


Review

# In Silico Studies of Drug Discovery and Design Against COVID-19 Focusing on ACE2 and Spike Protein Virus Receptors: A Systematic Review

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**Abstract:** The emergence of COVID-19 has prompted researchers worldwide to focus on developing drugs that specifically target ACE2 receptors and SARS-CoV-2 Spike Protein receptors. They have embraced an in-silico approach that employs virtual screening, molecular docking, and molecular dynamics to achieve this. This innovative method harnesses existing chemical and natural product databases to identify the most suitable ACE2 receptor blockers and SARS-CoV-2 Spike Protein inhibitors. By following the PRISMA statement guidelines, a thorough literature search yielded 21 relevant articles, forming the basis of this systematic review. The review provides a comprehensive summary and detailed description of the methodologies, protocols, software tools, and noteworthy drug candidates identified in these studies. Additionally, it sheds light on the crucial molecular interactions by presenting an overview of the interacting residues elucidated in the reviewed articles, offering valuable insights for effective therapeutic interventions. Furthermore, the review presents thought-provoking suggestions for future research directions, aiming to inspire and guide advancements in drug development efforts.

## 1. Introduction

Coronaviruses are enveloped RNA viruses that are widely distributed among humans and other animals and can cause respiratory, enteric, and hepatic diseases in most cases (1). Severe Acute Respiratory Syndrome – Corona Virus (SARS-CoV) was the primary cause of severe acute respiratory syndrome outbreaks in Guangdong Province, China, in 2002 and 2003. In 2004, several reports from laboratory and nonlaboratory cases highlighted the possibility of SARS re-emergence (2). In late December 2019, there was a report about emerging pathogens linked to coronaviruses. Local health facilities in Wuhan reported several clusters of patients with pneumonia of unknown origin linked to a seafood and wet animal wholesale market in Hubei Province, China (3). Previous research in viral diagnostics, isolation and identification has revealed that the novel virus shares more than 85% identity with a bat SARS-like CoV genome. Further analysis of the 2019-nCoV genome from clinical specimens reveals 86.9% nucleotide sequence identity to a previously published bat SARS-like CoV. Although it is similar to some beta coronaviruses found in bats, it is not the same as SARS-

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CoV (4). This study served as the foundation for many subsequent studies on novel coronavirus drug discovery and vaccine development.

Another critical research is on the matter of viral receptors in the human body. SARS-CoV-2 uses human Angiotensin-converting Enzyme 2 (ACE2) as an entry receptor with similar affinity to previous SARS-CoV isolates, according to Walls et al. (5), indicating that the virus could spread effectively in humans. Another study by Zhou et al. (6) found that SARS-CoV-2 can use all ACE2 proteins, except mouse ACE2, as an entry point to enter cells that express ACE2, but not those that do not, indicating that ACE2 is the cell receptor through which the virus can enter the cells. They also show that SARS-CoV-2 does not use previous SARS-CoV receptors such as aminopeptidase N (APN) and dipeptidyl peptidase 4 (DPP4).

Drug discovery and development is a complicated process that needs extensive research and also has high-risk factors of failure (7). Conventional drug discovery methods could take more than a decade to complete, and the cost of research could be prohibitively expensive, which could reach around \$70 million for each NME reaching the clinic (8). The failure that could occur in the process namely lack of clinical efficacy, unmanageable toxicity, poor drug-like properties and lack of commercial needs (9). Computational methods such as virtual screening and molecular docking could assist in the early stages of discovery (10). Modern computational-aided drug design is divided into two parts based on the molecular information source utilized, which are structure-based drug design and ligand-based drug design (11). Compounds from the existing database or new chemical entities could be screened more quickly to find the hit and lead compound, which could then be developed further to become the final drug (12).

In this article, the authors conducted a systematic review of drug discovery research and methods by using computational methods to find drug candidates from drug repurposing or a database of chemical entities. The authors systematically selected documentation on this subject and compared different computational analysis methods towards the ACE2 receptor or Spike Protein SARS-CoV-2 receptor in the current study.

## 2. Materials and Methods

### 2.1 Study Protocols

The Preferred Reporting Items for Systematic Reviews and Meta-Analysis (PRISMA) statement is used as a guideline to conduct the literature search (13). For the search strategy, the following descriptors were employed: "ACE2", "SARS-CoV-2", "Spike Protein", "*in silico*", "computational", "virtual screening", and "molecular docking" using the Boolean operator. The search was conducted in ScienceDirect. The potential studies were screened according to the inclusion and exclusion criteria.

### 2.2 Inclusion and Exclusion Criteria

The inclusion criteria for this article review were: (1) experimental studies written in English, (2) studies that use virtual screening or molecular docking as their drug discovery method, (3) databased used in the studies are existing drug databases or databases of natural product with non-peptide small molecule and (4) studies are published in the range year 2020 - 2023.

