Molecular Docking Analysis of Phytochemical Thymoquinone as a Therapeutic Agent on SARS-CoV-2 Envelope Protein

Abdul Khader Sultan Mohideen 1,2

1 Faculty of Zoology, The New College (Autonomous), Chennai-600014, Affiliated to University of Madras, Tamil Nadu, India
2 Correspondence: smdbio@yahoo.co.in

Abstract: The sudden outbreak due to severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) is responsible for causing acute, highly dreadful coronavirus disease (COVID-19). The pore-forming proteins in the SARS-CoV-2 envelope protein employ amphipathic α-helix for pore formation. The pore openings are essential for the transport of ions, toxins, and viroporin activity. Moreover, there is an insurgence to identify lead compounds to target the novel coronavirus for therapeutic purposes. Therefore in the present study, the SARS-CoV-2 envelope protein sequence was analyzed, constructed the three-dimensional homology model, and screened against the bioactive phytochemical Thymoquinone (TQ). Molecular docking was performed between the modeled E protein and TQ to study protein-ligand interactions using ArgusLab 4.0. The investigation revealed that the modeled E protein contains a single α-amphipathic helix identified for the first time across the Amino-terminal region of the transmembrane domain may contribute to pore formation of small membrane proteins. Molecular docking results showed the promising inhibitory potential of the ligand TQ and the binding free energy of the bound complex was found to be -9.01 kcal/mol. The in silico approach has explicitly demonstrated the significant inhibitory effects of the ligand TQ. Therefore it may be used effectively as an antagonist against the SARS-CoV-2 infection owing to its outstanding pharmacological properties.

Keywords: Coronavirus; SARS-CoV-2 Envelope protein; Phytoconstituent; Thymoquinone; Bioactive compound; Homology modeling; Molecular docking.

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1. Introduction

The sudden outbreak of novel coronavirus (CoV) has completely appalled across the world due to its pandemic nature. The coronavirus, also called severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) appeared in the month of December 2019 at Wuhan, China. This virus is highly pathogenic, which is responsible for causing serious, life-threatening pneumonia coronavirus disease 2019 (COVID-19). Infected patients show symptoms of respiratory discomfort and dyspnoea. However, these viruses are not only confined to the respiratory tract but may also affect the central nervous system [1].

CoVs are enveloped viruses consisting of a single-stranded RNA as its genome. The size of the genome ranges from 26 to 32 kilobases, regarded as one of the largest among retroviruses [2]. The most striking feature of coronavirus is the presence of club-shaped spikes

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that protrude from its surface [3]. CoVs infect a wide range of avian and mammalian species and can cause severe illness in humans [4].

Therefore, there is an insurgence to identify natural chemical compounds possessing pharmacological properties expressing a high degree of safety, stability, and efficacy. Medicinal plants have always been the natural source of different phytoconstituents, inevitably used as contemporary drugs that have found their presence in modern therapeutic approaches [5]. They are used to heal and cure human diseases due to the presence of phytochemical constituents [6].

Phytoconstituents occur naturally in various parts of medicinal plants like leaves, flowers, vegetables, bark, and roots that show defense mechanisms against various diseases [7]. The important phytoconstituents of plants include tannins, resins, lectins, alkaloids, flavonoids, phenolic compounds glycosides, etoposides, tanniposides, fatty acids, waxes, terpenoids, phenolic acids, and polypropanoids, etc., [8, 9]. These constituents also contain several moieties like naphthoquinone, anthraquinones, and benzoquinone, which facilitate biological or pharmacological interaction with its substrate, essential for drug design and development [10].

In the present study, Nigella sativa L. (N. sativa) was selected due to its outstanding pharmacological and biological activities. N. sativa is also known as black seed or kalonji, is an annual flowering plant in the family Ranunculaceae. It is native to the Indian Subcontinent and West Asia widely used as a medicinal plant throughout the world [11]. It is popularly adopted in traditional systems of medicine such as Unani, Ayurveda, and Siddha. The seeds and its oil have an age-old history of folklore applied in different systems of food and medicine [12]. In Islamic literature, it is recognized as one of the indispensable forms of holistic medicine. It has been recommended to use on a regular basis in Tibb-e-Nabwi, Prophetic Medicine [13]. In the Middle East, the seeds of N. sativa are used for dietary purposes. Moreover, the biological activities of N. sativa seeds are mainly due to the presence of essential oil constituent thymoquinone (TQ) ranging from 30–48%, which was first obtained by El-Dakhakhny [14].

