therapeutic drugs

The 2D structures of selected phytochemicals such as nigeplanine, nigellamine C, nigellamine D, nigellicine, nigellidine, and nigellone as ligands and selected therapeutic drugs such as ribavirin, favipiravir, remdesivir, and nirmatrelvir are given in Figure 4.



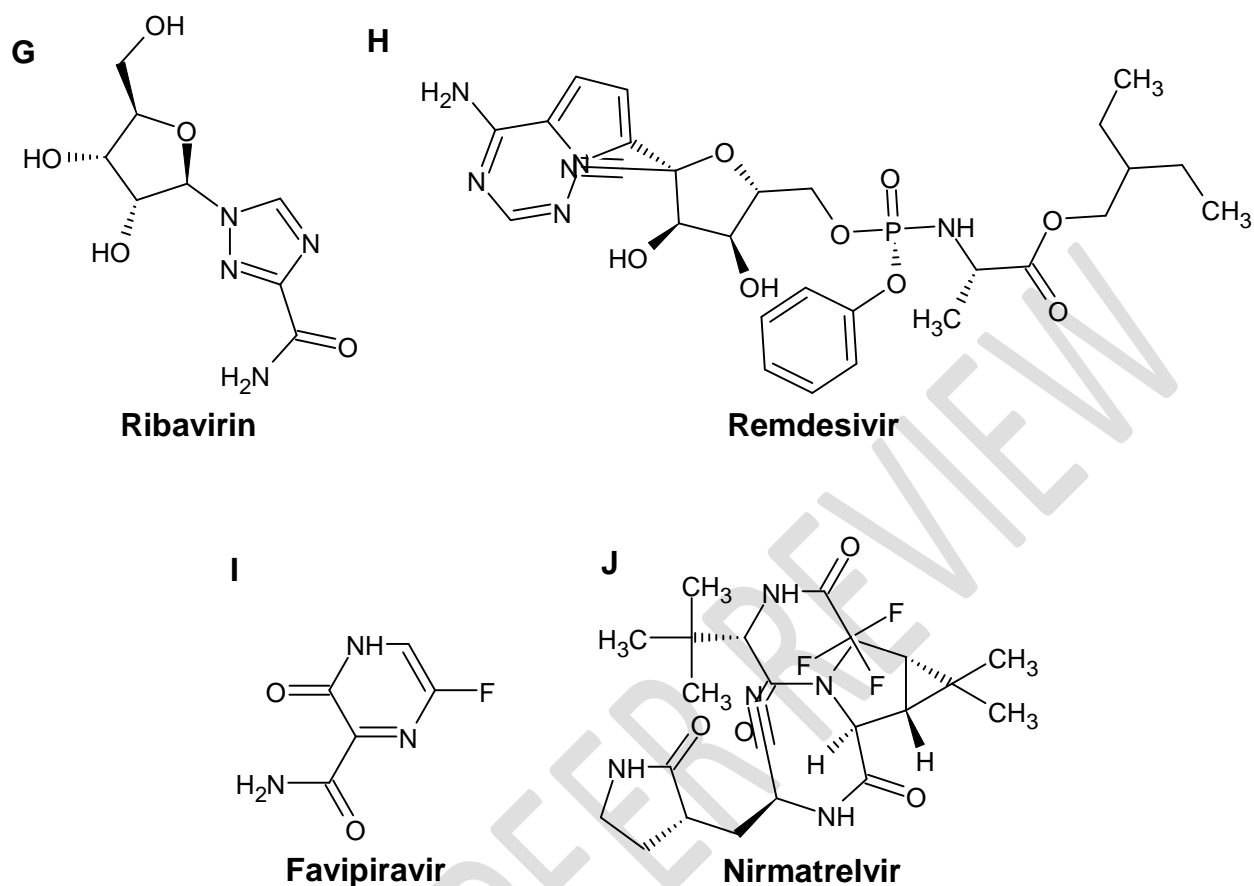


Figure 4. 2D chemical structure of different metabolites and commonly used COVID-19 therapeutics.

