In Silico Evaluation of Different Flavonoids from Medicinal Plants for Their Potency against SARS-CoV-2

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Abstract: The ongoing pandemic situation of COVID-19 caused by severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) poses a global threat to both the world economy and public health. Therefore, there is an urgent need to discover effective vaccines or drugs to fight against this virus. The flavonoids and their medicinal plant sources have already exhibited various biological effects, including antiviral, anti-inflammatory, antioxidant, etc. This study was designed to evaluate different flavonoids from medicinal plants as potential inhibitors against the spike protein (Sp) and main protease (Mpro) of SARS-CoV-2 using various computational approaches such as molecular docking, molecular dynamics. The binding affinity and inhibitory effects of all studied flavonoids were discussed and compared with some antiviral drugs that are currently being used in COVID-19 treatment namely favipiravir, lopinavir, and hydroxychloroquine, respectively. Among all studies flavonoids and proposed antiviral drugs, luteolin and mundulinol exhibited the highest binding affinity toward Mpro and Sp. Drug-likeness and ADME studies revealed that the chosen flavonoids are safe and non-toxic. One hundred ns-MD simulations were implemented for luteolin-Mpro, mundulinol-Mpro, luteolin-Sp, and mundulinol-Sp complexes and the results revealed strong stability of these flavonoid-protein complexes. Furthermore, MM/PBSA confirms the stability of luteolin and mundulinol interactions within the active sites of this protein. In conclusion, our findings reveal that the promising activity of luteolin and mundulinol as inhibitors against COVID-19 via inhibiting the spike protein and major protease of SARS-CoV-2, and we urge further research to achieve the clinical significance of our proposed molecular-based efficacy.

Keywords: SARS-CoV-2; spike protein; main protease; flavonoids; ADME/Tox; MD simulation

1. Introduction

COVID-19, manifested by severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) has found its place in the concept of a new emerging outbreak that was first recorded in patients suspected with symptoms of the possible common cold found in Wuhan city of China in late December 2019, followed by a rapid spread worldwide until now [1]. SARS-CoV-2 from the coronaviridae family possesses a single-stranded RNA molecule that infects both humans and mammals. The clinical manifestation covers a wide range of symptoms including lymphopenia, fever, dry cough, diarrhea, rhinorrhea, vomiting,
nausea, and fatigue [2,3]. The SARS-CoV-2 is comprised of four main viral proteins such as spike glycoprotein (S), envelope protein (E), membrane protein (M), and nucleocapsid protein (N) [4]. The human angiotensin-converting enzyme 2 (HACE2) is a receptor on the host cells vital for the incorporation of SARS-CoV-2 spike protein into the lung tissue [5,6]. Many studies have revealed the crystal structures of the main protease of SARS-CoV-2, including or excluding inhibitory molecules. Thus, it has been the most characterized target for novel drug discovery [7–9]. To design novel drugs against SARS-CoV-2, we have been provided with several efforts to obtain an understanding of the protein-ligand interaction on a molecular basis [10]. Currently, a potential therapeutic drug is yet to be discovered; however, it is reported that chloroquine, hydroxychloroquine, favipiravir, lopinavir, simprevir, darunavir, and remdesivir can act as a helpful medication against this virus [11–13]. Additionally, the combination of lopinavir and ritonavir with nebulized alfa-interferon is described as a sympathetic medication against SARS-CoV-2 [14–17]. The role of traditional medicine in COVID-19 treatment is studied and reported [18–20]. Medicinal plants containing bioactive compounds have always been of great interest to scientists as they play a significant role in curing human diseases. Therefore, they can reinforce the existing drugs to elicit improved effects against different diseases without negligible adverse effects [21]. Over 80% of the African population relies on traditional medicines against different pathogens such as primary health problems according to WHO [22]. Flavonoids originate from medicinal plants, such as phenolic compounds, and have several biological applications including antiviral, anti-inflammatory, antibacterial, antifungal, anti-aging, anti-Alzheimer’s disease, antitumor, antioxidant, etc. [23–31]. In addition, flavonoids exhibit antiviral activity against various viruses such as Sindbis virus, HCMV, HSV-1, HSV-2, DENV, parainfluenzavirus 3, chikungunya virus [32–38]. Among the naturally occurring flavonoids, anthranol, apigenin, cannflavin, derrisin, jaceidin, lupiwighteone, luteolin, methylglovanon, mundulinol, naringenin, rhamnetin, and tamarixetin, and due to their significant biological roles they are reported the most important flavonoids [39–42]. These flavonoids are abundant in various medicinal plants such as *Andrographis paniculata*, *Moringa peregrina*, *Santolina insularis*, *Melissa officinalis*, *Solanum nigrum*, and *Tamarix nilotica* and these plants are already used as traditional medicine in some countries [43–48]. Therefore, the present study aimed to depict the inhibitory potential of these flavonoids with variable pharmacological effects, targeting the spike protein (Sp) and main protease (M<sub>pro</sub>) of SARS-CoV-2 by in silico approaches. The inhibitory effects of all studied flavonoids were discussed and compared with some currently used antiviral drugs in COVID-19 treatment.

