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ABSTRACT

Angiotensin-converting enzyme 2 (ACE2), also known as peptidyl-dipeptidase A, belongs to the dipeptidyl carboxylyleptidases family has emerged as a potential antiviral drug target against SARS-CoV-2. Most of the ACE2 inhibitors discovered till now are chemical synthesis; suffer from many limitations related to stability and adverse side effects. However, natural, and selective ACE2 inhibitors that possess strong stability and low side effects can be replaced instead of those chemicals’ inhibitors. To envisage structurally diverse natural entities as an ACE2 inhibitor with better efficacy, a 3D structure-based-pharmacophore model (SBPM) has been developed and validated by 20 known selective inhibitors with their correspondence 1166 decoy compounds. The validated SBPM has excellent goodness of hit score and good predictive ability, which has been appointed as a query model for further screening of 11,295 natural compounds. The resultant 23 hits compounds with pharmacophore fit score 75.31 to 78.81 were optimized using in-silico ADMET and molecular docking analysis. Four potential natural inhibitory molecules namely D-DOPA (Amb17613565), L-Saccharopine (Amb6600091), D-Phenylalanine (Amb3940754), and L-Mimosine (Amb21855906) have been selected based on their binding affinity (−7.1, −7.0 kcal/mol), respectively. Moreover, 250 ns molecular dynamics (MD) simulations confirmed the structural stability of the ligands within the protein. Additionally, MM/GBSA approach also used to support the stability of molecules to the binding site of the protein that also confirm the stability of the selected four natural compounds. The virtual screening strategy used in this study demonstrated four natural compounds that can be utilized for designing a future class of potential natural ACE2 inhibitor that will block the spike (S) protein dependent entry of SARS-CoV-2 into the host cell.

Abbreviations: +ssRNA, positive single strand RNA; ICTV, International Committee on Taxonomy of Viruses; 2D, two dimensional; MD, molecular dynamics; 3D, three dimensional; MERS, Middle East Respiratory Syndrome; AA, amino acids; PBVS, pharmacophore based virtual screening; ACE2, angiotensin-converting enzyme 2; PDB, Protein Data Bank; ADMET, absorption, distribution, metabolism, excretion, and toxicity; RMSD, root mean square deviation; ADT, Auto Dock Tools; RMSF, root mean square fluctuation; COVID-19, coronavirus disease 2019; RO5, rule of five; DUDE, Database of Useful Decoys; SARS-CoV-2, severe acute respiratory syndrome coronavirus 2; HTS, High Throughput Screening; SB, structure-based; SBPM, structure-based pharmacophore model; TMPRSS2, transmembrane protease serine protease 2; T.E.S.T., Toxicity Estimation Software Tool; WHO, World Health Organization.

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1. Introduction

The ongoing novel coronavirus 2019 (nCoV-2019) outbreak has been recently taken place and hit almost all over the world. The disease has been named COVID-19 (coronavirus disease 2019) by the World Health Organization (WHO) recently after the outbreak started in Wuhan, Hubei province of China on 31 December 2019 [1]. The International Committee on Taxonomy of Viruses (ICTV) renamed the outbreak-causing virus as severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) [2], which is responsible for lower respiratory tract disease in humans [3]. COVID-19 pandemic is a great threat to both tropical and polar regions of the world [4], and WHO decreed the disease as a sixth public health emergency. There are rapidly growing numbers of cases globally, and most of the countries have already reported nationwide community transmission [5]. The transmissible and pathogenic virus infects an estimated >223,475,490 people and caused 4,611,347 confirmed deaths (September 09, 2021, 09:41 GMT) across 221 countries and territories around the world [6]. The emergence of new COVID-19 has led to increased demand for new antiviral strategies [7]. But, to date, no specific proven drugs, and therapeutics have been developed that can prevent or treat infections resulting from these pathogens [8,9].

SARS-CoV-2 is a morpohologically oval, round, or often polymorphic shape in nature with a diameter being 60-140 nm [10]. From a genomic perspective, the virus consists of a positive sense single-strand RNA (+ssRNA) and belongs to the lineage of β (beta)-coronavirus [11,12]. The +ssRNA genome of the virus carries a length of around 29.8 kilobases (kb) formed by 29.86% adenosine, 18.39% cytosine, 19.63% guanine, and 32.12% thymine’s [13]. Phylogenetically the virus belonging to the β-genus of coronavirus and their virion consists of the major surface spike (S) protein, integral membrane (M) and envelope (E) protein, and complexes genomic RNA forming nucleocapsid (N) proteins [14]. Early genomic sequence analyses of the SARS-CoV-2 indicated that the virus encodes similar structural proteins (such as spike glycoprotein) and enzymes (helicase, and RNA-dependent RNA polymerase) as of SARS and Middle East Respiratory Syndrome (MERS) virus [7,15]. Furthermore, the SARS-CoV-2 genome shares 79.6% and 96% sequences similarity to SARS-CoV and bat coronavirus, respectively [16], which previously utilizes the ACE2 protein as a receptor to get entrance into the host [17–19].