The exclusion criteria were: (1) studies written in other languages besides English, (2) non-related studies such as: (a) *in vitro* studies using cell line, (b) synthesis method, (c) computational using artificial intelligence, (d) network analysis method, (e) screening or docking using modified structure, (3) duplicate publications, (4) articles not available as full text.

## 2.3 Selection of Studies and Data Collection

All search results were collected and screened. All identified studies are assessed based on the title and abstract without using any specific data extraction form. The data are gathered according to the year of publication, computational method, and database used in the study.

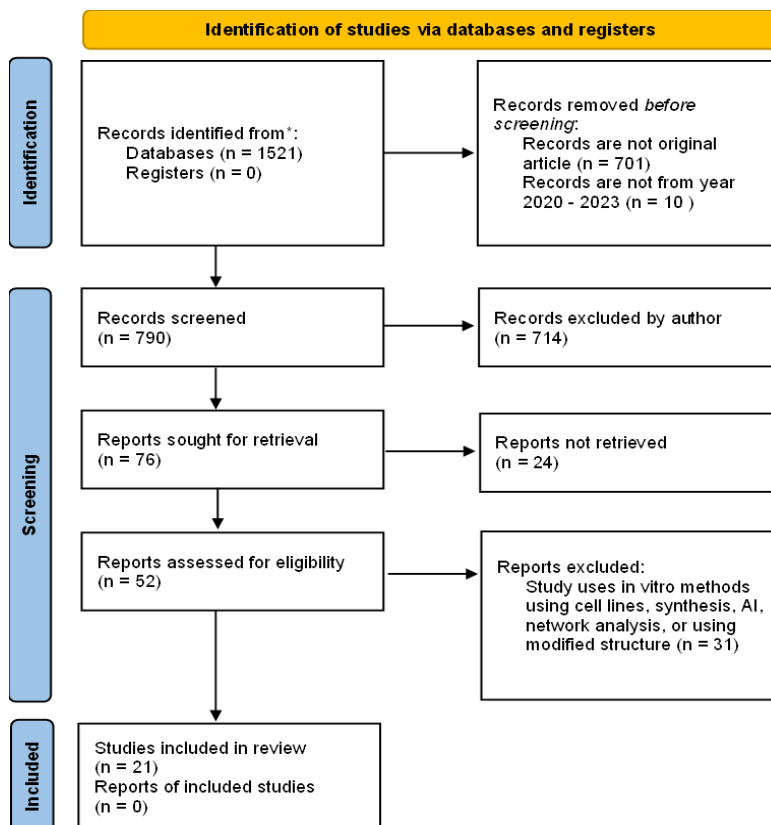
## 2.4 Quality of Article Assessment Study

The idea of bias in computational drug research studies is not well established, therefore it is hard to determine the risk of bias in this type of study. The authors decided to go with a tool designed for the assessment in a similar *in silico* review published by Mohamed et al. (14) with some refinement. The assessment encompassed five main aspects of the quality of studies included in the present systematic review: design (single-target or bi-target or multi-target), target template and crystal resolution, docking tools, molecular dynamics simulation (yes or no) and the resource of the database. The quality of each eligible article was appraised by the author.

## 3. Results and Discussion

### 3.1 Literature Collection Process

The keywords used in the literature search yielded 790 research articles published between 2020 and 2023. Only 76 of those articles met the criteria for related studies. We retrieved 52 potential literature articles from the 76 articles and excluded 24 articles because they were written in a language other than English, the method used was unclear, or it was a review article that was missed in the first screening. We evaluated the retrieved articles for eligibility and decided to exclude 31 records that used a modified structure for docking, an *in vitro* method, a synthesis method, AI simulation, and network analysis. As a result, 21 articles were identified as meeting the inclusion criteria and were given full consideration. The final 21 articles are processed to the Quality of Article Assessment Study.



**Figure 1.** PRISMA flowchart for the literature search process.

### 3.2 Quality of Article Assessment Study and Summary of Studies

We have summarized the quality of articles in our studies in Table 1. Bias in computational drug research studies is not well established, as stated in the methods. But one potential bias is the target's molecular plasticity. The target's flexibility is usually limited or ignored when only molecular docking is used in a study. In this case, MD simulation could play a key role in drug discovery and design, as well as pre-and post-docking simulation. It can be used as a generator for multiple target conformations for virtual screening, or as a validator for post-docking to distinguish between improper docking poses and meaningful ones (15). 13 of our reviewed papers have been validating further by using various molecular dynamics programmes such as GROMACS (16), UNRES (17), NAMD (18), Schrodinger Desmond (19), and AMBER (20).

Table 1. Quality of articles included in systematic review.

