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African derived phytochemicals may interfere with SARS-CoV-2 RNA capping machinery via inhibition of 2'-O-ribose methyltransferase: An *in silico* perspective

Gideon A. Gyebi^{a,*}, Oludare M. Ogunyemi^b, Adedotun A. Adefolalu^c, Alejandro Rodríguez-Martínez^d, Juan F. López-Pastor^d, Antonio J. Banegas-Luna^d, Horacio Pérez-Sánchez^{d,*}, Adegbenro P. Adegunloye^e, Olalekan B. Ogunro^f, Saheed O. Afolabi^g

^a Department of Biochemistry, Bingham University, Karu, Nigeria

^b Human Nutraceuticals and Bioinformatics Research Unit, Department of Biochemistry, Salem University, Lokoja, Nigeria

^c Department of Biochemistry, Federal University Lafia, Nigeria

^d Structural Bioinformatics and High Performance Computing Research Group (BIO-HPC), Universidad Católica de Murcia (UCAM), Spain

^e Department of Biochemistry, Faculty of Life Sciences, University of Ilorin, Ilorin, Nigeria

^f Department of Biological Sciences, KolaDaisi University, Ibadan, Nigeria

^g Department of Pharmacology and Therapeutics, Faculty of Basic Medical Sciences University of Ilorin, Ilorin, Nigeria

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ABSTRACT

Despite the ongoing vaccination against the life-threatening COVID-19, there is need for viable therapeutic interventions. The S-adenosyl-L-Methionine (SAM) dependent 2'-O-ribose methyltransferase (2'-O-MTase) of the severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) presents a therapeutic target against COVID-19 infection. In a bid to profile bioactive principles from natural sources, a custom-made library of 226 phytochemicals from African medicinal plants with especially anti-malarial activity was screened for direct interactions with SARS-CoV-2 2'-O-MTase (S2RMT) using molecular docking and molecular dynamics (MD) simulations as well as binding free energies methods. Based on minimal binding energy lower than sinefungin (a reference methyl-transferase inhibitor) and binding mode analysis at the catalytic site of S2RMT, a list of 26 hit phytochemicals was defined. The interaction of these phytochemicals was compared with the 2'-O-MTase of SARS-CoV and MERS-CoV. Among these compounds, the lead phytochemicals (LPs) viz: mulberrofuran F, 24-methylene cycloartenol, ferulate, 3-benzoylhosloppone and 10-hydroxyusambarensine interacted strongly with the conserved KDKE tetrad within the substrate binding pocket of the 2'-O-MTase of the coronavirus strains which is critical for substrate binding. The thermodynamic parameters analyzed from the MD simulation trajectories of the LPs-S2RMT complexes presented an eminent structural stability and compactness. These LPs demonstrated favorable druggability and *in silico* ADMET properties over a diverse array of molecular computing descriptors. The LPs show promising prospects in the disruption of S2RMT capping machinery *in silico*. However, these LPs should be validated via *in vitro* and *in vivo* experimental models.

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1. Introduction

The coronavirus disease-19 (COVID-19), was classified a worrisome global pandemic by the World Health Organization (WHO), following the virulent infection rate of Severe Acute Respiratory

Syndrome Coronavirus 2 – (SARS-CoV-2) in humans [1]. Recent epidemiological findings present a cumulative total of about over 246 million confirmed cases and 5 million deaths have been reported since the start of the outbreak [2]. The SARS-CoV-2 belongs to one of the two zoonotic coronaviruses, the other ones being the Middle East Respiratory Syndrome Coronavirus (MERS-CoV). MERS-CoV and SARS-CoV have engendered severe respiratory disorder in mankind since the 21st century commenced [3]. SARS-CoV-2 has been described to be part of the most virulent

* Corresponding authors.

E-mail addresses: gideonagyebi@gmail.com (G.A. Gyebi), hperez@ucam.edu (H. Pérez-Sánchez).

viruses of this century, with the most fatalities till date [4]. Coronaviruses are described as rapidly evolving viruses, with a high rate of genomic mutation [5]. Recently, several variants of SARS-CoV-2 have been identified: the United Kingdom (UK), South Africa and Brazil variants are cited in the several literatures as (B.1.1.7 for UK, 501Y.V2 or 20C/501Y.V2B.1.351 for South Africa and P.1 for Brazil variants) [6]. This, along with a high infection rate has made the development of drugs quite elusive. Like the earlier coronaviruses, SARS-CoV-2 makes use of its cell environment for its replication and survival [7]. The viral RNA maintains its integrity through the “cap”, a unique organization towards the 5' end of the RNA molecule which comprise of a C-2'-O-methyl-ribosyladenine and N-methylated guanosine triphosphate; an arrangement similar to the host cell's RNA [8,9]. The “cap” structure plays significant function in pre-mRNA splicing, mRNA export, RNA stability and escaping the cellular innate immune system [10]. However, in humans, the cap is established in the nucleus of the cell, on the newly transcribed RNA to which the virus has no access. Consequently, they have to own their specified cap-synthesizing enzymes [9]. The last methylation step that cumulates into the RNA cap requires two enzymes, non-structural proteins (nsp) 14 and 16. The nsp 14 for GTP nucleobase, N-7 methylation while the nsp 16 for C-2'-O methylation of the following nucleotide. Both enzymes are methyltransferases (MTases) that depend on S-adenosylmethionine (SAM) [11,12]. Nsp14 when complexed to nsp10 has been reported to reduce cases of mismatched nucleotides via its exoribonuclease domain (ExoN) [13]. The 2'-O-ribose methyltransferase (2'-O-MTase) activity of nsp16 is also influenced by the enzyme's association with nsp10 [13]. The activity of nsp16 functioning as a 2'-O-ribose methyltransferase (2'-O-MTase) is also influenced by its association with nsp10 [14,15]. These properties indicate nsp14 and nsp16 as promising therapeutic targets for SARS-CoV-2, especially nsp16 being a very promising molecular target for structural drug design. The 2'-O methyltransferase (MTase) is also essential for coronaviruses replication (in cell cultures) [16,17]. Identifying bioactive compounds with therapeutic activities against these targets is a necessary step to designing potent antiviral agents. Initial large-scale screening of bio-active agents capable of inhibiting target proteins, using bioinformatics tools have been variously reported [18–23]. The use of plants and their parts, ‘herbal remedies’, in traditional medicine has been well documented. These plants are used as concoctions, decoctions, infusions etc. Indeed, the efficacy of many of these remedies has been attributed to their bioactive phytochemicals [24]. Compounds derived from plants have been known to possess enormous structural diversity that has served as good starting points for investigating new drug [25].

There are several reports that focuses on the use of computation methods to screen different databases and libraries of natural compounds for potential inhibitors of several targets of SARS-CoV-2, this information have been compiled in some reviews [26,27]. Though there are few reports that targets 2'-O-MTase as a viable therapeutic target [28,29], there is no report on the repurposing of antimalarial compounds against SARS-CoV-2 2'-O-MTase

The inhibitory potential of phytochemicals against viral methyltransferases have been well documented [30,31]. This approach can be exploited in the quest for inhibitors of important targets against the novel SARS-CoV-2. In the wake of the ravaging (and still evolving) effect of the COVID-19 pandemic, the dearth of effective anti-viral drugs, and the relatively long process of drug discovery, computational simulation techniques has been a viable tool employed to study the evolving mutations [32,33], and for screening possible novel drug candidates [34–38]. In this study, we employ computational techniques to predict the interactions of a list of bioactive phytochemicals (BP) that were compiled from literature search and known to be derived from African medicinal plants against SARS-CoV-2 and other coronaviruses 2'-O-MTase.

Table 1
Binding site coordinates of nsp16 protease of coronaviruses.

Dimensions	SARS-CoV-2 (Å)	SARS-CoV (Å)	MERS-CoV (Å)
Center_x	89.26	54.25	89.26
Center_y	16.92	60.82	16.92
Center_z	26.44	65.17	26.44
Size x	31.00	26.34	31.00
Size y	29.63	25.41	29.63
Size z	31.34	20.81	31.34

2. Methods

2.1. Retrieval and preparation of protein structure for molecular docking

The 3D structure of nsp-16/10 of SARS-CoV-2 complexed with its native substrate (PDBID: 6WRZ), and previous viruses SARS-CoV (PDB ID: 3R24) and MERS-CoV (PDB ID: 5YNB), were retrieved from the Protein Data Bank (<http://www.rcsb.org>). Existing ligands and water molecules associated with the protein structures were removed and missing hydrogen atoms were added. Using MGL-AutoDockTools (ADT, v1.5.6), the Kollam charges were added as the partial atomic charge [39]. The non-polar hydrogens were merged while the polar hydrogens were added to the proteins. This procedure was applied to all proteins and then saved into a dockable pdbqt format for docking calculations.

2.2. Ligand preparation for molecular docking

The structure data format (SDF) of the reference inhibitors (sinefungin and S-adenosyl-L-homocysteine (SAH)) and 226 phytochemicals were downloaded from the PubChem database (www.pubchem.ncbi.nlm.nih.gov). These ligands were converted to mol2 using Open babel [40]. Compounds that were unavailable on the database were drawn using ChemDraw version 19, the same was converted to mol2 chemical format.

2.3. Virtual screening and active site targeted molecular docking of phytochemicals

The screening of the 226 bioactive compounds against SARS-CoV-2 2'-O-ribose methyltransferase (S2RMT) was performed using AutoDock Vina [41]. Based on the docking scores, interaction in the catalytic site and binding poses, 26 hit phytochemicals were selected. These hit phytochemicals were docked for interaction with the active pockets of the S2RMT of other strains (SARS-CoV and MERS-CoV). For all the docking

In OpenBabel that is incorporated into PyRx 0.8. the Universal Force Field (UFF) was used as the energy minimization parameter and conjugate gradient descent as the optimization algorithm. The energy of all the ligands were minimized using conjugate gradient descent as the optimization algorithm in OpenBabel that is incorporated into PyRx 0.8. The active sites of the three enzymes were defined by the grid boxes and presented in Table 1. All other parameters were kept as default.

2.4. Molecular dynamics simulation

Desmond module of Schrodinger 2019–4 was employed for the MD simulation of the LPS-S2RMT. Water boxes were added to the proteins subsequent to addition of the missing hydrogen atoms and removal of any ligand in the TIP3P molecules solvent system [42] under orthorhombic periodic boundary conditions for 10 Å, buffer region with OPLS3 force field. An isothermal–isobaric ensemble (constant number of particles N, constant pressure P and

constant temperature T) which is an ensemble of Nose-Hoover thermostat [43] and barostat was applied to maintain the constant temperature (310 K) and pressure (1 bar) of the systems, respectively. An energy minimization of 1000 steps with steepest descent followed by conjugate gradient algorithms was utilized. The Parameters such as temperature, salt concentration, and pH were set at the physiological values (310 K, 0.154 M NaCl and 7.0, respectively) during the simulation period. Multiple time step RESPA integration (Reference System Propagator Algorithms) was used in the dynamics study for bonded, near and far-bonded interactions with 2, 2 and 6 fs, respectively. The data were collected for every 100 ps, and the obtained trajectory was analyzed with Maestro graphical interphase (Schrödinger Release 2021–1: Maestro, Schrödinger, LLC, New York, NY, 2021). Various structural parameters, like Root Mean Square Deviation (RMSD), Root Mean Square Fluctuations (RMSF), Radius of Gyration (rGyr), Intramolecular Hydrogen Bonds (intraHB), Molecular Surface Area (MoSA), Solvent Accessible Surface Area (SASA) and Polar Surface Area (PSA) were calculated as a function of time to explore the structural behavior of the proteins, ligands and protein-ligand complexes. In order to estimate the free energy change that describes the binding of these LPs through the MD trajectories, MM-GBSA calculations were carried out, and free energy estimations were computed for 11 snapshots (one every 10 ns as shown in Fig. S19: supplementary data).

