In silico discovery of non-psychoactive scaffolds in Cannabis halting SARS-CoV-2 host entry and replication machinery

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Aim: Coronavirus disease still poses a global health threat which advocates continuous research efforts to develop effective therapeutics. Materials & methods: We screened out an array of 29 cannabis phytoligands for their viral spike-ACE2 complex and main protease (Mpro) inhibitory actions by in silico modeling to explore their possible dual viral entry and replication machinery inhibition. Physicochemical and pharmacokinetic parameters (ADMET) formulating drug-likeness were computed. Results: Among the studied phytoligands, cannabigerolic acid (2), cannabigerol (8), and its acid methyl ether (3) possessed the highest binding affinities to SARS-CoV-hACE2 complex essential for viral entry. Canniprene (24), cannabigerolic methyl ether (3) and cannabichromene (9) were the most promising Mpro inhibitors. Conclusion: These non-psychoactive cannabinoids could represent plausible therapeutics with added-prophylactic value as they halt both viral entry and replication machinery.

Graphical abstract:

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SARS-CoV-2 is the causative agent of raging pandemic of COVID-19 that is manifested by mild-to-severe respiratory tract inflammation leading to pneumonia, lung fibrosis and causing even death. Up till now there is no approved remedy for COVID-19, hence, many research endeavors have been directed toward finding therapeutic drug leads to combat the infection propagation. SARS-CoV-2 is a sense RNA virus bearing projections on its surface, i.e., envelope (E) and spike (S) glycoproteins with the latter initiating the infection via its attachment with host angiotensin-converting enzyme 2 (ACE2). ACE2 is an enzyme expressed in nasal and oral mucosa as well as lung and GI tract.
Nature provides a huge reservoir of anti-infectious phytochemicals with anti-inflammatory potential, from which many innovative phytotherapeutics were developed. Natural products and their metabolites in various plants were recently reported to possess striking inhibitory action against different SARS-CoV-2 proteins which could serve as valuable source of drug leads competent in combating COVID-19 pandemic [1]. Among which, polyphenols and flavonoids were reported to stop SARS-CoV-2 replication and thus, mitigating the infection propagation [2]. Natural therapeutics capable of blocking viral entry into host cells can represent a plausible therapeutic avenue with an added prophylactic action [3]. Emodin, tetra-O-galloyl-β-D-glucose and hesperidin are the only few S-protein-ACE2 binding inhibitors reported till now [4]. The virus releases its RNA into the host cells after its entry, then the viral genome codes for nonstructural proteins viz. main protease (M<sup>pro</sup>), RNA-dependent RNA polymerase, the structural glycoprotein and its helicase, and papain-like protease [4]. Inhibition of the M<sup>pro</sup> activity is regarded as a second most important class of antiviral therapeutics as being able to inhibit viral replication, hence halting the infection spread [5]. Viral replication in host cells is well evidenced to be associated with hyper-induction of proinflammatory cytokines, known as ‘cytokine storm’ and immune activation which are the main pathogenic features of COVID-19 [6]. Oolonghomobisflavan-A, theasinsensin-D, and theaflavin-3-O-gallate from tea were reported as M<sup>pro</sup> inhibitors [7].

*Cannabis sativa* L. biosynthesizes an arsenal of resorcinyl core decorated scaffolds with para-oriented terpenyl, pentyl and isoprenyl groups collectively known as phytocannabinoids that possess a wide therapeutic spectrum viz. anti-cancer, anti-epileptic and analgesic actions [8].

Cannabis plants are divided into three different phenotypes according to the cannabinoid content: drug-type (phenotype I; *C. indica* Lam.), containing a higher proportion of THC, intermediate-type (phenotype II; *C. ruderalis* Janisch.) and fiber-type or hemp (phenotype III; *C. sativa* L.) with cannabidiol (CBD) as the main phytocannabinoid. The fiber-type *Cannabis* contains a very low Δ9-THC level, however it is rich in CBD and its related nonpsychoactive phytochemicals. This type has been used for textile manufacturing and food purposes but with no therapeutic actions. Various non-psychoactive phytocannabinoids are present in fiber-type cannabis such as cannabinoic acids viz. cannabigerolic acid (CBGA) and cannabidiolic acid (CBDA), and their decarboxylated forms, i.e., cannabigerol (CBG) and cannabidiol (CBD), in addition to other minor cannabinoids viz. cannabichromene (CBC), cannabinol (CBN), cannabichromenic acid (CBCA), cannabionic acid (CBNA) and several phenolic compounds such as cannflavins A and B, and canniprene [9]. The phytocannabinoidial content decreases from the upper to the lower part of the plants, i.e., hemp flowers contain the highest level, followed by the upper leaves. The peak concentration of cannabinoids is attained during the growing periods of the plant (10–11 weeks after cultivation) [10].

In continuation to previous research work [4,11,12], we aimed in the current study to explore the relative dual prophylactic/therapeutic potential of 29 cannabis phytochemicals by *in silico* exploration of their inhibitory action on viral spike-ACE2 complex, i.e., essential for the viral entry, as well as the viral replication machinery via the M<sup>pro</sup>. Further, physicochemical and pharmacokinetic parameters (ADMET) formulating drug-likeness were computed.