It has been reported that ACE2 a member of the dipeptidyl carboxy-dipeptidases family has a great impact on supporting the SARS-CoV-2 viral entry into the human host cell [20–22]. The S proteins of the virus are the main target for the neutralization antibody, attach to the host cellular ACE2 receptor, and allow it to enter into the targeted host [12,20,23]. SARS-CoV-2 engages their S protein into two molecular subunits as S1 and S2 [24], where S1 is responsible for attachment to the entry receptor and S2 to viral particle infusions [25]. S1 subunit of the virus is composed of receptor-binding domain and receptor binding motif [26], and this organizational structure help to contacts the receptor ACE2 of the host [20]. The S proteins of the virus also known as a type of class I viral fusion proteins require protease cleavage for their activation. The activation of the S protein is a two-way process known as priming cleavage, efficiently cleaved at the boundary between the S1 and S2 subunits [24], and activating cleavage that activated at the boundary to the S2’ priming site [25]. Once S protein comes in contact with the ACE2, a serine protease called transmembrane protease serine protease-2 (TMPRSS2) helps to cleave the ACE2 for initiating the S protein priming [20], and the resulting activation of the cleavage site mostly leading to the viral fusion and infectivity to the pathogens [20,24,27]. Being an attentional host cellular target, different quantitative kinetics studies through surface plasmon resonance revealed that ACE2 has had 10-20 folds higher affinity to the S protein of the SARS-CoV-2 than other coronaviruses [28]. Additionally, the human ACE2 known as 1 integral membrane protein [29], holds 17 amino acids (AA) residue to the N-terminal domain, 22 AA in the C-terminal domain, and 43 AA residue in the cytoplasmic domain [4], which contains potential phosphorylation sites have major impacts on SARS-CoV-2 viral infection [30]. For example, first helix and lysine 353 and proximal residues of the N terminus of ACE2 extracellular portion, interacts with viral spike glycoprotein has a major role in the virus infectivity. Since host cell entry of the SARS-CoV-2 depends on the receptor ACE2 [20], and compounds responses against this enzyme could at least partially protect against the virus. Although ACE2 was suggested to be a novel SARS-CoV-2 target [18,32,33], only a few and effective inhibitory compounds were identified hitherto.

Natural products, often defined as compounds or substances produced by a living organism are derived from nature and historically use as active components of many traditional medicines [34]. The compounds derived from natural sources have great therapeutic value and represent more than half of FDA-approved drugs [35], which also received attention for their extensive pharmacological and biological activities [36]. However, the chemically synthesized compound has poor biological activities or obvious side effects [37,38], hence developing novel ACE2 inhibitors from natural resources for the treatment of COVID-19 is an urgent matter.

The conventional process of developing new drugs normally involves lengthy, expensive, and requires intense effort [39]. For example, high-throughput screening (HTS) is a process that aids drug discovery by assaying a large number of potential drug-like compounds, whereby the technology combines multiple-well microplate and robotic processing [40]. Nevertheless, HTS requires huge resources, as the cost to process a single HTS program is relatively high and necessitates advance robotic systems [37]. Computer-aided drug design also known as in silico drug design on the other hand is a relatively new method compared to a high-throughput approach, to screen large databases of compounds [41–43]. The in silico virtual screening process help to generate hits to lead compounds in the way to the discovery of new drugs at a faster time and at a lower cost [44]. Advance in silico drug design option tends to decrease the amount of time to develop, design, and optimize a new drug. Over the past few decades, the virtual screening process was engaged to identify the best lead compounds with different structural features for combination with a specific biological target [45–47]. Moreover, the computer-aided drug design by using pharmacophore-based virtual screening (PBVS), molecular docking and dynamics simulation approaches has been identified diverse promising drug targets and hits [14]. Therefore, this study aimed to screen potential natural anti-COVID-19 compounds by combining SBPM, virtual screening, molecular docking, ADMET (absorption, distribution, metabolism, excretion, and toxicity), and MD simulation approaches.

2. Materials and methods

2.1. Pharmacophore modeling

The X-ray-derived crystal 3D structures of human ACE2 protein (PDB ID: 1R4L) in complex with XXS (PDB ID) were obtained from the Protein Data Bank (PDB) [29], and an SBPM was generated automatically using the LigandScout 4.3 advance software [48]. The advanced version of the software was used to determine and visualize the main pharmacophore features of the protein-ligand interaction like hydrogen bond donor and acceptor, negative and positive ionizable area, and hydrophobic interactions of the compound. Multiple chemical features like the number of aromatic rings, binding location of zinc, magnesium, and manganese, halogen bond donor were also detected and mapped for the characterizing of all or specific pharmacophore patterns of ligands [49]. In addition, LigandScout’s excluded volume feature has been added to the 3D structure-based models to retain the steric circumstance of the macromolecule. These excluded volume features ensure and match the sterical requirements of the active site and help to identify desire compounds [48] and increase the selectivity of the virtual screen compounds.
2.2. Pharmacophore model validation

The pharmacophore model generated from complex protein-ligand interaction was validated using a set of known active inhibitors by evaluating their ability to distinguish between known potential active and inactive compounds [19]. A set of known active ACE2 inhibitors were extracted by combining ChEMBL (www.ebi.ac.uk/chembl/) database and an extensive literature search shown in Fig. 1. Compounds with large molecular weight and peptides like molecules have been removed as well as non-ACE2 inhibitors with false activity labels [45]. The remaining set of known active ACE2 inhibitors and their correspondence decoy compounds were retrieved from the enhanced Database of Useful Decoys (DUDE) (http://dude.docking.org), which have been used to validate the 3D-interaction feature model. The decoy set of compounds obtained from the DUDE database was assigned into a multi-conformational virtual screening library by utilizing the command line executable tools “idbgen” in LigandScout 4.3, advance. The main purpose of using the executable tools “idbgen” is to create compound databases for virtual screening, and the idbgen derived LDB file format also speeds up processes for compute clusters of the compounds and annotates each conformation with generated 3D SB-pharmacophore features. To evaluate the preferential efficiency of the pharmacophore, model a 3D molecular structural database screening process was performed [48]. Valuable parameters like active hits (A_h), decoy compounds (D_C), early enrichment factor (EF), the total compounds in the database (D), the total number of hits retrieved (T_H), and goodness of hit score (GH) has been considered to evaluate the performance of the model. The EF of a pharmacophore model is a widely used metric generated during randomly screening of compounds library, which describes the number of active hits found by utilizing an appointed PM model as inverse to the number hypothetically active compounds found [50].

The GH statistical hypothesis test used to compare how well do the observed hits correspond to fit with the assumed pharmacophore model. The GH value ranging from 0 to 1, where 0 indicates acceptance of the null hypothesis model with low or no fit with the model, and 1 indicates most significant with the highest fit to the pharmacophore model [51].