2.5. PCA and FEL analysis and covariance matrix generation

PCA and FEL analysis and covariance matrix generation were performed through covar and anaeig GROMACS modules with Desmond MD trajectories and represented by matplotlib Python library.

2.6. Physicochemical and pharmacokinetic study

The LPs for S2RMT were submitted for drug-likeness and ADMET filtering analysis. The SwissADME webserver (<http://www.swissadme.ch/index.php>) was used to analysis the drug-likeness using the Lipinski and Veber filtering tools [44]. Several molecular descriptors on the SuperPred webserver (http://lmm.d.ecust.edu.cn/admet_sar1/predict/) was used to analysis the predicted Absorption, Distribution, Metabolism, Excretion and toxicity (ADME/tox [45]). The canonical SMILES of the LPs were used for the analysis.

3. Results

3.1. Molecular docking of phytochemicals with the target protein

The virtual screening of 226 bioactive phytochemicals from African medicinal plants against S2RMT demonstrated varying degrees of estimated binding energies as presented in Table S2: supplementary data. From the results obtained, a hit list of 26 BP with binding affinities higher than the reference inhibitors, Sinefungin (−7.7 Kcal/mol) and SAH (−8.2 Kcal/mol), with notable interaction with the catalytic residues. The interactions of the top 26 ranked BP with S2RMT were further compared with those of SARS-CoV and MERS-CoV 2′-O-MTase. From these analyzes, the four lead phytochemicals (LPs) with the highest binding affinity to the S2RMT were further selected viz: mulberrofuran F, a flavonoid; 24-methylene cycloartenol ferulate, a pentacyclic triterpenes; 10′-hydroxyusambarensine, an indole alkaloid; and 3-benzoylhosloppone, an abietane diterpenes (Table 2) with quantified free binding energy of (−10.7, −10.1, −9.4 and −9.2 Kcal/mol, respectively). The LPs interacted with SARS-CoV and MERS-CoV 2′-O-MTase with binding affinities of (−9.4, −8.9, −10.5 and −9.6 Kcal/mol) and (−8.7, −8.9, −10.1, and

−9.9 Kcal/mol), respectively (Fig. 1). It was observed the Mulberrofuran F (−10.7 Kcal/mol) the topmost ranked phytochemical to the S2RMT displayed a lower binding affinity of −9.2 and 8.7 Kcal/mol to SAR CoV and MERS-CoV2′-O-MTase. On the other hand, 10-Hydroxyusambarensine demonstrated the highest binding affinity to SARS-CoV and MERS-CoV 2′-O-MTase (−10.5 and −10.1 Kcal/mol, respectively). Thus, the compounds displayed selectivity for different strain of the coronaviruses base on their affinity. The structural stability of the S2RMT complexed with the LPs was analyzed through MD simulations.

3.2. Molecular interactions between the lead phytochemicals and coronaviruses 2′-O-MTase

The interactions of the LPs with the amino acid residues of coronaviruses 2′-O-MTase is given in Table 3. The active site directed docking of sinefungin and SAH to SARS-CoV-2, SARS-CoV and MERS-CoV 2′-O-MTase revealed that it actively interacted with the catalytic site residues majorly through conventional hydrogen-bond, in most cases the reference compounds served as H-Donor from its hydroxyl or amino group. In the three coronaviruses 2′-O-MTase, sinefungin and SAH were docked into the substrate binding cavity as SAM (Fig. 2). In the same manner mulberrofuran F, 24-methylene cycloartenol ferulate, 10-hydroxyusambarensine and 3-benzoylhosloppone interacted with the catalytic residues of the substrate binding pocket (SBP) that is localized in a canyon of the three coronaviruses [46]. Mulberrofuran F interacted via a hydrogen bond to ASP⁶⁸⁹⁷ which is part of the amino acid involved in methionine binding in the pocket of the second subdivision of the SBP [46]. The benzofuran-2-yl ring of Mulberrofuran F was responsible for the hydrophobic interaction with the SBP via Pi-Pi, T-shaped, Pi-alkyl and alkyl interactions (Table 3 and Fig. 2). 24-Methylene cycloartenol ferulate interacted with both the nucleoside and amino acid (methionine binding) pocket sub unit of SAM binding cleft of S2RMT. The same binding pattern was observed with SARS-CoV and MERS-CoV 2′-O-MTase. The 3-O-feruloyl moiety was directed into the cleft of S2RMT, interacting via H-bonds, while the cycloartenol moiety interacted with the nucleoside binding residue via Pi-alkyl and alkyl hydrophobic interaction (Fig. 2). The hydroxyl and amino group of the 9H-pyrido[3,4-b]indol-6-ol moiety of 10-Hydroxyusambarensine interacted with catalytic residues of S2RMT such as ASP⁶⁸⁹⁷ and ASP⁶⁹¹² via hydrogen bonds while the 1H-indolo[2,3-a]quinolizine moiety formed most of the hydrophobic (P-alkyl and alkyl) interactions. A Pi-Sigma interaction with PRO⁶⁹³² alongside carbon hydrogen bonds were formed between the indolopyridocoline moiety and the SAM binding residues (ASN⁶⁸⁹⁹ and GLY⁶⁸⁷¹) of S2RMT (Fig. 2 and Table 3). For 3- Benzoylhosloppone the carbonyl and hydroxyl group were responsible for the conventional and carbon hydrogen bonds to the catalytic residues S2RMT. The benzoyl ring interacted via Pi-alkyl and alkyl interactions to MET⁶⁹²⁹ and LEU⁶⁸⁹⁸ residues belonging to the amino acid (methionine binding) pocket a subunit of the SAM binding cleft of S2RMT. In a similar binding mode as the reference compounds, it was further observed that the LPs interacted with SARS-CoV and MERS-CoV2′-O-MTase in similar binding pattern as with S2RMT. They were docked into the ligand binding cleft of the two proteins and interacted with the catalytic and substrate binding residues (Figs. 2–4). The interacting amino acids are represented in Table 3.

3.3. Result from molecular dynamic analysis

An in-depth 100 ns MD simulation was performed on the LPs complexed with S2RMT. In other to access the stability of the bound system and the structural integrity upon the binding of the

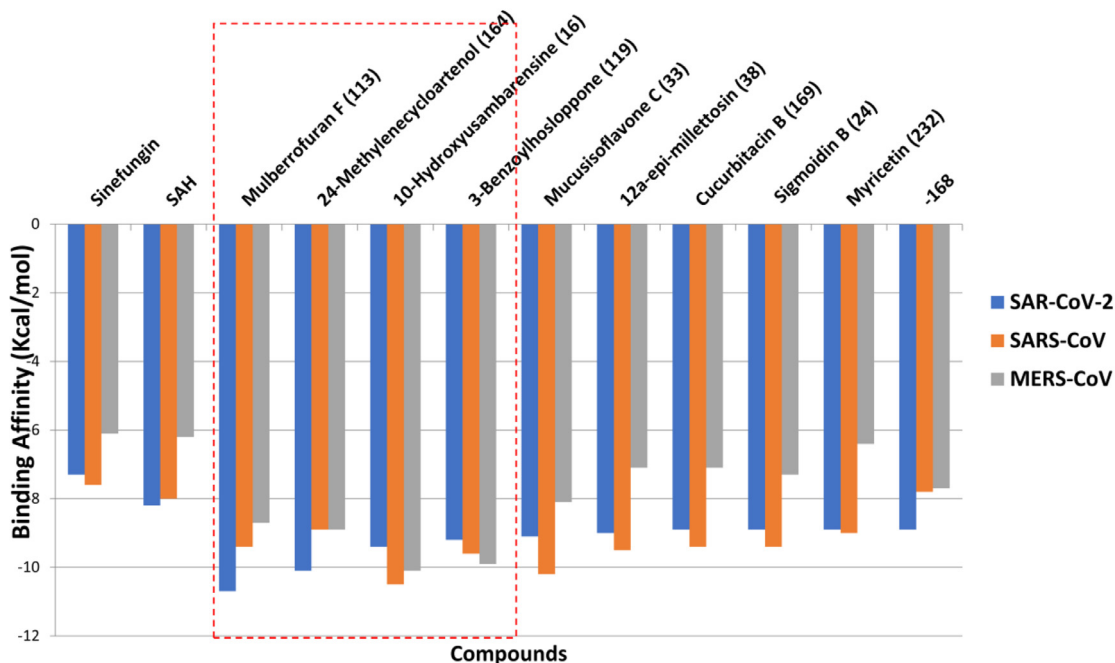


Fig. 1. Binding energies of the ten lead phytochemicals from the docking analysis of 226 phytochemicals and reference compounds docked to the active site of coronaviruses 2'-O-methyltransferase. The red dotted line shows the top 4 docked compounds. 168 = 2, 3, 19 -trihydroxy-urs-12-20-en-28-oic acid.

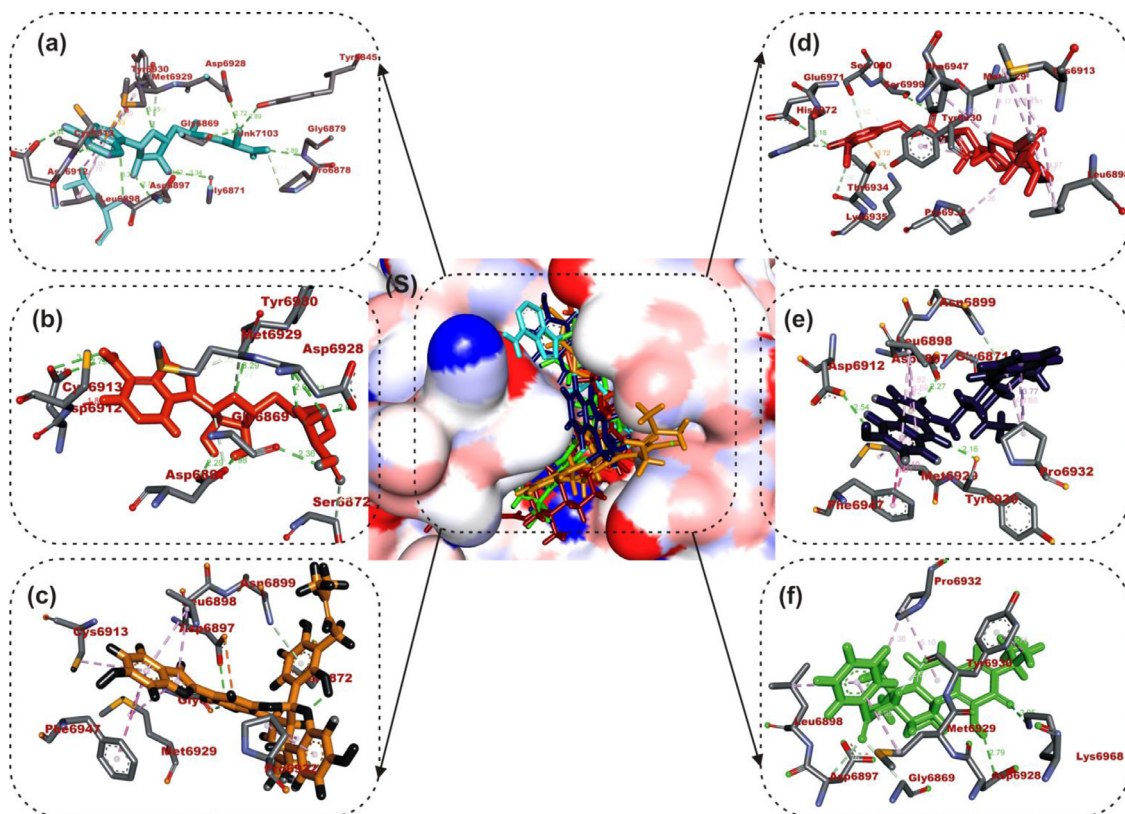


Fig. 2. Amino acid interactions of top lead phytochemicals from the docking analysis and reference inhibitors in substrate binding cavity SARS-CoV-2 2'-O-MTase. (S) solvent-accessible surface view. The top four ranked phytochemicals in sticks representation are represented by colors: (a) cyan: sinefungin (b) orange: SAM (c) gold: mulberrofuran F (d) red: 24-methylene cycloartenol ferulate (e) blue: 10-hydroxysambarensine (f) Green: 3-benzoylshosloppone. Types of interactions are represented by light purple-dotted line: Green-dotted lines: H-bonds; hydrophobic interactions (Pi-Alkyl, Alkyl and pi-stacking); yellow-dotted lines: purple-dotted line: Pi-Pi T Shaped; Pi-sulfur interactions, with three-letter abbreviations of amino acids.