**Materials & methods**

**Phytoligands under study**

An array of 29 phytoligands from *C. sativa* belonging to different classes viz. phytocannabinoids, stilbenoids and flavonoids were included in the study, i.e., cannabidiolic acid (CBD) (1), cannabigerolic acid (CBGA) (2), cannabigerolic acid monomethyl ether (3), cannabigerovarinic acid (4), cannabigerovarin (5), cannabidiol (CBD) (6), cannabidivarinic acid (7), cannabigerol (CBG) (8), cannabichromene (CBC) (9), cannabivaricin (10), Δ9-tetrahydrocannabivaricin (Δ<sup>9</sup>-THCV) (11), cannabidivarin (CBDV) (12), (+)-cannabichromenic acid (13), cannabionic acid (14), cannabidiocinodiol (CBND) (15), cannabinol methyl ether (16), cannabiol (17), cannabicyclolcic acid (18), cannabicyclol (19), cannielsoin A (20), cannabinol (21), cannflavin A (22), cannflavin B (23), canniprene (24), cannabifuran (25), dehydrocannabifuran (26), cannabicitran (27), cannabiripsol (28) and cannabimovone (29). 2D structures of these different phytocannabinoids were downloaded as structure-data file (SDF) from PubChem (https://pubchem.ncbi.nlm.nih.gov).

**Molecular docking of the phytoligands**

Molecular Operating Environment (MOE) software package version 2016.10 (Chemical Computing Group, Montreal, Canada) was used for molecular docking of the two viral proteins under study viz. SARS-CoV-2 S protein C-terminal domain (SARS-CoV-2-CTD) and M<sup>pro</sup>. SARS-CoV-2-CTD crystal structure complexed with human
ACE2 (hACE2) and the coordinates of Mpro in complex with its inhibitor 2-cyclohexyl-N-(3-pyridyl)acetamide were retrieved from the protein data bank (PDB ID 6LZG and 5R84, respectively) [11,13–15]. The proteins were prepared employing the default “structure preparation” module settings, where hydrogen atoms were added, hydrogen bonds were optimized, atomic clashes were removed, and the crystal structures were refined. The docking protocol was adopted as previously reported in [16]. Docking results were presented as a list based on the S-scores with RMSD values below 2 Å and graphically showing the phytoligands interactions.

In silico analysis of physicochemical properties & ADME profiling of the phytoligands

The physicochemical properties and ADME profiles of the studied phytoligands in human body were predicted using SwissADME calculation toolkit [17] and the online-server PreADMET (https://preadmet.bmdrc.kr/) [18].

Results

Molecular docking analysis of cannabis phytoligands within SARS-CoV-2 spike (S) protein–ACE2 complex

SARS-CoV-2 entry into the host cell is initiated by the spike (S) glycoprotein attachment [19] to hACE2 which proceeds by the cleavage of S protein by host cell proteases into S1 and S2 subunits for receptor recognition and cell membrane fusion, respectively. S1 is further subdivided into N-terminal domain (NTD) and C-terminal domain (CTD), which both function as a receptor-binding entity [20]. The S1 CTD of SARS-CoV-2 functions as the key region interacting with the hACE2 receptor. The crystal structure of SARS-CoV-2-CTD in complex with a single hACE2 molecule in asymmetric unit (PDB ID: 6LZG) was used in our study [13].

Docking simulations were used to explore the ability of the studied phytoligands in destabilizing the virus-enzyme complex or preventing its formation. We proposed that possible accommodation/fitting of such phytochemicals at the SARS-CoV-2-CTD-h2ACE interface and their possible interactions with the key amino acids of the complex may provide promising insights for the possible prophylactic action of the studied phytoligands.

The most suitable site for docking the phytoligands into the SARS-CoV-2-CTD-2hACE binding interface was localized using ‘site finder’ feature of MOE 2016.10. in the absence of a co-crystallized inhibitor at the interface of the studied complex by employing flexible docking mode using reference phytochemical viral entry inhibitor, i.e., hesperidin [21–23]. Most of the studied phytoligands were found to display moderate to promising binding affinities compared with hesperidin (30) (Table 1). Cannabigerolic acid (CBGA) (2) recorded the best binding affinity (-7.55 kcal/mol), followed by cannabigerol (CBG) (8), cannabigerolic acid monomethyl ether (3), cannabigerovarinic acid (4), cannabichromene (CBC) (9) and cannabifuran (25) showing slightly less binding affinities ranging from -7.21 to -7.01 kcal/mol. The remaining phytoligands showed a moderate fitting (-6.93 to -5.99 kcal/mol).

Molecular docking analysis of cannabis phytoligands within SARS-CoV-2 Mpro

Most of the investigated phytoligands displayed promising binding affinities to SARS-CoV-2 Mpro as compared with the reference (30) (Table 2). Canniprene (24) showed the best binding affinity (-7.02 kcal/mol) followed by cannabigerolic acid monomethyl ether (3), cannabichromene (CBC) (9), cannabigerolic acid (CBGA) (2), cannabinoids methyl ether (16), cannflavin B (23), cannabidinodiol (CBND) (15) and cannabinolic acid (14) which exhibited less binding affinities (-6.98 to -6.67 kcal/mol) but still better than the reference [30] that exhibited a binding affinity of -6.66 kcal/mol.

