The recent eruption and ensuing pandemic of coronavirus disease 2019 (COVID-19) caused by the severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) has presented unprecedented global public health and economic challenges. The disease was first detected in Wuhan, China, but has been identified in 212 countries and territories with 3,566,210 confirmed infected cases and 248,285 deaths worldwide as of May, 4, 2020. The World Health Organization (WHO) has estimated that the mortality rate of COVID-19 is 3.4%. In recent history, there have been 2 outbreaks of other coronavirus diseases, the SARS-CoV epidemic of 2003 that led to 1000 deaths [1] and the Middle Eastern respiratory syndrome (MERS-CoV) outbreak of 2012, which claimed 862 lives [2]. The WHO declared the current worldwide COVID-19 outbreak an international public health emergency. At the time of writing, it continues to spread beyond control. The SARS-CoV-2 is highly homologous to SARS-CoV and the fatality rate of SARS-CoV was 10% [3, 4]. SARS-CoV-2 is a single-strand positive-sense RNA virus of the Coronavirinae subfamily, Coronaviridae.

Objectives: The outbreak of coronavirus disease 2019 (COVID-19) caused by the severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) remains a serious global threat. At the time of writing, there are no specific therapeutic agents or vaccines to combat this disease. This study was designed to identify the SARS-CoV-2 main protease inhibitors using drug molecule information retrieved from DrugBank 5.0 (Wishart et al.)

Methods: A set of common pharmacophores were generated from a series of 22 known SARS-CoV inhibitors. The best pharmacophore used for virtual screening (VS) of DrugBank using the Phase module followed by structure-based virtual screening (VS) using Glide (Release 2020-1; Schrödinger LLC, New York, NY, USA) with SARS-CoV-2 main protease and 50 ns molecular dynamics (MD) simulation studies.

Results: Six hits were selected based on the fitness score, extra-precision Glide score, and binding affinity with the main protease (Mpro). The predicted inhibitor constant (Ki) values of the 3 best hits, DB03777, DB08634, and DB07456, were 0.8176, 0.2148, and 0.1006 μM, respectively. An MD simulation of DB07456 and DB13592 with the Mpro demonstrated stable protein-ligand complexes.

Conclusion: The selected inhibitors displayed a similar type of binding interaction with co-ligands and remdesivir, and the predicted Ki values of 2 inhibitors were found to be superior to remdesivir. These selected hits may be used for further in vitro and in vivo studies against the SARS-CoV-2 Mpro.

Keywords: COVID-19, DrugBank, molecular docking, molecular dynamics, pharmacophore, SARS-CoV-2, virtual screening

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family, Nidovirales order. Extensive research of human coronaviruses has been performed by numerous researchers, who have identified 4 Coronavirinae genera (α, β, γ, and δ). The human coronaviruses HCoV-HKU1, HCoV–OC43, MERS-CoV, SARS-CoV, and SARS-CoV-2 are members of the betacoronavirus genus. A detailed classification of human coronavirus is illustrated in Figure S1. The new SARS-CoV-2 has a 79.7% and 91.02% genomic similarity with SARS-CoV and the Malayan pangolin coronavirus (SARS-CoV-2 like), respectively. It also has 96% and 89.6% identity with the envelope and nucleocapsid protein of SARS-CoV, respectively. Similarities between SARS-CoV and SARS-CoV-2, such as the replication mechanism, may be useful to inhibit the SARS-CoV-2 infection cycle.