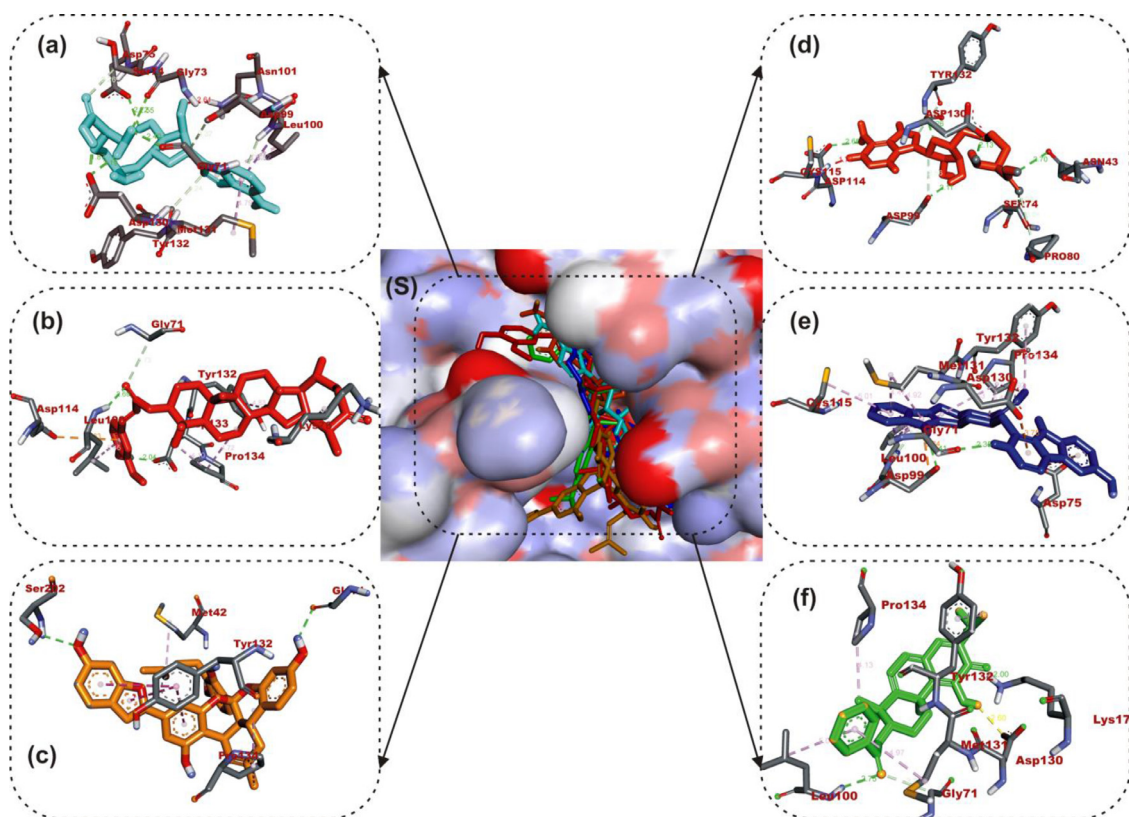


Fig. 3. Amino acid interactions of phytochemicals and reference inhibitors in substrate binding cavity SARS-CoV 2'-O-MTase. (S) solvent-accessible surface view. The top four ranked phytochemicals in sticks representation are represented by colors: (a) cyan: sinefungin (b) orange: SAM (III) gold: mulberrofuran F (d) red: 24-methylene cycloartenol ferulate (e) blue: 10-Hydroxyusambarensine (f) Green: 3-benzoylhosloppone.

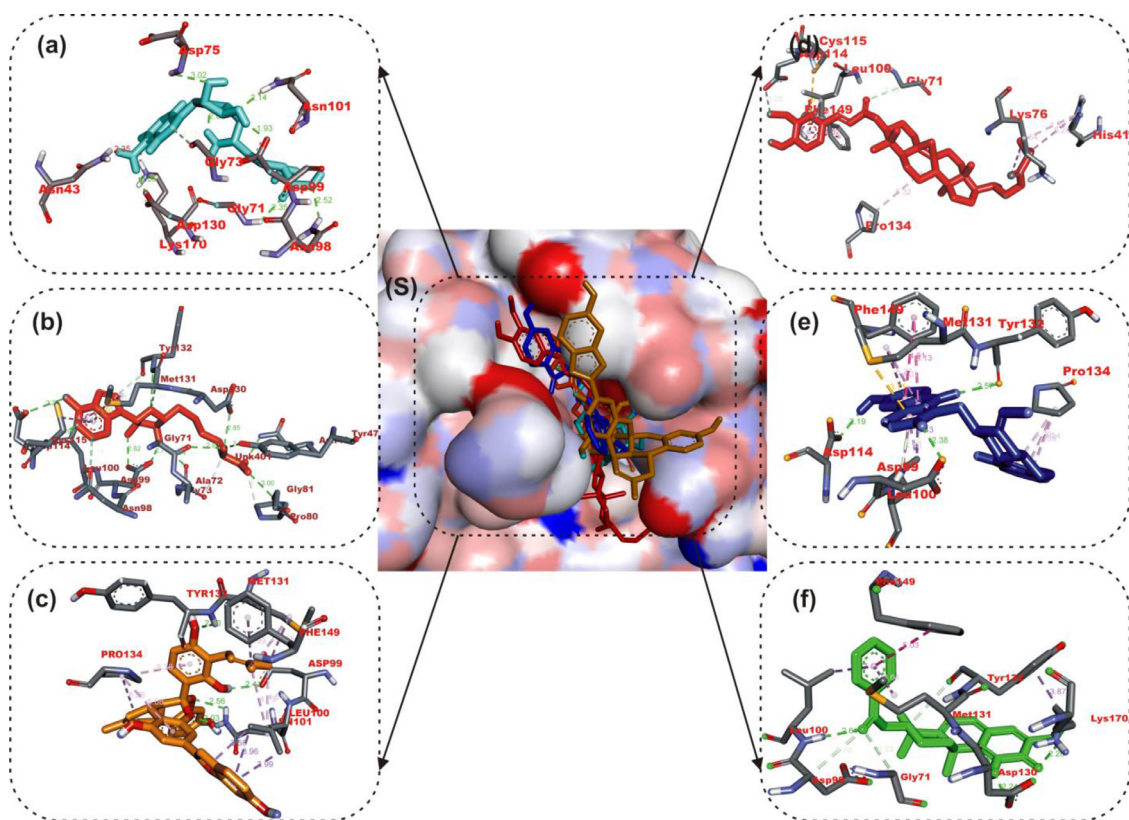


Fig. 4. Amino acid interactions of phytochemicals and reference inhibitors in substrate binding cavity MERS-CoV 2'-O-MTase. (S) solvent-accessible surface view. The top four ranked phytochemicals in sticks representation are represented by colors: (a) cyan: sinefungin (b) orange: SAM (c) gold: mulberrofuran F (d) red: 24-methylene cycloartenol ferulate (e) blue: 10-Hydroxyusambarensine (f) green: 3-benzoylhosloppone.

Table 2
Structure of reference inhibitors (sinefungin and SAH) and the top docked BP with the active site residues of SAR CoV-2 2'-O-MTase.

S/No	Bioactive Compounds	Class of compound	Plant species (Family)
S1	Sinefungin	Nucleoside	
S2	S-adenosyl-L-homocysteine(SAH)	Nucleoside	
1	Mulberrofuran F	Isoprenylated flavonoids	Morusmesozygia (Moraceae)
2	24-Methylene cycloartenol ferulate	Pentacyclic triterpenes	Entandrophrag maingolense (Meliaceae)
3	10-Hydroxyusambarensine	Indole alkaloids	<i>Strychnos ambarensis</i> (Loganiaceae)
4	3- Benzoylhosloppone	Abietane diterpenes	Hoslundiaopposita (Lamiaceae)

phytochemicals, the MD simulation trajectories of the complex systems were compared to that of the unbound systems. The following thermodynamic parameters (RMSD, RMSF, SASA, RoG, and number of H-bonds) protein secondary structure, ligand properties and protein-ligand contacts were computed from the trajectories, the plots were presented as a function of time frame.

3.3.1. Protein secondary structure

Protein secondary structure elements (SSE) of the S2RMT such as the alpha-helices and beta-strands were monitored throughout the simulation. Fig. 8a shows the SSE distribution by residue index. Fig. 8b summarizes the SSE composition, while Fig. 8c monitors each residue and its SSE assignment over time. The result of the

Table 3
Interactions of top docked compounds and reference inhibitors with active site residues of coronaviruses 2'-O-MTase.