Binding affinity and interaction between chain A and ligands

Identification of non-covalent interactions such as hydrophobic contacts, hydrogen bonds, salt bridges, etc., and the nature of the interaction of bioactive molecules with their target is very important in drug design because these are stabilizing factors of target-ligand interaction (Anighoro, 2020). All six selected compounds from *N. Sativa* showed potent binding affinity against chain A of spike protein S1 of SARS-CoV-2 (Table 2). Among them, nigellamine C showed the highest binding affinity of -7.9 kcal/mol where the contact residues were CYS18, ILE40, ALA45, VAL77, LEU195, VAL206, LEU50, ILE116, PHE24, PHE56, ALA45, LEU195, LEU195, TYR47 (Figure 5). Four different antiviral agents that had been used in the treatment of COVID-19 patients showed less binding affinity compared with these bioactive compounds of *Nigella*. Nigeglanine, nigellamine D, nigellicine, nigellidine, and nigellone showed binding affinity of -7.5, -7.3, -6.5, -7, and -6.7 kcal/mol, respectively whereas ribavirin, favipiravir, remdesivir, and nirmatrelvir showed -5.1, -4.8, -6.2, and -5.9 kcal/mol, accordingly (Table 2). Nigeglanine, nigellicine, and nigellidine showed mostly hydrophobic interaction while nigellamine C, nigellamine D, and nigellone showed conventional hydrogen bond interaction.

Table 2 Details of binding affinity and interaction between chain A and ligands

Compound name	Molecular formula	Molecular weight	Binding affinity (kcal/mol)	Residue in contact	Interaction type	Distance
Nigeglanine	C ₁₂ H ₁₄ N ₂ O	202.25	-7.5	CYS18	Alkyl	4.58597
				ILE40	Alkyl	4.80409
				ALA45	Alkyl	4.49616
				VAL77	Alkyl	4.34818
				LEU195	Alkyl	5.03289
				VAL206	Alkyl	5.3736
				LEU50	Alkyl	4.95935
				ILE116	Alkyl	4.8034
				PHE24	Pi-Alkyl	4.6959
				PHE56	Pi-Alkyl	5.38877
				ALA45	Pi-Alkyl	4.40149
				LEU195	Pi-Alkyl	4.80967
				LEU195	Pi-Alkyl	5.21649
				TYR47	Pi-Pi Stacked	5.54815
				Nigellamine C	C ₃₂ H ₃₈ N ₂ O ₅	530.7
PHE56	Hydrogen Bond	2.1157				
PRO66	Hydrogen Bond	2.94736				
TYR51	Hydrogen Bond	2.24315				
PHE59	Pi-Sigma	3.57235				
Nigellamine D	C ₃₉ H ₄₂ N ₂ O ₈	666.8	-7.3	SER55	Hydrogen Bond	2.35783
				PHE59	Hydrogen Bond	2.52482
				TYR51	Hydrogen Bond	2.47743
				THR54	Hydrogen Bond	2.82209
				PRO66	Hydrogen Bond	2.89098
Nigellidine	C ₁₃ H ₁₄ N ₂ O ₃	246.26	-6.5	THR54	Hydrogen Bond	2.31726
				PHE56	Pi-Pi T-shaped	5.00223
				PHE59	Pi-Pi T-shaped	4.95843
				PHE59	Pi-Pi T-shaped	5.60944
				TYR47	Pi-Alkyl	4.14739
Nigellone	C ₁₈ H ₁₈ N ₂ O ₂	294.3	-7	TYR51	Hydrogen Bond	2.83052
				TYR47	Pi-Pi Stacked	5.40681
				TYR51	Pi-Pi Stacked	5.33247
				PHE59	Pi-Pi T-shaped	4.73143
				PHE59	Pi-Pi T-shaped	5.44998
				TYR47	Pi-Alkyl	4.00084
				PHE59	Pi-Alkyl	4.11259
				TYR51	Hydrogen Bond	2.28764
				TYR51	Hydrogen Bond	2.28764
Ribavirin	C ₈ H ₁₂ N ₄ O ₅	244.20	-5.1	PHE56	Hydrogen Bond	1.94621
				THR54	Hydrogen Bond	2.35982
				THR54	Hydrogen Bond	2.35982
Favipiravir	C ₅ H ₄ FN ₃ O ₂	157.10	-4.8	CYS18	Pi-Sulfur	5.79321
				PHE74	Pi-Pi Stacked	4.52437
				ALA45	Pi-Alkyl	4.00721
				VAL77	Pi-Alkyl	5.27546
				LEU195	Pi-Alkyl	5.0039
Remdesivir	C ₂₇ H ₃₅ N ₆ O ₈ P	602.6	-6.2	TYR51	Hydrogen Bond	2.25127
				THR54	Hydrogen Bond	2.62554
				TYR51	Hydrogen Bond	2.15073
				TYR47	Pi-Pi Stacked	5.27052