2. Results and Discussion

2.1. Molecular Docking

The binding interaction obtained due to the interactions among all the studied compounds inside 6W63 and 6VYB has been displayed in Figures 1–4, respectively. Table 1 shows the result obtained from the molecular docking analysis for three proposed antiviral drugs (Favipiravir, Lopinavir, and Hydroxychloroquine) and the studied compounds against 6W63. The molecular docking analysis revealed that only luteolin and mundulinol have the highest binding energies (−13.97 and −13.12 kcal/mol, respectively) compared to favipiravir, lopinavir, and hydroxychloroquine in the state of 6W63 (−13.02, −12.98, and −12.45 kcal/mol, respectively). Different types of interactions such as hydrogen bond, Van der Waals, and pi-alkyl interactions were established between the amino acid residuals of 6W63 and the studied compounds with the highest binding energy. Luteolin compound interacts with ARG 131, LYS 137, TYR 239, LEU 287, GLU 288, and ASP 289 amino acid residuals of 6W63 through hydrogen bonds and the distance of these bonds are 2.56, 2.02, 2.17, 2.44, 2.11, and 2.27 Å, respectively. Additionally, it interacts with ASP 289 amino acid residue through Van der Walls interaction. Mundulinol interacts with ARG 188, and pi-alkyl interactions were established between the amino acid residuals of 6W63 and the studied compounds with the highest binding energy. Luteolin compound interacts with ARG 131, LYS 137, TYR 239, LEU 287, GLU 288, and ASP 289 amino acid residuals of 6W53 through hydrogen bonds and the distance of these bonds are 2.56, 2.02, 2.17, 2.44, 2.11, and 2.27 Å, respectively. Additionally, it interacts with ASP 289 amino acid residue through Van der Walls interaction. Mundulinol interacts with ARG 188, and GLU 55 amino acid residuals of 6W53 through hydrogen bonds with distance 2.29, and 2.01 Å, respectively. It also interacts with PHE 181, VAL 186, PRO 52, and TYR 54 amino acid residues through Van der Walls interactions, whereas in the state of 6VYB, the molecular
docking analysis showed that mundulinol, derrisin, and luteolin compounds displayed the highest binding energies (−11.08, −11.04, and −10.92 kcal/mol, respectively) compared to favipiravir, lopinavir, and hydroxychloroquine (−10.76, −10.72 and −10.35 kcal/mol, respectively) as shown in Table 2. Luteolin interacts with ASN 544, and GLY 545 amino acid residuals of 6VYB through hydrogen bonds and the distance of these bonds are 2.32, and 2.16 Å, respectively. It also interacts with ALA 522, and HIS 519 amino acid residues through Van der Walls interaction. The ASN 969, and ASP 867 amino acid residuals of 6VYB interacts with mundulinol through hydrogen bonds with a distance of 2.33, and 2.39 Å, respectively. It also interacts with PHE 970 amino acid residues through Van der Walls interactions. Basically, in the state of 6W63, the binding affinity of the studied compounds is comparatively higher than those of 6VYB. When we compare the docking results of docked ligands with 6W63 and 6VYB proteins, we have noticed that luteolin and mundulinol compounds have higher binding energies than currently used antiviral drugs in both cases (6W63 and 6VYB). Whilst derrisin failed to have higher binding energy than those of favipiravir, lopinavir, and hydroxychloroquine, derrisin is excluded from the promising drugs list as antiviral against SARS CoV-2. Based on this, only luteolin and mundulinol compounds exhibited the ability to work as dual inhibitors against 6VYB and 6W63 because both had binding affinities higher than those of selected drugs favipiravir, lopinavir, and hydroxychloroquine (as anti SARS-Cov-2 drugs) in two cases (6VYB and 6W63).

