Identification of potential inhibitors of SARS-CoV-2 from *Artemisia annua* compounds by *In silico* evaluation and their density functional theory (DFT)

Abdirahman ELMI*1, Ahmed Said MOHAMED1, Nazia SIDDIQUI2, Syad AL JAWAD3, Moustapha NOUR4, Idriss MIGANEH1 and Saleem JAVED5*

1 Medicinal Research Institute, Centre d’Etudes et de Recherche de Djibouti, IRM-CERD, Route de l’Aéroport, Djibouti
2USIC, Dayalbagh Educational Institute, Agra, India 282005
3International Islamic University Chittagong, Department of pharmacy, Bangladesh
4Centre d’Etudes et de Recherche de Djibouti, ISV-CERD, Route de l’Aéroport, Djibouti
5Department of Chemistry, Institute of H. Science, Dr. B. R. Ambedkar University, Agra, India 282002

Abstract

The genus Artemisia has recognized medicinal value and its use by humans dates back to centuries ago. With the appearance of the new coronavirus, end of 2019, several countries have recommended the use of herbal teas consisting mainly of *Artemisia*. The individual analysis of the constituents of this species is crucial to characterize and optimize its antiSARS-CoV-2 action. We evaluated by molecular docking the inhibitory action of major compounds of the *Artemisia* genus (*Artemisinin, Artemannin B, Alpha Thujone, P-Hydroxyacetophenone, Fisetin, Cirsimaritin, Capillin, β-Strosterol, and Quercetin*) against three targets namely SARS-CoV-2 main protease (Mp), SARS-CoV-2 receptor binding domain (RBD) and human furin protease (HF protease). The two flavonols, quercetin and fisetin, have the best binding energies with the three targets. Quercetin/Fisetin possesses binding energy of -7.17/-6.9, -6.3/-6.15 and ~5.98/- 5.49 kcal/mol with MP, RBD and HF protease respectively. Their physicochemical properties meet the requirements of an oral active principle and are not toxic according to predictive simulations. Thereby DFT calculation has been used to analyze the electronic and geometric characteristics of these two compounds. The gap energies were also deduced for the stable structure and their reactivity. The abundance of Quercetin in different plants may be another advantage in the use of this bio-compound in the treatment of coronavirus.

Keywords: *Artemisia annua*, DFT, Docking Molecular, SARS-CoV-2, Quercetin and Fisetin

1 INTRODUCTION

*Artemisia* is one of the most widespread genera in the Asteraceae family. It is a heterogeneous genus, composed of more than 500 different species distributed mainly in the temperate zones 1. A large number of studies have been carried out to determine precisely the chemical compositions of the *Artemisia* species, especially for the two species *A. annua* and *A. afra* which have high medicinal value. Sesquiterpene lactones, phenolic compounds and flavones have been identified in *A. annua* 2-7 and *A. afra* contains monoterpenoids, sesquiterpenes, glucolides, guaianolides, flavonoids 8. Many *Artemisia*-like plants exhibit strong antiviral properties. One of the first studies was carried out by M. M. Abid Ali Khan and al., in 1991 regarding the evaluation of inhibitory activity against tobamovirus. This work has shown promising results for some of the tested plant extracts such as *A. annua*. The active virus inhibitor effect has been shown to be the result of the presence of a mixture of low-molecular-weight sterols 9.

Also another study showed that *A. annua* has an antiviral potential greater than that of 20 other medicinal plants tested. The good antiviral activity of *A. annua* can be explained by the presence of sterols having better antiviral activity than artemisinin or artanuin-B, two characteristic molecules for this species 10.
Quercetin is also one of the naturally occurring substances found in *Artemisia A. annua* shown to be effective for a wide range of antiviral use.  

In the current context, the genus *Artemisia* is also one of the candidates for the fight against the global pandemic which has caused a major health crisis. The *A. annua* is considered as a possible treatment for Covid-19, but trials should be performed to assess its effectiveness and determine its adverse effects. Historically, in 2005, a study was carried out by the team of Shi-you Li, confirms that the ethanolic extract of *A. annua* exhibits anti-SARS-CoV activity with EC50 = 34 ± 2.6 µg/ml.  