Compounds	Coronavirus	Hydrogen bonds (Bond distance)	Other interactions
Sinefungin	SARS-Cov-2	TYR ⁶⁹³⁰ (3.36) ASP ⁶⁹¹² (3.00, 2.97) ASP ⁶⁸⁹⁷ (2.29) SER ⁶⁸⁷² MET ⁶⁹²⁹ (3.37) TYR ⁶⁹³⁰ (2.01) ASP ⁶⁹²⁸ (2.71, 2.83, 2.01) GLY ⁶⁸⁶⁹ (2.78) ASP ⁶⁸⁹⁷ (2.61, 2.70) CYS ⁶⁹¹³ (2.91)	CYS ⁶⁹¹³ MET ⁶⁹²⁹
S-Adenosyl-L-Homocysteine		TYR ⁶⁹³⁰ (3.29) ASP ⁶⁹¹² (3.00, 2.79) ASP ⁶⁸⁹⁷ (2.29) ASP ⁶⁹²⁸ (2.65, 2.83, 2.01) GLY ⁶⁸⁶⁹ (2.36) ASP ⁶⁸⁹⁷ (1.98)	SER ⁶⁸⁷² MET ⁶⁹²⁹ ASP ⁶⁸⁹⁷ TYR ⁶⁹³⁰
Mulberrofuran F		SER ⁶⁸⁷² (2.41) GLY ⁶⁸⁶⁹ (2.13) ASP ⁶⁸⁹⁷ (2.80)	ASP ⁶⁸⁹⁷ ASP ⁶⁸⁹⁹ PHE ⁶⁹⁴⁷ PRO ⁶⁹³² LEU ⁶⁸⁹⁸ MET ⁶⁹²⁹ CYS ⁶⁹¹³
24-Methylene cycloartenol ferulate		SER ⁶⁹⁹⁹ (3.31) GLU ⁶⁹⁷¹ (2.81) HIS ⁶⁹⁷² (3.37) TYR ⁶⁹³⁰ (3.56) THR ⁶⁹³⁴ (3.56) LYS ⁶⁹³⁵ (3.72) SER ⁷⁰⁰⁰ (3.50)	LYS ⁶⁹³⁵ LEU ⁶⁸⁹⁸ MET ⁶⁹²⁹ TYR ⁶⁹³⁰ HIS ⁶⁹⁷²
10 -Hydroxyusambarensine		ASP ⁶⁸⁹⁷ (2.27) TYR ⁶⁹³⁰ (2.16) ASP ⁶⁹¹² (2.54) ASN ⁶⁸⁹⁹ (3.60)	PRO ⁶⁹³² MET ⁶⁹²⁹ PHE ⁶⁹⁴⁷ LEU ⁶⁸⁹⁸
3- Benzoylhosloppone		LYS ⁶⁹⁶⁸ (3.05) ASP ⁶⁹²⁸ (2.79) GLY ⁶⁸⁶⁹ (3.61) ASP ⁶⁸⁹⁷ (3.76)	TYR ⁶⁹³⁰ PRO ⁶⁹³² LEY ⁶⁸⁹⁸ MET ⁶⁹²⁹
Sinefungin	SARS-COV	GLY ⁷⁵ (2.77) GLY ⁷¹ (2.39) ASP ⁷³ (2.54) ASP ¹³⁰ (2.18, 1.87) SER ⁷⁴ (3.25) ASP ⁹⁹ (3.32) TYR ¹³² (3.23)	LEU ¹⁰⁰ (2.95, 3.52,4.80)
S-Adenosyl-L-Homocysteine		TYR ¹³² (3.22) ASP ¹¹⁴ (2.65) ASP ¹³⁰ (2.13) ASN ⁴³ (2.70) ASP ⁹⁹ (2.11)	SER ⁷⁴ PRO ⁸⁰ ASP ⁹⁹ TYR ¹³²
Mulberrofuran F		SER ²⁰¹ (2.73) SER ²⁰² (3.21) GLY ⁷¹ (2.40)	TYR ¹³² PRO ¹³⁴ MET ⁴²
24-Methylene cycloartenol ferulate		ASP ¹³³ (2.04)	GLY ⁷¹ ASP ¹¹⁴ PRO ¹³⁴ TYR ¹³² LEU ¹⁰⁰
10 -Hydroxyusambarensine		ASP ⁹⁹ (2.41) GLY ⁷¹ (2.35)	ASP ⁷⁵ ASP ⁹⁹ ASP ¹³⁰ LEU ¹⁰⁰ PRO ¹³⁴ TYR ¹³² MET ¹³¹ CYS ¹¹⁵
3- Benzoylhosloppone		LYS ¹⁷⁰ (2.81) ASP ¹³⁰ (2.60)	GLY ⁷¹ PRO ¹³⁴ LEU ¹⁰⁰ MET ¹³¹
Sinefungin	MERS-CoV	ASN ¹⁰¹ (2.07) ASN ⁹⁸ (2.51) ASP ⁷⁵ (3.02) ASP ⁹⁹ (1.92, 3.36) ASP ¹³⁰ (3.72) GLY ⁷¹ (2.35) GLY ⁷³ (3.72, 3.133)	LYS ¹⁷⁰ ASN ⁴³
S-Adenosyl-L-Homocysteine		ASN ⁴³ (2.48) TYR ⁴⁷ (2.78) GLY ⁸¹ (2.99) CYS ¹¹¹ (2.96) GLY ⁷¹ (2.81) ASP ¹³⁰ (2.85) GLY ⁷³ (2.98) ASP ⁹⁹ (2.75)	MET ¹³¹ (3.60) LEU ¹⁰⁰ (3.60)
Mulberrofuran F		TYR ¹³² (2.74) ASN ¹⁰¹ (3.72) ASP ⁹⁹ (3.72)	PRO ¹³⁴ MET ¹³¹ ASP ⁷⁵ PHE ¹⁴⁹ LEU ¹⁰⁰
24-Methylene cycloartenol ferulate		ASP ¹¹⁴ (3.44) TYR ¹³² (2.74) LEU ¹⁰⁰ (3.25)	PRO ¹³⁴ PHE ¹⁴⁹ LEU ¹⁰⁰ LYS ⁷⁶ HIS ⁴¹ CYS ¹¹⁵
10 -Hydroxyusambarensine		TYR ¹³² (2.54) ASP ¹¹⁴ (3.07) ASP ⁹⁹ (2.37) LEU ¹⁰⁰ (3.75)	PRO ¹³⁴ MET ¹³¹ PHE ¹⁴⁹
3- Benzoylhosloppone		TYR ¹³² (2.74) GLY ⁷¹ (3.25) ASP ⁹⁹ (3.72) ASP ¹³⁰ (3.72) LEU ¹³⁰ (3.72) LYS ¹⁷⁰	MET ¹³¹ PHE ¹⁴⁹

analysis showed that 19.75% was Helix, 15.28% was strands, while 35.05% was Total SSE (Fig. 5).

3.3.2. Thermodynamic parameters

Root mean square deviation analysis. The RMSD plots for the five systems show that they were equilibrated before 10 ns. The systems exhibited the same progression of RMSD with minimal fluctuation with average RMSD values of 6.83043, 5.674218, 6.124726, 6.369042 and 5.989651 Å for the unbound enzyme, and S2RMT complexed to 10-Hydroxyusambarensine, Mulberrofuran F, 3-Benzoylhosloppone, 24-Methyono cycloartenol, respectively. The binding of the lead phytochemicals reduced the fluctuation in the phytochemical-enzyme complex system; this indicates a more compacted structure upon the binding of the phytochemicals (Fig. 6). The LP- S2RMT systems were further analyzed in Figs. S1-S4 (supplementary data) The C α shows the RMSD evolution

of a protein (left Y-axis). The ligand RMSD (right Y-axis) indicates how stable the LPs are with respect to the S2RMTP and its binding pocket.

Root mean square fluctuation analysis. The RMSF plots reveal the flexibility of the amino acid residues of the protein. Higher fluctuations are observed at the N and C terminal ends of the proteins due to terminal motions. The mean RMSF values for the systems are 2.37097, 2.642513, 3.20722, 2.304798 and 2.51605 Å for the unbound enzyme and S2RMT complexed to 10-Hydroxyusambarensine, Mulberrofuran F, 3-Benzoylhosloppone, 24-Methyono cycloartenol, respectively. The phytochemicals bound SARS-Cov-2 2'-O-MTase complexes displayed higher RMSF values when compared to the unbound enzyme (Fig. 7).

The RMSF plots of the LP- S2RMT systems were analyzed to reveal the secondary structure elements (alpha-helical and beta-

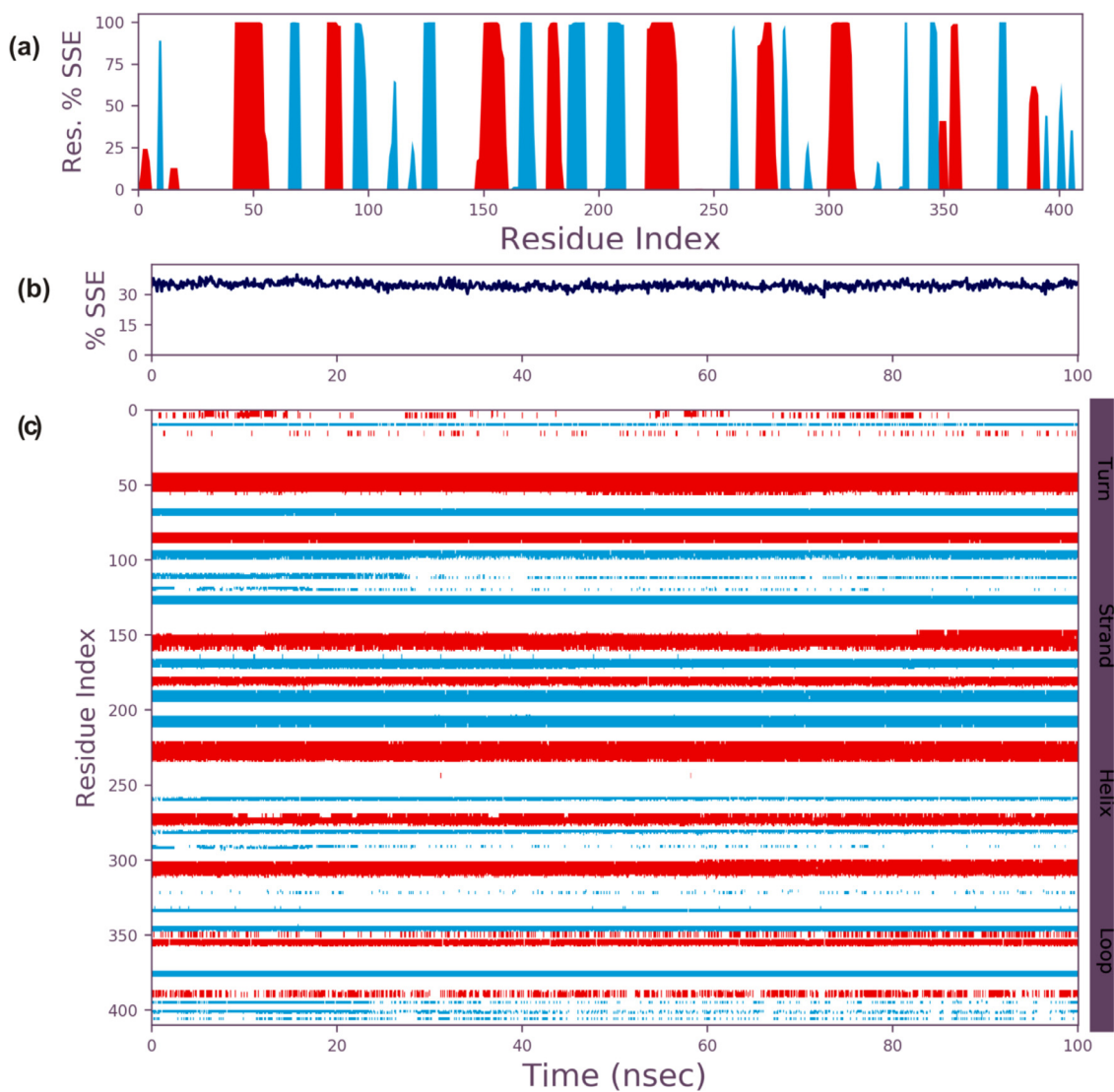


Fig. 5. Secondary structural analysis of SARS-Cov-2 2'-O-MTase during 100 ns MD simulation (a) SSE distribution by residue (b) summary of the SSE composition for each trajectory frame (c) residue and its SSE assignment over time.

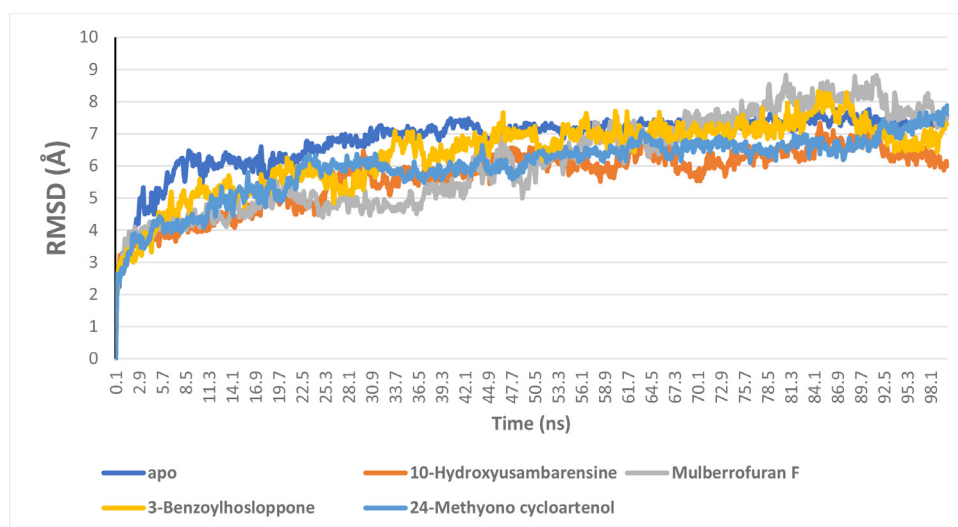


Fig. 6. The Backbone-Root Mean Square Deviation (RMSD) plots of molecular dynamics (MD) simulation of SARS-Cov-2 2'-O-MTase complexed to the four lead phytochemicals from the docking analysis.