Potential antiviral therapies can be grouped in 2 categories, namely improving the human immune system, in which interferon plays a vital function, and targeting virus replication through blocking the signal pathways. Potential alternatives to control or prevent COVID-19 infection include vaccines, interferon therapies, small molecule drugs, peptide or oligonucleotide-based therapies, and monoclonal antibodies, as shown in Figure S2. Of several options to prevent COVID-19 disease, one of the important is the use of small-molecule drugs. In the present exigent situation, given that there is no specific drug or vaccine yet approved for COVID-19, existing drug molecules are being analyzed using systematic bioinformatic tools as a potential means to block the replication cycle of SARS-CoV-2. Researchers have examined several existing drug molecules and natural compounds on the basis of structure-based virtual screening (VS) of various data-banks. In the present study, SARS-CoV-2 inhibitors were explored using resources from DrugBank (5.0; Wis-hart et al.) using a ligand-based and a structure-based VS virtual screening approach. For the ligand-based analysis, 22 SARS-CoV inhibitors were used to generate common pharmacophore hypotheses. The structures of the diverse SARS-CoV inhibitors are shown in Figure 1. The DrugBank database was used.

The structural organizations of the SARS-CoV genomes are displayed in Figure S3. The Mpro is a key protein for the proteolytic maturation of virus Nsps. The chymotrypsin-like protease of SARS-CoV-2 was selected as the target protein for the structure-based method VS. Various 3CLpro inhibitors of SARS-CoV-2 have been reported by several research groups to be potentially effective treatment options for COVID-19 based on a structure-based approach. The present work used the best 3CLpro hits from DrugBank for in vitro study.

**Methods**

**Collection of Materials**

The high resolution (2.16 Å) X-ray crystal structure of the SARS-CoV-2 Mpro or 3CLpro (PDB ID: 6LU7), consisting of 312 amino acid residues complexed with a peptide-like inhibitor (N3), was retrieved from the Research Collaboratory for Structural Bioinformatics Protein DataBank. The initial genomic sequences analyses of SARS-CoV-2 indicate that it has a high level of sequence similarity with SARS-CoV. In all, 22 known SARS-CoV inhibitors were collected based on the literature and used to build a common pharmacophore. The structures of the diverse SARS-CoV inhibitors are shown in Figure 1. The DrugBank database was used.

![Figure 1. The 22 structurally diverse severe acute respiratory syndrome coronavirus 2 inhibitors used to generate common pharmacophore hypotheses.](image-url)
for the ligand-based and structure-based VS.[22] The Mpro inhibitor remdesivir was also retrieved in the protein data bank (pdb) file format to compare with selected inhibitors. Windows 10 (Microsoft Corp., Redmond, WA, USA) and a 64-bit, Core 2 Duo CPU (Intel Corp., Santa Clara, CA, USA) microprocessor were used to perform the computations.

**Ligand Preparation**

All of the structures of the SARS-CoV inhibitors were drawn using ChemDraw Professional 15.1 (PerkinElmer Inc., Waltham, MA, USA) and saved in structure-data file (sdf) format. These inhibitors and DrugBank molecules were geometrically refined using the Schrödinger Ligprep module (Release 2020; Schrödinger LLC, New York, NY, USA).[23] During ligand preparation, the 2D structures were converted to the corresponding 3D structure, and added hydrogens and energy of ligands were minimized using the OPLS_2003 force field (Schrödinger LLC, New York, NY, USA) until it reached a root mean square deviation (RMSD) of 0.01 Å. Ligprep generates a single isomer with the original chiralities of each input. In order to perform molecular docking study using AutoDock 4.2 software (The Scripps Research Institute, La Jolla, CA, USA)[35] the selected molecules were converted to pdb format and then loaded in AutoDockTools 1.5.6 for conversion to pdbqt format (pdb with partial charges and AutoDock atom types).