Nirmatrelvir	C ₂₃ H ₃₂ F ₃ N ₅ O	499.5	-5.9	PHE56	Pi-Pi T-shaped	5.00422
				PHE59	Pi-Pi T-shaped	4.78039
				TYR51	Pi-Alkyl	4.95075
				THR58	Hydrogen Bond	2.34192
				ARG90	Hydrogen Bond	2.33035
				GLY86	Hydrogen Bond	1.80821

The 3D surface view, 3D tube pose view, H bond donor and acceptor scenario, 3D protein-ligand binding interaction, and 2D protein-ligand binding interaction of nigellamine C are given in Figure 6. Nigellamine C was selected for further analysis because of its highest binding affinity and the presence of hydrogen bonds to chain A of spike protein S1 of SARS-CoV-2.

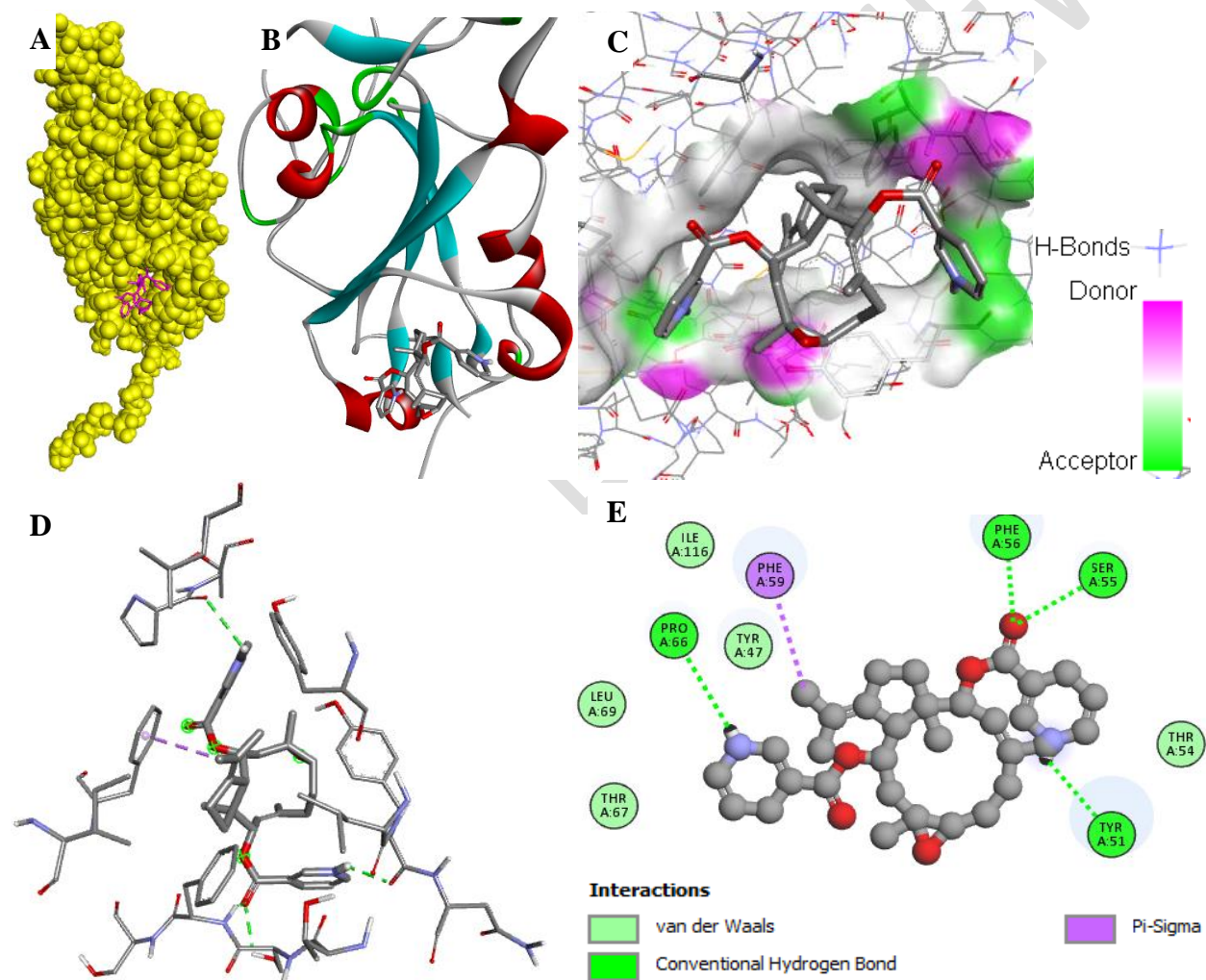


Figure 5. Binding modes for the nigellamine C within the active site of the chain A of the spike protein of coronavirus: (A) 3D surface view; (B) 3D tube pose view; (C) Hydrogen