In silico prediction of physicochemical properties, ADME & drug-likeness of cannabis phytoligands

SwissADME online server was employed to predict the most important physicochemical properties that formulate drug-likeness parameters of the most promising phytoligands [17]. The promising phytoligands exhibited drug-like bioavailability according to Veber’s [24] and Lipinski’s [25] parameters. Among the selected phytoligands, cannabigerovarinic acid (4), cannflavin B (23) and canniprene (24) were observed to be in full accordance to Lipinski’s and Veber’s parameters, whereas cannabigerolic acid (CBGA) (2), cannabigerol (CBG) (8) and cannabichromene (CBC) (9) showed one violation. Additionally, Pre-ADMET program [26] was employed for the calculation of ADME descriptors.
Table 1. Results of docking simulations of the studied cannabis phytoligands within SARS-CoV-2 spike (S) protein–ACE2 complex.

| No. | Name of phytoligands                      | Δ \( G^1 \) (kcal/mol) | Interactions at the binding interface | hACE2 residues | SARS-CoV-2-CTD residues |
|-----|------------------------------------------|-------------------------|--------------------------------------|----------------|-------------------------|
| 1   | Cannabidiolic acid (CBD)                 | -6.53                   | Glu37                                | No interaction |                         |
| 2   | Cannabigerolic acid (CBGA)               | -7.55                   | Glu37, Arg403, Glu406, Tyr505        |                |                         |
| 3   | Cannabigerolic acid monomethyl ether     | -7.21                   | His34                                |                |                         |
| 4   | Cannabigerovarinic acid                  | -7.21                   | Glu37                                | Glu406         |                         |
| 5   | Cannabigerovarin                         | -6.45                   | No interaction                       | Glu406         |                         |
| 6   | Cannabidiol (CBD)                        | -6.19                   | His34                                | No interaction |                         |
| 7   | Cannabidivarin                           | -6.42                   | No interaction                       | No interaction |                         |
| 8   | Cannabigerol (CBG)                       | -6.66                   | No interaction                       | No interaction |                         |
| 9   | Cannabichromene (CBC)                    | -7.11                   | No interaction                       | No interaction |                         |
| 10  | Cannabivarin                             | -6.29                   | His34                                | No interaction |                         |
| 11  | 9-Tetrahydrocannabivarin (9-THCV)        | -5.99                   | His34                                | No interaction |                         |
| 12  | Cannabidiavin (CBDV)                     | -5.99                   | No interaction                       | No interaction |                         |
| 13  | (±)-Cannabichromenic acid               | -6.33                   | No interaction                       | Glu409, Lys417 |                         |
| 14  | Cannabinolic acid                        | -6.64                   | His34                                | Arg403         |                         |
| 15  | Cannabinoliodiol (CBND)                  | -6.49                   | Glu37                                | No interaction |                         |
| 16  | Cannabinol methyl ether                  | -6.65                   | No interaction                       | No interaction |                         |
| 17  | Cannabinol                              | -6.53                   | Glu37                                | No interaction |                         |
| 18  | Cannabicyclic acid                       | -6.73                   | No interaction                       | No interaction |                         |
| 19  | Cannabicyclol                           | -6.75                   | His34                                | No interaction |                         |
| 20  | Cannabielsoin A                         | -6.81                   | No interaction                       | No interaction |                         |
| 21  | Cannabiriotriol                          | -6.93                   | No interaction                       | Glu406         |                         |
| 22  | Cannflavin A                             | -6.55                   | His34                                | No interaction |                         |
| 23  | Cannflavin B                             | -6.62                   | His34                                | No interaction |                         |
| 24  | Canniprene                               | -6.67                   | No interaction                       | No interaction |                         |
| 25  | Cannabifulurane                          | -7.01                   | His34                                | No interaction |                         |
| 26  | Dehydrocannabinol                       | -6.83                   | His34                                | Arg403         |                         |
| 27  | Cannabichitrin                           | -6.50                   | No interaction                       | No interaction |                         |
| 28  | Cannabiripsil                            | -6.80                   | His34                                | No interaction |                         |
| 29  | Cannabinomovone                          | -6.93                   | No interaction                       | No interaction |                         |
| 30  | Hesperidin                               | -7.51                   | His34                                | Lys417         |                         |

The key residues involved in the SARS-CoV-2-CTD-2hACE complex formation are listed in bold.

\( \Delta G^1 \) is the ligand–receptor complex binding free energy at RMSD ≤ 2 Å.

Discussion

As revealed from molecular docking of the phytoligands within SARS-CoV-2 S protein–ACE2 complex, the most promising phytoligands were cannabigerolic acid (CBGA) (2) and its monomethyl ether derivative (3), and cannabigerol (8), belonging to cannabigerol-type phytocannabinoids which were featured by the presence of a linear isoprenyl residue. These phytochemicals were found to be most enriched in cannabis varieties resulted from hybridization [27]. They are nonpsychoactive with low cannabinoid (CB) receptors potency however, they possess powerful antioxidant and anti-inflammatory properties which make them good candidates for managing the inflammatory conditions [28].