Since new coronavirus appeared, several studies have been carried out to discover natural substances extracted from species of the genus “Artemisia” and examine their potential as inhibitors of the virus.  

Recently, in April 2020, the Madagascan president promoted the name of an anti-Covid remedy, called “The Covid-Organics” (CVO): an improved traditional remedy based on *Artemisia* and medicinal plants from Madagascar. Combined with conventional therapies, herbal teas show an attenuation of respiratory symptoms. It is possible that molecules of *A. annua* play a role in blocking the two main receptors (serum protease, ACE2 receptor) of the cell membrane, i.e. the two main keys that allow the virus to access the cell (*Artemisia* and Covid-19: the remedy Madagascar boosts Africa, Paris Match, Published on 05/18/2020).

It is known also that the *A. annua* has always been regarded as a reference to clear heat and eliminate dampness. For this reason, *A. annua* acts as a pivotal medicine in the prescription of treatment of SARS. Similar, to SARS, Covid-19 is also a coronavirus that causes respiratory syndromes, which start with dampness, with a long latency and courses. The pathogenesis of Covid-19 is mainly characterized by the inward invasion of dampness towards the transformation into heat. Covid-19 treatment should focus on clearing heat, removing dampness and resolving phlegm. Considering the initial clinical manifestations of Covid-19 characterized by pathogen invading Shaoyang gallbladder meridian, *A. annua* is expected to be an essential drug for the treatment of Covid-19.

The natural product tested against this coronavirus reacted largely well against the spike glycoprotein (S) on the surface of SARS-CoV-2 precisely, by binding the domain of SARS-CoV-2 spike protein and SARS-CoV-2 main protease. In this present study we also targeted furin, a kind of proprotein convertases. Nine compounds from the *Artemisia annua* are tested by *Insilico* for their affinity with these three targets. Molecular simulation by density functional theory is carried out on the compounds with the best bond energies.

### 2 MATERIALS AND METHODS

#### 2.1 Compounds tested

The genus *Artemisia* is rich in phytocompounds and the presence of three major families of secondary metabolites has been reported. There are basically nine compounds found in the *Artemisia annua*. They have been stimulated to determine their anti-SARS-CoV-2 potential. These compounds are: Artemisinin, Arteannuin B, Alpha Thujone, P-hydroxycacetophenone, Fisetin, Cirsimaritin, Capillinn, β-Sitosterol and Quercetin.

#### 2.2 Docking targets

The targets are SARS-CoV-2 main protease (Mp), SARS-CoV-2 receptor binding domain (RBD) and human furin protease (HF protease). These targets have a crucial role in the virus propagation. In addition SARS-CoV-2 Mp is present in most viruses of the coronavirus family and therefore its functioning is widely described in the literature.

### 2.3 Docking protocol

The description of molecular docking, prediction of pharmacokinetic parameters as well as the toxiological property was carried out in our previous article.

#### 2.4 Density Functional Theory

Density Functional Theory (DFT) is currently the most successful and promising quantum chemical approach to calculate ground state properties of atoms and molecules. In this paper, all calculations were done by using the B3LYP method and 6-311++G(d,p) basis set. The computations for the present work has been performed using the Gaussian 03W and ORCA 4.0.1. All vibrational wavenumbers and geometrical parameters were computed as decrypted in. The topological parameters were determined using Multiwfn software. All the graphs were plotted using Origin 8.0 software or Multiwfn software.

### 3 RESULTS AND DISCUSSIONS

#### 3.1 Bioavailability parameters and Toxicity of selected compounds

When designing a drug, the study of the physico-chemical parameters of the candidate molecule is essential. These parameters predict the behavior of the compound during its oral administration and its bioavailability. Two rules govern the analysis of these parameters: Lipinski and Veber. According to the first rule, the candidate molecules must have a value equal or multiple of 5. The optimal properties and the corresponding values are: number of active hydrogen atoms (HBD > 5), number of binding sites for hydrogen atoms (HBA > 10), logarithm of the octanol / water partition (LogP < 5), molecular weight (MW < 500 Da) Beyond two unsatisfied conditions, a candidate molecule is discarded in the drug research process. Our nine compounds from *Artemisia annua* fulfill these two rules (Table A, Support information).

