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## **Article**

# **Pharmacoinformatics** molecular and docking simulation-based phytochemical screening of Artocarpus altilis against SARS-**CoV-2 by targeting Mpro**

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#### **Abstract**

A sudden onset of an emergent viral pathogen, SARS-CoV-2, manifested globally in late December 2019, originating in Wuhan, China, swiftly precipitating the COVID-19 pandemic. Subsequent to its rapid dissemination, intensive research efforts have been dedicated to identifying effective therapeutic modalities to counteract this formidable virus. In this context, a computational scrutiny of compounds derived from *Artocarpus altilis* (breadfruit) was undertaken to discern potential inhibitors of the SARS-CoV-2 main protease  $(M<sup>pro</sup>)$ enzyme, crucial for viral replication. Initiating our investigation, a comprehensive drug-likeness analysis was employed to discern compounds with optimal pharmacological attributes. Subsequently, molecular docking studies were conducted, focusing on the Mpro enzyme, with selected compounds from *A. altilis*. Nirmatrelvir, an FDA-approved drug in combination with ritonavir, served as the benchmark inhibitor in these analyses. The compounds under investigation, namely cycloartomunin, dihydrocycloartomunin, namely cycloartomunin, dihydrocycloartomunin, cycloartobiloxanthone, artomunoxanthentrione, and cycloartomunoxanthone demonstrated notable binding affinities of −7.6,  $-7.7, -7.7, -8.3,$  and  $-8.1$  kcal/mol, respectively, in the molecular docking studies. Comparative analysis revealed nirmatrelvir to exhibit an affinity of −8.1 kcal/mol in the same docking environment. Subsequently, a comprehensive pharmacological assessment was undertaken, juxtaposing the top five test compounds with the standard inhibitor. Concomitant with this, a computational toxicity analysis was integral to our assessment. Ultimately, the investigated compounds displayed promising docking outputs coupled with moderate pharmacological profiles. This study advocates further experimental validations to ascertain the inhibitory potential of these *A. altilis*-derived compounds against the SARS-CoV-2  $\rm \dot{M}^{pro}$  enzyme.

Keywords: COVID-19, SARS-CoV-2, M<sup>pro</sup> enzyme, breadfruit, *Artocarpus altilis*, molecular docking.

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# **Introduction**

The SARS-CoV-2 outbreak in late 2019 and its global propagation have had substantial global consequences on public health, economics, and social aspects (Miyah et al., 2022). SARS-CoV-2, also known as COVID-19, is a highly pathogenic virus that was discovered in Wuhan, China, near the end of 2019 (Hu et al., 2021; Ludwig & Zarbock, 2020). Since then, it has swiftly spread over the world, and the World Health Organization (WHO) designated it a global pandemic in March 2020 (Patnaik, 2021). As a result of the pandemic, many countries throughout the world imposed tight measures such as lockdowns and travel restrictions to prevent the spread of COVID-19 (Inoue & Todo, 2020; Zhong et al., 2021). Until November 22, 2023, the World Health Organization (WHO) had recorded over 772 million COVID-19 cases globally (https://covid19.who.int/). Regrettably, the aforementioned has led to a total of more than 6.98 million fatalities, which is a substantial figure in comparison to other diseases (WHO, 2023).

The causative agent of COVID-19, SARS-CoV-2, is a positive-sense, single-stranded RNA virus belonging to the family of Coronaviridae (Hu et al., 2020). The virus's surface has a special spike protein that binds to the human angiotensin-converting enzyme 2 (ACE2) receptor. This makes it easier for the virus to get into host cells and start copying itself (Deng et al., 2021). The SARS-CoV-2 virus has undergone significant mutations, leading to the emergence of several new variants (WHO, 2022).

The main protease (Mpro) enzyme in SARS-CoV-2 is very important for the virus to replicate and is thought to be a good target for drugs that could help treat COVID-19 (Huynh et al., 2021).  $M<sup>pro</sup>$ , also known as 3CL pro, is responsible for the proteolytic cleavage of the viral polypeptide during replication, which is essential for the production of functional viral proteins(Dharmashekara et al., 2021).The inhibition of  $\mathbf{M}^{\text{pro}}$  can potentially prevent viral replication and transmission, making it a valuable target for drug development against COVID-19 (Citarella et al., 2021).Recent years have seen a lot of research into the structure and function of Mpro. This has led to the creation of a number of inhibitors that target this protease (Hu et al., 2022). These inhibitors have shown effective inhibition of  $M<sup>pro</sup>$  in vitro and in vivo, demonstrating their potential as a therapeutic option for treating COVID-19 (Li et al., 2021).