Cannabichromene-type phytocannabinoids, represented by cannabichromene (9), possess isoprenyl moiety that is oxidatively fused to the resorcinyl ring. This type was reported to exhibit potent TRPA1 activation, anti-inflammatory and anti-nociceptive actions via the inhibition of cyclooxygenase enzyme and its associated prostaglandins but with no affinity to cannabinoid receptors, i.e., devoid of psychotropic activity [29]. These compounds are more abundant in the vegetative stage of hemp plant than in its reproductive stage [30]. Cannabifuran (25), an oxidatively cyclized analog of cannabidiol (15), is commonly isolated from aged samples of hashish with no reported biological actions till now.
### Table 2. Results of docking simulations of the studied cannabis phytoligands within SARS-CoV-2 M\(^{\text{pr}}\).

| No. | Name of phytoligands | \(\Delta G^1\) (kcal/mol) | Interactions at the active site of SARS-CoV-2-main protease protease |
|-----|----------------------|--------------------------|-------------------------------------------------|
| 1   | Cannabidiolic acid (CBDA) | -6.12 | No interaction |
| 2   | Cannabigerolic acid (CBGA) | -6.81 | No interaction |
| 3   | Cannabigerolic acid monomethyl ether | -6.98 | No interaction |
| 4   | Cannabigerovarinic acid | -6.57 | Glu166 |
| 5   | Cannabigerovarin | -6.62 | Gin189 |
| 6   | Cannabidiol (CBD) | -5.84 | No interaction |
| 7   | Cannabidivarinic acid | -6.38 | Cys145 |
| 8   | Cannabigerol (CBG) | -4.68 | Asn142 |
| 9   | Cannabichromene (CBC) | -6.87 | No interaction |
| 10  | Cannabivar 

Most of the promising phytoligands were able to accommodate into the interface and interact with the key amino acids (Figures 1 & 2). Accordingly, they can destabilize or halt the virus–receptor engagement which is commonly dominated by polar contacts mediated by these key hydrophilic amino acid residues [13]. Cannabigerolic acid monomethyl ether (3), cannabidiol (CBD) (6), cannabivar 

\(\Delta\gamma\)-tetrahydrocannabinovin (\(\Delta\beta\)-THCV) (11), cannabino acid (14), cannabicyclic acid (19), cannabionin A & B (22 & 23), cannabipicurin (25), dehydrocannabiphuran (26) and cannabiripsol (28) showed interactions with His34 of the hACE-2 involved in the complex formation (Figure 1), whereas (\(\pm\))-cannabichromenic acid (13) interacted with Lys417 which is the key residue of the SARS-CoV-2 CTD but did not exhibit any interactions with the hACE-2 side (Figure 1Q & R).

Cannabidiol (6) is a major phytocannabinoid in fiber hemp formed by spontaneous decarboxylation of its acidic form by the act of light and heat upon cannabis ageing. Despite having structural similarity with \(\Delta\gamma\)-THC, it exhibits a distinct pharmacological profile and lacks any psychoactive properties with a very low affinity for cannabinoids receptors. Cannflavins A and B (22 & 23), methylated isoprenoid flavones, are two of unique \(C. sativa\) flavonoids with a well-reported anti-inflammatory action via the inhibition of 5-Lipoxygenase (5-LO) and prostaglandin E 2 synthase (mPGE5-1) [9].

ACE2 inhibition was reported as being unfavorable in COVID-19 patients due to the consequent decrease in the production of angiotensin 1–7, that possess anti-inflammatory, antifibrotic and vasodilatory actions via the Mas...
Figure 1. Docking simulation of the studied phytoligands into the binding interface of SARS-CoV-2-CTD in complex with hACE2 (Protein Data Bank ID: 6LZG). (A & B) 3D and 2D binding modes of 1 (cyan sticks). (C & D) 3D and 2D binding modes of 2 (yellow sticks). (E & F) 3D and 2D binding modes of 3 (magenta sticks). (G & H) 3D and 2D binding modes of 4 (orange sticks). (I & J) 3D and 2D binding modes of 5 (light pink sticks). (K & L) 3D and 2D binding modes of 6 (green sticks). (M & N) 3D and 2D binding modes of 10 (violet sticks). (O & P) 3D and 2D binding modes of 11 (white sticks). (Q & R) 3D and 2D binding modes of 13 (red sticks). (S & T) 3D and 2D binding modes of 14 (deep yellow sticks). (U & V) 3D and 2D binding modes of 15 (pink sticks). (W & X) 3D and 2D binding modes of 17 (blue sticks). (Y & Z) 3D and 2D binding modes of 19 (deep cyan sticks). (AA & AB) 3D and 2D binding modes of 21 (grey sticks). (AC & AD) 3D and 2D binding modes of 22 (light brown sticks). (AE & AF) 3D and 2D binding modes of 23 (dark blue sticks). (AG & AH) 3D and 2D binding modes of 25 (dark green sticks). (AI & AJ) 3D and 2D binding modes of 26 (brown sticks). (AK & AL) 3D and 2D binding modes of 28 (dark red sticks). (AM & AN) 3D and 2D binding modes of 30 (light cyan sticks) in the binding interface of SARS-CoV-2-CTD in complex with hACE2 (PDB ID: 6LZG). The names of the phytoligands (1–30) are given in the Materials & methods section.
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Short Communication

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receptor [31]. The protective role of ACE2 was revealed in animal models of acute respiratory distress syndrome [32]. Patients who administer angiotensin-II inhibitors (ACE1) was reported to suffer from severe symptoms with a higher mortality rate than their counterparts who did not take these medications [33]. Thus, ligands not interacting with hACE2 side such as (+)-cannabichromenic acid (13) will be considered promising anti-SARS drug leads with a preventive potential.