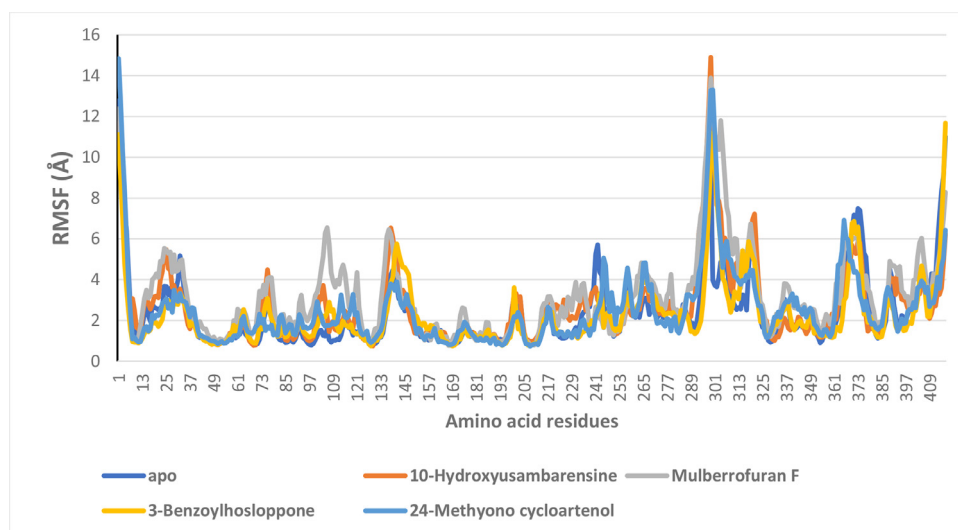


Fig. 7. Per residue Root Mean Square Fluctuations (RMSF) plots of molecular dynamics (MD) simulation of SARS-Cov-2 2'-O-MTase complexed to the four lead phytochemicals from the docking analysis.

strand) regions that interacted with the LP. For the 4 LP- S2RMT systems, the highest fluctuation was observed with the amino acid residues close to residue no. 300 and after residue no. 350. These residues weren't involved in interaction with the ligand. A minimal fluctuation was observed with the interacting amino acid residues before amino acid residue no. 150. The catalytic and substrate binding residue were stable throughout the simulation period. (Fig. S5: supplementary data)

The RMSF of the LP with respect to the S2RMT complexes was further analyzed. The atomic breakdown of the LP that corresponds to the 2D structure in the top panel (Fit Ligand on Protein) line shows the ligand fluctuations, with respect to the protein. A large degree of fluctuation was observed during the simulation period in Mulberrofuran F atoms especially around atom no. 33 with respect to the protein, though the internal atoms of the ligands experienced fewer fluctuations (Fig. S6a). For 24-Methylene cycloartenol ferulate-protein complex the highest fluctuation was around atom no. 34 and 44 (Fig. S6b). Atoms around the later interacted via hydrophobic contacts with the binding site residue. In the case of 10-Hydroxyusambarensine-protein complex the atoms were stable with a lesser degree of fluctuations at atom no. 34 a hydroxyl moiety (Fig. S6c). Atoms of the benzoyl ring moiety of 3- Benzoylhosloppone were the most stable, while the alkyl, carbonyl and hydroxyl group on hosloppone moiety caused some level of fluctuations (Fig. S6d).

The radius of gyration (RoG) analysis. The extent of the compactness of the enzyme upon binding of the ligands is measured from the RoG plots and values. A stably folded protein structure presents a steady RoG plot. Fig. 8 shows the RoG plots of the five systems. The plots for the systems show a steady progression with minimal fluctuations. The mean RoG values calculated for the S2RMT systems are 21.94529, 22.34246, 22.72907, 22.32701 and 22.66686 for the unbound enzyme and the enzyme complexed to 10-Hydroxyusambarensine, Mulberrofuran F, 3-Benzoylhosloppone, 24-Methyono cycloartenol, respectively. The unbound and the enzyme complexed to the lead phytochemicals displayed very close mean RoG values indicating compacted systems (Fig. 8).

The surface accessible surface area analysis. The measure of solvent accessible by the surface of the enzymes was computed

from the generated SASA values for the systems. Both RoG and SASA plots indicates the level of structural unfolding of proteins with reference to its original structure. Fig. 9 show the SASA plots for the enzymes systems. The average SASA values for the S2RMT systems are 20,326.16, 21,156.28, 21,112.91, 20,899.48 and 20,900.15 for the unbound enzyme and the enzyme complexed to 10-Hydroxyusambarensine, Mulberrofuran F, 3-Benzoylhosloppone, 24-Methyono cycloartenol, respectively (Fig. 9).

The changes in the number of H-bonds. The average number of hydrogen bonds for the unbound enzyme, 10-Hydroxyusambarensine, Mulberrofuran F, 3-Benzoylhosloppone, 24-Methyono cycloartenol complexes are 53.72927, 46.89011, 48.31968, 50.53147 and 48.3956. In the AChE systems, a slight reduction in average number of hydrogen bond was observed in the complexes when compared to the unbound protein (Fig. 10).

3.3.3. Protein-ligand contacts

The S2RMT interactions or contacts with the LPs were monitored throughout the simulation. The 2D-trajectory interaction diagram (Figs. 11a–14a) and the stacked bar interactions plots were categorized by type (Hydrogen Bonds, Hydrophobic, Ionic and Water Bridges) and summarized in Figs. 11b–14b. From the total of 21 amino acids contact made, CYS⁶⁹¹³ and ASP⁶⁸⁷³ maintained contact for about 50% and 20% of the simulation time, respectively, via H-Bonds to mulberrofuran F. TYR⁶⁹³⁰ maintained the highest contact via hydrophobic interaction mulberrofuran F. Almost all the residue maintained some level of contact during the simulation time via water bridges (Fig. 11b). 24-Methylene cycloartenol ferulate made contact with 25 amino acid residues. H-bond contacts were sparsely maintained, while residue like TYR⁶⁹³⁰, VAL⁶⁹³⁷ and LEU⁶⁸⁹⁸ interacting via hydrophobic interaction-maintained contact for ~40% 18% and 15% of the simulation time, respectively. Most residues interacting with water bridges maintained some level of prolonged contact (Fig. 12b). From the total of 18 contacts made, LYS⁶⁹³³, TYR⁶⁹³⁰ and ASP⁶⁹¹² interacting via H-bond maintained contact for at least 30% of the simulation time with 10-Hydroxyusambarensine. PHE⁶⁹⁴⁷ maintained the highest hydrophobic contact time (>50%) with 10-Hydroxyusambarensine Unlike the first 2 compounds 10-Hydroxyusambarensine maintained short ionic contact with ASP⁶⁸⁷⁹ and ASP⁶⁹¹². The 2D-trajectory interac-

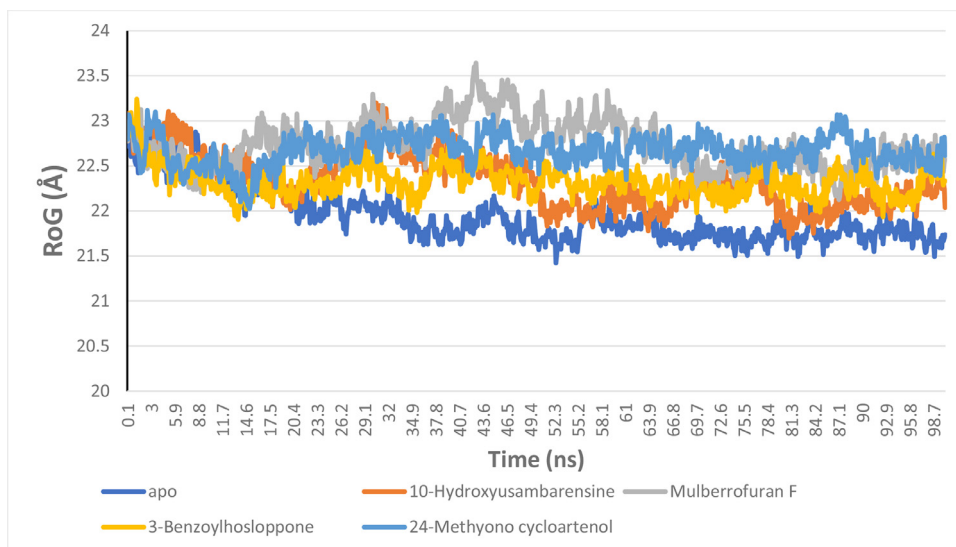


Fig. 8. The Radius of gyration (RoG) plots of molecular dynamics (MD) simulation of SARS-Cov-2 2'-O-MTase complexed to the four lead phytochemicals from the docking analysis.

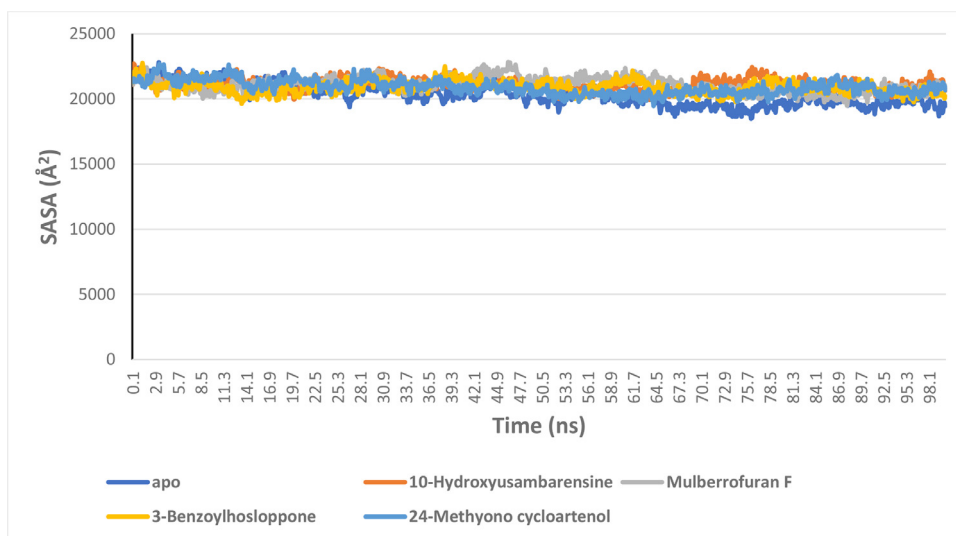


Fig. 9. The Surface Accessible Surface Area (SASA) plots of molecular dynamics (MD) simulation of SARS-Cov-2 2'-O-MTase complexed to the four lead phytochemicals from the docking analysis.

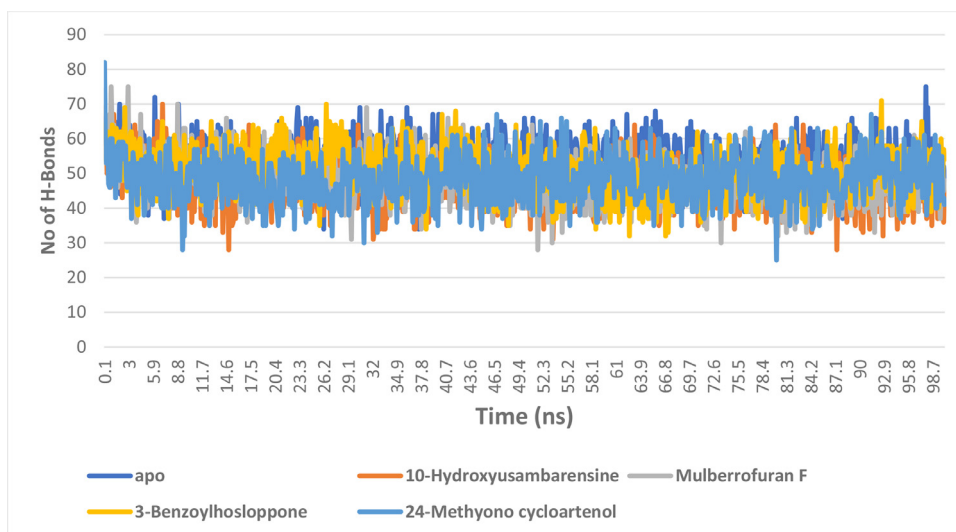


Fig. 10. The changes in the number of H-bonds during the MDS trajectory of SARS-Cov-2 2'-O-MTase complexed to the four lead phytochemicals from the docking analysis.