Table 1. Molecular docking analysis of studied compounds and some antiviral drugs against the main protease (PDB: 6W63) of SARS-CoV-2.

| Compounds       | Binding Energy | Residue       | Distance (Å) | Residue       | Hydrophobic  |
|-----------------|----------------|---------------|--------------|---------------|--------------|
| Anthranol       | −10.22         | GLU 189       | 2.11         | MET 165, MET 49 |
| Apigenin        | −12.32         | PHE 185, ASN 180 | 2.07, 2.22 | PHE 181, PHE 134 |
| Cannflavin      | −10.88         | ARG 188, THR 25 | 2.27, 2.58 | CYS 44, MET 49 |
| Derrisin        | −11.44         | ARG 188, GLU 166 | 2.44, 2.55 | MET 165, HIS 41, CYS 145 |
| Jaceidin        | −9.66          | LYS 137       | 2.06         | TYR 126, GLU 290 |
| Lupiwighteone   | −11.02         | ARG 188, ASN 142 | 2.07, 2.18 | PRO 168, MET 49 |
| Luteolin        | −13.97         | GLU 287, ASP 289 | 2.56, 2.02 | ASP 289 |
|                  |                | LYS 137, ARG 131 | 2.17, 2.44 |             |
| Methylglovanon  | −9.66          | -             | -            | 2.11, 2.27 |           |
|                |                |                |              | 2 PHE 134, GLU 240 |
| Mundulinol      | −13.12         | ARG 188, GLU 55 | 2.29, 2.01 | PHE 181, VAL 186, PRO 52, TYR 54 |
| Naringenin      | −12.72         | ASP 33, TYR 101 | 2.36, 2.18 | PHE 103 |
|                |                | GLN 83, LYS 88 | 2.19, 2.26 |             |
| Rhamnetin       | −10.15         | LEU 287       | 1.88         | TYR 239 |
| Tamarixetin     | −11.13         | PHE 103, TYR 101 | 2.11, 2.03 | LYS 102, GLU 178 |
| Favipiravir     | −13.02         | ASP 33, TYR 101 | 2.38, 2.16 | PHE 103, TYR 239 |
|                |                | LYS 88        | 2.24         |             |
| Lopinavir       | −12.98         | GLU 288, ASP 289 | 2.11, 2.32 | PHE 181, VAL 186 |
| Hydroxychloroquine | −12.45     | ARG 188, GLU 166 | 2.25, 2.34 | VAL 186, PRO 52, TYR 54 |
|                 |                |                |              |             |
Table 2. Molecular docking analysis of studied compounds and some antiviral drugs against 6VYB.