bond donor and acceptor; (D) 3D protein-ligand binding interaction and (E) 2D protein-ligand binding interaction. The superimposed structure of pre and post-molecular dynamics simulation of the protein-ligands complex is given in Figure 6A. Comparison between the surface view of initial

and final structures of the complexes from MD simulations at the snapshot of 4 psec are presented in Figure 6B.

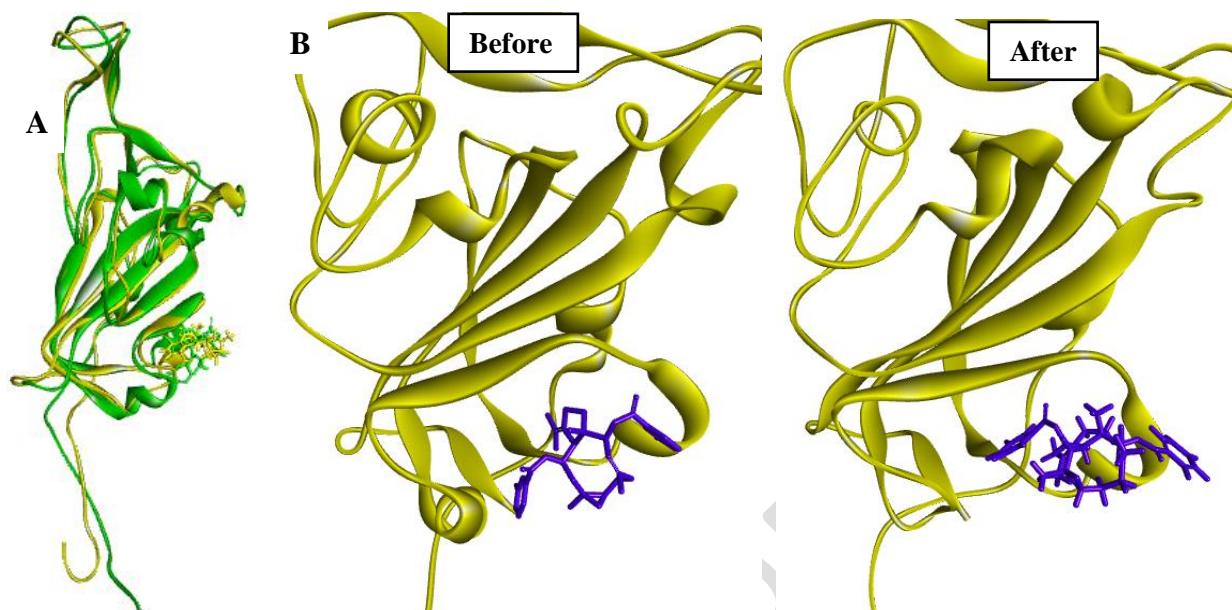


Figure 6. The superimposed structure of pre and post-molecular dynamics simulation nigellamine C. (A) The yellow color denotes the pre-molecular dynamics structure, and the green color denotes the post-molecular dynamics structure, and (B) comparing surface view between the initial and final structures of the complexes from MD simulations at the snapshot of 4 psec.

