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In Silico Analysis of 14-Deoxy 11, 12-Didehydro Andrographolide (AGP 2) From Sambiloto (Andrographis paniculata) As Drug Candidate Against SARS-CoV-2

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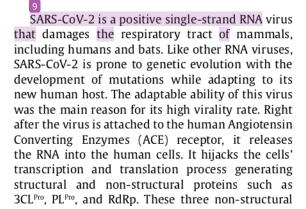
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ABSTRACT

The outbreak of the COVID-19 pandemic in the world has urged researchers to develop a vaccine or therapeutic drugs to fight this virus. This study aimed to assay 14 deoxy-11,12-didehydroandrographolide (AGP 2) ability as an inhibitor of 3-chymotrypsin like-protease (3CLPro), Papain-like protease (PLPro), and RNAdependent RNA-polymerase (RdRp), the viral proteins of SARS-CoV-2 and to evaluate it safeness as a drug candidate. In-silico technique was performed in this study to analyze the binding interaction, complex stability between protein and ligand, and drug-likeness properties. The proteins and ligands were obtained from Protein Data Bank (PDB) 3d PubChem web tools, then using PyRx to identify the binding affinity score, PyMoL to visualize the 3D binding interaction, and WebGro web tools to analyze the stability of each complex. A drug-likeness evaluation was done using SwissADME, pkCSM, and Way2drug web tools. The result of this study showed that the binding affinity score for each complex is; AGP 2-3CLPro (-6.7 kcal/mol), AGP 2- PLPro (-6.4 kcal/mol), and AGP 2-RdRp (-7.0 kcal/mol) where the AGP 2-RdRp and AGP 2-3 CLPro showed a stable form indicating the inhibitor ability of AGP 2. This study also demonstrates that the drug-likeness properties of AGP 2 are safe to use. Additionally, it has been proved that AGP 2 can be developed into a therapeutic drug with further studies.

1. Introduction



proteins play a role in virus pathology function by initiating proteolytic maturation and the formation of the virus Replication-Transcription Complex (RTC). With these functions, SARS-CoV-2 can easily infect the host cells faster and cause casualties in large numbers. According to (Murugan *et al.* 2021), it effectively fights SARS-CoV-2 by inhibiting these viral proteins. Until now, research on using nature drugs as an alternative to tackling this virus using *in silico* techniques is still scarce.

One study was reported in Indonesia and India about *in silico* using plant secondary metabolites fighting 3CL^{Pro}, PL^{Pro}, and RdRp. Fajrin *et al.* (2022.) found that quercetin can inhibit 3CL^{Pro} and PL^{Pro} reviewed on its favorable binding affinity and RMSD value. On the other hand, Pandeya *et al.* (2020) assert that three compounds from *Argemone mexicana*: protopine, allocryptopine, and

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6-acetonyldihydrochelerythrine can inhibit RdRp from SARS-CoV-2 effectively.

Plant secondary metabolites believed can be an effective option to propose as the alternative, as most of them are safer and more cost-effective that contain the natural antiviral compound (Koparde *et al.* 2019). Meanwhile, *Andrographis paniculata*, also known as Sambiloto, has been listed by the FDA as one of the medicinal herbs and was widely used for various diseases (Royani *et al.* 2014). In Indonesia, Sambiloto was used to treat various diseases, such as flu and fever, and to decrease blood glucose in diabetic patients (Bahi *et al.* 2020).

Andrographis paniculata was reported to contain a major compound of diterpenoids with various derivatives like andrographolide, 14 deoxy-11, 12-didehydroandrographolide (AGP 2), neoandrographolide, 14-deoxyandrographolide, 14-deoxy-14,15-didehydroandrographolide, andrographanin, and isoandrographolide (Yunita 2021). AGP 2 was believed to exert no toxicity and has higher oral bioavailability than andrographolide. Not just that, it also has many pharmacology activities such as anti-virus. An in vitro research from China found that AGP 2 strongly inhibited H5NI replication, and the partial mechanism underlying this effect has been elucidated. This research shows AGP 2 has low cytotoxicity and significantly decreased the protein level of non-structural proteins of the H5N1 virus (Cai et al. 2015).