| Compounds          | Binding Energy | H-Bonds | Hydrophobic Bonds |
|--------------------|----------------|---------|-------------------|
|                    | Number | Residue | Distance (Å) | Number | Residue |
| Anthranol          | −9.08  | 1       | THR 315      | 2.11   | CYS 301, ALA 292 |
| Apigenin           | −10.09 | 5       | MET 731, LYS 733, SER 730, HIS 1058, ASP 867 | 2.15, 2.26, 2.27, 2.31, 2.40 | 2 | VAL 860, PRO 863 |
| Cannflavin         | −9.11  | 1       | THR 315      | 2.46   | CYS 301, ALA 292 |
| Derrisin           | −11.04 | 2       | ARG 319, ASN 317 | 2.38, 2.49 | 2 | PHE 318, VAL 595 |
| Jaceidin           | −10.54 | 2       | LEU 48, ASN 856 | 2.14, 2.28 | 2 | LEU 966, SER 967 |
| Lupiweightone      | −9.92  | 1       | PRO 665      | 2.46   | ILE 312, LEU 303, TYR 313 |
| Luteolin           | −10.92 | 2       | ASN 544, GLY 545 | 2.32, 2.16 | 3 | ALA 522, HIS 519 |
| Methylglovanon     | −9.43  | 1       | LYS 964      | 2.56   | LEU 303 |
| Mundulinol         | −11.08 | 2       | ASN 969, ASP 867 | 2.33, 2.39 | 1 | PHE 970 |
| Naringenin         | −10.12 | 2       | LEU 517, ASN 544 | 2.31, 2.26 | 2 | ALA 522, ASN 544 |
| Rhamnetin          | −10.15 | 2       | GLN 516, Ala 520 | 2.28, 2.19 | 2 | LEU 546, HIS 519 |
| Tamarixetin        | −10.33 | 2       | ARG 765, GLN 762 | 2.15, 2.23 | 2 | ALA 766 |
| Favipiravir        | −10.76 | 2       | ASN 969, THR 315 | 2.24, 2.56 | 2 | PHE 318, ALA 292 |
| Lopinavir          | −10.72 | 2       | MET 731, LYS 733 | 2.19, 2.38 | 1 | SER 967 |
| Hydroxychloroquine | −10.35 | 1       | GLY 545      | 2.33   | ASN 544 |
Figure 1. The binding interaction of Anthranol, Mundulinol, Cannflavin, Derrisin, Jaceidin, and Lupiwighteone in the active site of 6W63.
Figure 2. The binding interaction of Luteolin, Methylglovanon, Mundulinol, Naringenin, Rhamnetin, and Tamarixetin in the active site of 6W63.
Figure 3. The binding interaction of Anthranol, Mundulinol, Cannflavin, Derrisin, Jaceidin, and Lupiwighteone in the active site of 6VYB receptor.
Figure 4. The binding interaction of Luteolin, Methylglovanon, Mundulinol, Naringenin, Rhamnetin, and Tamarixetin in the active site of 6VYB receptor.
2.2. ADMET Properties and Drug-Likeness Model Score

All the values obtained from the ADMET analysis for our studied compounds are presented in Table 3. Lipinski’s rule of five is a widely used method for drug development [49]. All compounds in our study complied with Lipinski’s rules of five. Based on BBB permeability, the compounds were non-BBB permeable. Additionally, all the studied compounds have high GI absorption, especially apigenin, luteolin, and mundulinol compounds with optimum solubility. The solubility and Caco-2 permeability are used to evaluate the drug absorption. The solubility ranges from $-5.86$ to $-1.49$ indicating the moderate to the high solubility of all studied compounds. The high-water solubility can be due to different hydroxy groups present in each compound capable of forming bonds with water molecules. The human colonic adenocarcinoma cell line Caco-2 permeability of all studied compounds have high permeation. The gastrointestinal absorption of all studied compounds. All ADMET properties of all studied compounds are in a good agreement of those calculated by Vijayakumar et al. [50]. Vijayakumar et al. calculated ADMET properties of 48 flavonides compounds to show their ability to act as SARS-CoV-2 inhibitors. In which they calculate the solubility and Caco-2 permeability of Luteolin ($-3.01$, $0.65$), Apigenin ($-2.77$, $1.03$), and Mundulinol ($-2.77$, $1.034$) their obtained values are in a good agreement with our values. Waqar et al. [51] studied the activity of some flavonoids compounds against chikungunya virus replication using molecular docking and DFT-based approach. They calculate the Pharmacokinetics and pharmacological properties of some flavonoids compounds, they elucidate that Apigenin and Luteolin have high solubility and GI absorption. Additionally, the toxicity and carcinogenicity of both compounds were also predicted for in order to determine the safe use of the selected compounds during in vivo analysis. Apigenin and Luteolin showed non-toxic and non-carcinogenic nature. Rasool et al. [52] studied the inhibitory potential of selective phytochemicals against Mpro of 2019-nCoV by computer-aided approaches, they study ADMET properties of Mundulinol, Derrisin, Luteolin, and Apigenin. They found that Mundulinol, Luteolin, Derrisin, and Apigenin have high solubility and GI absorption. Additionally, four compounds exhibited the properties of non-BBB permeability. Finally, four compounds exhibited non-toxic and non-carcinogenic nature. In a collective sense from all cited published paper in this section that, the ADMET profile of Luteolin, Mundulinol, and Apigenin compounds is satisfactory and hence suitable for in silico studies with SARS-CoV-2 proteins. Finally, estimations of the toxicity and carcinogenic properties, suggesting that all studied compounds are non-toxic and non-carcinogenic. So, all compounds have suitable ADMET profiles, especially compounds such as apigenin, luteolin, and mundulinol. The drug-likeness model score for the selected compounds was predicted from the Molinspiration server (http://www.molinspiration.com accessed on 10 June 2021) and the output values are shown in Table 4. We excluded the compounds with zero or negative value of drug-likeness score from nominated compounds to be drugs (derrisin and rhamnetin), while the highest drug-likeness score was found to be 0.98 for apigenin followed by luteolin (0.95), mundulinol (0.92), and finally naringenin (0.90). Joining together the docking results, ADMET estimations, drug-likeness results suggest that compounds, luteolin, and mundulinol are promising to be developed as leads drugs with good pharmacokinetic properties. Therefore, we nominated these two compounds for further investigations via running MD simulations.
Table 3. ADMET properties of all studied compounds [GIT = Gastrointestinal tract and BBB = Blood–brain barrier].