The EF and GH score were calculated from Eqs. (1) and (2) accordingly [50,51], to validate the model performance in our rational drug design approaches. The EF score Eq. (1) and GH score Eq. (2) is given below:

\[
\text{Enrichment factor (EF)} = \frac{A_H \times D}{T_H \times A}
\]  
(1)

\[
\text{The goodness of hit Score (GH)} = \left\{ \frac{A_H \times (3A + T_H)}{4T_H \times A} \right\} \times \left\{ 1 - \frac{(T_H - A_H)}{(D - A)} \right\}
\]  
(2)

where, A_H is the active hits in the database, T_H is the total number of hits retrieved, D is the total compounds in the database, A is the total number of actives in the database, (T_H – A_H) = F_C is the number of false positives compounds, and (D – A) = D_C is the decoy compounds of the database.

2.3. Dataset generation

Ambinter (www.Ambinter.com) is a brand and worldwide supplier of advanced chemicals that supporting the scientific community by providing active compounds for drug discovery. Ambinter is a public access database that contains 36 million purchasable compounds that are ready to dock. Initially, the library was designed and developed for easy access to molecules and drug-like compounds for virtual screening, now it is widely used for VS, PBVS, and force field development [14]. The Ambinter database supply compounds from different vendors by utilizing a compounds code known as SMILES key. For new compounds discovery, the database contains a targeted library of SARS-CoV-2 that has been retrieved for the further screening process. In the present study, chemical features that were generated from the SBPM were used as a query for searching the chemical library. The molecules that fit with the query pharmacophore features were retained and retrieved for further validation.

2.4. Virtual screening

The dataset generated from the Ambinter database was virtually screened based on the validated 3D SB-pharmacophores features. The load screening database features of the LigandScout 4.3 advanced use to convert the compounds into a (*.ldb) database file format, have been uploaded to the molecule database list for quick pharmacophore features based virtual screening [48]. The screening has been done with the relative pharmacophore-fit as a scoring function with a maximum of four omitted pharmacophore features. The hit compounds fitted with the geometry and features of the 3D model were ranked according to the pharmacophore fit score and retrieved for further validation.

2.5. Protein and ligand preparation

Pharmacophore based virtual screening process can generate many ‘hits’ compounds with disproportionate quality [52]. To optimize hits and sorting the better interaction features in comparison with the XX5 inhibitor, and the selected virtually screened compounds have been docked to the binding site of the ACE2 protein. The crystal X-ray structure of the ACE2 protein (PDB ID: 1R4L) in complex with the ligand XX5 was chosen as the positive control system for the study [29]. The ligand in a complex with the protein ACE2 was separated using the Biovia Discovery Studio visualizer version (16.1.0). The PDB structure of proteins was prepared by the following steps (i) bond order has been assigned, (ii), metal ions, cofactors, and water beyond 5 Å from het groups has been removed, (iii) het state has been generated by using Epik at pH 7 ± 2 [53], (ii) non-polar H has been merged and polar H-atoms has been assigned by using PROPKA at pH 7, (iv) the protein structure integrity has adjusted, and the missing side-chain atoms of the protein residues has been predicted by Prime, and (v) the default
2.6. Binding site identification and grid generation

The binding site of the protein was visualized and analyzed to predict the binding affinity and selectivity between the complex crystal protein-ligand interaction [38]. The binding position of the complex interaction was characterized based on hydrogen bond donor-acceptor features, hydrophobic interaction features, negative-positive ionizable area, and zinc chelation in addition to unspecific halogen bond donor-acceptor features of the compounds. Additionally, the PrankWeb (http://prankweb.ech/) is an online resource and server-based tool used to complex protein-ligand binding site prediction and an extensive literature search also confirmed the binding site position of the protein [56]. After identification, the binding sites of the protein a receptor grid was generated using the predicted binding site position of the complex protein-ligand structure.

2.7. Molecular docking (MD) simulation

The docking positions between the selected compounds and ACE2 proteins were predicted by PyRx software [57]. PyRx is open-source software that utilizes Auto Dock 4 (AD4) and Auto Dock Vina (ADV) tools for molecular docking simulation [58]. In this study, the PyRx tools Autodock vina (version 1.1.2) a commonly docking program for molecular docking simulation, has been used to predict the protein-ligand interaction [57]. The crystal structure of ACE2 was downloaded and prepared before molecular docking simulation, and docking positions were visualized by BIOVA Discovery Studio Visualizer Tool (16.1.0). The top 20% of compounds with the highest binding energy (negative sign value), were considered for further investigation.

2.8. Absorption, distribution, metabolism, and excretion (ADME) test

The drug design and development process involves the assessment of ADME to identify molecules with the highest chance to become an effective drug for a specific disease [39]. ADME of a compound provides information about their physicochemical, pharmacokinetics, metabolism, and excretion properties of molecules into urine and feces [59]. Due to numerous compounds with limited access to the physical samples, physicochemical and pharmacokinetic properties of compounds are critical for their initial selection [60]. Early-stage evaluation of ADME in the drug design and discovery process mitigates the fraction of pharmacokinetics-related failure during clinical trials [61]. Nowadays, computer-generated models have complied as a potent substitute for experimental methods for early-stage prediction of ADME. For the studies, a freely accessible Swiss-ADME server (http://www.swissadme.ch/) was used to predict the various pharmacokinetic and pharmacodynamics properties [62]. The results of ADME for all the ligand molecules are represented in Table 3.