TQ is a major bioactive compound present in black seed oil. The black seed oil is listed in the United States Food and Drug Administration (USFDA) and is “Generally Recognized as Safe.”[15] and has received tremendous significance for its pharmacological properties and therapeutic potential [16]. The major pharmacological activities expressed by TQ are antimicrobial, anti-cancer, anti-inflammatory, antioxidant, etc. [17]. Furthermore, the topical application of N. sativa oil was found to be effective with minimal side effects during the treatment and combat of allergic rhinitis [18]. Moreover, the miraculous power of healing has fetched N. sativa, a place among the top-ranked evidence-based herbal medicines. This is mostly attributed due to the presence of active ingredients of the essential oil TQ, which has therapeutic effects [12].

Many researchers have carried out extensive work in unraveling SARS CoV structural proteins. The structural proteins like envelope (E) protein, spike (S) protein, and membrane (M) protein can be used as suitable candidates to study drug-target interactions. The SARS-CoV E protein is made of 76 amino acids consisting of one alpha-helical transmembrane (TM) region. The TM domain is made of 25 amino acid residues [19]. The E proteins are integrated with the lipidic virion layer, along with the S and M proteins [20].

A recent study pertaining to the screening of medicinal plant compounds showed 32,297 potential antiviral phytochemicals, which may inhibit the SARS-CoV-2 3CLpro
enzyme, thereby facilitating rapid drug discovery against COVID-19 pandemic [21]. In the same manner, inhibition of the ion channel activity in SARS CoV E protein may be used as a suitable target for developing new antiviral drugs [22]. In fact, experimental studies also suggest that defects in the SARS CoV E gene in a mouse model expressing SARS-CoV human ACE2 (angiotensin-converting enzyme-2) receptor failed to grow in the cerebrospinal nervous system [23]. Similarly, the lack of E protein led to the obstruction of virus interaction in the secretory pathway and inhibition of virus maturation [24, 25].

Thus, the main objective of this study is to search for novel bioactive compounds with therapeutic properties. Though crystal structures of the E protein from different organisms have been determined earlier, there is still a need for structural information. Hence, the present investigation is focused on the homology modeling of SARS-CoV-2 E protein to gain insights on its structural and functional aspects. Molecular docking is performed between SARS-CoV-2 E protein and the phytoconstituent TQ to study the protein-ligand interactions by adopting in silico strategies to elucidate its antiviral properties for medicinal purposes.

2. Materials and Methods

2.1. Sequence retrieval and analyses.

The amino acid sequence of SARS-Cov-2 envelope protein was extracted from GenBank: QHD43418.1 [26] and stored in FASTA format for further analysis (Fig.1).

![Figure 1. The amino acid sequence of SARS-CoV-2 E protein (residues 1-75) extracted from GenBank (QHD43418.1) showing different domains.](https://biointerfaceresearch.com/)  

Physico-chemical properties of the SARS-CoV-2 envelope protein were computed by the ProtParam tool (http://web.expasy.org/protparam/), to analyze the molecular weight, theoretical pl, amino acid composition, and grand average of hydropathicity (GRAVY). The antigenicity of the E protein was predicted with AllerTOP v.2.0 [27]. SOPMA server was used to predict the secondary structure of E protein to evaluate the conformational information about the location of the α-helices, β-strands, turns, loops, random, and coils from the protein structure [28].

2.2. Homology modeling of SARS-CoV-2 E protein.

BLASTp search of SARS-CoV-2 E protein using the default parameter was conducted against the Brookhaven Protein Data Bank (PDB) to find a suitable template for comparative or homology modeling (http://blast.ncbi.nlm.nih.gov/). The PDB structures were selected based on sequence coverage and identity, and stored in PDB format for further analysis. The three dimensional (3D) structure of SARS-CoV-2 E protein was predicted using the GalaxyTBM server [29]. The software generated several models through an optimization-based refinement method. The model with the lowest energy was selected for further analysis. The predicted structure was visualized using PyMOL molecular graphics viewer v.2.2.0 (www.pymol.org). Further, the structural domain regions were predicted with the help of the PSIPRED server (http://bioinf.cs.ucl.ac.uk/psipred/).
2.3. Structural validation of the modeled SARS-Cov-2 E protein.