ADME/T and bioactivity analysis

Absorption, distribution, metabolism, excretion, and toxicity properties together are called ADME-Tox properties prediction and play an important role in drug development because they account for the failure of 60% of drug molecules during the drug development process (Mandlik, Bejugam, & Singh, 2016). Thus, cost-effective in-silico tools early prediction of ADME analysis of nigellamine C is given in Table 3 where the intestinal absorption property was 98%, and the total clearance amounts was 0.37 log ml/min/kg. Physiochemical properties and lipophilicity were also predicted via SMILES where the TPSA (the polar surface area) was 90.91 Å² and the consensus log $P_{o/w}$ was 4.80 for nigellamine C.

Biological activities, drug-likeness, and medicinal properties of nigellamine C are given in (Table 3) where positive drug-likeness properties were shown in all models Lipinski, Ghose, Veber, Egan, and Muegge and the bioavailability score was 0.55. Nigellamine C showed 0 alerts in PAINS but 3 alerts in the Brenk assay and a negative response in the lead likeness assay. The synthetic accessibility score was 6.05, nigellamine C. Nigellamine C belongs to the toxicity class 6. The LD₅₀ value was 10000 mg/kg.

Table 3 ADME properties of the selected phytochemicals

ADME properties		Other properties	
Absorption		Biological activities	
Water solubility (log mol/L)	-4.664	GPCR ligand	-0.00
Intestinal absorption in human (%)	98.377	Ion channel modulator	-0.19
Skin Permeability (log Kp)	-2.762	Kinase inhibitor	-0.39
P-glycoprotein substrate	No	Nuclear receptor ligand	0.31
Distribution		Protease inhibitor	0.04
VDss in humans (log L/kg)	0.12	Enzyme inhibitor	0.31
Fraction unbound in humans (Fu)	0	Druglikeness	
BBB permeability (log BB)	-0.799	Lipinski	Yes; 1 violation: MW>500
CNS permeability (log PS)	-2.957	Ghose	No; 4 violations: MW>480, WLOGP>5.6, MR>130, #atoms>70
Metabolism		Veber	Yes
CYP2D6 substrate	No	Egan	No; 1 violation: WLOGP>5.88
CYP3A4 substrate	Yes	Muegge	Yes
CYP1A2 inhibitor	No	Bioavailability score	0.55
CYP2C19, CYP2C9 and CYP2D6 inhibitor	No	Medicinal chemistry	
CYP3A4 inhibitor	Yes	PAINS	0 alert
Excretion		Brenk	3 alerts: Three-membered_heterocycle, isolated_alkene, more_than_2_esters ?
Total Clearance (log ml/min/kg)	0.37	Leadlikeness	No; 2 violations: MW>350, XLOGP3>3.5
Renal OCT2 substrate	No	Synthetic accessibility	6.05

Nigellamine C showed inactive in hepatotoxicity, mutagenicity, and cytotoxicity as well as inactive in important targets such as aryl hydrocarbon receptor (AhR), androgen receptor (AR), androgen receptor ligand binding domain (AR-LBD), aromatase estrogen Receptor Alpha (ER), estrogen receptor ligand binding domain (ER-LBD), peroxisome proliferator-activated receptor gamma (PPAR-Gamma), nuclear factor (erythroid-derived 2)-like 2/antioxidant responsive element (nrf2/ARE), heat shock factor response element (HSE), phosphoprotein (tumor suppressor) p53 and ATPase family AAA domain-containing protein 5 (ATAD5).

QSAR analysis

Nigellamine C did not produce any acute systemic and topical toxicity nature in the QSAR model where the confidence level of all tests was $\geq 50\%$ Figure 7.