2.9. Toxicity test

An early assessment of compound’s toxicity is very much important in the field of drug discovery and development [63]. In-silico evaluation of toxicity is exceptionally arising as an integral stage for the determination of toxicity of a chemical compound that could be potentially harmful to humans and animals [64]. The in-silico toxicity of the selected compounds was evaluated using the Toxicity Estimation Software Tool (T.E.S.T.) version 4.2.1 [65]. The software predicts the toxicity of selected compounds by using Quantitative Structure-Activity Relationships (QSARs) methodologies. This model can predict toxicity by comparing the physical characteristics features of a chemical compound entered by the user. The toxicity of a chemical compound can be assessed in terms of toxicity endpoints like mutagenicity, carcinogenicity, and other features, and this endpoint can be measured both quantitatively and qualitatively [64]. In our studies, the quantitative toxicity endpoints, and drug relevant properties like 96-hour fathead minnow LC50, 48-hour Daphnia magna LC50, 48-hour Tetrahymena pyriformis IGC50, oral rat LD50, bioaccumulation factor, have been evaluated using the T.E.S.T. software. The organ toxicity like hepatotoxicity and toxicity endpoints like carcinogenicity, immunotoxicity, mutagenicity, and cytotoxicity of the compounds were evaluated qualitatively using the ProTox-II (http://tox.charite.de/) web server. The ProTox-II is freely accessible virtual lab for the prediction of toxicities of small compounds and user can be accessed the server without registration. Various toxicity endpoints such as acute toxicity, hepatotoxicity, cytotoxicity, carcinogenicity, mutagenicity, immunotoxicity of the compounds can be retrieved from the server by knowing the two-dimensional structure (2D) of the input compounds. ProTox-II incorporates molecular similarity, fragment propensities and machine-learning features that helps to predict various toxicity endpoints of the compounds, indicating their possible applicability for other compound classes.

2.10. Molecular dynamics (MD) simulation

Molecular dynamics (MD) simulations were used to validate the structural stability and conformational flexibility of a protein-ligand complex [44,66]. MD simulations provide powerful tools for the prediction of each atomic movement of protein, ligands, and physics governing interatomic interactions over a specific time [67]. To validate the structural stability and conformational flexibility protein ligands complex generated from the molecular docking study were subjected to 250 (ns) nanoseconds of MD simulations [68]. The Desmond module of Schrödinger (Release 2020-3) software package with OPLS-2005 force field for generating system topology has been selected to simulate the complex protein-ligand structure [68,69]. A protein to box minimum distance of 0.8 nm was generated to solvate the systems and the protein was centered in the box. The solvent box was filled with the simple point charge water (SPC) type of solvent with a 3-point solvent model. Na+ and Cl− ions were used to neutralize the system to reach a 0.15 M molar concentration. Energy minimization was done by gradient optimization to reduce the net force on each movable atom with a minimization step size of 0.1 (ns) and a maximum force of 500 kJ/mol-nm² to achieve the static state of the system, and minimization graph was generated from the system declared that the optimized construction was solvent saturated and geometrically stable. Equilibration of protein water system ensemble for 10 ns was done for both NVT (constant number of particles, volume, and temperature) and NPT (constant number of particles, pressure, and temperature) ensemble process. The NPT of the system was set at a constant temperature of 300 K and NVT of the system pressure (1.01325 bar) to sustain the stability of the system. Well, an equilibrated dynamics system was performed in the NPT (constant pressure and temperature) ensemble involved for 250 ns production run with integrator time 2 fs and resulted in production were saved every picosecond for further analysis. Analysis of molecular dynamic simulation generated the root-mean-square deviation (RMSD), root-mean-square fluctuation (RMSF), and protein-ligand contact map was used to examine the interactions and stability between the ACE2 proteins and natural compounds.
2.11. Calculation of MM/GBSA

For calculating the binding free energy of ligands to the macromolecules “molecular mechanics generalized Born surface area” (MM/GBSA) methods have become popular methods [70]. The MM/GBSA has been used to estimate the binding free energy of the compounds by using the Maestro package that incorporated in Schrödinger (Release 2020-3) by using default parameters [71].

3. Results

3.1. Structure-based pharmacophore modeling

The first step of our rational drug design approach was the building of a 3D structure-based pharmacophore model. For this purpose, we utilize both theoretical bits of knowledge and experimentally validated 3D structural model storage in the different protein databases. To identify new inhibitor compounds, the crystal X-ray structure of ACE2 protein (PDB ID: 1R4L) in complex with ligand XX5 was retrieved from the PDB, and a 3D-SBPM to the enzymatic cavity was generated. The binding activity of the XX5 inhibitor with the ACE2 protein has confirmed experimentally IC_{50}: 0.44 nM (100), and validated by X-ray diffraction, resolution: 3.00 Å, R-value free: 0.337 R-value work: 0.253 [29], highlighted that this series of inhibitors could bind the active site of the enzyme, followed by inhibition of their biological activity. To determine the active structure of natural inhibitors with similar or better chemical features than XX5, a 3D-SBPM was generated by automatically derives key chemical features of LigandScout 4.3 advanced tools. The automatic key chemical features option of the advance tool determines and displayed a total of 16, including 12 main and four dedicated pharmacophore features (Fig. 2), where two hydrogen bond donors (HBD), five hydrogen bond acceptor (HBA), two negative ionizable (NI), one positive ionizable (PI) area, two hydrophobic (H) interactions features have defined as main, and two aromatic rings (AR), one zinc-binding location (ZNB), and one halogen bond donor (XBD) features have defined as dedicated pharmacophore features of the protein-ligand complex interaction. Fifteen excluded volumes feature, also known as the program, increases selectivity generated features, derived from the sterical circumference of the protein has not displayed in both Fig. 2(a) & (b), the excluded volumes feature has shown in Fig. S1.

Fig. 2. (a) The 3D structure-based pharmacophore model of ACE2 protein in complex with XX5 ligand derived from the X-ray derived crystal structure of the protein retrieved from the PDB (PDB code: 1R4L), (b) Merge pharmacophore features of the complex interaction, where two hydrophobic feature has indicated by yellow spherical shape ◆, one positive ionizable by blue star shape ◆, two negative ionizable by red star shape ◆, two hydrogen bond donor by green spherical or green arrow shape ◆, five hydrogen bond acceptor by red spherical or red arrow shape ◆, one zinc-binding location by royal blue star or cone shape ◆, and one halogen bond donor by violet spherical or arrow shape ◆ have represented within the protein-ligand complex interaction. Fifteen excluded volume areas generated by the pharmacophore model have not displayed in this figure. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

3.2. Pharmacophore model validation

Pharmacophore models generated with distinguished chemical features were subject to validation to evaluate the quality of the model. Validation is an important attribute of a reliable pharmacophore model, which portends the capability to exactly predict internal and particularly external data sets of molecules [19]. The structure-based pharmacophore models were validated using the testing set that included 20 actives known ACE2 inhibitors with correspondence 1166 decoy molecules retrieved from the enhanced database DUDE.