| Compounds       | Estimated Solubility Log Score | Estimated Solubility Class | GIT Absorption | Caco-2 Permeability | BBB Penetration | Lipinski Violations | Toxicity | Carcinogenicity |
|-----------------|--------------------------------|-----------------------------|----------------|---------------------|----------------|---------------------|----------|-----------------|
| Anthranol       | −2.63                          | High soluble                | High           | 1.14                | No             | 0                   | Non-toxic | Non-carcinogenic |
| Apigenin        | −2.74                          | High soluble                | High           | 1.01                | No             | 0                   | Non-toxic | Non-carcinogenic |
| Cannflavin      | −5.47                          | Moderately soluble          | High           | 1.21                | No             | 0                   | Non-toxic | Non-carcinogenic |
| Derrisn         | −2.72                          | High soluble                | High           | 1.03                | No             | 0                   | Non-toxic | Non-carcinogenic |
| Jaceidin        | −5.11                          | Moderately soluble          | High           | 1.28                | No             | 0                   | Non-toxic | Non-carcinogenic |
| Lupiwiweightone | −3.71                          | High Soluble                | High           | 0.79                | No             | 0                   | Non-toxic | Non-carcinogenic |
| Luteolin        | −3.04                          | High soluble                | High           | 0.69                | No             | 0                   | Non-toxic | Non-carcinogenic |
| Methylglovanon  | −5.86                          | Moderately soluble          | High           | 1.25                | No             | 0                   | Non-toxic | Non-carcinogenic |
| Mundulinol      | −1.49                          | High Soluble                | High           | 0.46                | No             | 0                   | Non-toxic | Non-carcinogenic |
| Naringenin      | −2.88                          | High Soluble                | High           | 1.07                | No             | 0                   | Non-toxic | Non-carcinogenic |
| Rhamnetin       | −4.54                          | Moderately soluble          | High           | 1.12                | No             | 0                   | Non-toxic | Non-carcinogenic |
| Tamarixetin     | −4.63                          | Moderately soluble          | High           | 1.15                | No             | 0                   | Non-toxic | Non-carcinogenic |

Table 4. Prediction of drug-likeness model score of all studied compounds.

| Compounds       | Drug-Likeness Model Score |
|-----------------|---------------------------|
| Anthranol       | 0.60                      |
| Apigenin        | 0.98                      |
| Cannflavin      | 0.67                      |
| Derrisn         | −1.08                     |
| Jaceidin        | 0.25                      |
| Lupiwiweightone | 0.36                      |
| Luteolin        | 0.95                      |
| Methylglovanon  | 0.18                      |
| Mundulinol      | 0.92                      |
| Naringenin      | 0.90                      |
| Rhamnetin       | −0.50                     |
| Tamarixetin     | 0.27                      |

2.3. Molecular Dynamic Simulation (MD)

MD simulation mimics the structural perturbations and actual movement of a receptor molecule within corresponding biological conditions. Our docking poses are validated through the RMSD, a well-established parameter for protein stability and equilibration, and examine the average behavioural pattern for the entire protein structure during the MD simulation analysis.