No.	First Author, Year	Targets				Target Template		Crystal Resolution (Å)		Docking Tools	MD Simulation		Resource of structure Database
		Sgl	Sgl+	Dbl	Dbl+	Spike Protein	ACE2	Spike Protein	ACE2		Yes	No	
1	Gurung, 2022 (21)					7WBL		3.40		Autodock 4.2	Gromacs		PubChem
2	Kulkarni, 2020 (22)					6M0J		2.45		Autodock Vina			PubChem
3	Rameshkumar, 2021 (23)					6VW1		2.68		Autodock Vina			Var.
4	Natesh, 2021 (24)					6W41	1R42	3.08	2.20	Autodock Vina			PubChem
5	Al-shuhaib, 2022 (25)						1R42		2.20	GLIDE	UNRES		PubChem
6	Hadni, 2022 (26)					6M17		2.90		Autodock 4.2	NAMD		PubChem
7	Kar, 2022 (27)					6M0J		2.45		Autodock Vina	Gromacs		PubChem
8	Vardhan, 2020 (28)					2GHV	6M17	2.20	2.90	Autodock Vina	CABS-flex 2.0		Var.
9	Kiran, 2022 (29)					6VSB		3.46		Cresset Flare Docking			PubChem
10	Jain, 2021 (30)					6M0J		2.45		Autodock Vina	Gromacs		PubChem
11	Khan, 2021 (31)						6M0J		2.45	Autodock Vina	Gromacs		PubChem
12	Benitez-Cardoza, 2020 (32)						1R42		2.20	Autodock + MOE			EXPRESS-pick
13	Gowrishankar, 2021 (33)					6VYB	1R42	3.20	2.20	Autodock 4.2	Gromacs		PubChem
14	Muhseen, 2020 (34)					6LZG		2.50		Autodock Vina	Gromacs		NPACT + MPD3
15	Akinlalu, 2021 (35)						6LZG		2.20	Autodock Vina			DrugBank
16	Baby, 2021 (36)						1R4L		3.00	GLIDE	Schrodinger Desmond		DrugBank
17	Pokhrel, 2021 (37)						1R4L		3.00	Autodock 4.2	Schrodinger Desmond		Ambinter
18	Yu, 2022 (38)					7DF4	7DF4	3.80	3.80	Autodock Vina			TCM
19	Singh, 2022 (39)					6LZG	6M0J	2.50	2.45	Autodock 4.2	Gromacs		PubChem
20	Mishra, 2021(40)					6M0J	6M0J	2.45	2.45	GLIDE	Schrodinger Desmond		PubChem
21	Nabati, 2022 (41)					6M0J		2.45		Autodock Vina			ChemDiv

The majority of the reviewed articles use GROMACS and Schrodinger Desmond as their methods of validation and use various configurations for the simulation. 9 articles used a single target as their receptors, which is either ACE2 or the SARS-CoV-2 Spike Protein (SP), while 6 of the articles use a single target but with addition other than ACE2 or the SARS-CoV-2 SP. One article used exactly double targets which are ACE2 and SARS-CoV-2 SP, and the last 5 use both targets with the addition of another target. These other targets are proteins which also contribute to the infection or multiplication process of the virus such as Main protease (Mpro)/3-chymotrypsin-like protease (3CLpro), papain-like protease (PLpro), RNA-dependent RNA polymerase (RdRp), and other non-structural protein (Nsp) for example Nsp-3 and Nsp-9. Crystal resolution is also one of the crucial parameters in molecular docking and dynamics studies. In this review, all the crystal structures used in the studies range from 2 Å to 4 Å, considered adequate.

Resolution is defined as a metric for assessing the quality of data collected on the protein or nucleic acid crystal. It evaluates the level of detail in the diffraction pattern as well as the level of detail that will be visible when the electron density map is computed (42). Higher resolution values, such as 1Å, are highly ordered and allow you to see every atom in the electron density map, whereas lower resolution values, such as 3Å or higher, only show the basic contours of the protein chain (43,44).

The docking tools mainly used in the reviewed literature are Autodock4 (45) and Autodock Vina (46), though other tools are being used such as GLIDE (47), Cresset Flare (48), or MOE (49). The structure of the database mainly comes from PubChem (50), but other databases such as DrugBank (51), Ambinter (52), and ChemDiv (53) are also being used.

Table 2 provides a comprehensive summary of the included studies, highlighting the methods, protocols, software, ligand candidates, targets, grid sizes in virtual screening and molecular docking, ADMET/physicochemical analysis software, and key interacting residues. Moreover, the table also delineates the important residues implicated in the molecular interactions under investigation, providing valuable insights into the specific amino acids or functional groups involved in driving the observed effects. This information serves as a valuable resource for understanding the research parameters and findings. In subsequent subsections, we discuss each parameter in detail, offering insights into the methodologies employed and their implications. This comprehensive analysis aims to provide a deeper understanding of the reviewed studies, shedding light on the approaches utilized and their impact on the field of research.