However, whether AGP 2 from Andrographis paniculata can exhibit se same result as the previous in silico study to fight 3CL^{Pro}, PL^{Pro}, and RdRp of SARS-CoV-2 still needs to be determined. Due to AGP 2's numerous bioactivities, it should be considered to invent therapeutics that can inhibit and interfere with the viral replication process of the Covid 19 virus. Thus, this study aim sto assay AGP 2 ability to possess as an inhibitor to 3-chymotrypsin like-protease (3CL^{Pro}), Papain-like protease (PL^{Pro}), and RNA-dependent RNA-polymerase (RdRp), the viral proteins of the virus and to evaluate the pharmacoproperties of 14-deoxy 11, 12-didehydroandrographolide (AGP 2) as a drug candidate using the in-silico technique.

2. Materials and Methods

2.1. Docking of Andrographolide 2 to SARS-CoV-2 Proteins

2.1.1. Ligand and Receptors Preparation

The 3D structure of SARS-CoV-2 non-structural proteins (nsp), which is a 3-Chymotrypsin-like protease (3CLPro), Papain-like protease (PLPro), and RNA-dependent RNA-polymerase (RdRp) were obtained from Protein Data Bank (PDB) with PDB ID of each are 3CLPro (6LU7), PLPro (7LFU) and RdRp (6NUS). The 14-deoxy 11, 12-didehydroandrographolide's (AGP 2) 3D structure was obtained from PubChem with ID (CID: 5318517) (Lim et al. 2021). In this step, the protein must undergo the purification process to eliminate the unwanted segments of the inhibitor before entering the molecular docking step. The purification process was performed using PyMol.

2.1.2. Molecular Docking

Protein-ligand binding docking analysis was performed using PyRx, and the 3D visualization of the interaction was using PyMol. PyRx application tools provide data about the active binding site of the protein receptor with a binding affinity score (Nuryady *et al.* 2021). This study has three binding protein-ligand complexes: AGP 2–3CL^{Pro}, AGP 2–PL^{Pro}, and AGP 2–RdRp. The visualized data using PyMol will show the amino acid residues that interacted with the ligand in precise length.

2.1.3. Molecular Dynamic Simulation

All the complexes of the protein-ligand result in this study have been used to perform the molecular dynamics (MD) simulations using the WebGro web tools (https://simlab.uams.edu/). The MD simulations were carried out to study the stability of the complexes formed in the ambient condition based on the RMSD and RMSF values (Murugan *et al.* 2021). The equilibration and MD run parameters were set to default in this web tool as the temperature was set to 300 K, pressure bar 1.0, and the time was set to 20 nanoseconds while the approximate number of frames was 1.000.

2.2. Drug-Likeness Prediction 2.2.1. ADME Prediction Analysis

ADME (Absorption, Distribution, Metabolism, and Excretion) of the drug's pharmacodynamics materials is imperative to analyze. This prediction aims to analyze the drug-likeness of the proposed ligand and whether it is safe to consume by human or animal models. The analysis was done using SwissADME web tools (https://www.swissadme.ch). The data obtaiged contains information on ligand drug-likeness, such as physicochemical properties, lipophilicity, water-solubility, pharmacokinetics, drug-likeness, and medicinal chemistry (Enmozhi et al. 2020a).

The percentage of oral absorption (%ABS) were calculated using the equation from (Zhao *et al.* 2002):

 $%ABS = 109 - (0.345 \times TPSA)$

2.2.2. Toxicity Prediction Analysis

Toxicity prediction is essantial to predict the drug candidate's tolerability before a eing ingested into the human and animal models. pkCSM (http://biosig.unimelb.edu.au/pkcsm/prediction) is an online database used in this toxicity prediction. The web tools provide details of ligand's toxicology effects

on AMES toxicity, human maximum tolerance dose (HMTD), hERG-I inhibitor, hERG-II inhibitor, LD₅₀, and hepatoxicity (Enmozhi *et al.* 2020b).

2.2.3. Quantitative Structure-Activity Relationship (QSAR) Analysis

The QSAR analysis was then performed to check the feasibility of the ligand's bioactivity using Way2drug web tools (http://way2drug.com/passonline). The ligand's canonical smiles were inserted into the website's command box and examined. The result was shown in the list with the expectation that the ligand possesses antiviral activities against a positive single-stranded RNA virus with indications of passing the threshold of Pa > 0.3.