**Protein Preparation, Grid Generation and Docking Validation**

The crystal structure of the Mpro (PDB ID: 6LU7) was used to prepare the protein using Protein Preparation Wizard (Schrödinger LLC, New York, NY, USA).[24] The protein structure was processed by assigning the bond orders, adding hydrogen, creating zero-order bonds to metals, filling side chains, and adding missing loops using the Prime protein structure prediction program (Schrödinger LLC, New York, NY, USA).[25-27] Finally, the protein was minimized using heavy atoms to RMSD 0.30 Å with OPLS3 force field. The receptor grid was prepared by selecting the coligand ligand of length 15Å from the center of the coligand. The coligand was split from its crystal structure and then docked with the prepared receptor grid. The best docked pose of the coligand was then superimposed on the original crystallographic bound conformation Figure S6 and the RMSD was calculated. The measured RMSD value was 2.0147Å. An RMSD value of <3Å is considered acceptable; therefore, the reproducibility of the docking protocol was good. In order to perform the molecular docking study using AutoDock 4.2, the protein was loaded in AutoDock Tools 1.5.6 and water molecules were removed, polar hydrogens bonded to heteroatoms were added, and the result was saved in pdbqt format.

**Generation of Common Pharmacophore Hypothesis**

The Phase[28-30] module (Release 2020-1; Schrödinger LLC, New York, NY, USA) was used to create common pharmacophores. Phase contains 6 in-built pharmacophore features: hydrogen bond acceptor (A), hydrogen bond donor (D), hydrophobic group (H), negatively charged group (N), positively charged group (P), and aromatic ring (R). The 22 prepared SARS-CoV inhibitors were imported and common pharmacophore hypotheses were created by selecting multiple active ligands and using a minimum (Table 1) of 4 pharmacophore features. The best pharmacophore hypotheses are shown in Figure 2.

**Ligand Based Virtual Screening**

The best pharmacophore, HHRR.1, was used to search DrugBank[22] with VS to retrieve SARS–CoV pharmacophore-matched inhibitors. The VS workflow used to identify SARS–CoV pharmacophore-matched inhibitors is presented in Figure 3. The database compounds matching at

| HypoID | Survival | Site | Vector | Volume | Select | Matches |
|--------|----------|------|--------|--------|--------|---------|
| HHRR_1 | 4.6674   | 0.5523 | 0.8589 | 0.5774 | 1.5326 | 15      |
| HHRR_2 | 4.6390   | 0.5503 | 0.8555 | 0.5665 | 1.5527 | 13      |
| HHRR_3 | 4.4058   | 0.4738 | 0.798  | 0.4202 | 1.5377 | 14      |
| HHRR_4 | 4.3177   | 0.5575 | 0.6887 | 0.4032 | 1.5221 | 14      |
| HHRR_5 | 4.2837   | 0.4788 | 0.7121 | 0.3945 | 1.5222 | 15      |
| HHRR_6 | 4.2764   | 0.5704 | 0.6129 | 0.4088 | 1.5381 | 14      |

**Figure 2.** The Phase-generated best pharmacophore hypothesis, HHRR.1, illustrating hydrophobic (H2, H4: green) and aromatic rings (R5, R6: orange). In all, 22 known severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) inhibitors were used for this model.
least 4 pharmacophoric features were identified and 1000 hits were kept in reserve for structure-based VS.

**Structure Based Virtual Screening**

The output pharmacophore-matched hits were utilized for VS with Glide.[31-33] The input hits were filtered with QikProp (Schrödinger LLC, New York, NY, USA)[34] and the Lipinski rule, and reactive functional groups were removed. The filtered hits were used as an input for high-throughput VS and 10% of the best output hits were subjected to standard precision (SP) docking. Finally, 10% of the best SP outputs were used for extra precision (XP) docking. The VS workflow of structure-based VS is shown in Figure 3.

**Molecular Dynamics Simulation (MD)**

Molecular dynamics (MD) simulations to predict the stability of the protein-ligand complex and interaction analysis were performed using Desmond (Schrödinger LLC, D.E. Shaw Research, NY, NY, USA).[35] The best 3 ligand- Mpro complexes were placed in the orthorhombic box with a buffer distance of 10 Å to add water and TIP3P was used to generate water models.[36] The temperature and pressure of the system was kept at 300.15 K and 1.01325 bar, respectively. The cut-off for the van der Waals radius and electrostatic interactions was 9.0 Å and the time step was 2.0 fs. The 50 ns root-mean-square fluctuation (RMSF) MD simulations for the 3 complexes were performed with the isothermal-isobaric (NPT) ensemble using OPLS3e force field. The RMSD, RMSF, and interaction diagram were generated using the Maestro 12 interface (Schrödinger LLC, New York, NY, USA). The methods of simulation were conducted as described by Yoshino et al.[37]