Also, a predictive evaluation of the possible toxicity of these compounds was made with four different parameters namely Ames toxicity, Carcinogens, Acute oral toxicity and Rat acute toxicity. They have low toxicity whatever the parameters considered (Table B, Support information).

#### 3.2 Molecular docking

The search for therapeutic molecules is carried out in different stages. Molecular modelling makes it possible to pre-select the candidate molecules for a given target. The hydrogen bond between the ligand (therapeutic molecule) and the protein/enzyme of the pathogen is studied and a binding energy is measured. A low binding energy corresponds to a good affinity between the ligand and the biological target.

Binding energies range from -2.519 for Artemisinin at the HF protease site to -7.169 for Quercetin at the SARS-CoV-2 Mp site (Table 1). On the SARS-CoV-2 RBD target site, six compounds present in *Artemisia sp* (66% of compounds tested) have better binding energies than hydroxychloroquine which is very widely used against Covid-19. And this SARS-CoV-2 RBD target is interesting since this part allows this virus to attack to the host cell. By blocking this receptor with inhibitors like these biomolecules, the virus will not be able to attach itself to...
cells. Recalling, moreover, that the virus is essentially made up of proteins surrounding genetic material and needs the cellular tools to multiply. Compounds with a better binding energy than one of the reference drugs are shown in 2D to better visualize their interactions with the amino acid residues of the targets (Figure 1). Quercetin, having a better score than the two references, is represented in 3D (Figure 2).

The first two targets (SARS-CoV-2 Mp and SARS-CoV-2 RBD) located on the virus  have a better binding energy, twice as small as that of the third human furin protease target, -4.825, -4.916 and -2.519 kcal/mol respectively. The action of this active compound in A annua would be directed towards the virus rather than host cell protection. This observation is applicable to its derivative Arteannuin B which has binding energies -5.075 and -5.604 and -3.342 kcal/mol at the SARS-CoV-2 Mp, SARS-CoV-2 RBD and human furin protease sites respectively. Against SARS-CoV-2 RBD these two compounds are better than hydroxychloroquine (Table 1).

However, on the three target sites, Quercetin and Fisetin have the best affinities (Table 1). The antiviral action of Quercetin is already obtained against HIV, against influenza virus, herpes virus and against the dengue virus. A recent study has shown that this compound can interfere with 85% of the functions of the SARS-CoV-2 viral proteins inside human cells. This value rises to 93% if Quercetin is combined with vitamin D.
Figure 1: 2D visualization of molecular interaction of SARS-CoV-2 Mp (A), SARS-CoV-2 RBD (B) and human furin protease (C) with the biomolecules having at least better binding energy than one of standard.
Table 1: Molecular docking of biomolecules with SARS-CoV-2 Mp, SARS-CoV-2 RBD and human furin protease.

<table>
<thead>
<tr>
<th>Compound</th>
<th>SARS-CoV-2 Mp</th>
<th>SARS-CoV-2 RBD</th>
<th>human furin protease</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>BE (kcal/mol)</td>
<td>BE (kcal/mol)</td>
<td>Amino acid interaction (H-bond)</td>
</tr>
<tr>
<td>Artemisinin</td>
<td>-4.825</td>
<td>-4.916</td>
<td>-</td>
</tr>
<tr>
<td>Arteannuin B</td>
<td>-5.075</td>
<td>-5.604</td>
<td>TYR363</td>
</tr>
<tr>
<td>Alpha Thujone</td>
<td>---</td>
<td>---</td>
<td>---</td>
</tr>
<tr>
<td>P-Hydroxyacetophenone</td>
<td>-6.182</td>
<td>-5.918</td>
<td>LEU390; TYR365</td>
</tr>
<tr>
<td>Fisetin</td>
<td>-6.907</td>
<td>-6.152</td>
<td>LEU390; ASN388</td>
</tr>
<tr>
<td>Cirsimaritin</td>
<td>-6.107</td>
<td>GLU166</td>
<td>SER393; TYR365; ASN388</td>
</tr>
<tr>
<td>Capillin</td>
<td>-5.489</td>
<td>-4.324</td>
<td>-</td>
</tr>
<tr>
<td>β-Sitosterol</td>
<td>-3.646</td>
<td>---</td>
<td>-3.148</td>
</tr>
<tr>
<td>Quercetin</td>
<td>-7.169</td>
<td>GLU166; HIE163; GLU166; HIE164; GLN189</td>
<td>ILE358; ASN388</td>
</tr>
</tbody>
</table>