3. Results

3.1. Molecular Docking

Molecular docking analysis and visualization were done using PyRx and PyMoL databases. Based on Figure 1, the three protein-ligand binding complexes' results are as follows, the AGP 2-RdRp (-7.0 kcal/mol) complex comes with the highest binding affinity score, followed by AGP 2-3CLPro (-6.7 kcal/mol), then AGP 2-PLPro (-6.4 kcal/mol).

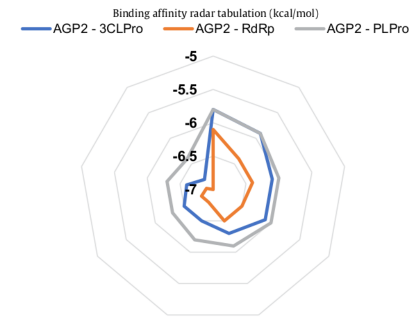


Figure 1. Binding affinity radar tabulation between AGP 2-RdRp, 3CLPro, and PLPro

Figures 2, 3, and 4 show this study's visualization of each complex. Each figure shows the binding interaction between AGP 2 and the proteins where the pink structure is the ligand, and the blue stick is the residues. The yellow line represents the binding interaction with bond-length information.

3.2. Molecular Dynamic Simulation

Figure 5 shows the Molecular Dynamic simulation results. The data on complex stability was reviewed from the Root Mean Square Deviation (RMSD) and Root Mean Square Fluctuation (RMSF). Meanwhile, Table 1 shows the detailed data of the molecular

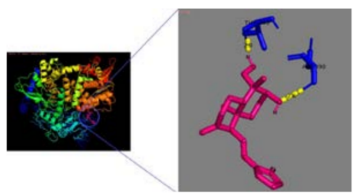


Figure 2. AGP 2-RdRp binding interaction between protein's residues and the ligand

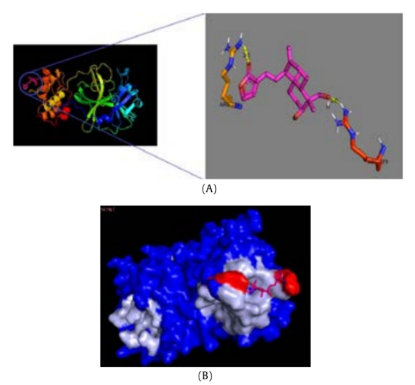


Figure 3. (A) AGP 2–3CL^{Pro} binding interaction between protein's residues and the ligand, (B) AGP 2–3CL^{Pro} binding interaction in 3D surface visualization. The red area represents the polar binding interaction; meanwhile, the light blue area is the non-polar binding interaction, and the blue area represents the protein structure

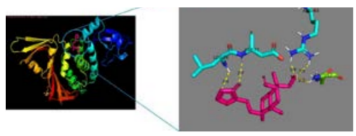


Figure 4. AGP 2-PL^{Pro} binding interaction between protein's residues and the ligand

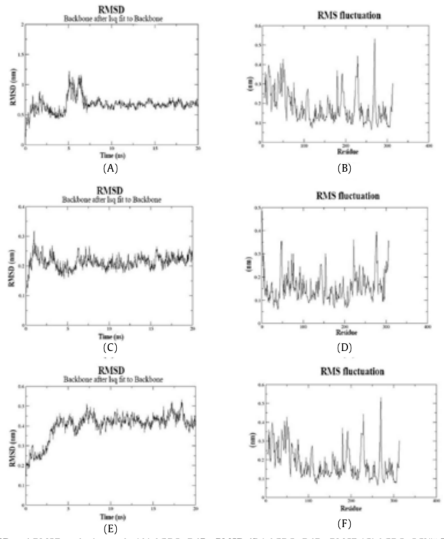


Figure 5. RMSD and RMSF analysis graph: (A) AGP 2-RdRp RMSD, (B) AGP 2-RdRp RMSF, (C) AGP 2-3CL^{pro} RMSD, (D) AGP 2-3CL^{pro} RMSF, (E) AGP 2-PL^{pro} RMSD, (F) AGP 2-PL^{pro} RMSF

dynamic simulation that shows the AGP 2–PL^{Pro} RMSD value did not obey the threshold, as it is above the standard value. The other two complexes obey the standard threshold of RMSD and RMSF itself.