*Artocarpus altilis*, commonly referred to as breadfruit, belongs to the family Moraceae (Turi et al., 2015). It is a notable plant with a substantial therapeutic profile (Sikarwar et al., 2014). Many ongoing studies are looking at the pharmacological effects of *Artocarpus*  *altilis*. Anti-inflammatory activity, antioxidant activity, antimicrobial potential, immunomodulatory potential, anti-diabetic activity, anti-cholinergic effect, chelating activity, nutritional assessment, as a cosmetic agent, ACE inhibitors, and other studies are being conducted on these plants (Ajiboye et al., 2020; Jalal et al., 2015; Riasari et al., 2019; Tara Kamal, 2012). Surprisingly, the activity of this plant against SARS-CoV-2was previously undocumented in the literature. As a result, it was chosen for this investigation to investigate its action against the targets of Omicron variant SARS-CoV-2.

Nirmatrelvir–ritonavir (Paxlovid™), which has received emergency use authorization by the Food and Drug Administration (FDA) for the outpatient treatment of COVID-19 infection in adults (Gui et al., 2023). Nirmatrelvir is a SARS-CoV-2  $M<sup>pro</sup>$  inhibitor, which actively inhibits the viral replication process by blocking the virus from cleaving viral polyproteins into their functional parts, thus limiting the infection's spread in the body (Park et al., 2022). Multiple studies have shown that nirmatrelvir–ritonavir has potent inhibitory activity against SARS-CoV-2 proteases, making it a promising candidate for COVID-19 treatment (Rodrigues et al., 2022). A clinical trial by Pfizer, the developer of nirmatrelvir/ritonavir, reported that nirmatrelvir/ritonavir treatment reduced the risk of hospitalization or death by 89% in patients with mild to moderate COVID-19 symptoms (Huang et al., 2022). Thus, we have considered nirmatrelvir as a standard inhibitor in our assessment.

Computer-aided drug design (CADD) is a computational method that uses various software tools and algorithms to create, optimize, and test new drugs before they can be synthesized and tested in the lab (Yu & MacKerell, 2017). In this computational assessment, we implemented molecular docking, pharmacokinetic and pharmacodynamic property studies to discover the best possible drug candidate. Molecular docking provided us with information on each ligand's binding affinity, direction, and kind of interaction with the appropriate target proteins. The pharmacokinetic profiles were obtained in order to analyze data on the absorption, distribution, metabolism, and excretion (ADME) of chemicals that occur inside the body following medication delivery. To determine the LD50 values and toxicity classes of the various ligands, a toxicology scrutiny was conducted.

In the current work, we used a screening method to filter out 10 phytochemical compounds from *A. altilis* based on drug-likeness criteria. Through preliminary computational analysis, the compounds were examined across a wide spectrum of studies. As a result, this

framework evaluates and offers potential promising *A. altilis* medication candidates against the SARSCoV-2 Omicron B.1.1.529 strain.

# **Materials and Methods**

## **Selection and Preparation of Ligands**

A total of ten phytochemicals derived from *Artocarpus altilis* were chosen for this investigation based on their drug-likeness. The molecules were chosen using Lipinski's rule of five and the Ghose filter (Khan et al., 2019; Nogara et al., 2015). Only molecules that followed both rules were chosen for this investigation. The chosen ligands' 3D conformers were obtained in SDF format from the online databases PubChem (https://pubchem.ncbi.nlm.nih.gov/) and IMPPAT 2.0 (Indian Medicinal Plants, Phytochemistry, and Therapeutics; https://cb.imsc.res.in/imppat/) (Kim et al., 2016; Vivek-Ananth et al., 2023).

#### **Retrieval and Preparation of Target Protein**

The crystal structure of the M<sup>pro</sup> enzyme (PDB ID: 8D4L) of SARS-CoV-2 was retrieved in PDB format from the RCSB Protein Data Bank database (https://www.rcsb.org/) (Dejnirattisai et al.,2022; Rose et al.,2016). The resolution of the downloaded spike protein was 1.70°A. The protein structures were cleaned by eliminating unwanted atoms and molecules (including ligands) with PyMOL version 2.5.2 software (Schrodinger, LLC) (Lill et al., 2011). The spike protein's receptor-binding domain (RBD) was taken out of the crystal structure, and PyMOL was used to get rid of the extra protein chains. The proteins' chains were stored in PDB formats for molecular docking (Morris & Lim-Wilby, 2008).

#### **Molecular Docking**

The CB-Dock2 server (https://cadd.labshare.cn/cbdock2/php) was used to execute molecular dockings on the chosen ligands against the target proteins (Liu et al., 2022). The binding affinity (kcal/mol) for each protein-ligand combination, as well as noncovalent interactions and docking orientations, were examined using the Dassault Systmes BIOVIA Discovery Studio 2021 Client version 21.1.0 software (Baroroh, S.Si., M.Biotek. et al., 2023). The 2D and 3D schematic drawings of the protein-ligand docking complexes were obtained from BIOVIA Discovery Studio (BIOVIA, Dassault Systèmes, 2023).