2.3.1. Root Mean Square Deviation (RMSD)

The RMSD score determines the direct changes in a protein from its primary coordinates. The RMSD values for the protein backbone coupled with its potential inhibitors were estimated using the initial structure as a frame reference (0 to 100 ns). RMSDs of free 6W63 receptor, mundulinol + 6W63, and luteolin + 6W63 complexes are shown in Figure 5. Additionally, RMSDs of free 6VYB receptor, mundulinol + 6VYB, and luteolin + 6VYB complexes are presented in Figure 6. The RMSD values gradually augmented from 0 to 20 ns and retained the values constant after reaching an equilibration throughout the simulation period for all the studied systems. The RMSD values for all studied complexes were fluctuating between 20 to 40 ns indicating that the compounds were familiarizing a new conformation within the binding pocket. The average RMSD values for the last 20 ns for the systems, were 0.430 ± 0.06, 0.435 ± 0.09, 0.440 ± 0.14, 0.525 ± 0.03, 0.530 ± 0.07, and 0.540 ± 0.09 for 6W63 receptor, mundulinol + 6W63, luteolin + 6W63, 6VYB receptor, mundulinol + 6VYB, and luteolin + 6VYB complexes, respectively. Lower RMSD values of all studied complexes indicate that mundulinol and luteolin inhibitors are stable inside 6W63 and 6VYB and provide a significant base for our study.
Figure 5. The plot of root mean square deviation (RMSD) of receptors backbone vs. simulation time for solvated 6W63 receptors in complex with Mundulinol and Luteolin during molecular dynamics simulations at 100 ns.

Figure 6. The plot of root mean square deviation (RMSD) of receptors backbone vs. simulation time for solvated 6VYB receptors in complex with Mundulinol and Luteolin during molecular dynamics simulations at 100 ns.
2.3.2. Root Mean Square Fluctuation (RMSF)

RMSF was calculated in respect of the backbone atom per amino acid residues and the fluctuations at residue level were demonstrated by the RMSF plot. Figure 7 shows RMSFs of 6W63 receptor, mundulinol + 6W63, and luteolin + 6W63 complexes. Figure 8 shows RMSFs of 6VYB receptor, mundulinol + 6VYB, and luteolin + 6VYB complexes. Both the free receptors (6W63 and 6VYB) and the complexes with low RMSF values showed the nearly same behaviour of residue fluctuation report. So, the RMSF plot confirmed that the binding of all studied compounds is stable inside the receptors also, there is no major consequence found during the simulation analysis regarding the flexibility of the receptor. To gain better understandings of the receptor flexibility, the time average of RMSF values of the 309 amino acids of 6W63 and 975 amino acids of 6VYB receptors in the absence and presence of the studied compounds during the simulation period was computed. In the state of 6W63, the RMSF values for the complexes reported less fluctuation for the catalytic dyad residues (His41 and Cys148). The average RMSF values were 0.33 ± 0.04, 0.40 ± 0.07, and 0.47 ± 0.07 Å for 6W63 receptor, mundulinol + 6W63, and luteolin + 6W63 complexes, respectively. In the state of 6VYB, the RMSF values suggested that the following residues (HIE-41, LEU-141, and GLN-189) showed less fluctuation in the system. The average RMSF values were 0.50 ± 0.06, 0.53 ± 0.04, and 0.55 ± 0.05 Å for 6VYB receptor, Mundulinol + 6VYB, and Luteolin + 6VYB complexes, respectively.

Figure 7. The root mean square fluctuation (RMSF) values of 6W63 protease alone and in complex with the Mundulinol and Luteolin were plotted against residue numbers.
2.3.3. Hydrogen Bonds Analysis

Hydrogen bonds analysis is used to gain more insights into the molecular recognition, molecular interactions, and selectivity of the studied compounds inside the receptors. During MD simulation, the protein-ligand interactions obtained from the changes in the secondary structures were controlled by the analysis of hydrogen bonds. There are different conformations obtained from MD simulation in which each conformation of a receptor has its specific model of interaction with the ligand. We analysed the number of hydrogen bonds formed of MD simulations of all trajectories for the complexes. As showed in Figure 9, the number of hydrogen bonds formed during MD simulation time for mundulinol + 6W63 and luteolin + 6W63 complexes, the average number of conformations were nearly three hydrogen bonds for mundulinol – 6W63 complex and two hydrogen bonds for luteolin – 6W63 complex. Figure 10 represents the mean number of hydrogen bonds generated during MD simulations time for mundulinol – 6VYB complex and luteolin – 6VYB complexes was higher than the number of established hydrogen bonds with SARS-CoV-2 main protease. Further, the number of hydrogen bonds of mundulinol – 6VYB is six while the number of hydrogen bonds of luteolin – 6VYB is five. These results showed that the mundulinol and luteolin compounds possessed a large number of hydrogen bonds with 6VYB during the simulation than 6W63. So, mundulinol and luteolin were capable of maintaining a strong interaction with the binding pocket of 6VYB and 6W63 through the simulation period.
2.4. MM-PBSA Binding Free Energy