3.3. ADME Prediction Analysis

Drug candidate screening was done using the Swiss ADME database (Table 2). Based on Lipinski's Rule of Five, there are no violations detected for this drug candidate, as all the parameters showed favorable results.

Figure 6 shows AGP 2 analysis using the Boiled-Egg model. The egg-shaped plot is divided into three parts, including a yolk (BBB access), the white egg (HIA access), and the grey region (no HIA or BBB access). AGP 2 was plotted in the white egg region with the blue color of the dot, indicating that this drug candidate is suitable for oral consumption and acts as the positive substrate of P-glycoprotein (PGP+) in a cellular mechanism.

The bioavailability radar (Figure 7) showed a hexagonal structure, with each vertices representing

Table 1. RMSD and RMSF results

Complexes	F	Result		Threshold	
Complexes	RMSD (Å)	RMSF (Å)	RMSD (Å)	RMSF (Å)	
AGP 2–RdRp	0.75	1.4			
AGP 2-3CLPro	2.2	1.3	3.0	2.5	
AGP 2-PLPro	4.3	1.0, 1.3, 1.4, 1.8			

Table 2. ADME prediction result

Complexes	Pharmacoproperties	Result	Threshold (Yasin et al. 2020)
AGP 2	Molecular weight (g/mol)	350.45	500
	Lipophilicity (cLogP)	2.30	<5
	Water solubility (cLogS)	Soluble	Soluble
	Topological polar surface area (TPSA) (Å2)	86.99	40-120
	Number of rotatable bond	3.00	<10
	Hydrogen-bond acceptor	5.00	<10
	Hydrogen-bond donor	3.00	<5
	Oral absorbtion (%ABS)	78.99	Higher is good

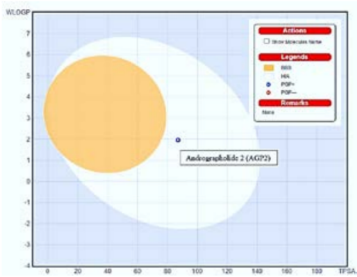


Figure 6. Boiled-egg model

a parameter that defines a bioavailable drug. The pink area within the radar indicates the optimal range for each property (Lipophilicity: XLOGP3 between -0.7 and 5.0, size: MW between 150–500 g/mol, polarity: TPSA between 20–130 Å2, Solubility: Log S (ESOL) not higher than 6, Saturation: fraction of carbons in the sp3 hybridization not less than 0.25 and Flexibility: not more than nine rotatable bonds).

Table 3 shows the bioavailability result of AGP 2. All these results show that each parameter was in the optimal range. Bioavailability is the extent and rate to which the active drug ingredient or active moiety from the drug product is absorbed and becomes available at a site of drug action.

3.4. Toxicity Prediction Analysis

The toxicity prediction of the drug candidate was made using pkCSM web tools. This screening of toxicity for the drug includes AMES toxicity, human maximum tolerated dose (HMTD), hERG I and hERG II inhibition, LD_{50} , and hepatoxicity analysis (Table 4). The results showed no violation in this screening, indicating that the toxicity of AGP 2 is good.

3.5. QSAR Analysis

The QSAR analysis (Table 5) showed that three related bioactivities are necessary for this research objective's success, including antiviral activities and RNA synthesis inhibitors. The Probable Active

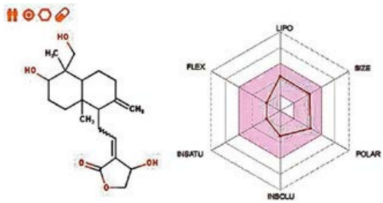


Figure 7. Bioavailability radar of AGP 2

Table 3: Bioavailability radar's result

LIPO	FLEX	INSATU	INSOLU	POLAR (Å2)	SIZE (g/mol)
2.16	3	0.75	-3.18	86.99	350.45

Lipophilicity (LIPO), flexibility (FLEX), insaturation (INSATU), insolubility (INSOLU), and polarity (POLAR)