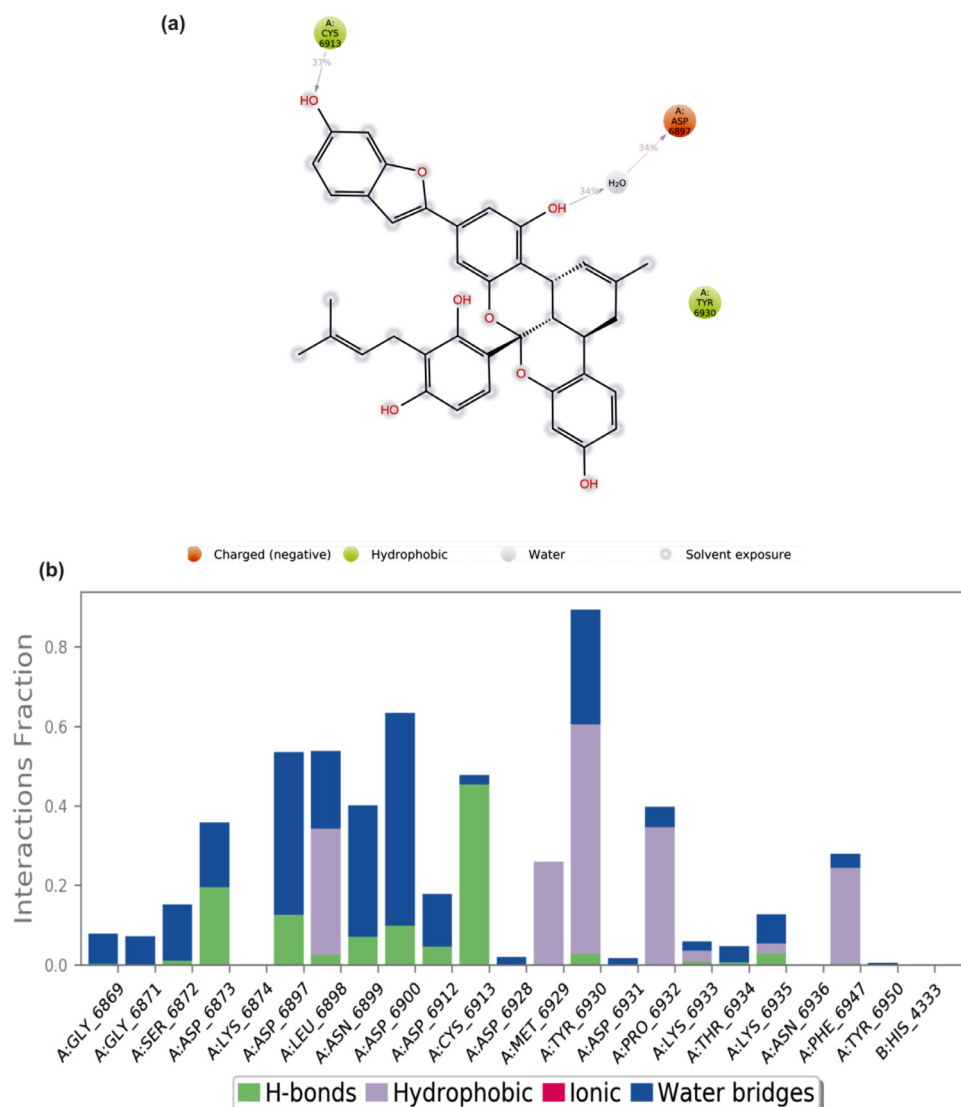


Fig. 11. (a) A schematic details of binding groups of mulberrofuran F interacting with the amino acid residues of SARS-Cov-2 2'-O-MTase (S2RMT) during the period of 100 ns MD simulation analysis. Interactions that occurred more than 30.0% of the simulation time in the selected trajectory (0.00 through 100.00 ns), are shown (b) simulation interactions plot showing categorized S2RMT- mulberrofuran F interactions.

tion diagram (Fig. 13a,b) depicts that 3- Benzoylhosloppone maintained a H-bonding with TYR⁶⁹³⁰ for 66% of the simulation time, while it maintained a hydrophobic contact for about 33% of during the period of simulation with PHE⁶⁹⁴⁷ (Fig. 14a,b).

A timeline representation of the interactions and contacts (H-bonds, Hydrophobic, Ionic, Water bridges) summarized in Figs. S11–S14 is presented in the supplementary data.

3.3.4. Ligand properties

The LPs properties analyzed on its reference conformation. From the plots (Figs. S7a–10a), we observed stable RMSD fluctuations (<3.0 Å) for most cases indicating no huge dynamical alterations during the course of simulations. All the compounds in ligand-protein systems showed a stable rGyr profile, suggesting no conformational alterations (expansion or compression) (Fig. S7b–S10b). Except for mulberrofuran F the other three displayed no intramolecular hydrogen bond during the simulation run (Figs. S7c–S10c). The MolSA, SASA and PSA plots for all the four compounds during the simulation run showed minimal fluctuations, indicating an impressively stable complex upon the binding of compounds

to the active sites of the protein. Other ligand properties such as the Ligand Torsion Profile (LTP) was analyzed, the results are presented in the Figs. S15–S18 (supplementary data). The ligand torsions plot summarizes the conformational evolution of every rotatable bond (RB) in the ligand throughout the simulation trajectory (0.00 through 100.00 ns). The 2D schematic of a ligand, rotatable bonds, conformation and torsion are represented in the Figs. S15–S18 (supplementary Data).

3.3.5. MM-GBSA method for estimating phytochemical binding free energy

The computed free energy estimations for 11 snapshots (one every 10 ns) are summarized in average values and their standard deviation in Table 4. All the LPs yielded dG values that corroborates the docking analysis, with Mulberrofuran F possessing the highest binding free energy (dG). The evolution of the binding free energy for the four systems is graphically presented as a function of the time during the simulation (Fig. S19: supplementary data).

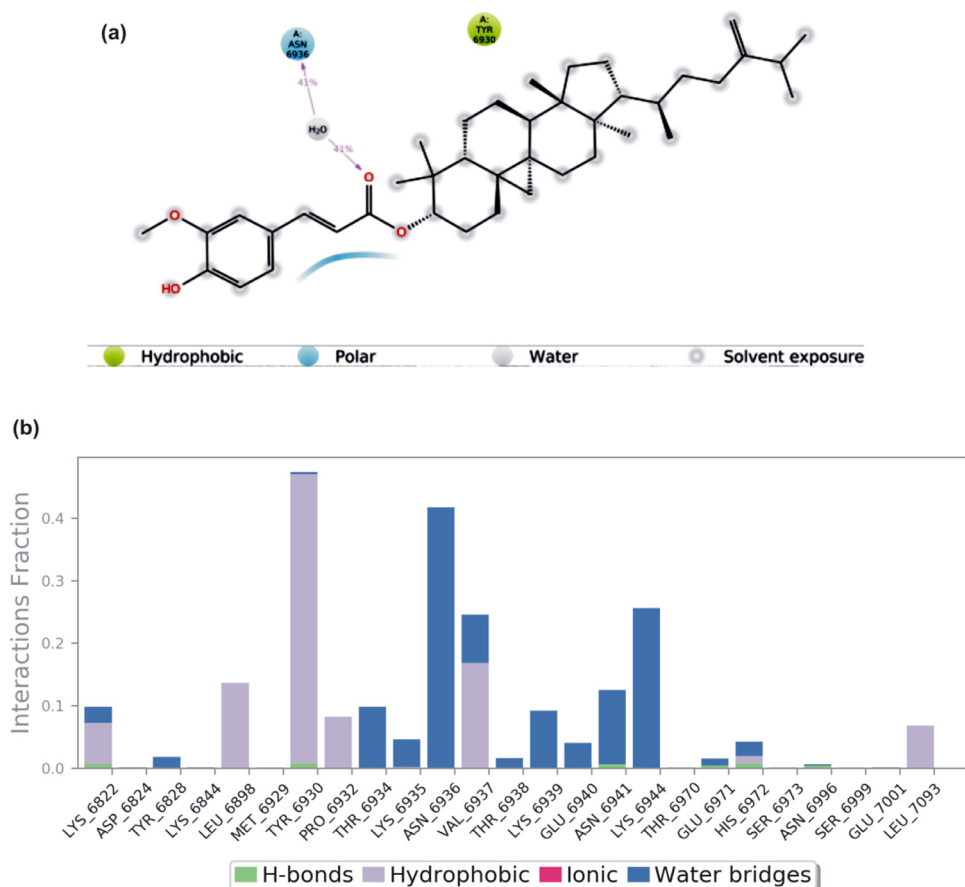


Fig. 12. (a) A schematic details of binding groups of 24-Methylene cycloartenol ferulate interacting with the amino acid residues of SARS-Cov-2 2'-O-MTase (S2RMT) during the period of 100 ns MD simulation analysis. Interactions that occurred more than 30.0% of the simulation time in the selected trajectory (0.00 through 100.00 ns) are shown (b) simulation interactions plot showing categorized S2RMT- 24-Methylene cycloartenol ferulate interactions.

Table 4

MMGBSA obtained dG average values and their standard deviation for the four studied compounds.

Compound	RMSD value at 100 ns (Å)	dG Average (kcal/mol)	dG Standard deviation
10-Hydroxyusambarensine	9.618	-112.4300034	21.67643475
Mulberrofuran F	6.735	-140.1412904	18.02256363
3-Benzoylhosloppone	4.481	-132.1901051	12.52935498
24-Methyono cycloartenol	6.782	-139.3845749	26.35954092

4. Principal component and free energy landscapes analysis

The free energy landscape representations generated by the two first principal components (PC1 and PC2) of the complexes with each one of the inhibitors show similar PCA distribution in the SARS-Cov-2 2'-O-MTase-ligand complexes. Additionally, all of them have differences with the PCA distribution for free protein system. It was observed more different metastable conformations with low-energy states, represented as free energy basins in the blue regions, for those complexes with inhibitor respect to the observed in the free protein. Besides, only one region near to the minimum energy was detected, while rest of complex show more of one metastable region with the minimum value (Fig. 15).

Regarding the traces of covariance matrix, the most relevant evidence is the difference of trace between free protein (10.7905 nm²) and protein binds with Mulberrofuran F (8.35602 nm²). Thus, these results suggest SARS-Cov-2 2'-O-MTase structure obtains a greater compaction when is binding of Mulberrofuran F due to trace decrease in the complex. The rest of ligands don't show a considerable increase in complex compaction respect protein (Table 5).

Table 5

Trace of the covariance matrix for each SARS-Cov-2 2'-O-MTase-compound complex.