MM-PBSA calculations for two selected complexes were estimated to analyse the biophysical basis [53] of the molecular recognition of Mundulinol and Luteolin as dual targets inhibitors. It is evident from Table 5 that the electrostatic interactions ($\Delta E_{\text{Electrostatic}}$) values of Luteolin are more favoured than Mundulinol. Further, the available data in Table 5 reveals that $\Delta E_{\text{Van der Waals}}$ values (intermolecular van der Waals) are similar for Luteolin and Mundulinol when they are targeted SARS-CoV-2 main protease M\text{Pro} (PDB ID: 6W63) and
Mundulinol is a little more favoured than Luteolin when they bind to SARS-CoV-2 spike protein Sp (PDB ID: 6VYB). The Lower binding free energy determines the better receptor and ligand interaction. Based on mentioned parameters, Table 5, Luteolin compound showed better binding free energy when it binds to 6VYB and 6W63 when compare with Mundulinol. Additionally, the binding free energies of both compounds in the state of 6VYB are greater than those of 6W63. This indicated the compounds as stable and had a negligible effect on the flexibility of the 6VYB and 6W63 during the simulation periods and reinforced the significance of our study. ∆E_{binding} can offer a more realistic view concerning the affinities of Mundulinol and Luteolin against the selected targets.

### Table 5. MM-PBSA calculations of binding free energy for two selected complexes.

| Complexes          | ΔE_{Electrostatic} (kJ/mol) | ΔE_{Van der Waal} (kJ/mol) | ΔE_{polar} (kJ/mol) | SASA (kJ/mol) | ΔE_{binding} (kJ/mol) |
|--------------------|----------------------------|----------------------------|---------------------|---------------|-----------------------|
| Mundulinol + 6W63  | -114.54 ± 1.33             | -33.65 ± 2.18              | 35.55 ± 2.16        | -36.32 ± 2.18 | -148.96 ± 3.55        |
| Luteolin + 6W63    | -117.54 ± 2.06             | -33.65 ± 2.36              | 35.55 ± 2.65        | -36.32 ± 3.34 | -150.96 ± 2.64        |
| Mundulinol + 6VYB  | -115.54 ± 2.32             | -37.65 ± 2.98              | 38.55 ± 2.39        | -37.32 ± 4.05 | -151.96 ± 3.55        |
| Luteolin + 6VYB    | -125.21 ± 2.64             | -36.23 ± 3.49              | 35.15 ± 2.18        | -37.65 ± 3.54 | -163.94 ± 2.63        |

### 3. Materials and Methods

#### 3.1. Ligand and Protein Preparation

The structures of studied flavonoid derivatives were downloaded from the PubChem database and saved in SDF format. The energy minimization and structural optimization of the compounds were performed by using the Molecular Mechanics 2 (MM2) force field. Next, the Autodock tool was applied for saving the energy minimized structures in pdbqt format. The 3D crystal structure of spike protein (PDB ID: 6VYB) and main protease (PDB ID: 6W63) of SARS-CoV-2 were obtained from the protein data bank (https://www.rcsb.org/ (accessed on 10 June 2021)). Elimination of water and the small molecules from crystal structures was accomplished via BIOVA Discovery Studio software. Following, the joining of the polar hydrogen atoms and processing the atomic charges were completed by Kollman and Gasteiger method. Finally, the 3D structure of proteins was kept in pdbqt format.

#### 3.2. Assessment of Pharmacokinetics and Pharmacological Properties

Absorption, distribution, metabolism, and excretion along with the toxicity prediction (ADME/Tox) were analysed for filtering the studied compounds and evaluate the pharmacokinetic profile. The prediction of ADME/Tox and the drug-likeness were determined by using swissadme (http://www.swissadme.ch/index.php (accessed on 10 June 2021)) and admetSAR database (http://lmmd.ecust.edu.cn/admetsar1/predict (accessed on 10 June 2021)) [56]. Pharmacokinetics and pharmacological properties of the phytochemicals, i.e., gastrointestinal (GI) absorption, solubility (ESOL), blood–brain barrier (BBB) penetration, and Lipinski’s rules violations were investigated. The properties set for compound screening were: Lipinski’s violations = 0; Solubility = High; Glabsorption = High or Moderate; BBB-permeability = No; and Toxicity = Zero/Nil.