The predicted 3D structure of the model was determined with the help of PROCHECK, Structural analysis, and verification server (SAVES) (http://nihserver.mbi.ucla.edu/SAVES) which estimates the stereochemical aspects besides the main chain and side chain parameters with detailed analysis [30].

The quality of the model was assessed for various parameters, which include Root Mean Square Deviation (RMSD) using TM-score (https://zhanglab.ccmb.med.umich.edu/TM-score/), Errat for estimating the progress of crystallographic model building and refinement, ProSA for the prediction of Z-score value using a suitable template [31].

2.4. Functional assessment of SARS-Cov-2 E protein.

The functional assessment of the modeled E protein was performed using the DIAL server [32], and the active site amino acid residues were predicted with the help of the Galaxy server [29].

2.5. Ligand preparation and molecular docking.

The molecular structure of the ligand TQ was extracted in SDL format from the Brookhaven protein data bank using the PDB ID: 4HCO [33]. The ligand was removed from the complex and then converted into a standard PDB format by applying OpenBabel v.2.3.2 (http://openbabel.org) (Fig. 2) for docking the catalytic site of the modeled SARS-Cov-2 E protein by applying ArgusLab software v4.0 [34]. In the process of docking, the ligand-binding sites of the modeled E protein were prepared for TQ based on the Galaxy server results obtained before molecular docking. The ArgusLab program has a built-in empirical scoring function (AScore) and docking engine (ArgusDock). The energy minimization was carried out using the universal force field (UFF) molecular mechanics. The calculations were adjusted to dock mode and flexible model with a grid resolution of 0.40 Å. The pose with the lowest binding energy was considered for calculating the scoring function, which shows the modeled E protein-ligand interactions. The different binding modes between the bound complexes were observed through PyMOL V.2.3.3 (www.pymol.org). The docking was further confirmed by uploading TQ and the modeled E protein to the PatchDock server [35].

Figure 2. Molecular structure of the phytoconstituent Thymoquinone (TQ) obtained from PDB ID: 4HCO is used effectively as an inhibitor owing to its biological and pharmacological activities.
3. Results and Discussion

3.1. Physico-chemical properties and amino acid composition of SARS- Cov-2 E protein.

The SARS- Cov-2 E protein has a molecular weight of 83.6 kDa, and the theoretical isoelectric point (pI) was found to be 8.57. Some substances like viruses and certain synthetic polymers have very high molecular weights. In fact, the molecular weight of protein plays a vital role in the prediction of protein structure [36]. The theoretical pI exhibits its alkaline nature (pI >7) due to the presence of basic amino acids, lysine, and arginine. These basic amino acids are involved in membrane protein activity and recognition of membrane voltages [37]. The GRAVY value was found to be 1.128, which is ≥1, indicating its hydrophilic nature and affinity towards water molecules [38]. The modeled E protein identified as a non-allergen and resembles with C-C chemokine receptor type 1, C-C chemokines do not have an amino acid between the first two cysteines of the characteristic four-cysteine motif [39]. Furthermore, it binds to the human macrophage inflammatory protein 1 alpha/RANTES receptor [40]. Subsequently, it transduces a signal by increasing the intracellular calcium ions level, responsible for affecting stem cell proliferation.

The secondary structure predicted by GOR4 (with default parameters) from the SARS-Cov-2 E protein demonstrated that the modeled E protein having the composition of Helix = 33.33%, Coil = 53.33%, Strand =13.33% and Beta-turn = 0.00% respectively. It is quite obvious from this secondary structure prediction that the modeled E protein belongs to the α-helical class of proteins.

3.2. Modeled structure of SARS-Cov-2 E protein.

The 3D structure of SARS-Cov-2 E protein was predicted by homology modeling. BLASTp analysis (http://www.ncbi.nlm.nih.gov/) of the amino acid sequence of the modeled E protein against the PDB database revealed good alignment with the crystal structure of microbial origin, PDB ID: 5X29_A [41]. The best template PDB ID: 5X29_A was assessed based on query coverage (82)% , an E-value (1e-30), and sequence identity (89%). The model with the least energy (-6.61) predicted by the GalaxyTBM server was considered for further analysis. The 3D structure of the modeled E protein mainly composed of α-helices, loops, or β turns (Fig.3). Interestingly, the predicted 3D structure of SARS-Cov-2 E protein has the characteristic features of the coronavirusidae family, designated as an integral small membrane protein.