Canniprene (24), a unique dihydrostilbenoid to C. sativa, recorded the highest binding affinities to SARS-CoV-2 M\(^{\text{pro}}\), which possesses anti-inflammatory action via the inhibition of pro-inflammatory eicosanoids production. Canniprene was reported to exhibit more potent action than cannflavin A as 5-LO inhibitors, however, it is less effective in inhibiting mPGES-1 [34]. Moderate fitting was observed in the rest of the cannabis phytoligands (ranging from -6.62 to -4.31 kcal/mol). Most of the phytoligands were able to accommodate into the active site and showed strong interactions with the key aminoacids (Figure 1). Cannabigerovarinic acid (4), cannabinolic acid (14), cannabidiol (17), cannabitrile (21) and canniprene (24) interacted with Glu166 that is involved in the reference inhibitor–active site complex. Additional interactions were observed in the phytoligands, i.e., cannabidivar (CBDV) (12) and cannabiripsol (28) with Glu189, however, phytoligands, i.e., cannabinodiol (CBND) (15) and cannabifuran (25) exhibited interactions with Cys145. Cannabicyclol (19), cannabielsoin A (20) and cannabicitran (27) showed interactions with the neighboring Met165. Cannflavin B (23) interacted with the active site through its key residue His163, whereas cannabigerol (CBG) (8) and cannabimovone (29) interacted through Asn142. A comparative display of the energy binding value of \(\Delta G\) (-kcal/mol) of Spike–ACE2 complex and M\(^{\text{pro}}\) with the studied cannabis phytoligands is illustrated in Figure 2.

All the investigated phytoligands displayed acceptable aqueous solubility with canniprene (24) appearing at the top of the list with an excellent intestinal absorption (>90%), medium to high BBB penetration, moderate permeability through CaCo-2 cells model and low permeability through MDCK ones (Table 3). Most of the phytoligands were predicted to possess moderate to high plasma proteins binding profiles and expected to be more bound to plasma proteins (PPB \(\approx 96–100\%\)). The investigated phytoligands were also predicted to be incapable of inhibiting cytochromes P450 2D6 (CYP2D6) but not CYP3A4, except cannabigerolic acid monomethyl ether (3) that was devoid of inhibiting both CYP2D6 and CYP3A4 (Table 3).

Indeed, oropharynx and nasopharynx are the main entry ports for SARS-CoV-2 as well as sources of transmission. Both saliva and nasopharyngeal secretions harbor a significant viral load in asymptomatic or presymptomatic virus carriers, and hence playing crucial role in both the pathogenicity and the transmission of SARS-CoV-2. Accordingly, an antiviral oral and nasopharyngeal rinses can be considered as one of the most efficient intervention in combating SARS-CoV-2 transmission [35]. Previous studies reported for the efficacy of mouthwashes in inhibiting viruses such as HIV, herpes simplex virus (HSV) and Middle East respiratory syndrome coronavirus (MERSCoV) [36–38]. Since, canniprene (24) possessed the best optimal ADMET profile, among the studied phytoligands, in addition to its most M\(^{\text{pro}}\) inhibitory action, this phytoligand might serve as promising drug candidate to be administered orally as being devoid of any psychoactivity.

Cannabigerol-type phytocannabinoids, i.e., cannabigerolic acid (2), cannabigerol (8) and its acid methyl ether (3), possessed the most potent inhibitory action on the viral entry machinery in our in silico modeling study which makes them promising candidates to be administered via topical application in the form of mouthwash/gargles or intranasally. Cannabigerolic acid monomethyl ether (3) was revealed to exhibit a dual promising inhibitory action against both the viral entry and the replication machinery. However, the efficacy and tolerability of the identified phytoligands need to be confirmed for its effective integration into the clinical practice. Table 3.

**Conclusion**

There is an urgency in finding new anti-SARS therapies with an added prophylactic value to be employed side by side to the developed vaccines for an efficient curtailment of COVID-19. The current study posed new drug leads from cannabis with low toxicity and in silico dual inhibitory action against viral entry and replication machinery that is essential for a more efficient management of COVID-19. The phytoligands ‘cannabigerolic acid (2), cannabigerolic acid monomethyl ether (3) and cannabichromene (9)’ possessed the highest inhibitory potential against both viral entry and replication machineries as illustrated in the comparative histograms shown in Figure 3. It seems crucial now to develop dosage forms for local application in the oral cavity to halt the SARS-CoV-2 load for prophylaxis [44]. Intranasal delivery of the proposed drug leads is then suggested as an additional option to minimize COVID-19
Figure 2. Docking simulation of the studied phytoligands into the active site of SARS-CoV-2-main protease (Protein Data Bank ID: 5R84). (A & B) 3D and 2D binding modes of 4 (cyan sticks). (C & D) 3D and 2D binding modes of 5 (yellow sticks). (E & F) 3D and 2D binding modes of 7 (magenta sticks). (G & H) 3D and 2D binding modes of 8 (orange sticks). (I & J) 3D and 2D binding modes of 12 (light pink sticks). (K & L) 3D and 2D binding modes of 14 (green sticks). (M & N) 3D and 2D binding modes of 15 (violet sticks). (O & P) 3D and 2D binding modes of 17 (white sticks). (Q & R) 3D and 2D binding modes of 18 (red sticks). (S & T) 3D and 2D binding modes of 19 (deep yellow sticks). (U & V) 3D and 2D binding modes of 20 (pink sticks). (W & X) 3D and 2D binding modes of 21 (blue sticks). (Y & Z) 3D and 2D binding modes of 23 (deep cyan sticks). (AA & AB) 3D and 2D binding modes of 24 (grey sticks). (AC & AD) 3D and 2D binding modes of 25 (orange sticks). (AE & AF) 3D and 2D binding modes of 26 (dark blue sticks). (AG & AH) 3D and 2D binding modes of 27 (dark green sticks). (AI & AJ) 3D and 2D binding modes of 28 (brown sticks). (AK & AL) 3D and 2D binding modes of 29 (dark red sticks). (AM & AN) Overlay of the docked (light cyan sticks) and the co-crystallized inhibitor 2-cyclohexyl-N-(3-pyridyl)acetamide 30 (green sticks) in 3D and 2D binding modes, in the active site of SARS-CoV-2-main protease (Protein Data Bank ID: 5R84). The names of the phytoligands (1–30) are given in the Materials & methods section.
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Table 3. Computational physicochemical, ADMET and drug-likeness parameters of the selected phytochemicals.