#### **ADME and Toxicity Prediction**

The top-docking ligands' canonical SMILES were taken from the PubChem and IMPPAT 2.0 databases and entered into the SwissADME website (https://www.swissadme.ch/) (Daina et al., 2017; Kim et al., 2019). SwissADME provided the ADME (absorption, distribution, metabolism, and excretion)

statistics for each ligand. Following that, the ProTox-II service (https://tox-new.charite.de/protox\_II/) was used to estimate the toxicity profile of each ligand (Banerjee et al., 2018). These two sources were used to record the physicochemical, pharmacokinetic, and pharmacodynamic aspects of each ligand. During the ADME and toxicity prediction, each ligand's topological polar surface area (TPSA), lipophilicity (MLogP), water solubility (LogS), bioavailability score, blood-brain barrier (BBB) permeability, interaction with P-glycoprotein (P-gp), LD50 value, and toxicity class were explored.

## **Results and Discussion**

The docking score, or binding affinity, of a ligand denotes the level of attraction at which the ligand is supposed to bind to the target (Figure 1) (Pantsar  $\&$ Poso, 2018). The docking conformations show at which orientation the ligands bind to the respective targets (Figure 2). A complete docking operation includes the bond types, bond lengths, and a complete overview of the ligand's static interactions with the target (Guedes et al., 2014). Artomunoxanthentrione displayed the highest binding affinity, which is −8.3 kcal/mol. It was found to exhibit almost 20 interactions with the amino acid residues of  $M<sup>pro</sup>$ . The interactions obtained from the CB-Dock2 server for each molecule have been mentioned in Table 1. The binding affinities of cycloartomunin, cihydrocycloartomunin, cycloartobiloxanthone, artomunoxanthentrione, cycloartomunoxanthone, and nirmatrelvir (standard) in complex were  $-7.6$ ,  $-7.7$ ,  $-7.7$ ,  $-8.1$ , and  $-8.1$ kcal/mol, respectively.



**Figure 1:** Binding affinity chart of each compound in complex with  $M<sup>pro</sup>$  enzyme of SARS-CoV-2.

**Table 1:** The binding affinities and noncovalent (hydrogen bonds and hydrophobic) interactions of the test compounds and the standard inhibitor (nirmatrelvir).







**Figure 2:** 3D conformations (left) and 2D view of the docking outputs of (A) cycloartomunin, (B) dihydrocycloartomunin, (C) cycloartobiloxanthone, (D) artomunoxanthentrione, (E) cycloartomunoxanthone, and (F) nirmatrelvir in complex with  $M<sup>pro</sup>$ .

The physicochemical, pharmacokinetic, and pharmacodynamic aspects of the compounds were analyzed using the data obtained from Protox-II and SwissADME. The ADME profiles give a detailed overview of the molecular weights (MWs), topological polar surface areas (TPSAs), lipophilicity, water solubility, gastrointestinal absorptions, bioavailability scores, blood-brain barrier (BBB) permeability, and certain parameters for the ligands. These factors reflect how well molecules will be absorbed, distributed, metabolized, and eventually eliminated once they reach the human body. All of the molecules under investigation have been found to conform to Lipinski's rule of five and the Ghose rule, indicating their potential for oral bioavailability.

In particular, the relationship between the TPSA value and the permeability of the blood-brain barrier has been considered, with TPSA below 90  $\AA^2$  being associated with higher permeability and those above 140  $\AA^2$  being linked to lower permeability (Hitchcock & Pennington, 2006; Pajouhesh & Lenz, 2005). Among the test compounds, cycloartomunoxanthone exhibits the lowest TPSA of  $96.36 \text{ Å}^2$ , while dihydrocycloartomunin and cycloartobiloxanthone shows the highest TPSA, that is 109.36  $\AA^2$ . However, artomunoxanthentrione and cycloartomunin showed TPSA values of 103.04  $\AA^2$  and 98.36  $\AA^2$ , respectively. Normatrelvir, the standard inhibitor exhibits a TPSA of 131.40  $\AA^2$ .

According to Lipinski's rule of five, drugs intended for oral administration should have a lipophilicity value below 5.0. All molecules investigated in this study were found to have lipophilicity (MLogP) below this threshold. Artomunoxanthentrione exhibited the lowest MLogP value as a test compound, which is 1.06, whereas that of cycloartomunoxanthone is 1.84 (highest among the test compounds). Cycloartomunin, dihydrocycloartomunin, and cycloartobiloxanthone have MLogP values of 1.77, 1.77, and 1.63, respectively (as shown in Table 2). The standard inhibitor, nirmatrelvir, has a value of 0.41. Water solubility (LogS (ESOL)) was also investigated for each compound. Cycloartomunin, dihydrocycloartomunin, cycloartobiloxanthone, artomunoxanthentrione, and cycloartomunoxanthone have LogS (ESOL) values of −6.10, −6.39, −5.46, −6.05, and −5.67, respectively. Nirmatrelvir occupies a LogS (ESOL) of −3.58. All five of the test compounds were found to have poor water solubility except cycloartobiloxanthone and cycloartomunoxanthone (both are moderately soluble). However, all the test compounds and the standard inhibitor show high gastrointestinal absorption, and each has a bioavailability score of 0.55, except artomunoxanthentrione (0.56). Cycloartobiloxanthone and Cycloartomunoxanthone were found to be substrates of the P-glycoprotein (P-gp).