Figure 3. Three-dimensional structure of SARS- Cov-2 E protein is made of four α-helices, loops, or β turns. The active sites of the residues are present between α1 and α2-helices. The N and C-terminal shown in blue and red color occupy at the extracellular region.
3.3. Description of SARS-CoV-2 E protein structure.

The modeled SARS-CoV-2 E proteins are a short integral membrane protein, which contains three domains, namely N-terminal domain, transmembrane domain (TMD), and C-terminal domain (residues 1-75). The modeled E protein is consisting mainly of four right-handed α-helices connected in between loops or β-turns. The α-helices are the simplest and most common form of protein structural motifs that occurs in a helix bundle. This type of arrangement helps in DNA binding motifs, including helix-turn-helix motifs and zinc finger motifs [42].

The N-terminal domain with α-1 (residues 1-8) situated on one side that forms the extracellular space known as accessible molecular surface or solvent-exposed area. The TMD consists of 2 subunits, namely S1 and S2 made of α-2 (residues 9-38) and α-3 (residues 42-57) helices, aligned parallel and connected by a short loop or β turn (residues 39-41). The loop residues are polar and hydrophobic, lies within the cytoplasmic region—the α-3 shares partly by a transmembrane helix and pore-lining helix of the TMD. The presence of a short loop or β turn is a characteristic feature of thermophilic organisms that provides the extra protein stability at high temperatures, thereby preventing unfolding and denaturation [43]. The N-terminus of the TMD contains a single α-amphipathic helix made of alternative hydrophobic/hydrophilic residues, identified as a structural motif. The structural motif has the amino acid sequence VSEETG (residue number 5-10) where V and G are hydrophobic and in between the polar residues are arranged. While, in TLIVN (residue number 11-15), TN are hydrophilic and in between non-polar residues are present. The α-4 helix (residues 58-75) is positioned extracellular on the C-terminal domain of the SARS-CoV-2 E protein (Fig. 4).

Figure 4. The SARS-CoV-2 E protein is comprising of three domains: the N-terminal domain, the transmembrane domain (TMD), and the C-terminal domain. The TMD consists of two subunits, namely S1 (residues 9-38) and S2 (residues 42-57) connected by a loop.

Moreover, the predicted single amphipathic α-helix at the N-terminus of TMD is located very close to the surface of the lipid membrane, essential for protein-ligand interactions. Moreover, the secondary structural elements (α-helices) may be involved in transmembrane pore formation or pore-forming proteins (PFPs), which employs amphipathic α-helix for pore formation. The pore openings allow the movement of ions and other components like toxins via the cell membrane barrier [44]. This is in agreement with the pore-forming domain consisting of hydrophobic helical hairpin buried between two amphipathic layers of α-helices, as observed in Bax and colicins [45]. Similarly, it is also seen in the CoV E protein TMD consisting of one amphipathic α-helix that oligomerizes to form an ion-conductive pore in membranes [46]. Moreover, they can affect the host cell’s physiology, tissue integrity, and inflammatory responses that may inhibit antimicrobial treatment [47, 48].
Furthermore, computational approaches and spectroscopic studies demonstrate that the SARS-CoV E TMD goes through oligomerization, which is a characteristic feature of ion-channeling proteins [49]. In addition, the synthetic peptides of SARS-CoV E viroporin are capable of transporting Ca2+ associated with the inflammatory response mostly seen in acute respiratory distress syndrome [50]. A similar mechanism was also observed with other viral proteins that might affect or modulate the oligomerization of Infectious bronchitis virus E protein. Furthermore, the M2 protein amphipathic helix motif is necessary for the release of Influenza A virus particles needed for the formation of budding, scission, and viroporin activity [51].

3.4. Protein model validity.

The geometrical and structural consistency of the modeled SARS-Cov-2 E protein was evaluated at the SAVES server. The phi and psi angle distributions of the Ramachandran map using PROCHECK revealed that 92.9% and 5.7% amino acid residues are located in the most favored and additionally allowed regions, while 1.4% and 0.0% amino acid residues in the generously allowed and disallowed regions. This represents the geometric fitness of the modeled E protein (Fig. 5).