| Compound no. | Physiochemical parameters | ADME | Bioavailability and drug-likeness |
|--------------|--------------------------|------|----------------------------------|
|              | LogP† | M.Wt | HBA | HBD | NROTB | TPSA†† | S (mg/L) | HIA§ | PPB§ | BBB# | Caco2‡ | CYP3A4 inhibitor | CYP2D6 inhibitor | Lipinski§§ | Veber¶¶ |
| 2            | 4.19  | 360.49 | 4   | 3   | 10    | 77.76  | 1.07    | 93.79 | 100  | 6.57 | 19.45 | 10.54 | Yes | None | 1 violation; MLOGP > 4.15 | Yes |
| 3            | 4.40  | 374.51 | 4   | 2   | 11    | 66.76  | 0.50    | 96.19 | 99.88 | 6.95 | 23.47 | 2.62 | Non | None | 1 violation; MLOGP > 4.15 | 1 violation; Rotors > 10 |
| 4            | 3.74  | 324.32 | 4   | 3   | 8     | 77.76  | 5.92    | 93.24 | 100  | 4.53 | 20.23 | 3.82 | Yes | None | Yes | Yes |
| 8            | 4.70  | 316.48 | 2   | 2   | 9     | 40.46  | 0.18    | 93.53 | 100  | 17.70 | 45.92 | 49.42 | Yes | None | Yes | Yes |
| 9            | 4.31  | 314.46 | 2   | 1   | 7     | 29.46  | 0.08    | 97.63 | 100  | 16.28 | 25.53 | 27.14 | Yes | None | Yes | Yes |
| 23           | 1.31  | 368.38 | 6   | 3   | 4     | 100.13 | 2.46    | 90.89 | 95.64 | 0.66 | 7.80  | 0.22 | Yes | None | Yes | Yes |
| 24           | 3.34  | 342.43 | 4   | 2   | 7     | 58.92  | 21.91   | 93.83 | 100  | 6.80 | 33.86 | 0.12 | Yes | None | Yes | Yes |

8BBB: Blood-brain barrier penetration; Hydrogen blood acceptor; HBD: Hydrogen blood donor; M.Wt: Molecular weight; NROTB: Number of rotatable bonds; PPB: Plasma protein binding; S: Aqueous solubility (mg/L).
†Log P: logarithm of compound partition coefficient between n-octanol and water.
††TPSA: polar surface area. Drug-like TPSA ≤ 140–150 Å².
‡MDCK: permeability through Madin-Darby Canin kidney cells. PMDCK values < 25 nm/sec (low permeability), values ≥ 25–500 nm/sec (medium permeability) and values > 500 nm/sec (high permeability) [42].
§HIA: human intestinal absorption. HA values < 20% (poorly absorbed), values > 20–70% (moderately absorbed) and values > 70% (well absorbed) [39].
¶PPB values ≤ 90% (poorly bound) and values > 90% (strongly bound) [18].
#BBB values ≤ 0.1 (low CNS absorption), values > 0.1–2 (medium CNS absorption) and values > 2 (high CNS absorption) [40].
‡‡Caco2: permeability through cells derived from human colon adenocarcinoma. PCaco2 values < 4 nm/sec (low permeability), values ≥ 4–70 nm/sec (medium permeability) and values > 70 nm/sec (high permeability) [41–43].
§§Lipinski rule: log P ≤ 5, M.Wt ≤ 500 Da, HBA ≤ 10 and HBD ≤ 5 [26].
¶¶Veber rule: NROTB ≤ 10 and TPSA ≤ 140 [25].
propagation and this represents an important area in viral respiratory diseases research. However, this is worthy of further exploration for the sake of identifying optimal parameters for intranasal delivery.

The remarkable in silico anti-SARS entry and replication machinery potential of cannabis phytoligands has shed the light on revisiting the cannabis bright side. This further stimulates the exploration of the possible biological actions of other cannabis phytochemicals than the psychoactive cannabinoid motif.

Summary points

- In silico screening of 29 cannabis phytoligands against SARS spike-ACE2 complex and Mpro.
- Cannabigerolic acid and cannabigerol were revealed as inhibitors of SARS-CoV-hACE2 complex.
- Canniprene and cannabigerolic methyl ether possessed anti-Mpro activity.
- The identified phytoligands could serve as anti-SARS therapies with an added prophylactic value.