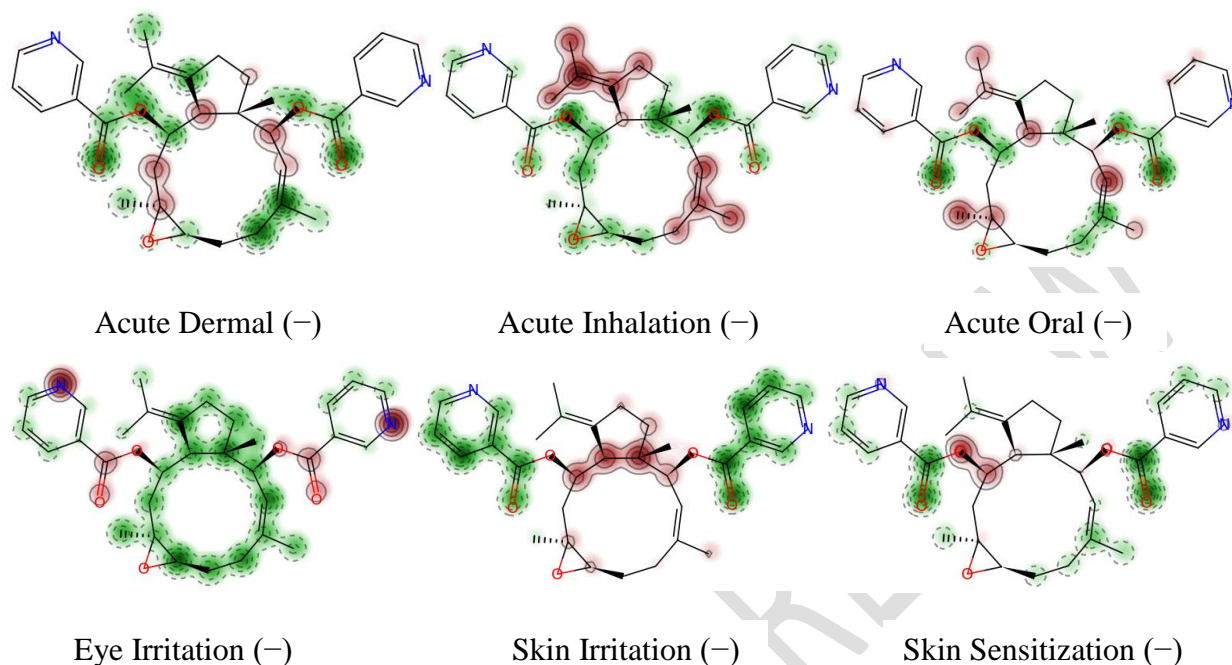


Figure 7. Maps of fragment contributions and predictions of Nigellamine C. (Red regions with continuous lines indicate the fragment is predicted to increase toxicity, the green regions with dashed lines indicate the fragment is predicted to decrease toxicity and the gray isolines define the frontier between the positive (red) and the negative (green) contributions. (+) denotes the toxic and (-) denotes the non-toxic nature).

Summary of results and probable mode of action of Nigella compounds

Compared with the COVID-19 drugs ribavirin, favipiravir, remdesivir and nirmatrelvir with nigellamine C showed more binding affinity against chain A of the spike protein of coronavirus in molecular docking analysis. Nigellamine C also proved itself as a probable potential drug candidate in ADME/T and QSAR analysis. It can be considered for inhalation and oral drug development. Generally, coronavirus binds to ACE-2 (Angiotensin-converting enzyme 2), heparan sulfate, sialic acid receptor of ciliated cells in nasal tissue cells, CD147, GRP78 (glucose-regulating protein 78), lectins (CD209L/CD209), and integrins of epithelial cells of lung tissue, DPP4 (dipeptidyl peptidase-4) /CD26 endothelial cells of the blood vessel, vimentin of pneumocyte of alveoli, NRP-1 (neuropilin-1) in epithelial cells in the olfactory tissue and bone marrow-derived macrophages, AXL (tyrosine-protein kinase receptor) of pulmonary epithelia (Eslami et al., 2022). A probable mode of action of this compound is given in Figure8.