![Ramachandran Map](https://biointerfaceresearch.com/)

**Figure 5.** The Ramachandran map for SARS-Cov-2 E protein displays the phi-psi torsion angles of all the amino acid residues. The color shading on the map shows different regions. The red color conforms to the core regions exhibiting the most favorable combinations of phi-psi values.

This is in conformity with the SARS Coronavirus E protein pentameric ion channel, which exhibited 83.3% and 15.6% residues in the most favored and additionally allowed regions, while 1.1% and 0.0% residues were found in the generously allowed and disallowed regions (PDB: 5X29_A). Furthermore, PROCHECK analysis showed both the main and side chain parameters are within the acceptable range suggesting the reliability of the SARS-CoV-2 E protein structure.

Further, the calculation of overall RMSD and TM score between the modeled SARS-Cov-2 E protein and 5X29_A was found to be 1.24 Å and 0.7, respectively. It confirmed a pair of best-aligned chains between the template and predicted model, indicative of a perfect match between two structures. ERRAT map showed an overall quality factor of 74.24%, which is indicative of a good resolution structure. The Z score was found to be -0.35 in SARS-CoV-2 E protein and 0.59 in PDB ID: 5X29_A (Fig. 6 ).
Figure 6. ProSA-web z-scores of the SARS-CoV-2 E protein and the template 5X2N_A in PDB calculated by X-ray crystallography (light blue) or NMR spectroscopy (dark blue) corresponding to their length. The plot exhibits only chains <1000 residues and a z-score ≤ 10. The z-scores of modeled E protein and template are highlighted as large dots.

The predicted 3D model of SARS-Cov-2 E protein after validation was deposited in the Protein Model Database (PMDB). The modeled E protein can be retrieved using the Accession No. PM0082981 (https://bioinformatics.cineca.it/PMDB).

3.5. Functional characterization of the predicted structure.

The active site residues of the modeled SARS-Cov-2 E protein were found to be S6, T9, T11, I13, and V17. The monomeric form of modeled E protein contains three functional motifs, which include two N-glycosylation sites located from the residues 48-51 (NVSL), 66-69 (NSSR) and one Protein kinase C phosphorylation site from the residues 67-69 (SSR) respectively. The modeled E protein also represented two N-glycosylation sites along the subunit S2 and C-terminal region where carbohydrates may be linked to proteins with the help of asparagine (N-linked). In fact, asparagine residue can derive an oligosaccharide only if the residue is an integral part of L-Asn-X-Ser or L-Asn-X-Thr amino acid sequence where ‘X’ can be any amino acid residue except proline. Glycosylation is responsible for maintaining protein stability in the blood. On the contrary, unglycosylated protein contains only 10% of the biological activity of the glycosylated form, and it gets easily removed from the system by the kidneys. Apart from that, potential glycosylation sites play an important role in adding sugars, enabling the protein to be more hydrophilic to take part in interactions with other proteins [52].

3.6. Docking of SARS-Cov-2 E protein to ligand TQ.

When docking runs were executed, the SARS-Cov-2 E protein showed tremendous binding affinity and a remarkable preference to its ligand TQ from the N-terminal domain (Fig 7). The binding energy was found to be -9.01 kcal/mol using Argus Lab 4.0. The findings were further confirmed by the PatchDock server showing similar binding features. The estimated global energy was found to be -32.03 kcal/mol. In fact, negative free binding energy value confirms the modeled E protein and TQ complex to be more stable.

The structure-activity relationship is well illustrated in the present study between the SARS-Cov-2 E protein and ligand TQ interactions. In fact, a number of distinctive characteristics are linked with the active residues of the modeled E protein. TQ access to the binding residues is aided through the N-terminal domain through α1 and α2 helices of the modeled E protein.
The surrounding residues of SARS-Cov-2 E protein and TQ are displayed within 8 Å radius sphere. The modeled E protein residues, S6, T9, T11, I13, and V17 are close to its ligand TQ (central residue) along the N and C direction contribute to short-range interactions with the bound TQ (Fig. 7).

**Figure 7.** Docked complex of SARS-Cov-2 E protein and phytoconstituent Thymoquinone (TQ) obtained from Argus Lab v.4.0 demonstrating protein-ligand interactions. The central residue TQ is in close contact with the active site residues of the modeled E protein.