Author contributions

A Khattab: conceptualization, methodology, writing – original draft preparation; M Teleb: methodology, software, formal analysis, data curation. All authors read and approved the final manuscript.

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Availability of data & material
Data sharing is not applicable to this article because no datasets were generated or analyzed in this study.

References
1. Singh Rahul, Bhardwaj Vijay Kumar, Sharma Jatin, Dinesh Kumar RP. Identification of potential plant bioactive as SARS-CoV-2 Spike protein and human ACE2 fusion inhibitors. Comput. Biol. Med. 136(2021), 104631 (2021).
2. Singh R, Bhardwaj VK, Sharma J, Purohit R, Kumar S. In silico evaluation of bioactive compounds from tea as potential SARS-CoV-2 nonstructural protein 16 inhibitors. J. Tradit. Complement. Med. 12(1), 35–43 (2021).
3. Sharma J, Kumar Bhardwaj V, Singh R, Rajendran V, Purohit R, Kumar S. An in silico evaluation of different bioactive molecules of tea for their inhibition potency against non structural protein-15 of SARS-CoV-2. Food Chem. 346, 128933 (2021).
4. Sayed AM, Khattab AR, Asmaa M, AboulMagd HMH, Rateb Zaid H, Abdelmohsen UR. Nature as a treasure trove of potential anti-SARS-CoV drug leads: a structural/mechanistic rationale. RSC Adv.: 10(34), 19790–19802 (2020).
5. Rossi F, Tortora C, Argenziano M, Di Paola A, Punno F. Cannabinoid receptor type 2: a possible target in SARS-CoV-2 (CoV-19) infection? Int. J. Med. Sci. 21(11), 3809 (2020).
6. Das S, Sarmah S, Lyndem S, Singha Roy A. An investigation into the identification of potential inhibitors of SARS-CoV-2 main protease using molecular docking study. J. Biomol. Struct. Dyn. 39(9), 3347–3357 (2020).
7. Bhardwaj VK, Singh R, Sharma J, Rajendran V, Purohit R, Kumar S. Identification of bioactive molecules from tea plant as SARS-CoV-2 main protease inhibitors. J. Biomol. Struct. Dyn. 59(10), 3449–3458 (2020).
8. Sholler DJ, Huestis MA, Amendolara B, Vandrey R, Cooper ZD. Therapeutic potential and safety considerations for the clinical use of synthetic cannabinoids. Pharmacol. Biochem. Behav.: 199, 173059 (2020).
9. Nocerino E, Amato M, Izzo AA. Cannabis and cannabinoid receptors. Fitoterapia 71(Suppl. 1), S6 (2000).
10. Nahler G. Pure cannabidiol versus cannabidiol-containing extracts: distinctly different multi-target modulators. Altern. Complement. Integr. Med. 4(1), 1–11 (2018).
11. El Hawary SS, Khattab AR, Marzouk HS, et al. In silico identification of SARS-CoV-2 spike (S) protein–ACE2 complex inhibitors from eight Tecomia species and cultivars analyzed by LC-MS. RSC Adv.: 10(70), 43103–43108 (2020).
12. Khattab AR, Teleb MKM. In-silico study of potential anti-SARS cell entry phytoligands from Phlomis aurea: a promising avenue for prophylaxis. Future Virol.: 16(11), 761–775 (2021).
13. Wang Q, Zhang Y, Wu L et al. Structural and functional basis of SARS-CoV-2 entry by using human ACE2. Cell 181(4), 894–904.e9 (2020).
14. Fearon D, Powell AJ, Douangamath A., PanDDA Analysis Group. Deposition–crystal structure of COVID-19 main protease in complex with Z219104216. Protein Data Bank 10, (2020).
15. Pratama MRF, Poerwono H, Siswodihardjo S. Molecular docking of novel 5-O-benzoylpinostrobin derivatives as wild type and L858R/T790M/V948R mutant EGFR inhibitor. J. Basic Clin. Physiol. Pharmacol.: 30(6), 20190301 (2020).
16. Elhawary SS, Younis IY, El Bishbishy MH, Khattab AR. LC–MS–based chemometric analysis of phytochemical diversity in 13 Ficus spp. (Moraceae): correlation to their in vitro antimicrobial and in silico quorum sensing inhibitory activities. Ind. Crops Prod.: 126, 261–271 (2021).
17. Daina A, Michielin O, Zoete V. SwissADME: a free web tool to evaluate pharmacokinetics, drug-likeness and medicinal chemistry friendliness of small molecules. Sci. Rep. 7, 1–13 (2017).
18. Shady NH, Khattab AR, Ahmed S et al. Hepatitis c virus ns3 protease and helicase inhibitors from red sea sponge (Amphimedon) species in green synthesized silver nanoparticles assisted by in silico modeling and metabolic profiling. Int. J. Nanomed.: 15, 3377–3389 (2020).
19. Lu G, Wang Q, Gao GF. Bat-to-human: spike features determining “host jump” of coronaviruses SARS-CoV, MERS-CoV, and beyond. Trends Microbiol.: 23(8), 468–478 (2015).
20. Zhou P, Yang X Lou, Wang XG et al. A pneumonia outbreak associated with a new coronavirus of probable bat origin. Nature 579(7798), 270–273 (2020).