The active test set with inhibitor constant Ki values (0.13 nM to 10,000 nM) were merged with the decoy compounds, to observe how well the pharmacophore model can distinguish the active compounds from inactive compounds. A 3D molecular structural database screening process was performed and subsequently, the EF value and GH value of the compounds were calculated (Table 1). The GH score of 0.75 (Out of 1) and the EF score of 1.1 were found from Eqs. (1) and (2) accordingly, AUC (area under the ROC curve) value found 0.75 (Fig. S3), indicating that the model was very good and is rational for virtual screening.

3.3. Dataset preparation

The Ambinter database contains a mix of natural products with synthetic and/or semisynthetic compounds along with their chemical structures and physicochemical properties. A total of 11,295 natural compounds small molecules were obtained from the Ambinter natural compounds database library (Ambinter and Greenpharma). Lipinski’s rule of five (ROS) also known as Pfizer’s rule of five is used to evaluate drug-likeness properties of compounds [44], which were applied for the subsequent phase of screening.

Table 1

Validation of pharmacophore model using GH scoring method includes enrichment factor, goodness of hit score of the model.

| Parameter | Symbol | Calculation | Model |
|-----------|--------|-------------|-------|
| Total compounds in database | D | – | 12,561 |
| Total actives in database | A | – | 11,295 |
| Decoy compounds | DC | D-A | 1166 |
| Total hits retrieved | TH | – | 23 |
| Active hits retrieved | AH | – | 23 |
| % yield of active hits | – | (AH/TH)×100 | 100% |
| % ratio of active hits | – | (AH/A)×100 | 0.20% |
| Enrichment factor | EF | Eq. (1) | 1.1 |
| False negatives | PF | A – AH | 11,272 |
| False positives | FP | TH – AH | 0 |
| Goodness of hit score | GH | Eq. (2) | 0.75 |
3.4. Pharmacophore-based virtual screening

Compounds with no violation of RO5 were used for pharmacophore-based virtual screening. The final key pharmacophore interaction features generated from the protein-ligand complex were applied to the 11,295 natural compounds, and a total of 23 derivatives was generated with a pharmacophore fit score ranging between 78.81 and 75.31 (Table S1). During pharmacophore-based virtual screening not all the features of the model could be matched, therefore four features of the model have been omitted, resulting in lower pharmacophore fit scores of some derivatives [48]. A higher fit score generated by measuring the geometric fit of the features of a molecule indicates a better fit for the 3D-SBPM. As the higher fit score indicates a better fit for the model, therefore molecules that fit the pharmacophore model should also show activity to the ACE2 protein.

3.5. Binding site identification and receptor grid generation

Identifying binding sites within a protein is an important part of molecular docking, and the virtual screening process [72]. The pocket on a target protein is a combination of the different binding sites that present compatible size, shape, and suitable chemical complementarity with ligands. To determine the binding site of the protein the co-crystal structure of ACE2 bound to XX5 was analyzed through the BIOVA Discovery Studio Visualizer Tool (16.1.0). The Discovery studio visualizer tool revealed that the XX5 of ACE2 formed 4 conserved hydrogen bonds with ARG273, THR371, PRO346, and TYR515 shown in Fig. 3. Another 4 residues namely CYS344, MET360, LYS363, and PHE504 formed an alkyl bond by interacting with the ligand XX5. Attractive charge and salt bridge bonding have also been observed in ARG 514, ARG518, NZN03, and ARG273 positions accordingly.

TYRS10 formed both Pi-Sigma and Pi-Alkyl bonds with the protein, where only one halogen bond with Cl has formed in the protein ASP368 position. Unfavorable acceptor-acceptor and Pi-Pi-T-shape bonds both have formed in the protein HIS345 residual position. The PrankWeb binding site finder and an extensive literature search were also used to confirm the binding site residue and position of the protein. Protein-ligand complex analysis from the server also revealed another 2-binding site position of the protein. Atomic coordinates of the protein and ligand were obtained from the discovery studio and PrankWeb analysis tools used for receptor grid generation. A grid box comprised of X = 57.30, Y = 51.20 and Z = 25 points spaced dimension by 0.350 Å was centered on the ACE2 binding site with a position of X = 40.03, Y = 0.29, and Z = 22.45.

3.6. Molecular docking simulation

Molecular docking is an increasingly important key tool in structure-based virtual screening and computer-aided drug design approaches [73]. The molecular docking approach can be used to predict the predominant binding mode(s) of a ligand with a protein at the atomic level [4]. The goal of ligand-protein docking is to perform virtual screening, rank the results according to their binding energy, understanding the protein-ligand mechanism of action and propose a structural hypothesis of how the ligands inhibit the target [58]. To understand the binding interaction of the targeted protein with the selected 23 compounds, the PyRx tools Autodock vina (version 1.1.2) molecular docking program has been run with an exhaustiveness value 8 [38,57]. The top four compounds have been chosen based on their binding energies ranging between −7.0 kcal/mol and −7.5 kcal/mol (Table 2). Molecular docking scores of selected 4 compounds Amb17613565, Amb6600091, Amb3940754, and Amb21855906 were found −7.5 kcal/mol, −7.1 kcal/mol, −7.1 kcal/mol, and −7.0 kcal/mol respectively, which have better binding energy than the control compounds XX5 (-6.4 kcal/mol).

3.7. Interpretation of protein-ligand binding interaction

Amb17613565, Amb6600091, Amb3940754, and Amb21855906 displayed better pharmacophore fit scores compared to the other compounds and molecular docking simulation determines the binding energy −7.5 kcal/mol, −7.1 kcal/mol, −7.1 kcal/mol, and −7.0 kcal/mol, respectively. Amb17613565 formed 3 conventional hydrogen bonds with ARG273 (2.73 Å distance) and 2 with ASP269 (2.09 Å and 2.03 Å distance) positions, one pi-alkyl bond with a distance 5.21 Å has also been observed in ALA153 residual position shown in Fig. 4.