In the protein-ligand interactions, the side chains of the SARS-CoV-2 modeled E protein residues S6, T9, T11, I13, and V17 showed a close affinity to the bound ligand TQ. The distance between the hydrogen bond lengths was measured between the modeled E protein and ligand TQ (Fig. 8). It was observed from the docked complex that the O atom of the ligand TQ could act as a nucleophile on the side chain residues of modeled E protein. Therefore, the O atom of TQ when it comes in contact with the HG atom of S6 may trigger a chemical reaction leading to the reduction of oxygen and oxidation of hydrogen molecules. This may result in the formation of water or hydrogen peroxide as end products [53].

**Figure 8.** Molecular interactions between SARS-Cov-2 E protein and ligand TQ. The distance between the hydrogen bond lengths was calculated between the modeled E protein and ligand TQ. The O atom (oxygen) of the TQ (Central residue) is in close association with the side chain HG atom (Gamma hydrogen) of the residue S6, may act as a potential nucleophile.

In addition, the HG1 and HG2 atoms of protein residues T9 and T11 base pairs with O atom of ligand TQ, which are polar and capable of forming H bonds as proton donors or acceptors, normally found in close proximity to the surface of protein or lipid membranes. In contrast, the CD1 atom of the protein residue I13 interacts with the O atom of ligand TQ and establishes Carbon-Oxygen hydrogen bonding (CH-O) between them. Klaholz and Moras
demonstrated protein-ligand binding mechanism in nuclear retinoic acid receptor bound to retinoid SR11254 to identify CH-O hydrogen bonds in ligand recognition. Subsequently, they discovered several CH-O bonds among different moieties of the ligand with hydrogen bond donors/acceptors at different positions, showing multiple CH-O interactions may significantly facilitate ligand binding affinity [54]. Thus, it is clearly demonstrated that CH-O bonding significantly contributes to the ligand-protein binding. Furthermore, the C atom of the protein residue V17 interacts with the C atom of ligand TQ. It should be noted that C-C interactions are considered to be weak. However, polarization provides stability to the bound complex [55]. Based on these correlations, it was identified that the modeled E protein residues have a definite role in protein-ligand interactions (Table 1).

| S.No. | Ligand | Residue and atom | Ligand atom | Bond length(A°) |
|-------|--------|-----------------|-------------|----------------|
| 1     | TQ     | S6 & HG         | O           | 2.1            |
| 2     | TQ     | T9 & HG1        | O           | 3.4            |
| 3     | TQ     | T11 & HG1 & CG2 | C           | 3.0            |
| 4     | TQ     | I13 & CD1      | O           | 2.4            |
| 5     | TQ     | V17 & CG2      | C           | 3.9            |

Thus it is explicitly demonstrated from the docked complex that the modeled E protein displayed tremendous affinity to its ligand TQ and can be used effectively as a potential inhibitor against SARS-CoV-2 E protein. Moreover, TQ can enhance the synthesis of reactive oxygen species (ROS) in cells [56] that can be deleterious to DNA, RNA, and proteins, which may eventually cause cell death [57]. Above all, TQ has been clinically tested for its pharmacological activities against various types of ailments, including human carcinomas, as it shows negligible toxicity against normal cells [58]. Therefore, the search for novel antiviral agents to block COVID-19 is an immediate requirement, as there is no approved medicine for COVID-19 to date [59].

4. Conclusions

To sum up, the modeled 3D structure of the SARS-CoV-2 E protein provides comprehensive structural and functional details. More importantly, the structural motif identified for the first time across the N-terminal region of the TMD contains a single α-amphipathic helix that may play a significant role in pore formation of small membrane proteins essential for the transport mechanism of ions, toxic substances, and viroporin activity. The docked complex between the SARS-CoV-2 E protein and ligand TQ displayed tremendous interactions suggesting potential inhibitory effects of the phytoconstituent. Thus, this in silico based approach has explicitly demonstrated the bioactive phytochemicals may inhibit the SARS-CoV-2 E protein ion channel, pore-forming protein activity. The present investigation may be used as a baseline to further develop TQ into a potential antiviral drug against the COVID-19 through suitable in-vitro and in-vivo models for drug transport through oral and intranasal medications.

Funding

This research received no external funding.
Acknowledgments

The author would like to thank the Management, Principal, and Head, PG & Research Department of Zoology, The New College (Autonomous), Chennai-600014, Affiliated to the University of Madras, India for their support and encouragement.

Conflicts of Interest

The authors declare no conflict of interest.

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