Table 4: Toxicity prediction results

AMES	HMTD (log mg/kg/day)	hERG I	hERG II	LD_{50} (mg/kg)	Hepatotoxicity
No 0.128		No	No	2,162	No
Mutagenesis (AMES), human maximum tolerated dose (HMTD), and median lethal dose (LD ₅₀)					

Table 5. QSAR analysis result

Table 5. Q5 in analysis result					
Bioactivity	Pa Score				
Antiviral (Rhinovirus)	0.367				
Antiviral (General)	0.424				
RNA synthesis inhibitor	0.436				

(Pa) score of all the related bioactivities is shown in Table 5. The threshold of the ligand Pa score based on QSAR analysis should be Pa >3.0, indicating the ligand is close to active.

4. Discussion

4.1. Molecular Docking

Figure 1 shows a binding affinity radar tabulation, where each non-structural protein has nine active binding sites as the results of PyRx analysis. The outer number of the nonagon-structure vertices represents the active binding site, with sequence number one hitting the highest binding affinity complex score and number nine being the lowest score. Based on the results, the AGP 2-RdRp complex showed the highest binding affinity with a score of -7.0 kcal/mol, followed by AGP 2-3CLPro with a score of -6.7 kcal/mol and AGP 2-PLPro with a score of -6.4 kcal/mol. In this study, all the complexes have a binding affinity range from -6.4 kcal/mol to -7.0 kcal/mol. In comparison with the existing drugs (remdesivir and molnupiravir) with a binding affinity range from -6.6 kcal/mol (Murugan et al. 2021) up to -7.5 kcal/mol (Sharov et al. 2022) AGP 2 showed an excellent binding affinity since the score range was not significantly different.

AGP 2–3CL^{Pro} binding interaction between protein's residues and the ligand; (b) AGP 2–3CL^{Pro} binding interaction in 3D surface visualization. The red area represents the polar binding interaction; meanwhile, the light blue area is the non-polar binding interaction, and the blue area represents the protein structure.

The active binding site of AGP 2-RdRp was located in the palm domain (Figure 2), aligning with the previous research from (Zhu et al. 2020). In this interaction, two hydrogens formed with two residues, namely Thr-393 and Asp-390, where Thr-393 acted as the hydrogen-bond acceptor and Asp-390 as the hydrogen-bond donor. Meanwhile, the AGP 2-3CL^{Pro} complex (Figure 3) shows the formation of two hydrogen bonds and a polar binding with the ligand at the protein's surface. This active binding site is linear with the study from (Mody et al. 2021), which claims the active binding site in his study is located at the protein's surface. However, the interacted residues in this study (Arg-222, Arg-279) acted as the hydrogen-bond donor to the ligand. It differs from Liu et al. (2020) mentioned

that the dyad residues (His-41 and Cys-145) are the main elements forming a hydrogen bond with the ligand.

Moreover, the AGP 2-PLPro complex (Figure 4) showed five hydrogen bonds bonded to four residues, Asp-77, Arg-83, Asn-157 and Leu-76, acting as the hydrogen-bond donor to the ligand. Uniquely, Leu-76 formed a hydrogen bond with the ligand even though this residue is classified as a non-polar element. However, based on (Matthews 2001), leucine (Leu) has a side chain with a polar atom that can interact with other polar molecules, and these abilities are known as amphiphilic.

Based on all the complex results in this step, all their interactions were categorized as strong bonds due to their bond-length interaction being less than 3.0 Å2 (Arba *et al.* 2020). Yan *et al.* (2017) also stated that a hydrogen bond less than 3.0 Å2 exhibits a strong bonding interaction and most likely exhibits a promising inhibitor characteristic.