**Prediction of Ki Values of Selected Hits**

The receptor was again prepared using AutoDock Tools 1.5.6 for docking with AutoDock 4.2. During the preparation, the water molecules were removed, and polar hydrogens were added, followed by computation of the Gasteiger charge and adding the Kollman charge. The best hits identified on the basis of the fitness score and XP Glide score were used to predict Ki values using AutoDock 4.2.[38]

**Results and Discussion**

A set of 6 pharmacophore hypotheses; HHRR_1, HHRR_2, HHRR_3, HHRR_4, HHRR_5, and HHRR_6, were generated from 22 SARS-CoV inhibitors. The survival score of HHRR_1 was the highest, 4.6674, which matched 15 SARS-CoV known inhibitors nicely as shown in Figure S4. The finding that 22 SARS-CoV inhibitors, 15 fit with the best pharmacophore, indicates the quality of the model. DrugBank contains 10,156 compounds and the best 1000 pharmacophore-matched hits were subjected to structure-based VS. During VS, the hits were filtered using QikProp and the Lipinski rule, and reactive functional group compounds were removed. The best hits: DB06829, DB07456, DB13592, DB06834, DB11903, and DB03777, were selected based on a high XP Glide score. The 2D structures of these hits are shown in Figure 4. The range of fitness score of the selected hits was 1.816-1.459, and the range of the XP Glide score was -8.254 to -6.608. The fitness score, XP Glide score, and interacting active site amino acid residues and distance are shown in Table 2.

The fitness score of DB06829 was 1.753 and its XP Glide score was the highest, -8.254. This hit interacted with 2 key interacting amino acid residues, HIE-41 (2.10 Å) and CYS-145 (2.37 Å). The other interacting amino acid residues were HIS-163 (1.91 Å) and ARG-188 (2.05 Å). The 2D protein-ligand interaction (A), 3D protein-ligand interaction (B), pharmacophore match (C), and position in the active site of DB06829 are shown in Figure 5a.

![Figure 3](image-url) The virtual screening workflow to identify SARS-CoV-2 inhibitors. HTVS: High-throughput virtual screening; SP: Standard precision; VS: Virtual screening; XP: Extra precision.

![Figure 4](image-url) Selected severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) inhibitors with DrugBank ID, fitness with pharmacophore HHRR.1, and extra precision (XP) Glide score.
Table 2. Fitness score, XP Glide Score and different type of interactions

| Drug Bank ID | Fitness XP | GScore | Interacting active site amino acid residues with interacting distance (Å) |
|--------------|------------|--------|---------------------------------------------------------------------|
| DB06829      | 1.753      | -8.254 | HIE-41 (n-n), GLY-143 (2.68), SER-144 (2.71), CYS-145 (2.58), GLU-166 (1.75), ARG-188 (2.03) |
| DB07456      | 1.816      | -8.167 | HIE-41 (two n-n), LEU-141 (2.35), CYS-145 (2.46)                      |
| DB13592      | 1.707      | -7.037 | HIE-41 (2.36), SER-144 (2.23), CYS-145 (2.39)                        |
| DB06834      | 1.573      | -6.814 | HIE-41 (two n-n), HIS-163 (2.03)                                     |
| DB11903      | 1.459      | -6.748 | GLY-143 (2.19), GLN-189 (2.16)                                       |
| DB03777      | 1.778      | -6.608 | HIE-41 (three n-n), SER-144 (2.42)                                   |
| Remdesivir   | -          | -8.061 | HIE-41 (2.13), HIS-164 (1.98), GLU-166 (2.47), THR-190 (1.98)         |

Figure 5a. Docking poses of compound DB06829 (a) 2D ligand interaction diagram of hydrogen bond donor-hydrogen bond acceptor (purple line) and π-π stacking (green line); (b) 3D ligand interactions of hydrogen bond donor-hydrogen bond acceptor (purple dotted line) and π-π stacking with distance (light blue dotted line); (c) Pharmacophore-match structure; (d) Position of selected hits in the active site.