### 3.3 Virtual Screening

There are millions of chemical 'libraries' that a trained chemist could hope to synthesise. Combinatorial chemists have already demonstrated in several prototype systems that libraries containing 1,000-100,000 compounds can be assembled (54). Virtual screening help chemist decides what compound should be synthesised (55). Virtual screening can be done by docking method or pharmacophore method.

Pharmacophores are the "refined" essence of what makes an effective ligand-receptor interaction, explicitly three-dimensional, and represent fundamental physicochemical aspects of ligand-receptor interactions, and are extremely useful when experimental structural data is unavailable and homology models are unreliable. In that case, a good pharmacophore model could give powerful insight and screen more effectively (56,57). Since the conformer is only compared against a three-point pharmacophore model, the method of virtual screening using pharmacophore could be very useful for a large number of compounds when compared to virtual screening using the docking method (54). This is due to the memory used for each conformation is not as large relative to when docking is required.

This method of virtual screening using pharmacophore could be seen in the work of Pokhrel et al. (37) which screened more than 11 thousand from the Ambinter database. The complicated steps in the pharmacophore model method are in the step of model validation. The literature used the GH scoring method for pharmacophore model validation and also includes enrichment factor and goodness of hit score. This validation needs to use a set of other databases called decoy compounds, which are usually available in the Database of Useful Decoys (DUDe) (58). This database has to be compared to active compounds in this matter they use known active ACE2 inhibitors from ChEMBL (59) and a literature search. After the model is validated, then it can be used as the parameter for the screening. This could be a problem when there is no available decoy database, or the number of active compounds is inadequate.

**Table 2.** Summary of studies included in the systematic review.

Ref.	Method, Protocol	Software	Ligand	Target	Grid Size (Å)	MD	ADMET	Candidate Drugs (ΔG) (kcal/mol)	Std. Ref.	Important Residue
(21)	PISA, MolDoc, PhysProp, MD	PDBsum, ADT 1.5.6, AD4.2, DW 4.6.1, GROMACS 2019.2, LigPlot+ v1.4.5	36 compounds with a preclinical or clinical trial against previous variants	<b>SP Omicron - hACE2 (7WBL)</b>	36 x 52.875 x 57.75	100ns 300K	DW 4.6.1	Abemaciclib (-10.08), Dasatinib (-10.06), Spiperone (-9.54)	-	SP : Phe338, Asp339, Asp364
(22)	Act.Site, MolDoc, DFT	PyRx, ADV	Major components of essential oils	<b>ACE2-RBD (6M0J)</b>	(adjusted according to active site residues that are selected)	-	-	Anethole (-5.2), Cinnamaldehyde (-5.0), Carvacrol (-5.2), Geraniol (-5.0), Cinnamyl acetate (-5.2), L-4-terpineol (-5.1), Thymol (-5.4), Pulegone (-5.4)	-	SP : Arg454, Ser459, Glu471, Tyr505
(23)	VS, MolDoc, PhysProp, DL	ADV, AD4.1, CoDockPP, SA	458 flavonoid compounds	<b>SP Omicron - hACE2 (6VW1)</b> , Mpro (6LU7), RdRp (6M71)	40 x 40 x 40	-	SA	Albireodelphin (-11.2), Amentoflavon (-10.2), Cupressuflavon (-10.0), Agathisflavone (-9.9)	-	SP: Thr319, Thr394, Phe396, Arg553, Lys621, Asn628, Asp760, Asp761
(24)	MolDoc, ADMET	ADT, ADV, ADMETLab, PT2, OSIRIS Property	Standard drugs and spices	<b>Spike Protein (6W41), ACE2 (1R42)</b> , Mpro (6LU7)	(Interacting critical residues in Spike Protein and ACE2 complex)	-	AL, PT2, OSIRIS	Bioactives in asafoetida and sesame seed	Remdesivir	ACE2 : His34, Glu37, Asp38, Arg393 SP : Arg403, Gln493, Ser494
(25)	MolDoc, ADMET, MD	GLIDE, SA, UNRES online server	3392 compounds from Iraqi medicinal plants	<b>ACE2 (1R42)</b>	30 x 30 x 30	250ns 300K	SA	Epicatechin (-6.05)	-	ACE2 : Asp30, Asn33, His34, Glu37
(26)	MolDoc, ADMET, MD	ADT, AD4.2, DSU.	Bioactive flavonoids compounds	<b>Spike Protein RBD (6M17)</b> , 3CLpro (6LU7)	60 x 60 x 60	100ns 310K	ADMET parameters	Herbacetin (-8.03), Morin (-8.46), Silibinin (-9.03), Tomentin E (-8.32), Amentoflavone (-10.19), Bilobetin (-8.89), Baicalein (-8.19), Quercetin (-8.26)	-	SP: Glu35, Asp38, Lys353, Glu406, Try453, Ser494, Gly496, Asn501, Tyr505
(27)	MolDoc, ADMET, MD, PCA	ADV, ADT, SA, pkCSM, LigPlot+, PLIP, GROMACS 2018.3	300 compounds from 25 Indian medicinal plants	<b>ACE2-RBD (6M0J)</b> , Mpro (6LU7)	50 x 50 x 50	100ns 300K	SA, pkCSM	Oleanderolide (-8.3), Proceragenin A (-8.3), Balsaminone A (-8.3)	-	SP : Cys336, Gly339, Asn343, Ala348, Arg355, Ser373, Asp428, Thr430, Phe515,

