Screening of phytochemical compounds of *Tinospora cordifolia* for their inhibitory activity on SARS-CoV-2: an *in silico* study

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**ABSTRACT**

In the present study, we explored phytochemical constituents of *Tinospora cordifolia* in terms of its binding affinity targeting the active site pocket of the main protease (3CL pro) of SARS-CoV-2 using molecular docking study and assessed the stability of top docking complex of tinosponnone and 3CL pro using molecular dynamics simulations with GROMACS 2020.2 version. Out of 11 curated screened compounds, we found the significant docking score for tinosponnone, xanosporic acid, cardiofolioside B, tembetarine and berberine in *Tinospora cordifolia*. Based on the findings of the docking study, it was confirmed that tinosponnone is the potent inhibitor of main protease of SARS-CoV-2 with the best binding affinity of −7.7 kcal/mol. Further, ADMET along with toxicity analysis was studied to predict the pharmacokinetics and drug-likeness properties of five top hits compounds. The molecular dynamics simulation analysis confirmed the stability of tinosponnone and 3CL pro complex with a random mean square deviation (RMSD) value of 0.1 nm. The computer-aided drug design approach proved that the compound tinosponnone from *T. cordifolia* is a potent inhibitor of 3CL main protease of SARS-CoV-2. Further, the *in vitro* and *in vivo*-based testing will be required to confirm its inhibitory effect on SARS-CoV-2.

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**Introduction**

Coronaviruses are positive-sense RNA viruses with non-segmented genomes and overlapping Open reading frame (ORF 1a and 1b) with a size genome structures ranging from 28 to 32 KB. The polyproteins are processed into non-structural proteins (NSP) and structural proteins. Coronaviruses regularly undergo recombination after infecting host cells (Lauber et al., 2013). The proteases of novel coronavirus share high percentage of sequence and structure similarity with SARS coronavirus which has the main role in the processing of viral polyprotein. These proteins perform stripping ubiquitin to help in evading immune response of the host (Shanker et al., 2020). Small molecules and drug inhibitors can be designed to block the cascade signaling pathways to protect the host system. Viral proteins have multiple functional domains such as a largest NSP-3 protein and proteases like protease and papain-like protease (3CLpro and PLpro) are involved in polyprotein processing (Báez-Santos et al., 2015). Novel coronavirus infection leads to over activation of CD4 effector T-cells resulting in excessive production of inflammatory cytokines termed as cytokine storm that recruits excess immune cells at the site of infection and reduces tissue capacity which leads to organ failure (Felsenstein et al., 2020). Plant-based immuno-modulatory agents would reduce the severity of hyperactivation of inflammatory response and reduce cytokine storm. Several medicinal plant extracts and their constituents with immuno-modulatory activities proved their ability to reduce the over inflammatory response of cytokines (Liu et al., 2016). *Tinospora cordifolia* which is a medicinal plant native to India commonly called as Guduchi, heart-leaved moonseed and used in Ayurvedic formulations as a medicine to treat several diseases. *Tinospora cordifolia* is found to have antineoplastic, anti-diabetic, hypolipidemic, antioxidant and anti-inflammatory effects. Phytochemical studies of *T. cordifolia* showed the presence of numerous components such as tinosponnone, tinosporin, berberine, palmitine, choline, tembetarine, isocolumbin and tetrahydropalmatine and other alkaloids, steroids, lactones, glycosides and sesquiterpenoids. *Tinospora cordifolia* has an effect on the expression of cyclooxygenase-1, tumor necrosis factor-α, iNOS genes, cytokines (IL-6, IL-1β and PGE2) and NF-κB activation and has the potential to reduce the expression of the COX-2 enzyme and thereby reducing the expression of TNF-α, synthesis of IL-1β and IL-6 and other pro-inflammatory inter interleukins (Tiwari et al., 2018). The phytosterols such as stigmasterol and β-sitosterol found in *T. cordifolia* are found to be responsible for the inhibition of cyclooxygenase-2 synthesis and adhesion molecules in TNF-α stimulated by cells (Philip et al., 2018). Stigmasterol inhibits prostaglandin E2 and other matrix degradation mediators (Joshi & Kaur, 2016; Sharma et al., 2019; Singh & Chaudhuri, 2017; Tiwari...
The phytochemicals of medicinal plants have anti-viral properties and hence, in the present study, we explored the inhibitory effect of phytochemicals of *T. cordifolia* against the SARS-CoV-2 by computational analysis.

**Methodology**

**Protein preparation for docking**

The main protease (3CLpro) in complex with an inhibitor N3 was selected from PDB for the present study. The ligands bound to the protein were removed using Chimera. Structure-based virtual screening was performed using PyRx, Main protease (Mpro, aka 3CLpro, 3-chymotrypsin-like protease, (PDB ID: 6lu7) was used as the macromolecule (receptor). The amino acids interacting with the old ligand were considered as the active site in the generated protein and receptor grid (Peele et al., 2020).