Amb6600091 has formed four conventional hydrogen bonds with GLY268 (2.39 Å), ASN277 (2.67 Å), and two with both ASP367 (2.23 Å and 2.80 Å) residues. A carbon-hydrogen bond (C–H) and an unfavorable donor-donor bond have also been established in ASP269 (3.53 Å) and THR371 (1.25 Å) residues, accordingly, depicted in Fig. 5.

Amb3940754 also formed four conventional hydrogen bonds with ACE2 protein, 2 with ASP350 (2.34 Å and 3.03 Å), and another two with ARG393 (2.49 Å and 2.82 Å) residue. One van-der-Waals, one pi-pi stacked and one pi-pi T-shaped bond also have been found to formed with GLY352 (3.61 Å), PHE390 (4.06 Å), and PHE40 (5.08 Å) respectively Fig. 6.

Another natural compound Amb21855906 have interacted with the target protein residues by forming four conventional hydrogen bonds with LYS363 (2.11 Å), THR365 (2.67 Å), ASN277 (2.46 Å), and ASN149 (2.72 Å) residues, where one attractive charge bond with ASP368 (5.39 Å) and one salt bridge bond with ASP367 (2.66 Å) have also noticed to formed Fig. 7. Conventional hydrogen bonds also are known as classical bonds are formed when hydrogen covalently bound to elements or a molecular fragment X–H (X = N, O, or F; X is more electronegative than H) forms a second bond to another electronegative atom. All the natural compounds Amb17613565, Amb6600091, Amb3940754, and Amb21855906 formed hydrogen bonds with the desired protein in agreement with the previous binding interaction analysis of our protein-ligand complex studies.

Ion-ion interactions are an attractive force that occurs between two oppositely charged ions that hold together ionic molecules that have been found to form in Amb21855906. In pi-alkyl interactions z-hold a bond over an aromatic group and electron group of an alkyl group that stays around the ring, leading to holding strong stability of the protein-ligand complex.

Table 2

| Ambinter ID  | Formula     | Molecular weight | Docking score (kcal/mol) | Pharm.-Fit score |
|--------------|-------------|------------------|--------------------------|-----------------|
| Amb17613565  | C9H13N3O3   | 197.188          | -7.5                     | 78.81           |
| Amb6600091   | C11H22N6O6  | 276.286          | -7.1                     | 78.75           |
| Amb3940754   | C9H13N3O3   | 165.189          | -7.1                     | 77.71           |
| Amb21855906  | C9H13N3O3   | 198.176          | -7.0                     | 77.12           |
| XX5 (PDB ID) | C7H12Cl2N3 | 428.31           | -6.4                     | N/A             |

A higher docking score indicates better stability of the ligand with the target protein. The pharmacophore fit score of each compound is provided for comparison.

Fig. 3. Describing the protein binding sites generated from the protein-ligand complex structure (PDB ID: 1R4L).
compounds have found to form in the compounds Amb17613565 and Amb21855906.

### 3.8. Pharmacophore features analysis

Pharmacophore features of a compound play a significant role in identifying the specific active site of the protein. The pharmacophore of a compound can be described based on the H, AR, HBA or HBD, PI, NI features that helps design a new drug candidate against a specific disease. These features retain the necessary geometric arrangement of atoms requires to producing a specific biological response. Therefore, the pharmacophore features of the selected four compounds include Amb17613565, Amb6600091, Amb3940754, and Amb21855906 compounds have been analyzed and compared with the query pharmacophore features shown in Fig. 8. Each of the compounds has similar or better pharmacophore properties than the query pharmacophore features. Therefore, the selected compounds should be effective for our target protein.

### 3.9. ADME prediction

The collaboration between drugs and the human body is a bidirectional procedure, drugs influence the human body, bringing about receptor inhibition or activation and the human body disposes of drugs by absorption, distribution, metabolism, and excretion (ADME) [39,50]. Influence and disposes of drugs are interactional that occur simultaneously in the human body after administration and can lead to desired pharmacological function or may lead to undesirable side effects [62]. Drug design research and development (R&D) is an expensive, slow, and risky process that is generally faced with some unrehearsed even catastrophic failures in various stages of drug discovery [60]. The efficacy and safety deficiencies during drug design are thought to be the main cause of R&D-related failures, which depend mainly on compounds’ ADME properties [50]. Therefore, the ADME of the compounds needs to evaluate to minimize the pharmacological failures in the drug discovery process. In this study, the Swiss-ADME an in silico ADME predictions server was used to evaluate the pharmacokinetics and drug-likeness properties of the selected 4 compounds [62]. The ADME profiles like lipophilicity known as partition coefficient between n-octanol and water (Log $P_{o/w}$), water solubility an important property influencing absorption, drug-likeness that determines the chance of a molecule to become an oral drug and medicinal chemistry that evaluates the synthetic accessibility of the compounds were evaluated, and all of the 4 compounds were found to be suitable as listed in Table 3.

### 3.10. Toxicity test

During drug development toxicity evaluation is an important part and should be assessed in preclinical and clinical trial phases. Toxicity and adverse effects of a drug can be evaluated using in vitro and in vivo
compound being synthesized. To access the toxicity and adverse effects of selected 4 compounds the computational methods have been used in rapidity, accessibility and most importantly it can be done before a toxicity prediction have shown great advantages due to their accuracy, ADME properties like physicochemical properties, lipophilicity, water solubility, pharmacokinetics, drug likeness, and medicinal chemistry of selected 4 compounds.