(28)	MolDoc, MD, ADMET, PLI	ADV, pkCSM, Mi, CABS-flex 2.0 online,	154 phytochemicals in Limonoids and Triterpenoids class	<b>SP (2GHV), ACE2 (6M17), 3CLpro (6LU7), Plpro (4MM3), RdRp (6M71)</b>	-	10ns	pkCSM Mi	SP : Maslinic acid (-9.3), Glycyrrhizic acid (-9.3), Corosolic acid (-9.4) ACE2 : Glycyrrhizic acid (-9.5), Maslinic acid (-8.5), Obacunone (-8.1)	-	ACE2 : Arg273, His345, Arg393
(29)	MolDoc, SAP, ADMET.	Cresset Flare Docking, LigPlus, pkCSM	37 phytoconstituents from K. Kudineer Chooranam and JACOM	<b>Spike Protein (6VSB)</b>	(based on trial and error)	-	pkCSM	Chrysoeriol (-11.39), Luteolin (-11.15), Quercetin (-11.47)	-	SP : Cys336, Phe338, Gly339, Phe342, Asn343, Thr345, Asp364, Val367,
(30)	BSP, MolDoc, MD	CASTp, Rampage, PyRx 0.8, GROMACS 2020, LigPlot+, VMD	10 dietary flavonoid compounds	<b>Spike Protein (6M0J)</b>	44.34 x 70.98 x 44.58	1ns 300 K	-	Naringin (-9.8)	Dexamethasone	*SP : Asp367, Thr371, Glu406, Ser409, Lys441
(31)	MolDoc, MD	UCSF Chimera, Autodock Vina, GROMACS 5.1	24 drug molecules	<b>ACE2 (6M0J)</b>	-	10ns 300 K	-	Cefpiramide (-9.1)	-	-
(32)	VS, MolDoc, FE	MOE, AD, PT2	500,000+ small molecules from Chembridge Corp.	<b>ACE2 (1R42)</b>	-	-	PT2	Chem7781334 (-5.87), Chem7676800 (-5.84), Chem7956590 (-5.83)	-	ACE2 : Gln24, Asp30, His34, Tyr41, Gln42, Met82, Lys353, Arg357
(33)	MolDoc, PK, MD	DSV, AD4.2, Mi, admetSAR, GROMACS 5.1,	57 phytochemicals from Indian Herbs	<b>ACE2 (1R42), Spike Protein (6VYB)</b>	ACE2 : 22.5 x 22.5 x 22.5 SP : 30 x 28.125 x 25.5	25ns 300 K	Mi, admet SAR	ACE2: Apigenin-7-O-glucuronide (-8.8), Ellagic acid (-8.4), Vasicolinone (-7.5), SP: Apigenin-7-O-glucuronide (-7.2), Ellagic acid (-6.2), Vasicolinone (-6.4)	-	ACE2 : Lys26, Gln89  SP: Tyr28, Tyr269, Asp290
(34)	MolDoc, ADMET, MD, PCA, FE	ADT ADV, SA, pkCSM, GROMACS 5.1	1000 plant bioactive terpenes compound	<b>Spike Protein (6LZG)</b>	45 x 45 x 45	50ns 300 K	SA, pkCSM	NPACT01552 (-11.0), NPACT01557 (-10.3), NPACT00631 (-9.5)	-	Tyr449, Tyr453, Glu484, Gly496, Gln498, Asn501
(35)	Act.Site, MolDoc, ADMET.	DSV, UCSF Chimera, ADT CASTp, PyRx, ADV, SA, AdmetSAR, GraphPad Prism	791 FDA-approved drugs	<b>ACE2 (6LZG), 3CLpro (6LU7), ADP ribose phosphatase of NSP3 (6VXS), NSP9 RNA binding protein (6W4B)</b>	(residues on active site included)	-	SA, Admet SAR	Ethinodiol diacetate (-15.6), Methylnaltrexone (-15.5), Ketazolam (-14.5), Naloxone (-13.6),	Lopinavir, Remdesivir, Hydroxychloroquine	Ile291