The fitness score of DB07456 was 1.816 and the XP Glide score was the next highest at -8.167. This hit also interacted with 2 key amino acid residues of the SARS-CoV-2 Mpro, HIE-41 (2, π-π) and CYS-145 (2.46Å). The other interacting amino acid residue was LEU-141 (2.35Å). The 2D protein-ligand interaction (A), 3D protein-ligand interaction (B), pharmacophore match (C), and position in the active site of DB07456 are shown in Figure 5b.

The fitness score of DB13592 was 1.707 and the XP Glide score was -7.037. This hit interacted with HIE-41 (2.36Å), SER-144 (2.23Å), and CYS-145 (2.39Å). The 2D protein-ligand interaction (A), 3D protein-ligand interaction (B), pharmacophore-match structure (C), and position in the active site of DB07456 are shown in Figure 5c.

The fitness score of compound DB06834 was 1.573 and its XP glide score was –6.814. This hit interacts with active site amino acid residues were HIE-41 (two, π-π), HIS-163 (2.03Å). The 2D protein-ligand interaction (A), 3D protein-ligand interaction (B), pharmacophore match (C), and position in the active site of DB06834 are shown in Figure 5c.

Figure 5b. Docking poses of compound DB07456 (a) 2D ligand interaction diagram of hydrogen bond donor-hydrogen bond acceptor (purple line) and π-π stacking (green line); (b) 3D ligand interactions of hydrogen bond donor-hydrogen bond acceptor (purple dotted line) and π-π stacking with distance (light blue dotted line); (c) Pharmacophore-match structure; (d) Position of selected hits in the active site.

Figure 5c. Docking poses of compound DB13592A (a) 2D ligand interaction diagram of hydrogen bond donor-hydrogen bond acceptor (purple line) and π-π stacking (green line); (b) 3D ligand interactions of hydrogen bond donor-hydrogen bond acceptor (purple dotted line) and π-π stacking with distance (light blue dotted line); (c) Pharmacophore-match structure; (d) Position of selected hits in the active site.
teraction (B), pharmacophore-match structure (C) and position in the active site of DB06834 are shown in Figure 5d.

The fitness score of DB11903 was 1.459 and the XP Glide score was -6.748. This hit interacted with GLY-143 (2.19 Å) and GLN-189 (2.16 Å). The 2D protein-ligand interactions (A), 3D protein-ligand interaction (B), pharmacophore-match structure (C), and positions in the active site of DB11903 are shown in Figure 5e.

The fitness score of DB03777 was 1.778 and the XP Glide score was -6.608. This hit interacted with active site amino acids HIE-41 (3, π-π) and SER-144 (2.42 Å). The 2D protein-ligand interaction (A), 3D protein-ligand interaction (B), pharmacophore-matched structure (C), and position in the active site of DB03777 are shown in Figure 5f.

The coronavirus has more than a dozen proteins for viral entry and replication, including a papain-like protease (PLpro), a 3-chymotrypsin-like protease (3CLpro), an RNA-dependent RNA polymerase (RdRp), and a spike protein. SARS-3CLpro is a cysteine protease crucial to viral replication in the infection cycle. Here, we targeted SARS-CoV-2 3CLpro (6LU7) to identify potential inhibitors. All 6 hits: DB06829, DB07456, DB07458, DB13592, DB06834, DB11903, and DB03777, had a common pharmacophore with SARS-CoV and good binding affinity with the active site of the Mpro. The active site of the 3CLpro has a Cys-His catalytic dyad (CYS-145 and HIS-41). There was a covalent interaction between CYS-145 and the coligand (N3) in the original crystal structure of 6LU7 (Fig. S5). The interactions of remdesivir, an important SARS-CoV-2 ongoing clinical trial inhibitor, were HIE-41, HIS-164, GLU-160, and THR-198.