Table 3

| Properties                      | Amb6600091 | Amb17613565 | Amb3940754 | Amb21855906 |
|---------------------------------|------------|-------------|------------|-------------|
| Physico-chemical properties     |            |             |            |             |
| MW (g/mol)                      | 276.29     | 197.19      | 165.19     | 198.18      |
| Heavy atoms                     | 19         | 14          | 12         | 14          |
| Arom. heavy atoms               | 0          | 6           | 6          | 6           |
| Rotatable bonds                 | 11         | 3           | 3          | 3           |
| H-bond acceptors                | 8          | 5           | 3          | 5           |
| H-bond donors                   | 5          | 4           | 2          | 3           |
| Lipophilicity                   | Log P<sub>ow</sub> | 0.85 | 0.72 | 1.08 | 0.69 |
| Water solubility                | Log S (RSOL) | High | High | High | High |
| Pharmacokinetics                | GL absorption | Low | High | High | High |
| Drug likeness                   | Lipinski, violation | No | No | No | No |
| Medi. chemistry                 | Synth. accessibility | Very easy | Very easy | Very easy | Very easy |

respectively, where oral rat LD<sub>50</sub> indicate the number of chemical compounds (mg/kg body weight) that causes 50% of rats to die after oral ingestion were predicted using the T.E.S.T. software listed in Table 4. To estimate the value and average of the predicted toxicities from all the QSAR methods mentioned in the T.E.S.T. tools called consensus method was applied in this study.

3.11. Molecular dynamics simulation

Molecular dynamics (MD) simulation was performed to understand the dynamic behavior of our compounds within the protein. The MD simulation also determined the effect of explicit solvent molecules on the ACE2 protein and their fluctuations and conformational changes to obtain time-averaged features of the complex system in different time scales [44]. In this study, the results of MD simulation were analyzed based on three major physical properties comprising root mean square deviation (RMSD), root mean square fluctuation (RMSF), and protein-ligands contact mapping of the compounds in a specific time.

3.12. RMSD analysis

The protein-ligand complex structure variation generated from molecular docking was predicted by RMSD values obtained from 0 to 250 ns simulation run. The success in molecular dynamics simulation can be evaluated based on RMSD value when the RMSD value within an arbitrary threshold rang 0.3 nm or 3 Å the true pose is considered as a hit [50]. The RMSD of the complex structure indicated the stability and deviation in the average distance of C<sub>α</sub>-atoms as a function of simulation time [75]. The average RMSD values of Amb3940754, Amb6600091, Amb17613565, and Amb21855906 were found 2.35 Å, 2.75 Å, 2.5 Å, and 2.45 Å, respectively compared to control compound XX5 in complex with ACE2 protein (Fig. 9). The compound Amb6600091 showed maximum fluctuations from 200 ns to 235 ns and smoothly turned into a stable phase after the 240 ns run. The RMSD value obtained from the MD simulation showed minimal fluctuation throughout the 250 ns run and remaining stable until the end of the simulation, where higher fluctuations have observed only at the starting points during the simulation run of the complex structure.

3.13. RMSF analysis

The RMSF is a measure of the deviation between the position of residues contributing to protein structure and the binding site residues of the complex structure. The AA residues with low or no RMSF values are considered more stable because of their limited movement capabilities during the MD simulations [76]. The change of RMSF (ΔRMSF) value within an arbitrary threshold range of >3 Å is considered tangible and an important change in AA residue-specific flexibility [50]. The RMSF graph was calculated for ACE2 in complex with XX5 compound from 0 to 570 AA residue of C<sub>α</sub>-atoms responsible for 50% of fathead minnow, D. magna and T. pyriformis to die after 96, 48 and 48 h, respectively.
structure seen roughly in RMSF plots exhibited an advanced fluctuation level over the 250 ns timeframe. The RMSF graph demonstrated averaged low and significant values of the ACE2-Amb3940754 complex (2-2.3 Å), ACE2-Amb6600091 complex (2-2.1 Å), ACE2-Amb17613565 complex (2-2.2 Å), and ACE2-Amb21855906 complex (1.9-2.1 Å) compared to the reference compound XX5, commencing the natural compounds were closely bound to ACE2 protein concerning their average positions shown in Fig. 10.

### 3.14. Protein-ligand contact mapping

Protein interactions with the selected four compounds Amb3940754, Amb6600091, Amb17613565, and Amb21855906 have been monitored throughout the simulation interaction diagram (SID) of the Schrödinger (Release 2020-3). The hydrogen bonds, hydrophobic, ionic, and water bridge interactions found during the interaction analysis have been shown in the stacked bar charts (Fig. 11). Different types of bonding play an important role in stable binding to the targeted protein, where hydrogen bonding helps to determine influence drug specificity, metabolism, and adsorption. The hydrogen bonding interaction found for all the four compounds was observable until the last AA

### Table 4

| Endpoint                  | Target | Amb6600091 | Amb17613565 | Amb3940754 | Amb21855906 |
|---------------------------|--------|------------|-------------|------------|-------------|
| Organ toxicity            | Hepatotoxicity | Inactive    | Inactive    | Inactive   | Inactive    |
| Toxicity Endpoints        | Carcinogenicity | Inactive   | Inactive    | Inactive   | Inactive    |
|                           | Immunotoxicity  | Inactive    | Inactive    | Inactive   | Inactive    |
|                           | Mutagenicity    | Inactive    | active      | Inactive   | Inactive    |
|                           | Cytotoxicity    | Inactive    | Inactive    | Inactive   | Inactive    |
| LD_{50} (mg/kg)            |        | 5500       | 1460        | 2400       | 2000        |
| Toxicity class            |        | 6          | 4           | 5          | 4           |
| 96-hour fathead minnow LC_{50} | mg/L | 103.64     | 57.05       | 141.40     | 758.01      |
| 48-hour D. magna LC_{50}   | mg/L   | 464.57     | 6.26        | 31.02      | 19.6        |
| 48-hour T. pyriformis IGC_{50} | mg/L | 389.25     | 410.07      | 572.19     | 212.98      |
| Oral rat LD_{50}           | mg/kg  | 1768.26    | 3205.31     | 1631.99    | 1747.49     |
| Bioaccumulation factor     | Log10  | -1.38      | -0.48       | 0.09       | -1.49       |