(36)	MolDoc, MD	Protein Prep Wizard, Schrodinger's HTVS, Glide SP, Schrodinger Desmond platform	2800 FDA-approved drugs	<b>ACE2 (1R4L)</b> TMPRSS2 (Hom. Mod.)	(Glide grid tool)	50ns 300 K	-	Valrubicin (-8.59), Lopinavir (-7.89), Fleroxacin (-7.73), Alvimopan (-8.51), Arbekacin (-7.74), Dequalinium (-7.73)	-	Arg273, Lys363, Asp367, Thr371,
(37)	VS Pharmacophore MolDoc, ADMET, MD, FE	LigandScout 4.3, DSV 16.1, PyRx, AD4, ADV, SA	11,295 compounds from the database	<b>ACE2 (1R4L)</b>	20.05 x 17.92 x 8.75	250ns 300 K	SA	Amb17613565 (-7.5), Amb6600091 (-7.1), Amb3940754 (-7.1), Amb21855906 (-7.0)	XX5	Asn149, Gly268, Asp269, Arg273, Asn277, Asp350, Lys363, Thr365, Asp367, Arg393
(38)	MolDoc	ADV, M2M	28 natural plants from TCM	<b>ACE2 (7DF4), Spike Protein (7DF4)</b>	(Interacting critical residues in Spike Protein and ACE2 complex)	-	-	ACE2: Oleanolic acid (-7.1), Tryptanthrin (-6.55), Chrysophanol (-6.16), Rhein (-4.69) SP: Oleanolic acid (-3.74), Tryptanthrin (-4.26), Chrysophanol (-4.07), Rhein (-3.66)	-	ACE2: His34, Lys94, Gln102, Lys562, Trp566 SP: Arg403, Tyr449, Tyr453, Gln493, Ser494, Tyr495, Gly496
(39)	ADMET, MolDoc, Qua.Calc, MD,	SA, admetSAR, UCSF Chimera, AD4.2, DSV Gaussian16 suite, GROMACS 2015,	586 phytochemicals from 47 medicinal plants	<b>ACE2 (6M0J), Spike Protein (6LZG),</b> Plpro (6W9C), Mpro (6LU7), Importin $\alpha$ -5 (2JDQ), Importin $\beta$ -1 (1F59)	33.75 x 33.75 x 33.75	50ns 300 K	SA, Admet SAR	ACE2: Hetisinone (-8.46) SP: 14-deoxy-11,12-didehydroandrographolide (-7.76)	Arbidol (SP), Hydroxychloroquine (ACE2)	ACE2: Ala348, Trp349, His378, His401 SP: Arg23, Ser182, Phe183, Leu185
(40)	MolDoc, FE, MD.	Schrodinger Drug Discovery Suite, Pymol 2.4.1	85 antiviral and antimicrobial flavonoid compounds	<b>ACE2 (6M0J), Spike Protein (6M0J),</b> Mpro (6LU7), NSP12 (7BV2), NSP15 (6WXC)	30 x 30 x 30	100ns 300 K	-	SP: Isosilybin (-5.19) ACE2: Legalon (-2.78)	-	SP: Tyr453, Gly496, Gln498, Tyr505 ACE2: Lys26, Glu37, Asn90, Gln96, Ala387, Arg393
(41)	VS, MolDoc, ADMET	Chimera 1.13, Molegro Virtual Docker v6.0, PyRx 0.8, DSV, ADV, ADT1.5.6, OpenBabel SA, ADMETlab 2.0	100,000+ compounds from ChemDiv	<b>Spike Protein (6M0J),</b> Mpro (7AMJ), RdRp (7B3D), Plpro (6WX4)	23.60 x 45.00 x 21.88	-	SwissADME, ADMETlab 2.0	8008-2051 (-9.0) K279-0710 (-8.8)	Nebivolol	Arg403, Tyr449, Tyr453, Ser494, Gly496, Gln498, Asn501,



### 3.4 Molecular Docking

The docking method is another option for virtual screening. It can also further be used to validate virtual screening results. Molecular docking studies are mainly used to predict the ligand-receptor complex's binding affinity, preferred binding pose, and interaction with the least amount of free energy. Docking studies also can reveal the interaction between protein-ligand, protein-nucleotide, and also protein-protein interactions (PPIs). Noncovalent interactions can include ionic bonds, hydrogen bonds, and van der Waals interactions (60). In addition to the software mentioned previously in the summary of studies, several other software options are widely used in many molecular docking studies. RosettaLigand (61), Surflex (62), and Ligandfit (63) are some of the other popular software.

The docking mechanism is a two-step mechanism. It started with sampling conformation of the ligand in the receptor then followed by ranking these conformations using a scoring function. The effectiveness of a docking programme is determined by two major factors: search algorithms and scoring functions (64).