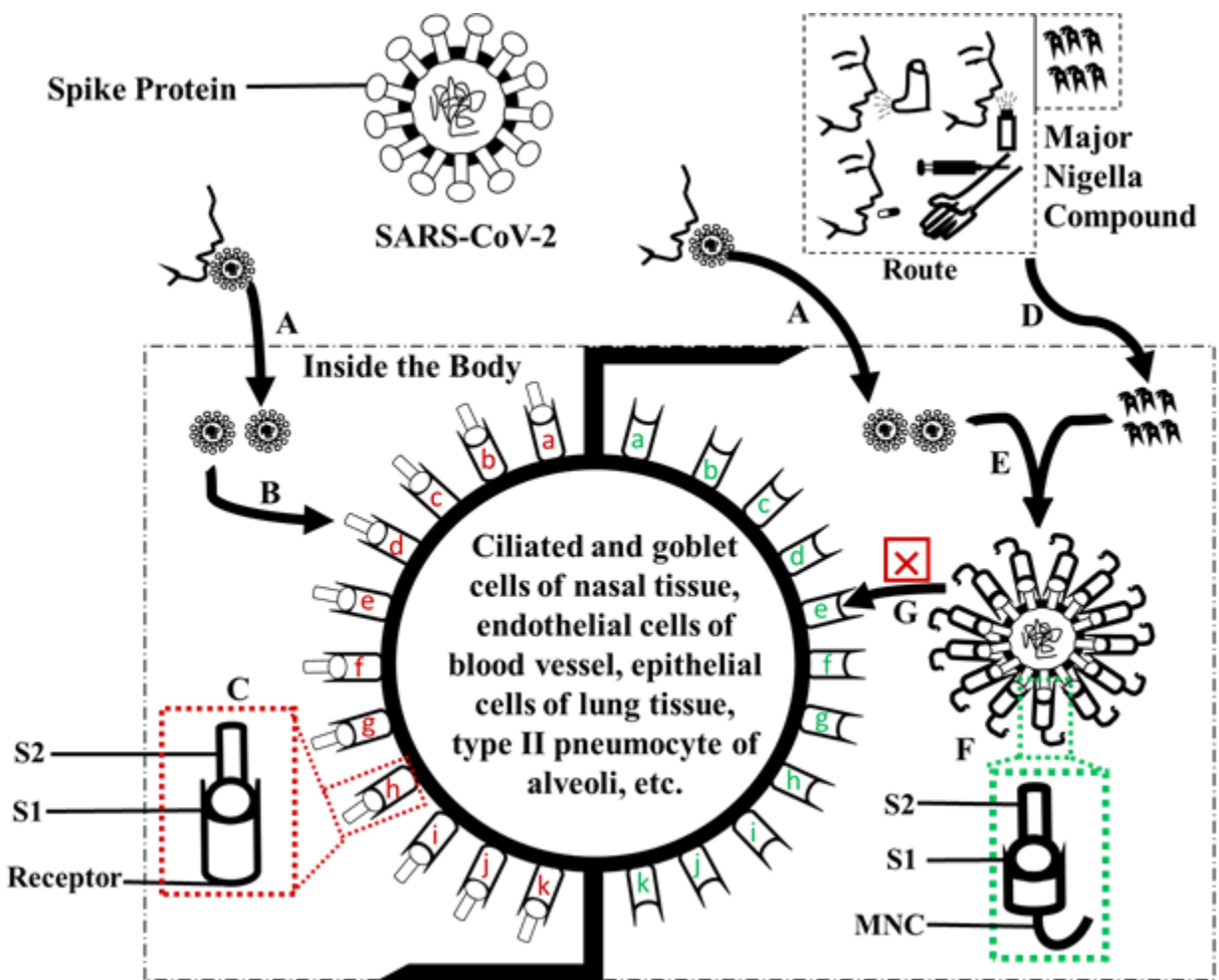


Figure 8. Probable mode of action of Nigellamine C as a drug candidate.

CONCLUSION

Pharmaceutical industries, food and ornamental industries, medicine, chemical industries, and research laboratories give attention to traditional medicinal plants because these are the sources of bioactive natural compounds. Nigellamine C has proved itself as a more potent antiviral drug candidate among other compounds based on molecular docking, ADME/T, and QSAR analysis. This study will help in advancing phytochemical-dependent antiviral drug development in the future.

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