**Ligand preparation and virtual screening**

The phytochemicals, namely phytosterols, berberine, cardiofolioside B, chasmanthin, colubmin, cordifolide A, palmarin, tembetarine, tinosponnone, tinosporinone, xanosporic acid of *T. cordifolia* was downloaded from Indian Medicinal Plants, Phytochemistry and Therapeutics (IMPPAT) database (https://cb.imsc.res.in/imppat/home) in 3D-SDF format. IMPPAT is a curated database of Indian medicinal plants, phytochemical and therapeutic associations (Mohanraj et al., 2018). Energy minimized was performed using Chimera (http://www.cgl.ucsf.edu/chimera). Hydrogen atoms were added before docking and the top-ranked two ligand poses of *T. cordifolia* phytochemical compounds bound to the active site pocket of main protease were used for further analysis. The top hits were analyzed for finding ADME properties using the pkCSM online server (http://biosig.unimelb.edu.au/pkcsm/prediction).

**Molecular dynamics simulations**

The complex of main protease and the top-ranking phytochemical drug candidate with best docking score was prepared for MD simulations and run for 10 ns using GROMACS 2020.2 using CHARMM 36 force field. Topology generation and energy minimization of ligand were performed using the CgenFF server. The trajectories files were generated and saved. The random mean square deviation (RMSD) of the complex with respect to time frame was plotted in the graph (Abraham Peele et al., 2020).

**Results and discussion**

**Molecular docking for phytochemicals of *T. cordifolia***

The 3CL main protease is essential for SARS-CoV-2 replication and considered as crucial target for developing the drug (Peele et al., 2020; Wu et al., 2020; Zhang et al., 2020). A total of 11 phytochemicals of *T. cordifolia* was downloaded from Indian Medicinal Plants, Phytochemistry and Therapeutics (IMPPAT) database (https://cb.imsc.res.in/imppat/home) in 3D-SDF format. IMPPAT is a curated database of Indian medicinal plants, phytochemical and therapeutic associations (Mohanraj et al., 2018). Energy minimized was performed using Chimera (http://www.cgl.ucsf.edu/chimera). Hydrogen atoms were added before docking and the top-ranked two ligand poses of *T. cordifolia* phytochemical compounds bound to the active site pocket of main protease were used for further analysis. The top hits were analyzed for finding ADME properties using the pkCSM online server (http://biosig.unimelb.edu.au/pkcsm/prediction).

![Figure 1. (A) Docked pose of tinosponnone with 3CL main protease. (B) Ligand interaction of tinosponnone with 3CL main protease.](image)
respectively and the interactions were analyzed using 2D diagram. Based on the lowest docking energy value of docked complex, tinosponone was selected as potent inhibitor molecule (Table 1). The compound tinosponone showed hydrogen bond interactions with GLU166 and ASN 142 (Figure 1(a,b)). Xanosporic acid showed hydrogen bond interactions with HIS 41, HIS 163 and GLN 189 (Figure 2(a,b)). Cardiofolioside B showed interactions of hydrogen bond with SER 46, HIS 41, CYS 145, THR 24 (Figure 3(a,b)). Tembetarine showed hydrogen bond interactions with GLU166, HIS 163 (Figure 4(a,b)). Berberine showed hydrogen bond interactions with PHE 140 and ASN 142 (Figure 5(a,b)).

**ADME prediction**

The top 5 phytochemicals of *T. cordifolia* shown better binding affinity with 3CL main protease was subjected to ADME prediction.
The compound, namely tinasponone has satisfied Lipinski’s rule of five conditions and the data were represented in Table 1. Tinasponone possessed good pharmacokinetic properties of druggability with better permeability through membranes and does not interfere with the sites of CYP450 metabolism of other drugs. The physicochemical properties of tinosponone have revealed that the molecular weight 330.38 g/mol with five hydrogen bond acceptors and one hydrogen bond donor, topological polar surface area (TPSA), 76.74Å² and lipophilicity, LogP 2.81. AMES test for toxicity prediction of tinasponone by pkCSM server has shown that the tinasponone has no toxicity.

**Molecular dynamics simulations**

The nature of binding was studied using molecular dynamics simulation study. The top-ranked drug, tinosponone with main protease complex has shown the stability conformation throughout the running of simulation for 10 ns and the RMSD value was obtained below 0.2 nm. The average deviation was found to be negligible, i.e. 0.12 nm (Figure 6). Hence, from the present study on in silico approach, it is suggested that the best docked phytochemical compounds could be tested biologically to develop potent drugs against novel coronavirus.

**Conclusion**

Based on virtual screening and molecular docking analysis, the phytochemical compounds, namely tinosponone, xanosporic acid, cardiofolioside B, tembetarine and berberine of *T. cordifolia* were identified as possible lead molecules to fight against SARS-CoV-2. The present in silico study proved that tinosponone as potent, selective and nontoxic inhibitor of 3CL protease of SARS-CoV-2.

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**Disclosure statement**

No potential conflict of interest is reported by the authors.

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**Figure 5.** (A) Docked pose of berberine with 3CL main protease. (B) Ligand interaction of berberine with 3CL main protease.

**Figure 6.** RMSD analysis for top-ranked lead tinosponone and main protease complex.