There are 2 main algorithms in molecular docking, which is the stochastic algorithm when where the search is carried out by modifying the ligand conformation or population of ligands. Example algorithms for this method are Monte Carlo and Genetic Algorithms (65). On the other hand, systematic search methods promote minor modifications in structural parameters, which gradually change the conformation of the ligands. The algorithm examines the energy landscape of the conformational space and, after many search and evaluation cycles, converges on the lowest energy solution corresponding to the most likely binding mode. The systematic algorithms are presented in GLIDE and DOCK (66).

Scoring functions are used to predict the target-ligand complex's binding free energy, which is a measure of the small molecule's binding potency for the biomolecular target. Scoring functions are classified into three types: force-field-based scoring, empirical-based scoring, and knowledge-based scoring (67). AutoDock scoring is an example of force-field-based scoring, which is derived from the classic force field and evaluates the binding energy as a sum of nonbonded interactions. Empirical-based scoring, such as GlideScore, is a weighted sum of various types of receptor-ligand interactions. DrugScore is a knowledge-based scoring system that penalises repulsive interactions while favouring preferred contact between each of the atoms in the protein and ligand within a given cut-off (64).

### 3.5 Grid Size and Parameter

In our reviewed studies, we examined several variations in the grid size parameter. Several studies reported the grid size qualitatively, stating that they used active site residues or critical interacting residues as grid size (22,24,35,38). Because of the lack of quantitative data, this approach of disclosures would, of course, reduce the reproducibility of the studies. In other studies, the grid size is determined differently for each receptor target (33). In others, it is the same size for all tested receptors (23,25–27,39–41). Understandably, a specific PDB will have a set of specific grid sizes that can be used in it, but the grid size should always be bigger than the ligand that is docked. According to Feinstein and Brylinski's research, the highest accuracy is obtained when the dimensions of the search space are 2.9 times larger than the radius of the gyration of a docking compound (68). They developed a procedure based on this discovery to customise the box size for individual query ligands to maximise docking accuracy. This finding essentially reduces the number of scoring failures caused by overly generous box sizes while also avoiding sampling failures caused by a too-narrow search space.

### 3.6 Database of Chemicals

Although the majority of the chemical structure is obtained from PubChem, the dataset used for each study is unique to each author. Several studies are being conducted to investigate whether already approved drugs can be repurposed from their original purpose to become ACE2 blockers and SARS-CoV-2 inhibitors (31,35,36). Another method for preparing the dataset for testing is to look for compounds found in traditional medicine or plant constituents as drug candidates (22,24,39,40,25,27–30,33,34,38). The final notable approach is to select a chemical suitable as a candidate from a large dataset ranging from 10,000 to 500,000 compounds (32,37,41). However, these differences in approach may provide beneficial insight from different perspectives, bringing us closer to real drug candidates ready for development.

### 3.7 Target Receptors

We may find a range of PDB IDs for ACE2 and SARS-CoV-2 Spike Protein receptors in this review study. The same PDB ID was used in several articles for the same receptors. We can see this in four publications (24,25,32,33) for ACE2, which used PDB 1R42 (69) as the receptors. Some other interesting point is that a PDB file can be used as a receptor target for ACE2 or SARS-CoV-2 because it contains both receptors in one file. We can see the PDB 6M0J (70) can be used as a SARS-CoV-2 Spike Protein target in 5 articles (22,27,30,40,41) and for ACE2 in 3 other articles (31,39,40). Both mentioned receptors are good in crystal resolution (2.20 and 2.45 respectively)

### 3.8 Interacting Residues

We summarised the common residues that appear more than once in the reviewed literature from the reported residues that interact with the ligands. Tyr453 and Gly496 appear in four different articles in the SARS-CoV-2 Spike Protein, while Arg403, Ser494, and Tyr505 appear in three distinct articles. Several residues appear in two articles: Cys336, Phe338, Gly339, Asn343, Asp364, Ser373, Tyr449, Gln493, and Asn501.

The binding residues in the ACE2 receptor that appear more than once in the reviewed article are as follows. His34 was mentioned in five different articles. Glu37 and Arg393 were mentioned in four different articles. Three different articles featured Asp30, Arg273, and Asp367. While Lys26, Asn33, Lys363, and Thr371 were found in two separate articles.

These residue similarities could help researchers find more specific binding sites, as well as serve as a reference for future drug discovery and development aimed at ACE2 and SARS-CoV-2.

### 3.9 Molecular Dynamics

Advances in software and hardware performances allowed researchers to adopt molecular dynamics to address drug discovery issues, especially protein–ligand stability (71). The dynamic nature of the receptor has been largely demonstrated and conformational changes have been related to ligand binding (71).