21. Wu C, Liu Y, Yang Y et al. Analysis of therapeutic targets for SARS-CoV-2 and discovery of potential drugs by computational methods. Acta Pharm. Sin. B. 10(5), 766–788 (2020).
22. Haggag YA, El-Ashmawy NE, Okasha KM. Is hesperidin essential for prophylaxis and treatment of COVID-19 infection? Pharmacol. Biochem. Behav.: 15, 3377–3389 (2020).
23. Mahdian S, Ebrahim-Habibi A, Zarrahi M. Drug repurposing using computational methods to identify therapeutic options for COVID-19. J. Diabetes Metab. Disord.: 19(2), 691–699 (2020).
24. Veber DF, Johnson SR, Cheng HY, Smith BR, Ward KW, Kopple KD. Molecular properties that influence the oral bioavailability of drug candidates. J. Med. Chem.: 45(12), 2615–2623 (2002).
25. Lipinski CA, Lombardo F, Dominy BW, Feeney PJ. Experimental and computational approaches to estimate solubility and permeability in drug discovery and development settings. Adv. Drug Deliv. Rev.: 64(Suppl.), 4–17 (2012).
26. PreADMET. https://preadmet.bmdrc.kr/adme/
27. Hanusi LO, Meyer SM, Muñoz E, Tagliatela-Scafati O, Appendino G. Phyto- cannabinoids: a unified critical inventory. Nat. Prod. Rep. 33(12), 1357–1392 (2016).
28. Ligresti A, Moriello AS, Starowicz K et al. Antitumor activity of plant cannabinoids with emphasis on the effect of cannabidiol on human breast carcinoma. J. PharmacoL Exp. Ther. 318(3), 1375–1387 (2006).
29. Hartsel JA, Eades J, Hickory B, Makiyannis A. Cannabis sativa and Hemp. In: Nutraceuticals: Efficacy, Safety and Toxicity, 735–754 (2016).
30. Berman P, Futoran K, Lewitus GM et al. A new ESI-LC/MS approach for comprehensive metabolic profiling of phytocannabinoids in cannabis. Sci. Rep. 8(1), 1–5 (2018).
31. Abassi ZA, Skorecki K, Heyman SN, Kinanee S, Armaly Z. COVID-19 infection and mortality: a physiologist’s perspective enlightening clinical features and plausible interventional strategies. Am. J. Physiol. - Lung Cell. Mol. Physiol. 318(5), L1020–L1022 (2020).
32. Imai Y, Kuba K, Rao S et al. Angiotensin-converting enzyme 2 protects from severe acute lung failure. Nature 436(7047), 112–116 (2005).
33. Jakovac H. COVID-19: is the ACE2 just a foe? Am. J. Physiol. Lung Cell. Mol. Physiol. 318(5), L1025–L1026 (2020).
34. Pellati F, Borgonetti V, Brighenti M, Benvenuti S, Corsi L. Cannabis sativa L. and nonpsychoactive cannabinoids: their chemistry and role against oxidative stress, inflammation, and cancer. Biomed. Res. Int. 2018, 1691428 (2018).
35. Hensel A, Bauer R, Heinrich M et al. Challenges at the time of COVID-19: opportunities and innovations in antivirals from nature. Planta Med. 86(10), 659–664 (2020).
36. Eggers M, Eickmann M, Zorn J. Rapid and effective virucidal activity of povidone-iodine products against middle east respiratory syndrome coronavirus (MERS-CoV) and modified vaccinia virus Ankara (MVA). Infect. Dis. Ther. 4(4), 491–501 (2015).
37. Baqui AAMA, Jabra-Rizk MA, DePaola LG, Falkler WA, Meiller TF. In vitro effect of oral antiseptics on human immunodeficiency virus-1 and herpes simplex virus type 1. J. Clin. Periodontol. 28(7), 610–616 (2001).
38. Carrouel F, Gonçalves LS, Conte MP et al. Antiviral activity of reagents in mouth rinses against SARS-CoV-2. J. Dent. Res. 100(2), 124–132 (2020).
39. Yee S. In vitro permeability across Caco-2 cells (colonic) can predict in vivo (small intestinal) absorption in man – fact or myth. Pharm. Res. 14(6), 763–766 (1997).
40. Ma XL, Chen C, Yang J. Predictive model of blood–brain barrier penetration of organic compounds. Acta Pharmacol. Sin. 26(4), 500–512 (2005).
41. Yamashita S, Furubayashi T, Kataoka M, Sakane T, Szehi H, Tokuda H. Optimized conditions for prediction of intestinal drug permeability using Caco-2 cells. Eur. J. Pharm. Sci. 10(3), 195–204 (2000).
42. Irvine JD, Takahashi L, Lockhart K et al. MDCK (Madin-Darby canine kidney) cells: a tool for membrane permeability screening. J. Pharm. Sci. 88(1), 28–33 (1999).
43. Yazdanian M, Glynn SL, Wright JL, Hawi A. Correlating partitioning and Caco-2 cell permeability of structurally diverse small molecular weight compounds. Pharm. Res. 15(9), 1490–1494 (1998).
44. Cegolon L, Javanbakht M, Mastrangelo G. Nasal disinfection for the prevention and control of COVID-19: a scoping review on potential chemo-preventive agents. Int. J. Hyg. Environ. Health 230, 113605 (2020).