Remdesivir also interacted with HIE-41, one of the important residues (Fig. S7). Therefore, HIE-41 and CYS-145 were crucial target amino acid residues for inhibition. Three selected hits, DB06829, DB07456, and DB13592, had close nonbonding interactions with CYS-145. The hits DB06829, DB07456, DB13592, DB06834, and DB03777 had close

Figure 5d. Docking poses of compound DB06834 (a) 2D ligand interaction diagram of hydrogen bond donor-hydrogen bond acceptor (purple line) and π-π stacking (green line); (b) 3D ligand interactions of hydrogen bond donor-hydrogen bond acceptor (purple dotted line) and π-π stacking with distance (light blue dotted line); (c) Pharmacophore-match structure; (d) Position of selected hits in the active site.

Figure 5e. Docking poses of compound DB11903 (a) 2D ligand interaction diagram of hydrogen bond donor-hydrogen bond acceptor (purple line) and π-π stacking (green line); (b) 3D ligand interactions of hydrogen bond donor-hydrogen bond acceptor (purple dotted line) and π-π stacking with distance (light blue dotted line); (c) Pharmacophore-match structure; (d) Position of selected hits in the active site.

Figure 5f. Docking poses of compound DB03777A (a) 2D ligand interaction diagram of hydrogen bond donor-hydrogen bond acceptor (purple line) and π-π stacking (green line); (b) 3D ligand interactions of hydrogen bond donor-hydrogen bond acceptor (purple dotted line) and π-π stacking with distance (light blue dotted line); (c) Pharmacophore-match structure; (d) Position of selected hits in the active site.
nonbonding interactions with HIS-41. The hits DB06829, DB07456, and DB13592 interacted with both CYS-145 and HIS-41. The hit DB06829 showed 1 π-π interaction and 3 closed, H-bond donor-acceptor interactions with an interacting distance of <3.0 Å. The hit DB07456 showed 2 π-π interactions, and 4 H-bond donor-acceptor interactions. Hit DB13592 demonstrated 3 H-bond donor-acceptor interactions from a close distance. The hits DB06834 and DB111903 revealed 2 H-bond donor-acceptor interactions at a distance of <3.0Å. All of the selected hits had a common pharmacophore with SARS-CoV, good binding affinity, and were well packed inside the active site pockets of the SARS-CoV-2 Mpro. The targets of all of the inhibitors are listed in Table S1.

The binding energy values of the selected hits B06829, DB07456, DB13592, DB06834, DB111903, DB03777, and remdesivir predicted by AutoDock were -8.22, -9.55, -6.04, -9.10, -7.89, -9.67, and -9.05, respectively. The predicted Ki values of B06829, DB07456, DB13592, DB06834, DB111903, DB03777, and remdesivir were 0.9400, 0.1006, 37.300, 0.2148, 1.6500, 0.0817, and 0.2334, respectively, as illustrated in Table 3. The binding energy and Ki values of DB07456, DB06834, and DB03777 were greater than those of remdesivir. DB07456 interacted with both CYS-145 and HIS-41; its XP Glide score was the second greatest, and its Ki value was 0.1006. Therefore, the hit DB07456 may be the best SARS-CoV-2 protease inhibitor. On the basis of fitness score, XP Glide score, predicted Ki values, interacting amino acid residues, and comparison with remdesivir, the 3 best hits, DB06829, DB07456, and DB13592, were selected for 50 ns MD simulation.