#### 3.3. Molecular Docking Analysis and Binding Energy Estimation

The inhibitory role of our studied flavonoid compounds against the spike protein (PDB ID: 6VYB) and main protease (PDB ID: 6W63) of SARS-CoV-2 was analysed by the molecular docking study. The molecular docking study, in respect of binding of these flavonoids with receptors and their reactivity, following methods applied in various studies [58–60]. All the parameters were set at default values for docking analysis [53]. The docking grid covered all the active sites of the spike protein and main protease receptors with a grid size of 70 Å × 70 Å × 70 Å and a spacing value of 0.375 Å. The docking analysis used the Lamarckian Genetic Algorithm (LGA); all the calculations for docking analysis...
were carried out using the autodock 4.2 software [61]. Further evaluation of the docking output was completed using the discovery studio program. After the docking is completed, the best pose was chosen based on the lowest binding energy. Later on, we aligned our docked poses with that of co-crystallized structure, and finally RMSD lower than 1.0 Å was calculated.

3.4. Molecular Dynamic (MD) Simulation

Molecular dynamic simulation is broadly utilized to evaluate the biological molecules at the atomic level [62, 63] including structural stability and conformational changes. The compounds luteolin and mundulinol exhibit the lowest docking energy with the main protease and spike protein of SARS-CoV-2. The lowest energy of compounds denotes favourable binding affinity [64], therefore it can be said that these the compound could be treated as an inhibitor. Herein, top-ranked compounds bound complexes obtained from molecular docking analysis (mundulinol + 6W63, luteolin + 6W63, mundulinol + 6VYB, and luteolin + 6VYB complexes) were subjected to NAMD software for MD simulation [65]. The temperature and pressure were set to 303 K and 1 atm, respectively. All complexes were solvated using the transferable intermolecular potential water molecules (TIP3P) model. The CHARMM force field was used for a short energy minimization (1000 numsteps) by the Steepest Descent method. Following, the MD simulations were accomplished in two steps: equilibration under NVT ensemble for 5 ns timescale, and production for 100 ns timescale under the NPT ensemble. All theoretical background of MD simulations was performed according to Al-Anazi et al. [66].

3.5. MM-PBSA Calculation

The free energy for binding ($\Delta G_{\text{bin}}$) was estimated via the calculation of the Free Energy (CaFE) program exploiting the Molecular Mechanics-Poisson Boltzmann Surface Area (MM-PBSA) method [67, 68]. It handles simulation trajectories produced from numerous force fields and is enforced by VMD and NAMD. Three distinct energy terms are estimated through the MM-PBSA method that includes energy variations between the complexes and keeps the receptor and ligand molecule apart from each other in the gas phase from NAMD. Then, the APBS program was used to measure the polar solvation free energy [69]. Lastly, the difference of solvent-accessible surface area (SASA) and the free energy of nonpolar solvation was also estimated. The energies needed for binding of the ligand-receptor complexes were calculated throughout the whole MD simulation (100 ns).

4. Conclusions

Current research endeavours against COVID-19 involve numerous antiviral and conventional drugs to fight this pandemic, yet no ultimate therapeutic interventions are available so far. Using the molecular docking analysis, we found the interaction of luteolin and mundulinol with SARS-CoV-2 main protease, spike protein has the highest binding energy when compared to all studied compounds and some proposed antiviral drugs. So, the molecular docking study revealed that luteolin and mundulinol compounds can act as potential inhibitors for Mpro and Sp. The ADME/Tox properties strongly suggest all the studied compounds were non-toxic and non-carcinogenic. Finally, the RMSD value of the receptor-ligands complexes calculated for both receptors exhibited stability at around 0.6 Å and RMSD of two compounds (luteolin and mundulinol) complexed inside 6W63 and 6VYB remained at a favourable range of 0.3 Å and retained its stability throughout the simulation process. The free receptor and backbone atoms of the complex exhibit comparable RMSF values, showing the stability of the studied compounds inside 6W63 and 6VYB receptors. On the whole, the favourable effects of luteolin and mundulinol led to a conclusion that these may be considered as promising treatment strategies against COVID-19 after further clinical trials.

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