4.2. Molecular Dynamic Simulation

Based on the Molecular Dynamic Simulation result in Figure 5 and Table 1, these complexes, AGP 2-RdRp and AGP 2-3CLPro, exert good stability after the binding interaction occurs. The RMSD and RMSF value for both complexes obey the standard threshold of this study. According to (Fuglebakk et al. 2012), the RMSD value should be below 3.0 Å. The lower value of RMSD means the conformational changes in the complex are scarce; hence it is a stable binding. As for the RMSF, Fatriansyah et al. (2022) stated that if the value is below 2.5 Å, the fluctuation of the residues is low, which means the complex binding is stable. Since both complexes showed a value below all the criteria of RMSD and RMSF, the complexes manifest a strong and stable interaction. As for the AGP 2–PL^{Pro} complex, the RMSD result showed a high value above the standard threshold. Even if the RMSF value of this complex is below the threshold, this complex is classified as not stable binding because the conformational change of this complex is high.

4.3. ADME Prediction Analysis

Based on the ADME prediction analysis results (Table 2), all the pharmacoproperties of AGP 2 showed a good result, as all the criteria of Lipinski's Rule of Five were obeyed. From the Boiled-egg model evaluation (Figure 6), AGP 2 was indicated to

be suitable for oral consumption through intestinal absorption (Suhud *et al.* 2019).

The molar refractivity and %ABS score of AGP 2 showed a significant number of 95.21 and 78.99%, which indicates that AGP 2 has good membrane permeability (Guan *et al.* 2019). The favorable condition justifies that AGP 2 is permeable through particular membranes and can remain constant even amid strong or weak solvent-solvent or solute-solvent interactions (Enmozhi *et al.* 2020b). (Tong *et al.* 2019) stated that high bioavailability is associated with a lower molecular weight and TPSA. Linearly with AGP 2 physicochemical properties in this study (Figure 7 and Table 3).

The pharmacokinetics data predicted no violation of metabolism for CYP1A2, CYP2C19, CYP2D9, CYP2D6, and CYP3A4. These Main cytochromes (CYP) refer to various metabolism and interaction inside the human body (Durán-Iturbide et al. 2020) AGP 2 does not stop liver metabolism by in biting CYP1A2 (Sugiono et al. 2018); it also does not interfere with the metabolism of several therapeutic drugs, especially anti-convulsants, antiulcer, anti-malarial, anesthetic, and sedative drugs by inhibiting CYP2C19 (Leg et al. 2018). As for the metabolism of β blockers, anti-hypersensitive drugs, anti-arrhythmic drugs, and anti-depressants will not stop by AGP 2 as it does not inhibit CYP2D6 (Taylor et al. 2020). AGP 2 also does not inhibit CYP2C9 which results in the metabolism of anti-clotting agents, anti-hypersensitive, anti-hypertensive, management of Type II Diabetes and Nonsteroidal Anti-Inflammatory (NSAIDs) (Park et al. 2021). The P3A4 was not inhibited by AGP 2, which caused the oxidation of fatty acids, steroids, xenobiotics, and hormone synthesis (Samuels and Sevrioukova 2021).

4.4. Toxicity Prediction Analysis

AMES toxicity prediction (Table 4) showed that no mutagenesis/DNA damage could happen because of the AGP 2 intake. AGP 2 is classified as a nongenotoxic carcinogen that can act through DNA-damaging mechanisms (Arriaga-alba *et al.* 2012). This prediction also provides a human maximum tolerated dose (HMTD); it is predicted that HMTD was 0.128 log mg/kg/day. This dose aimed to maximize the benefit of the drug and minimize the harm. It should be known that the HMTD varies between patients' acute, maintenance, and

preventive therapy. The proper dose secures the effective maintenance treatment, minimization of adverse side effects, tolerability, and long-term adherence to the drug intake (Stampfer et al. 2019). The ion-gated channels hERG I and hERG II were predicted to be secure from AGP 2 inhibition; thus, they can perform their task safely. LD₅₀ value in this study was 2.162 mg/kg, which indicates AGP 2 is classified in class IV of toxicity (500-5,000 mg/kg). The highest the value of LD₅₀, the more tolerance to poisoning it is (Worasuttayangkurn et al. 2019). The AGP 2 was predicted to not cause oxidative stress in the liver due to DNA damage, lipid peroxidation, and protein dysfunction (Fikry and Ahmed 2019). In overall toxicity analysis, AGP 2 was predicted not to cause and lead to health issues and was safe to consume.