*Remdesivir and Hydroxychloroquine used as references.
The two flavonols which have the best scores in docking, Quercetin and Fisetin, are classified in category II for Acute oral toxicity with an LD50 between 50 mg/kg and 500 mg/kg (Table 1). Although the two compounds have very similar structures, the OH group at the 5th position on the A ring and the B ring as well as the hydroxyl number differ. Quercetin has five hydroxyl groups and 4 for Fisetin. Also, the latter has an OH at 5′ of the B ring. The structure-activity study of different flavonoids has shown that the presence of an OH at the 5′B ring decreases the inhibitory action of polyphenols. On the other hand, OH in the A ring, as is the case with Quercetin, is beneficial in the action against rhinovirus.

![Figure 2: 3D visualization of docking analysis of human furin protease binding with quercetin, better binding energy than the two standards.](image)

On the first two targets, the amino acids that the two flavonols interact with are similar. ASN142; HIE163 bond with Fisetin and for Quercetin ASN142; HIE163; GLU166; HIE164; GLN189 on SARS-CoV-2 Mp (Table 1). Likewise, Fisetin bond with LEU390 and ASN388 on SARS-CoV-2 RBD and Quercetin bond with ILE358 and ASN388. On the other hand, against human furin protease, the binding amino acids are totally different for the two flavonols: ALA292, PRO256, ASP258 for Fisetin and LEU227, GLU236, ASH264 for Quercetin. In the latter target, the two compounds can be complementary. On this same target, Quercetin is better than the reference drugs, remdesivir and hydroxychloroquine (Table 1).

The structural study in particular Density Functional Theory of these two flavonols is crucial to have the possible link of their action with the SARS-CoV-2 virus.

3.3 Density Functional Theory of Quercetin and Fisetin

3.3.1 Optimized molecular geometry for quercetin and fisetin

The B3LYP /6-311++G(d,p) optimized structure of both the molecules Fisetin and Quercetin along with their atom numbering are presented in Figure 3. Optimized bond lengths and angles are mentioned in Table C and D (Support Information). The studied compounds retain a C1 point group. The optimized geometry of the studied compounds is in the range of expected bond lengths and bond angles. The C-C, C-O, C-H and O-H bond lengths are in the regular order. C-C bond angles in the ring and C-C-O along with C-O-H showed well expected bond angles as given in Table C and D (Support Information).

![Figure 3: Optimized geometric structure with atom numbering of Fisetin (left) and Quercetin (right), brown-coloured are carbon and red are nitrogen.](image)
3.3.2 Vibrational spectral analysis

The vibrations frequencies, Raman and IR spectrum analysis of both Fisetin and Quercetin compounds have been obtained by using B3LYP/6-311++G(d,p) basis set along with small scaling factor 0.961. The fact that no imaginary frequencies were found in prediction implies that the optimized geometry is located at the local lowest point on the potential energy surface.

<table>
<thead>
<tr>
<th>Parameters</th>
<th>B3LYP/6-311++G(d,p)</th>
<th>Parameters</th>
<th>B3LYP/6-311++G(d,p)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Fisetin</td>
<td>Quercetin</td>
<td>Fisetin</td>
</tr>
<tr>
<td>μx</td>
<td>-1.7573</td>
<td>4.3252</td>
<td>βxxx</td>
</tr>
<tr>
<td>μy</td>
<td>-0.5470</td>
<td>-5.8199</td>
<td>βyxx</td>
</tr>
<tr>
<td>μz</td>
<td>-0.0008</td>
<td>-1.1814</td>
<td>βyy</td>
</tr>
<tr>
<td>μ(D)</td>
<td>1.8405</td>
<td>7.3467</td>
<td>βyyy</td>
</tr>
<tr>
<td>αxx</td>
<td>-92.2677</td>
<td>-93.7067</td>
<td>βxxx</td>
</tr>
<tr>
<td>αxy</td>
<td>17.4231</td>
<td>-0.5036</td>
<td>βyxx</td>
</tr>
<tr>
<td>αyy</td>
<td>-125.1757</td>
<td>-136.0758</td>
<td>βyy</td>
</tr>
<tr>
<td>αxz</td>
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<td>-8.3828</td>
<td>βxxz</td>
</tr>
<tr>
<td>αyz</td>
<td>-0.0032</td>
<td>4.7018</td>
<td>βyzz</td>
</tr>
<tr>
<td>αz0 (e.s.u)</td>
<td>-1.696x10^{-23}</td>
<td>-1.896x10^{-23}</td>
<td>βtot (e.s.u)</td>
</tr>
</tbody>
</table>