Compound	Trace of covariance matrix (nm ²)
No compound (Free protein)	10.7905
10-Hydroxyusambarensine (16)	10.9227
Mulberrofuran F (113)	8.35602
3-Benzoylhosloppone (119)	10.9818
24-Methyono cycloartenol (164)	18.2716

4.1. Drug-likeness and pharmacokinetic properties of selected compounds

The result for the predictive druglikeness and ADMET filtering analyzes for the LPs presented in Table 6. For ADMET analyzes, the molecular descriptors used for the filtering included blood brain barrier (BBB) penetration, this who compounds could cross the blood brain barrier, the aqueous solubility (AS), predicts the solubility of each LPs in water at 25 °C. The various cytochrome P450 descriptors were used to assess the cytochrome P450 inhibitory ac-

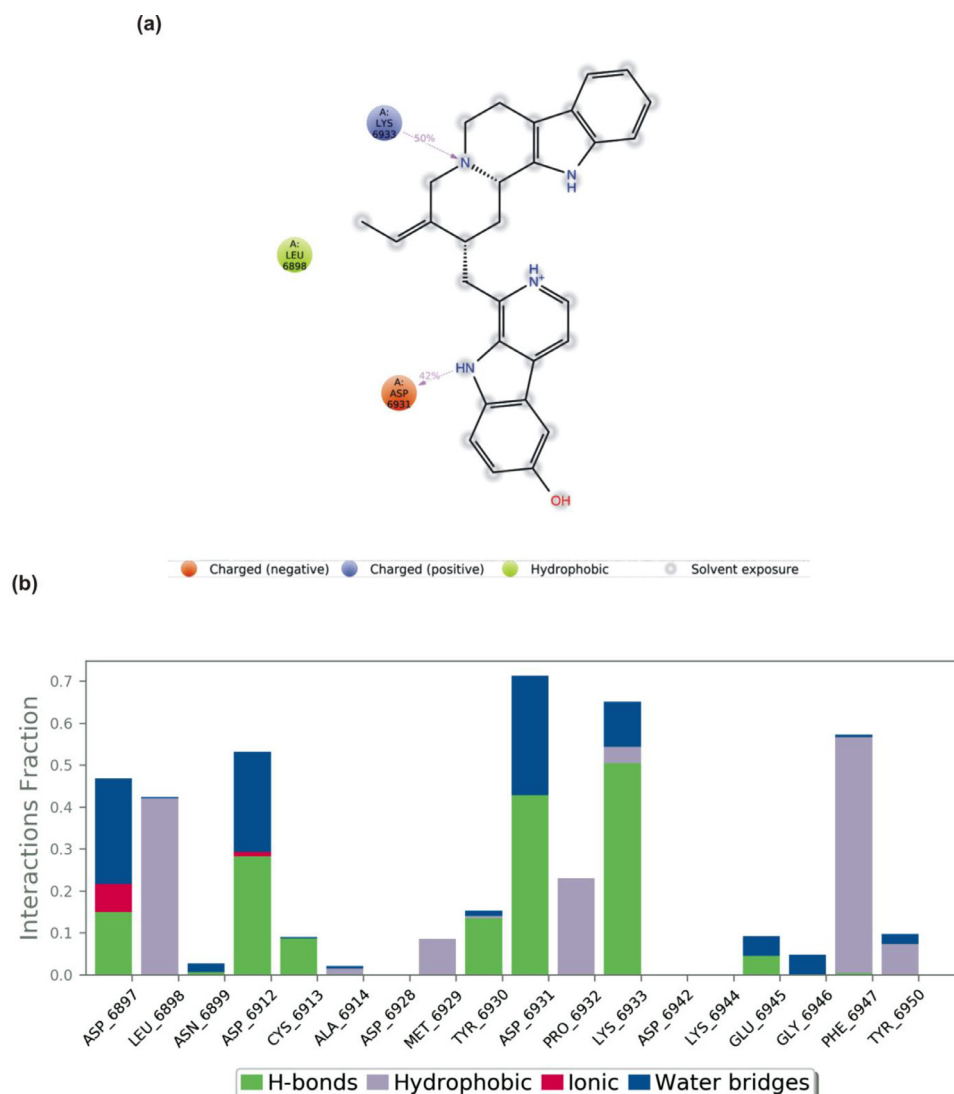


Fig. 13. (a) A schematic details of binding groups of 10-Hydroxyusambarensine interacting with the amino acid residues of SARS-CoV-2 2'-O-MTase (S2RMT) during the period of 100 ns MD simulation analysis. Interactions that occurred more than 30.0% of the simulation time in the selected trajectory (0.00 through 100.00 ns) are shown (b) simulation interactions plot showing categorized S2RMT-10-Hydroxyusambarensine interactions.

tivities of the LPs. The human intestinal absorption (HIA), predicts the intestinal absorption of the LPs after oral administration. The drug likeness and ADMET analysis of 10-hydroxyusambarensine have been reported in our previous paper [18] while the 3 reported herein (mulberrofuran F, 24-methylene cycloartenol ferulate, and 3-benzoylhosloppone) fulfilled the all the requirement for Lipinski analysis with corresponding favorable predicted ADMET parameters. The *in silico* druglikeness and ADMET properties suggested mulberrofuran F to have low GI absorption, while 24-methylene cycloartenol ferulate, and 3-benzoylhosloppone have high GI absorption. The three compounds had high probability of absorption, subcellular distribution, and low toxicity [47]. The ADMET analysis shows that the LPs have the ability to be absorbed in the human intestine, high aqueous solubility, low acute oral toxicity with a good bioavailability score (Table 6).

5. Discussion

SARS-CoV-2 is a virulent and highly evolving virus, whereas the drug discovery process has not matched the increasing therapeutic need of this viral infection [48]. Naturally existing phytochemicals from plants are potential bioactive repositories, including

antiviral activity, which, if adequately explored, could provide affordable, accessible and available use as therapeutic agents against coronavirus infections [49]. Like other coronavirus, the SARS-CoV-2 evades host immune detection and reduces the chance of immune response in the incubation period of 2 to 14 days. This evasion of immune detection is projected to be achieved through the modification of viral mRNA by 2'-O-methyltransferase activity of nsp16/nsp10 which enables the virus to escape detection by the host's innate immune mechanism [50]. Compounds that block viral immune evasion through the suppression of viral RNA 2'-O-methylation, will encourage early expression of interferon-stimulated genes which in turn will serve to impede SARS-CoV-2 replication [51]. The 2'-O-Methyl transferase activity of nsp-16 is S-adenosyl-methionine (SAM)-dependent and regulated by nsp10 binding. The binding of SAM induces essential conformational changes, in the enzyme, that favors RNA affinity and methylation [52]. Hence it is expected that compounds that interacts with the SAM binding site may elicit host response against the virus. Structure based drug design has employed molecular docking to predict the binding-conformation of ligands in the binding site of target receptor and the strength of association (binding affinity) [53]. In the present study, we screened 226 bioactive phytochemicals

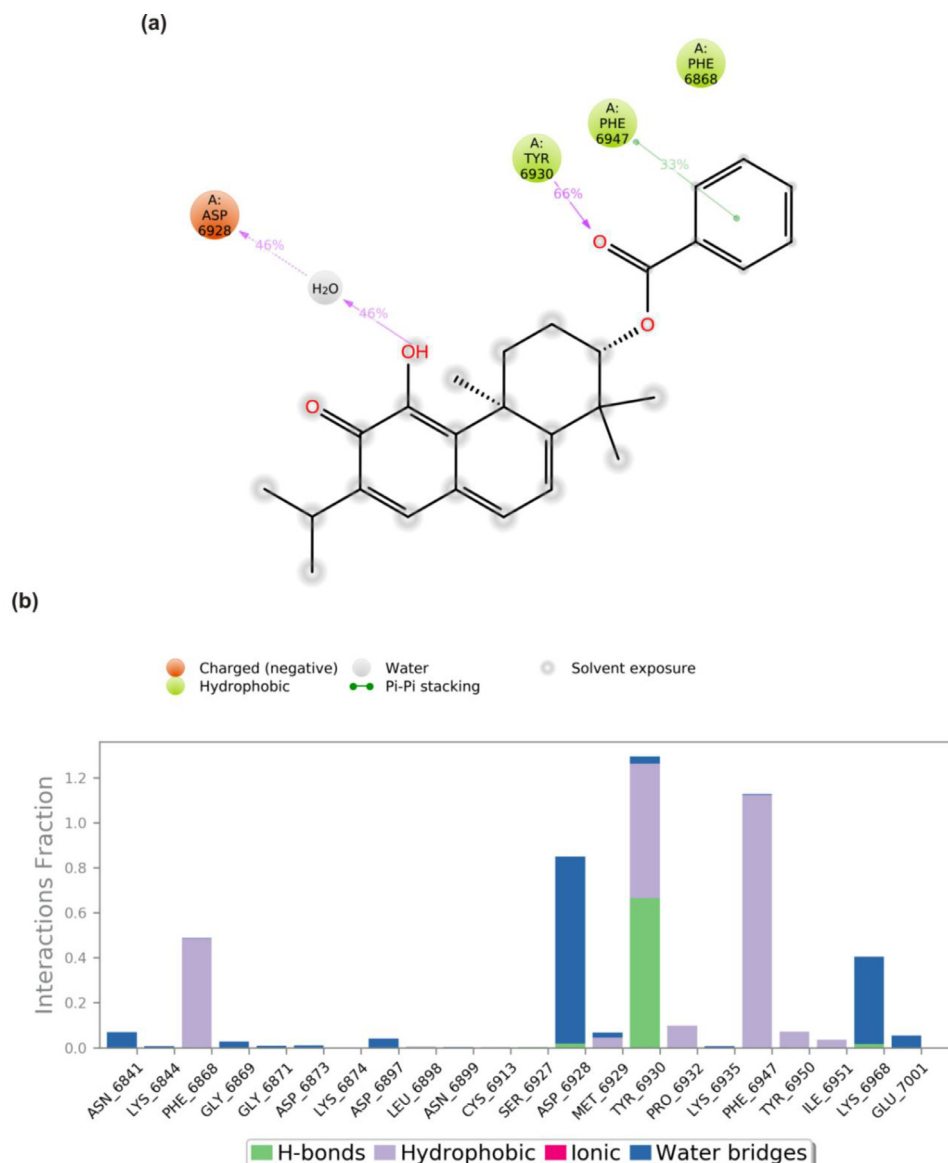


Fig. 14. (a) A schematic details of binding groups of 3-Benzoylhosloppone interacting with the amino acid residues of SARS-Cov-2 2'-O-MTase (S2RMT) during the period of 100 ns MD simulation analysis. Interactions that occurred more than 30.0% of the simulation time in the selected trajectory (0.00 through 100.00 ns) are shown (b) simulation interactions plot showing categorized S2RMT-3-Benzoylhosloppone interactions.

pounds from various African plants against nsp16 of SARS-CoV 2. The docking, interactive and binding free energy analysis identified the LPs (Mulberrofuran F, 24-Methylene cycloartenol ferulate, 10-Hydroxyusambarensine and 3-Benzoylhosloppone) with high potential and selective inhibition of the coronaviruses nsp16 protein. Hydroxyusambarensine is an alkaloid from the roots of *Strychnosambarensis*, previously reported as an antimalarial [54]. Mulberrofuran F isolated from *Morus alba*, has been used to treat hypotension [55,56]. 24-Methylene cycloartenol ferulate, also called γ -Oryzanol (OZ) has been identified in various cereals, including barley, rice bran and corn [57]. It has been reported to exhibit antioxidant, anti-lipidemic, anti-diabetic and neuro-modulatory properties [57,58]. These LPs interacted with the surface residues (Lys-46, Asp-130, Lys-170 and Glu-203) at the bottom of the central groove, that catalysis the transfer SAM methyl group within the substrate binding pocket [46]. In all strains of CoV, the catalytic tetrad (Lys46, Asp130, Lys170) and Glu203 are conserved [59], this may have been responsible for the high binding potential to the three CoV understudied. Though the LPs interacted with the cat-

alytic residues in a similar binding pattern as the SAH (the product of methylation of SAM) and sinefungine (a known inhibitor), they interacted with a stronger binding affinity than these compounds. Thus, these compounds may be able to bind to the S2RMT tightly and hence compromise the RNA methylation function of the enzyme, this will in turn, disrupt the capping machinery, prevent evasion of recognition by the host innate immune system [60–62] and preclude the viruses from resisting the IFN-mediated antiviral response [10,16]. To further understand the dynamic behavior of the LPs at the binding site of S2RMT, MD simulation was employed [63]. The binding patterns and per-residue amino acid interactions of the LPs-S2RMT complexes in the dynamic state collaborated with those done from the static docking analysis. The various thermodynamics parameters that were analyzed from the 100 ns atomistic MDS trajectory files of the LPs-S2RMT complexes revealed stable complexes that can be adapted into other forms of experiments. The comparison of the RMSD plots for the complex systems shows that the binding of the LPs to S2RMT did not cause any structure deformation in the protein [64]. From the RMSF plots