4.5. QSAR Analysis

Renatha et al. (2022) stated that the threshold of the ligand Pa score based on QSAR analysis should be Pa > 3.0. Based on the result from the QSAR analysis in Table 5, AGP 2 is expected to have many bioactivities. In this study, the ligand should be predicted to possess the ability as an antiviral against a positive single-strand RNA virus. As the target proteins used in this study play an important role in the SARS-CoV-2 replication process, thus the proposed ligand must fight the RNA synthesis mechanism of the virus. From a previous in-silico study, the diterpenoids and their derivatives (andrographolide, 14 deoxy-11, 12-didehydroandrographolide (AGP 2), neoandrographolide, 14-deoxyandrographolide) effectively inhibited the spike protein from SARS-CoV-2 with considerable binding affinity from the range -6.7 kcal/mol to -7.8 kcal/mol (Murugan et al. 2021). As the proposed ligand's bioactivity exceeds the threshold, it can be against the non-structural proteins of SARS-CoV-2, including 3CLPro, PLPro, and RdRp.

In conclusion, based on our findings, AGP 2 possesses the capability as a suitable inhibitor candidate in fighting RdRp, 3CL^{Pro}, and PL^{Pro} from SARS-CoV-2 as its binding affinity score were -7.0 kcal/mol, -6.7 kcal/mol, and -6.4 kcal/mol respectively. The molecular dynamic simulation analysis then uncovers that the AGP 2-RdRP and AGP 2-3CL^{Pro} complexes are steady, barring AGP 2-PL^{Pro}. As for the drug-likeness analysis and evaluation, it showed that AGP 2 heeds all the benchmarks of

the safety indicator for drugs, including absorption, distribution, metabolism, excretion, and toxicity. Since the binding affinity, stability, and drug-likeness of the formed complexes and AGP 2 compound itself is adequate, it verified that AGP 2 could be materialized as a drug candidate for combating Covid 19 through extensive research in the future.

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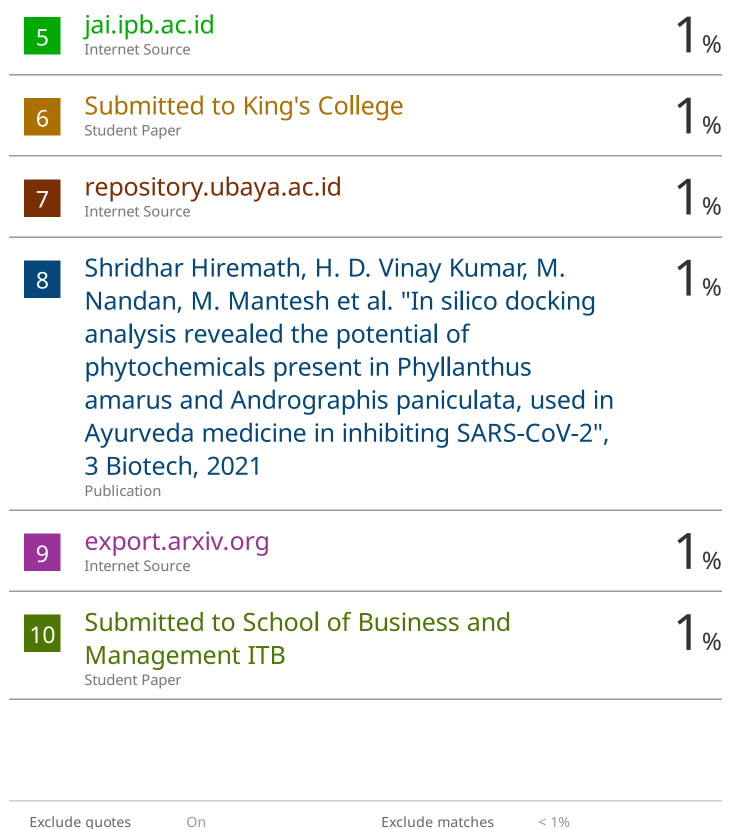
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In Silico Analysis of 14-Deoxy 11, 12-Didehydro Andrographolide (AGP 2) From Sambiloto (Andrographis paniculata) As Drug Candidate Against SARS-CoV-2

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