Fisetin has 31 atoms and 87 fundamental vibrations while Quercetin has 32 atoms and 90 fundamental vibrations, both showed Cs and C1 point group symmetry. The calculated IR and Raman frequencies together with relative intensities are presented in Table E and F (Support Information). C=O in Fisetin and Quercetin appears at 1612 and 1622 cm\(^{-1}\) respectively. All C-H and C-C bonds vibrations are in the expected range in both the molecules.

The IR spectrum of quercetin shows an intense and thin band at approximately 3300 cm\(^{-1}\), characteristic of bound –OH groups. However, the signatures of –OH bound in Fisetin appear at approximately 3500 cm\(^{-1}\). These differences in intensity and vibration are due to the diversity in the number of –OH groups and the chemical environment of the two molecules respectively. These bands can be explained by the formation of intramolecular hydrogen bonds, generally of the type C = O---HO– and –OH---OH. The number of hydroxyl groups as well as their positions can be an influencing factor for the anti-Covid activity against intermolecular interactions (hydrogen-bridge) between Fisetin or Quercetin and virus. Calculated IR and Raman Spectra are displayed in Figure. 4 and 5 for Fisetin and Quercetin.
Figure 4: Infrared spectra of Fisetin (left) and Quercetin (right) using DFT/6-311++G(d,p).

Figure 5: Calculated computational Raman spectra of Fisetin (left) and Quercetin (right) using DFT/6-311++G(d,p).
### 3.3.3 Molecular electrical potential surface

The contribution of charges may be presented in 3D by utilizing molecular electrical potential (MEP). This prediction facilitates the recognition of interactions of molecule and chemical bond nature. It displays the molecular size, shape, electrostatic potential charge through colour grading. Thus, various physicochemical properties of a molecule can be determined by MEP, like, interaction of nucleic acids with their constituent bases; enzyme-ligand or protein-ligand interactions among others. The different colour in Figure 6 shows distinct values of the electrostatic potential at the surface of the compounds.

The electrostatic potential is arranged from smallest (red) to largest (blue). Here, we are showing deepest red for the negative charge and deepest blue for the positive charge. The colour code of the maps is found to be in the range of (deepest red) -1.70 to +1.70 eV (deepest blue), for the Fisetin and -1.381 (deepest red) to + 1.381 eV (deepest blue) for the Quercetin molecule. The red colour corresponds electrophilic attack and the blue colour shows the nucleophilic attack.

![Figure 6: Molecular electrostatic potential (MEP) of Fisetin (left) and Quercetin (right)](image)

### 3.3.4 Non-Linear Optical (NLO) property analysis

The NLO properties of Fisetin and Quercetin were investigated by the DFT/B3LYP technique to establish the NLO character from polarizability, hyperpolarisability and dipole moment calculations. The value of hyperpolarisability have a strong sensitivity to the electron.

The dipole moment shows highest value at 1.84 and 1.734 D for Fisetin and Quercetin respectively. The values are tabulated in Table 2. The value of the dipole moment increases with the intramolecular interactions.

The hyperpolarisability value for the Fisetin and Quercetin are $9.26 \times 10^{-31}$ and $1.89 \times 10^{-30}$ esu and this large value of the two molecules implies that they have significant NLO properties.