Overall information about the stability of the protein backbone after the formation of a protein-ligand complex can be analyzed according to the RMSD parameters. MD simulation of both apoprotein and the 3 best Mpro-ligand complex hits obtained from VS were monitored and analyzed for 50 ns to understand the dynamic behavior and stability of the protein-ligand complexes. The RMSD of the apoprotein and protein-ligand complexes of DB07456, DB06829, and DB13592 are shown in Figure 6. Important MD parameters, such as RMSD, RMSF, 2D-ligand-protein interaction diagram, and a timeline representation of interactions of the ligand with different active site amino acid residues, were calculated from the 50 ns MD trajectory. The protein-ligand complexes of DB07456, DB06829, and DB13592 exhibited an average RMSD value of 1.90 Å, 1.95 Å, and 1.68 Å, respectively. The RMSD of DB13592 was similar to the RMSD value of the apoprotein (1.68 Å). The average RMSD value of the ligand-Mpro complexes of compounds DB07456, DB06829, and DB13592 and the ligand-free protein are given in Table 4.

| Drug Bank ID | Binding Energy | Predicted Ki (μM) |
|--------------|----------------|------------------|
| DB06829      | -8.22          | 0.9378           |
| DB07456      | -9.55          | 0.1006           |
| DB13592      | -6.04          | 37.300           |
| DB06834      | -9.1           | 0.2148           |
| DB111903     | -7.89          | 1.6500           |
| DB03777      | -9.67          | 0.0817           |
| Remdesivir   | -9.05          | 0.2334           |

| Ligand No | Average RMSD (Å) | Average RMSF (Å) |
|-----------|------------------|------------------|
| Apo protein (6lu7) | 1.68          | 1.80             |
| DB07456   | 1.90            | 1.25             |
| DB06829   | 1.95            | 1.80             |
| DB13592   | 1.68            | 1.35             |

Figure 6. Root mean square deviation (RMSD) plot of apoprotein [main protease (Mpro), black], DB07456-Mpro complex (green), DB06829-Mpro complex (red), and DB13592-Mpro complex (blue) during 50 ns molecular dynamics simulation.
indicates the high stability of the protein-ligand complexes. Therefore, the RMSD of the Mpro-DB07456 and Mpro-B13592 systems might indicate that they probably did not undergo significant conformational changes during the MD simulation. The acceptable range for an RMSD value is <3.0 Å \cite{ref}, therefore, all 3 protein-ligand complexes were within the acceptable range.

The average RMSF value of the apoprotein was 1.80 Å and the average RMSF value of DB07456 and DB13592 was 1.25 Å and 1.35 Å, respectively, which was less than that of the apoprotein. The RMSF value of DB06829 was 1.80 Å, which was similar to that of the apoprotein. The average RMSF values of the protein-ligand complexes of DB07456, DB13592, DB06829, and the apoprotein are shown in Figure 7 and Table 4. As seen in Figure 7, it is clear that there were no significant fluctuations of amino acid residues after binding in the active site. The greater RMSF values suggest that the protein structure is more flexible. The marginal flexibility of the protein-ligand system resulted in lower RMSF values.

The active site amino acid residues of the Mpro contributing to binding interactions with DB07456 were HIS-41 (π-π), THR-26 (hydrogen bond acceptor), ASN-142 (hydrogen bond acceptor), and GLU-166 (water bridge H-bond) produced 54%, 91%, 48%, and 44%, respectively. The interaction of DB06829 with HIS-41 yielded 44% and DB13592 with THR-190 resulted in 34%. The DB07456-Mpro, DB06829-Mpro, and DB13592-Mpro interaction patterns observed during the 50 ns MD simulations are shown in Figure 58. The contacts formed in the DB07456-Mpro complex over the course of the simulation were THR-26, HIS-41, ASN-142, and GLU-166. The contacts made by DB06829 with apoprotein the Mpro residues were THR-26, HIS-41, ASN-142, GLY-143, GLU-166, ARG-188, and THR-190. The DB13592 hit contacts with Mpro residues were HIS-41, MET-49, GLU-166, and GLN-189. All of the contacts formed by DB07456, DB06829, and DB13592 are presented in the top and bottom panels in Figure 8, illustrating how the residues interacted with the ligand in each trajectory frame during the simulation time. Wang\cite{ref} proposed that 1 hot spot residue, HIS-41, in the Mpro is a conserved residue across many viruses, including SARS-CoV, SARS-CoV-2, MERS-CoV, and hepatitis C virus (HCV). The selected hits DB07456 and DB06829 also showed fair interactions with HIS-41. It was observed that DB07456 and DB13592 had stable binding affinity throughout the MD simulation.