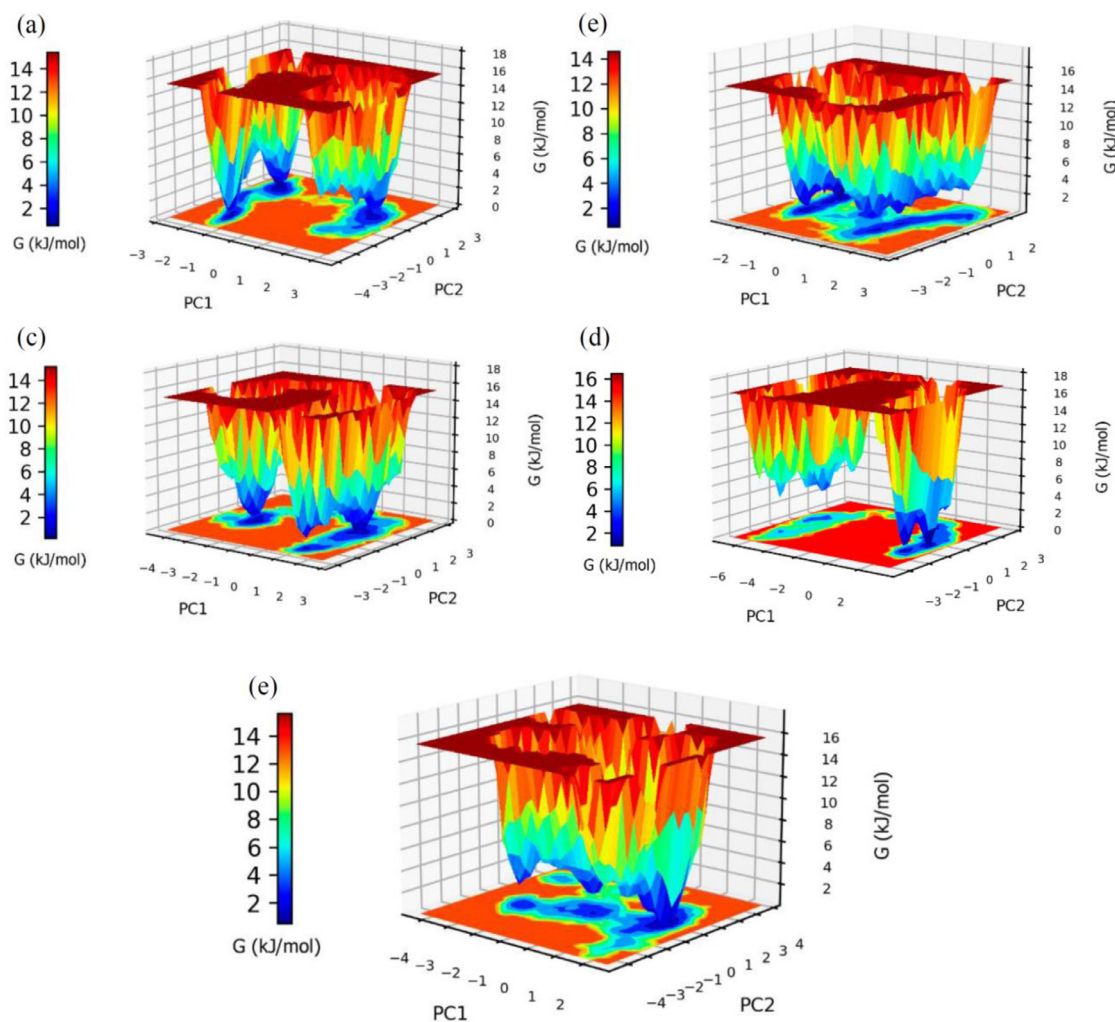


Fig. 15. Free energy landscape (FEL) between first and second principal components (PC1, PC2) graph representation for SARS-Cov-2 2'-O-MTase complexed with (a) mulberrofuran F (b) 24-methylene cycloartenol ferulate (c) 10-hydroxyusambarensine (d) 3-benzoylhosloppone and (e) without any compound systems.

analysis of the four systems, the higher fluctuation that was observed with the interacting residues is consistent with previous reports, where higher structural fluctuations occurred in ligand binding sites of catalytic loop regions [65]. The RoG and SASA plots of all the systems did not show fluctuation above the optimum of $>2 \text{ \AA}$ further indicating that the structural integrity of the proteins was preserved [66]. The binding free energy that is measured from the simulation trajectories provides more accurate computation of ligand binding affinities than the static docking analysis [67]. These results were calculated based on the total binding free energy of the complex. In these calculations, the binding free energy (ΔG_{bind}) measures the affinity of a ligand to its target protein. Thus, the ΔG_{bind} calculations are important to gain in-depth knowledge about the binding modes of the hits in drug design [68]. The results from the binding free energy calculation (MM-GBSA) agreed with that from the docking analysis; further establishing Mulberrofuran F as the most potent phytochemical. Also, from the predictive drug-likeness, pharmacokinetic and ADMET filtering analyzes, the top docked phytochemicals were predicted to be druggable and nontoxic. The result from the filtering analyzes showed descriptors that suggests a favorable ADMET and pharmacokinetic properties. This further indicates the druggable potential of the LPs [69,70]. The LPs displayed properties that suggest their ability to cross the BBB, hence their potential to ensure overall viral clearance in the brain cells [47]. Also, the LPs expressed high pos-

sibility of human intestinal absorption and not susceptible to the permeability-glycoprotein (P-gp, a drug efflux pump). Therefore, it is suggested to be well absorbed into the blood stream, subverting the restraining effect of the P-gp to pump compounds back into the intestinal lumen [71].

6. Conclusion

Herein, we have virtually screened a list of 226 bioactive phytochemicals compiled from a literature search of compounds from African medicinal flora with reported bioactivity against infectious diseases (including viral infection). Altogether the top four docked compounds demonstrated higher binding affinity than the reference inhibitors to the coronaviruses 2'-O-MTase, nevertheless, they displayed similar binding pattern as the reference inhibitors. These phytochemicals were identified to interact with important catalytic residue in the substrate binding site of SARS-CoV-2, SARS-CoV and MERS-CoV 2'-O-MTase as the reference inhibitors, hence, they may disrupt the RNA capping machinery, the replication and survival of the viruses. These potential inhibitors of SARS-CoV-2 2'-O-MTase were stable in a simulated dynamic condition and exhibited positive drug-likeness in the ADMET studies, thus, they are well adaptable for a recommended *in vitro* and *in vivo* experimental studies as anti-COVID-19 agents.

Table 6
Physicochemical properties of the top-binding phytochemicals from African plants to SARS-CoV-2 2'-O-MTase.

a) Physicochemical properties	Mulberrofuran F	24-Methylene cycloartenol ferulate	3- Benzoylhosloppone
Molecular weight (g/mol)	630.68	630.68	418.52
Num. heavy atoms	47	44	31
Num. arom. heavy atoms	27	6	6
Num. rotatable bonds	4	9	4
Num. H-bond acceptors	8	4	4
Hydrogen bond donor	5	1	1
cLogP	4.55	4.55	2.61
Molar Refractivity	179	179	101.11
TPSA (Å ²)	132.75	55.76	
Lipinski violation	1	1	0
Drug likeness			
Lipinski	Yes	Yes	Yes
Veber	Yes	Yes	Yes
Bioavailability Score	0.55	0.17	0.55
(b) ADMET SAR	Absorption (Probability)		
Blood-Brain Barrier	BBB+ (0.565)	BBB+ (0.649)	BBB+ (0.835)
Human Intestinal Absorption	HIA+ (0.984)	HIA+ (0.973)	HIA+ (0.974)
Caco-2 Permeability	Caco2+ (0.577)	Caco2+ (0.745)	Caco2+ (0.678)
P-glycoprotein Substrate	Non-substrate (0.727)	Substrate (0.795)	Substrate (0.752)
P-glycoprotein Inhibitor	Non-inhibitor (0.656)	Non-inhibitor (0.606)	Non-inhibitor (0.806)
Renal Organic Cation Transporter	Non-inhibitor (0.910)	Non-inhibitor (0.797)	Non-inhibitor (0.814)
Distribution (Probability)			
Subcellular localization	Mitochondria (0.786)	Mitochondria (0.802)	Mitochondria (0.838)
	Metabolism	Metabolism	Metabolism
CYP450 2C9 Substrate	Non-substrate (0.780)	Non-substrate (0.758)	Non-substrate (0.777)
CYP450 2D6 Substrate	Non-substrate (0.852)	Non-substrate (0.812)	Non-substrate (0.912)
CYP450 3A4 Substrate	Non-substrate (0.567)	Non-substrate (0.822)	Non-substrate (0.813)
CYP450 1A2 Inhibitor	Non-inhibitor (0.5154)	Non-inhibitor (0.5814)	Non-inhibitor (0.5814)
CYP450 2C9 Inhibitor	Non-inhibitor (0.8197)	Non-inhibitor (0.500)	Non-inhibitor (0.539)
CYP Inhibitory Promiscuity	Low CYP Inhibitory Promiscuity (0.8818)	Low CYP Inhibitory Promiscuity (0.729)	Low CYP Inhibitory Promiscuity (0.815)
Toxicity			
AMES Toxicity	Non-AMES toxic (0.506)	Non-AMES toxic (0.50)	Non-AMES toxic (0.882)
Carcinogens	Non-carcinogens (0.934)	Non-carcinogens (0.9712)	Non-carcinogens (0.912)
Acute Oral Toxicity	III (0.429)	IV (0.607)	III (0.749)
Rat Acute Toxicity LD ₅₀ , mol/kg	3.3280	1.4139	1.9882
Aqueous solubility (LogS)	-4.3480	-5.8146	-4.7646
Pharmacokinetics			
GI absorption	low	High	High
Log K _p (skin permeation) cm/s	-4.53	-1.42	-1.42

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Informed consent statement

Not applicable.

Data availability

All data supporting the findings of this study are available within the article and its supplementary materials.

Ethical approval

Not required.

Declaration of Competing Interest

The authors declare that they have no competing interests.

CRediT authorship contribution statement

Gideon A. Gyebi: Conceptualization, Visualization, Writing – original draft, Methodology. **Oludare M. Ogunyemi:** Methodology, Writing – review & editing. **Adedotun A. Adefolalu:** Writing – review & editing. **Alejandro Rodríguez-Martínez:** Methodology. **Juan F. López-Pastor:** Methodology. **Antonio J. Banegas-Luna:** Methodology. **Horacio Pérez-Sánchez:** Supervision. **Adegbenro P. Adegunloye:** Writing – review & editing. **Olalekan B. Ogunro:** Writing – review & editing. **Saheed O. Afolabi:** Writing – review & editing.

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Supplementary materials

Supplementary material associated with this article can be found, in the online version, at doi:10.1016/j.molstruc.2022.133019.

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