In brief, MD simulation begins with the selection and preliminary analysis of protein structure. Protein structures must be in 3D conformation and can be downloaded from the Protein Data Bank (PDB). If a 3D conformation is not available, the structure can be obtained using homology modelling. There must be no missing residues in the structure, and any that are missing must be added using the Modeller software. After the structure has been complexed with the best ligand, the input file for MD simulation must be prepared. A complex is formed in a system by adding water, after which the protein is minimised in a vacuum and a short MD is performed, restraining the protein. The system is then equilibrated and configured for the parameter being used. After all of the preparations have been completed, the long MD simulation can begin.

13 of the 21 articles reviewed are undergoing molecular dynamics testing to further validate the findings. Simulation times range from 1 ns to 250 ns, with an average of 50 ns to 100 ns. 11 of the 13 performed the simulation at 300 K, while only one performed the simulation at 310 K, and the other did not specify the simulation temperature.

#### 4. Summary

Following our review of these articles, we made several recommendations to improve research in this area. To begin, it is critical to select target receptors of high quality for *in silico* experiments. This quality can be attributed to the structure having a lower resolution (below 2.5 Å) and no missing residues. This minor adjustment can significantly improve the quality of the research.

Second, a protein-protein interaction could help to ensure that the docking site is in the correct position. This allows the interacting residues of both receptors to be determined and provides additional evidence that the experiment is being carried out in the correct location.

Third, it is encouraged to use multiple receptor conformations for virtual screening and molecular docking. This can be accomplished by processing the same PDB file through MD simulation and capturing the conformation for each several ns. This gives virtual screening more consideration for what is matched with the receptors and what is not, while also semi-accommodating the protein's flexibility, which is non-rigid in nature. Another approach is to use multiple PDB files for the same receptor. This could provide insights into the consistency of the virtual screening results, such as whether the experiment used a different structure for the same macromolecule and still produced similar results.

Fourth, as discussed in the grid size section, it is recommended to set the grid size to the relative size of the ligands, which is 2.9 times larger than the radius of gyration of the docked molecule, according to the reference. As a result, when using a really big molecule for virtual screening and docking, this needs to be considered, because if it is too small as well, the search space becomes too narrow, making it ineffective.

Last but not least, molecular dynamics simulation should always be included in the research steps. This experiment could greatly aid the researcher's understanding of the ligand-receptor interaction, specifically how they interact with each other over time, which could be deduced as the stability of the ligand docked in the receptors. For each complex, 50 ns is a good starting point for the simulation duration, but longer simulations are preferable because they can provide more insights into the interaction. Trajectories analysis for MD simulation is also useful for research, and it is recommended that it be processed until the free energy calculation step using the MMGBSA/MMPBSA method.

#### 5. Conclusions

We've shown several drug discovery research related to the ACE2 and SARS-CoV-2 Spike Protein by using the in-silico method. Aside from virtual screening, molecular docking, ADMET prediction, and molecular dynamic simulation, the research is carrying out several experiments. Several publications include active site determination, quantum chemical calculation, synthetic accessibility prediction, principal component analysis, and free energy calculation. These additional experiments could be extremely beneficial in ensuring the results of the lead compound or drug candidates. We've also highlighted the key interacting residues for each receptor reported in each article and summarised them in the order of how often they appeared in the reviewed articles. Finally, we've made several recommendations on how to make future research on this topic more elaborate and of higher quality, so that it can provide more precise results.

## List of Abbreviations

Act. Site	: Active Site Prediction	MOE	: Molecular Operating Environment
AD	: AutoDock	MolDoc	: Molecular Docking
ADMET	: ADMET Prediction	PCA	: Principal Component Analysis
ADT	: AutoDockTools	PhyProp	: Physical Properties
ADV	: AutoDockVina	PISA	: Protein Interface Stat. Analysis
AL	: ADMET Lab	PK	: Pharmacokinetics
BSP	: Binding Site Prediction	pkCSM	: pkCSM Server
DFT	: Density Functional Theory	PLI	: Protein-Ligand Interaction
DL	: Drug-likeness	PT2	: ProTox-II
DSV	: Discovery Studio Visualizer	Qua.Calc.	: Quantum Chemical Calculation
DW	: DataWarrior	SA	: SwissADME
FE	: Free Energy Calculation	SAP	: Syntetic Accesibility Prediction
MD	: Molecular Dynamics	SP	: Spike Protein
Mi	: Molinspiration Server	VS	: Virtual Screening

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## Conflict of Interest

The authors declare no conflict of interest.

## Authors contribution

Conceptualization	: Matheus Prayoga Claus; Arry Yanuar
Investigation	: Matheus Prayoga Claus
Supervision	: Matheus Prayoga Claus; Arry Yanuar
Administration	: Matheus Prayoga Claus
Writing and Editing	: Matheus Prayoga Claus; Masteria Yunovilsa Putra; Arry Yanuar

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