**Conclusion**

In the emergency situation of a COVID-19 pandemic, repurposing existing drugs may be helpful in the effort to...
combat the disease. In-silico study resulted in 6 hits with fitness to a common pharmacophore of SARS-CoV inhibitors, well-positioned in the pockets, and good binding affinity with the Mpro of SARS-CoV-2. The DrugBank identifiers of these hits are DB06829, DB07456, DB07458, DB13592, DB06834, DB11903, and DB03777. The binding energy values of DB07456, DB06834, and DB03777 were greater than that of remdesivir, and their corresponding Ki values were also favorable compared with remdesivir. The inhibitors of DB07456 showed good binding affinity with the Mpro predicted by the software. This molecule also interacted with the crucial amino acid residues HIS-41 and CYS-145. The 50 ns MD simulation study of the 3 best hits revealed DB07456 and DB13592 as potential SARS-CoV-2 Mpro inhibitors. These are commercially available and therefore, these hits may be useful to the scientific community for further in vitro and in vivo study to combat SARS-CoV-2. The VS studies performed thus far by different groups were based on the structure-based method only, but in this study, we employed pharmacophore-based VS as well as structure-based VS, and a MD simulation study, which may eliminate false-positive results.

**Disclosures**

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**Conflict of Interest:** None declared.

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In silico identification of potential inhibitors of the main protease of severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) using a combined ligand-based and structure-based drug design approach.

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Figure S5. The original crystallographic-bound pose of carbon monoxide (CO) ligand (N3) of severe acute respiratory syndrome 2 (SARS-CoV-2) main protease protein (PDB ID: 6LU7). (a) 2D ligand interaction, (b) 3D interaction diagram. The CO ligand is light blue. The covalent bond with CYS-145 was broken before taking the image.

Figure S6. The best docked pose of the carbon monoxide (CO) ligand (light blue) superimposed on the original crystallographic-bound conformation (green).

Figure S7. The 3D ligand interaction diagram of remdesivir with severe acute respiratory syndrome 2 (SARS-CoV-2) main protease protein (PDB ID: 6LU7). Purple dotted line indicates H-bond donor-acceptor interactions and distance.

Figure S8. The contacts between DB07456, DB06829, and DB13592 and amino acid residues during molecular dynamics simulation.
| Compound ID | Target protein |
|-------------|----------------|
| DB06829     | Tyrosine-protein phosphatase which acts as a regulator of endoplasmic reticulum unfolded protein response. Mediates dephosphorylation of EIF2AK3/PERK; inactivating the protein kinase activity of EI. |
| DB07456     | Serine/threonine kinase which acts as a master kinase, phosphorylating and activating a subgroup of the AGC family of protein kinases. Its targets include: protein kinase B (PKB/AKT1, PKB/AKT2, PKB... |
| DB13592     | Cytochromes P450 are a group of heme-thiolatemonooxygenases. In liver microsomes, this enzyme is involved in an NADPH-dependent electron transport pathway. It oxidizes a variety of structurally un... |
| DB06834     | Non-receptor tyrosine kinase which mediates signal transduction downstream of a variety of transmembrane receptors including classical immunoreceptors like the B-cell receptor (BCR). Regulates seve... |
| DB11903     | This drug entry is a stub and has not been fully annotated. It is scheduled to be annotated soon. |
| DB03777     | Serine/threonine kinase which acts as a master kinase, phosphorylating and activating a subgroup of the AGC family of protein kinases. Its targets include: protein kinase B (PKB/AKT1, PKB/AKT2, PKB... |