### 3.3.5 UV-Vis spectral analysis and Frontier molecular orbital

The UV-Vis spectrum of the Fisetin and Quercetin as represented in Figure 7 was computed in gas phase. The electronic transition values for both the molecules are gathered in Table 3. In gas phase, the absorption wavelengths of Fisetin and Quercetin are detected at 352, 303 and 356, 307 nm. These values are similar in gas and solvent phase. Therefore the solvent has no effect on the optical activity of the molecule.

![Figure 7: UV-Vis spectra of Fisetin (left) and Quercetin (right) (Theoretical-Gas phase)](image)
The HOMO and LUMO energy diagram is represented in Figure 8. The HOMO and LUMO energy gap informs the molecular chemical stability \(^{39}\). The energy difference between the HOMO and the LUMO for Fisetin is 3.835 eV and 4.00 eV for Quercetin as seen in Figure 8. This gap energy is an essential parameter for electron conductivity characterisation \(^{40}\). Parameter like chemical hardness (CH) also contributes to the chemical stability. The CH of Fisetin is 1.9175 and 2.00 for Quercetin as shown in Table G (Support Information).

![Energy Gap Diagram](image)

*Figure 8: Atomic orbital HOMO - LUMO composition of the frontier molecular orbital of Fisetin (Left) and Quercetin (Right).*

Similarly, electronegativity corresponding to the attraction of electron towards the atom through the bond was calculated 3.5785 and 4.068 for Fisetin and Quercetin respectively. The electrophilicity index for both molecules with B3LYP method was found 3.3391 and 4.137 for Fisetin and Quercetin respectively. These intermediate values suggested the large energy conversion between the HOMO and LUMO \(^{36-38,41}\).

Both the compounds showed very low chemical softness of 0.5215 and 0.500 for Fisetin and Quercetin respectively, indicating non-toxicity in nature (Table G, Support Information).

### Table 3: Comparison of the electronic properties of Fisetin and Quercetin.

<table>
<thead>
<tr>
<th></th>
<th>Gas phase Fisetin</th>
<th>Gas phase Quercetin</th>
</tr>
</thead>
<tbody>
<tr>
<td>(\lambda_{cal}(\text{nm}))</td>
<td>Band Gap (eV)</td>
<td>Energy (cm(^{-1}))</td>
</tr>
<tr>
<td>352</td>
<td>3.5244</td>
<td>28426.8</td>
</tr>
<tr>
<td>303</td>
<td>4.0921</td>
<td>33005.7</td>
</tr>
</tbody>
</table>

In addition, this computer simulation gives leads in the search for inhibitors against this virus. The affinity between the therapeutic molecules and the pathogen is not, however, the only decisive parameter in the choice of candidate drugs, especially in the case of Covid-19. In fact, as described in the literature, the action of the immune system and therefore of its regulation during this condition is crucial \(^{42}\).

Finally, the result between a plant extract and its separately tested compounds is different in some cases due to synergistic action (group action). Even if the two
characteristic compounds of *A. annua* don’t have the best score, however, *In vitro* tests, the ethanolic extract of this plant has been shown a good activity against the coronavirus.  

4 CONCLUSION

Since the appearance of the new coronavirus at the end of 2019, several types of treatment have been proposed, including herbal medicines. Despite the urgency of the situation, compliance with the protocol for identifying active molecules against this virus remains the only winning way. The flavonoids are the secondary metabolites most exploited for this purpose. In the process of discovering therapeutic molecules, *In silico* investigations accelerate the search for candidate molecules. In this present study, nine bio compounds detected from the *Artemisia annua* (Artemisinin, Arteannuin B, Alpha Thuione, P-Hydroxyacetophenone, Fisetin, Cirsimaritin, Capillin, β-Sitosterol, and Quercetin) are evaluated for their inhibitory effect through molecular docking on three targets essential in the spread of Sars-cov-2. Quercetin and fisetin give promising results. Thereby, a molecular simulation is carried out on these two flavonoids (DFT, HOMO, LUMO and gap energies) to evaluate their stability and their activity. Quercetin is present in many plants and this relatively good availability makes it a serious candidate in the fight against the coronavirus. *In vitro* tests have shown good anti-SARS-Cov-2 activity but due to its low solubility in water, its administration to living beings remains a challenge.

Declaration of competing interest

The authors declare that they have no conflict of interest.

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