Fig. 9. Depicted the RMSD values extracted from the Cα atoms of the selected four compounds in complex with the ACE2 protein. Herein, showing the RMSD of reference ACE2-XXS (blue) protein complex with the compounds (A) Amb3940754 (orange), (B) Amb6600091 (gray), and (C) Amb17613565 (yellow), where (D) Amb21855906 (red) of the compounds. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

Fig. 10. Depicted the RMSF values extracted from the Cα atoms of the selected four complex structures. Herein, showing the RMSF of ACE2-XXS protein (blue) in complex with the compounds (A) Amb3940754 (orange), (B) Amb6600091 (gray), (C) Amb17613565 (yellow), and (D) Amb21855906 (red) of the compounds. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)
residue of the protein. Hydrogen bonds and their relative strength in aqueous ionic solutions at ambient conditions are necessary to initiate the protein-ligand binding interaction [51,77]. A hydrogen bond is a weak type of dipole-dipole traction between molecules forms when a strongly electronegative atom in H-bond acceptor exists in the vicinity of another electronegative atom with a lone pair of electrons known H-bond donor [78]. In addition, other bonding interactions like hydrophobic, ionic, and water bridges bond at the same residue position of the protein. In this study, intermolecular hydrogen bonding interactions and other bonding interactions like hydrophobic, ionic, and water bridges bonds of the protein-ligands complex were determined and depicted in Fig. 11. During the MD simulation, the ACE2-Amb17613565 complex, ACE2-Amb6600091 complex, ACE2-Amb3940754 complex, and ACE2-Amb21855906 complex provided the highest number of hydrogen-bonding interaction for all the four protein-ligand complexes until the last residue of the simulation run. The analysis of the number of H-bonds formed in the ACE2-Ligands complexes indicating the improved stability of ligands to the binding site of the protein.

3.15. MM/GBSA analysis

MM/GBSA methods have been used in this study to estimate the ligand-binding free energy to the desired protein. The MM/GBSA of the protein-ligand complex structure has been calculated from the few snapshots (~200) of the MD simulations trajectory. The analysis of the complex structure found higher net negative binding free energy values -38.47 kcal/mol, -33.75 kcal/mol, -32.54 kcal/mol, and -35.18 kcal/mol for the selected four compounds Amb3940754, Amb6600091, Amb17613565, and Amb21855906, respectively with the targeted protein (Fig. 12). Additionally, physico-chemical components analysis of the four compounds identified a significant contribution in the GBind Coulomb (Coulomb energy) and GBind vdW (Van der Waals interaction energy) shown in Fig. 12. Therefore, it can be considered that the selected compounds will be able to maintain a long-term interaction with the desired ACE2 protein and result in inhibition of the desired protein.

4. Discussion

SARS-CoV-2 is a recently originated pandemic virus, against which vaccines and new drugs are constantly being tested [75,76]. For thousands of years, natural products and their derivatives isolated from various sources have been demonstrated to be effective therapeutic agents, and thus play an important role in treating diverse infectious diseases [34,79]. The chemical structure and extensive biological activities of these compounds vary comprehensively, that’s why natural compounds incessantly offer inspiration to innovations in drug discovery and medical sciences [35]. Therefore, we aim to identify potential natural ACE2 inhibitors through computational approaches such as pharmacophore modeling, virtual screening, molecular docking, ADME, and MD simulation to overcome the present demonic situation originated through the SARS-CoV-2.

Initially, a validated SBPM was applied to the virtual screening of 11,295 natural compounds that retrieved 23 similar scaffolds as hits with a maximum fit value of 78.81. The filtered compounds, which contain all the chemical features attendant in the SBPM, were retrieved for molecular docking simulation to avoid false-positive hits generated from the structure-based pharmacophore screening. Molecular docking simulation was performed to observe the complex structure of the small natural compounds with ACE2, calculate the binding energy of the complex interaction, and finding the best geometrical arrangements. The best four compounds with binding affinity range between ~7.5 and ~7.0 kcal/mol have been chosen for further evaluation.

Hereafter these four compounds have been submitted for in silico ADME, where ADME properties like lipophilicity, water-solubility, drug-like effectiveness, pharmacokinetic and physicochemical properties were evaluated and found optimum. After that, the in-silico toxicity
properties like acute toxicity, hepatotoxicity, cytotoxicity, carcinogenicity, mutagenicity, immunotoxicity along with LD50 score were also evaluated. Based on evaluation we found that all four compounds were nontoxic to the host. The MD simulation was performed on the selected four compounds and demonstrated good stability and affinity to the protein binding site.

Then we performed a molecular dynamics simulations study for investigating the stability of these four compounds with ACE2 protein. Because if the ligands do not form stable interaction with protein, then the inhibition of protein may hinder. By using the Desmond module of Schrödinger, we run the MD simulation for 250 ns for the selected four natural compounds. Here we observed the RMSD, RMSF, and protein-ligand contact of the complex system. The RMSD, RMSF, and protein-ligand contact study found for all the selected compounds showed enhance stability and optimized fluctuations with the ACE2 proteins.

5. Conclusions

Structure-based drug design is becoming an essential, efficient, and exterior approach to identify inhibitory compounds against a specific target protein. In this study, we describe the quick and successful identification of novel natural ACE2 inhibitors by a computer-aided drug design approach. The CADD approaches include pharmacophore modeling, virtual screening, molecular docking, ADME, and MD simulation, which identified four natural compounds Amb17613565, Amb6640091, Amb3940754, and Amb21855906 can potentially inhibit the activity of ACE2 and resulting blocking the entry of SARS-CoV-2 into the human host cell.

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FA, AS, MSR designed the project; TAB, AS, FA, MMIT, and ZN performed the experiments; FA, SP, TAB, and AS evaluated and interpreted the data; SP, TAB, FA, SMN, MA-AL-M, RA, RRI, MEK, MSH, MMIT, IQ, TMK, and MSR prepared the draft manuscript; FA, AS, ZN, and MSH finalized the manuscript. All authors approved the final version of the manuscript.

Declaration of competing interest

The authors declare no conflict of interest.

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Appendix A. Supplementary data

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