In terms of toxicity, the lethal dose 50 (LD50) for cycloartomunin, dihydrocycloartomunin, cycloartobiloxanthone, artomunoxanthentrione, and cycloartomunoxanthone are 5000, 5000, 2500, 120, and 5000 mg/kg, respectively. All test compounds except artomunoxanthentrione belong to toxicity class 5. The toxicity class of artomunoxanthentrione is 3. Nirmatrelvir, on the other hand, has a LD50 of 3000 mg/kg and belongs to toxicity class 5. It is important to mention that the higher the toxicity class, the safer the molecule would be while considering the amount administered. The bioavailability radars of the molecules are shown in Figure 3. The computer-generated estimates of the phytochemicals' hepatotoxicity, carcinogenicity, immunotoxicity, mutagenicity, and cytotoxicity profiles have been meticulously recorded (Table 2). The toxicity data affirms that the most of the compounds, as indicated in green and light green, are safe in terms of their hepatotoxic, carcinogenic, mutagenic and cytotoxic profiles. Nonetheless, certain immunotoxicity concerns are indicated in red. The selected drug, Nirmatrelvir, is safe according to all of the toxicity profiles and is indicated in green and light green. A higher number of greens indicate more safety in terms of toxicity.



**Figure 3:** The bioavailability radars of (A) cycloartomunin, (B) dihydrocycloartomunin, (C) cycloartobiloxanthone, (D) artomunoxanthentrione, (E) cycloartomunoxanthone, and (F) nirmatrelvir retrieved from SwissADME. The colored zone is the suitable physicochemical space for oral bioavailability. LIPO (Lipophilicity):  $-0.7 < XLOGP3 < +5.0$ ; SIZE: 150g/mol < MV  $<$  500g/mol; POLAR (Polarity):  $20\text{\AA}^2$  < TPSA  $<$  130 $\text{\AA}^2$ ; INSOLU (Insolubility):  $-6$   $<$  Log S (ESOL)  $<$  0; INSATU (Insaturation):  $0.25 <$  Fraction Csp3 < 1; FLEX (Flexibility):  $0 <$  Num. rotatable bonds < 9.

**Table 2:** The physicochemical, pharmacokinetic, and pharmacodynamic properties of the molecules with the top 6 docking scores retrieved from SwissADME and Protox-II.



MW: molecular weight; TPSA: topological polar surface area; MLogP: lipophilicity; LogS (ESOL): water solubility; ESOL class: water solubility class; GI absorption: gastrointestinal absorption; bioavailability score: Abbott bioavailability score; BBB permeant: blood-brain barrier permeability; P-gp substrate: interaction with P-glycoprotein; Lipinski Vio: number of violations of Lipinski's rule of Gve; Ghose Vio: number of violations of Ghose's rule; LD50 (mg/kg):lethal dose 50; toxicity class: class based on LD50 value, various colors primarily indicated the toxicity profiles of each chemical; green denoted safety or non-toxicity, light green signified a moderate level of safety, red indicated harm or toxicity.

# **Conclusion**

The goal of this study was to look at compelling compounds from breadfruit (*Artocarpus altilis)* in order to trace possible inhibitors of the  $M<sup>pro</sup>$  enzyme of SARS-CoV-2. Artomunoxanthentrione and cycloartomunoxanthone were found to have affinities above and equal to the standard inhibitor. Artomunoxanthentrione had the highest binding affinity when docked with the target. Moreover, cycloartomunin, dihydrocycloartomunin, and cycloartobiloxanthone also displayed good interactions with the target. This study, however, solely investigates the *in-silico* characteristics and profiles of the phytochemicals from *Artocarpus altilis*. Additional validations are demanded to confirm the efficacy and potentiality of the test compounds as potential inhibitors of the SARS-CoV-2 target enzyme. As a result of this, our study findings suggest cycloartomunin, dihydrocycloartomunin, cycloartobiloxanthone, artomunoxanthentrione, and cycloartomunoxanthone as promising candidates against the M<sup>pro</sup> enzyme of SARS-CoV-2.

# **Conflicts of Interest**

The authors declare no conflicts of interest.

#### *Supplement*

Figures: Go to the following link to download high resolutions of the figures:

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