Graph Convolutional Network-Based Screening Strategy for Rapid Identification of SARS-CoV-2 Cell-Entry Inhibitors

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ABSTRACT: The cell entry of SARS-CoV-2 has emerged as an attractive drug development target. We previously reported that the entry of SARS-CoV-2 depends on the cell surface heparan sulfate proteoglycan (HSPG) and the cortex actin, which can be targeted by therapeutic agents identified by conventional drug repurposing screens. However, this drug identification strategy requires laborious library screening, which is time consuming, and often limited number of compounds can be screened. As an alternative approach, we developed and trained a graph convolutional network (GCN)-based classification model using information extracted from experimentally identified HSPG and actin inhibitors. This method allowed us to virtually screen 170,000 compounds, resulting in ~2000 potential hits. A hit confirmation assay with the uptake of a fluorescently labeled HSPG cargo further shortlisted 256 active compounds. Among them, 16 compounds had modest to strong inhibitory activities against the entry of SARS-CoV-2 pseudotyped particles into Vero E6 cells. These results establish a GCN-based virtual screen workflow for rapid identification of new small molecule inhibitors against validated drug targets.

INTRODUCTION

Since the outbreak of the COVID-19 pandemic, global communities have suffered a significant loss of lives and economic growth. Although the development of COVID vaccines can significantly inhibit the spreading of SARS-CoV-2, the virus is constantly evolving into more infectious and transmissible variants (e.g., the delta strain), resulting in frequent breakthrough infections among vaccinated people.1−3 The constant increase of hospitalized patients in the USA and around the world despite the rollout of the vaccination programs has summoned the need to develop potent small molecule therapeutics for COVID patients.

The cellular entry of SARS-CoV-2 is one of the key steps in the viral life cycle that represents a hot target for small molecule inhibitors.5,7 To target SARS-CoV-2 viral entry, the most popular target is the viral spike protein given that drugs targeting spike are less likely to disrupt cellular processes in the host cells. On the host side, a variety of proteins are being evaluated as potential anti-SARS-CoV-2 targets, which include angiotensin-converting enzyme 2 (ACE2), the major receptor for spike on the cell surface,8−12 and transmembrane serine protease 2 (TMSSPR2), the protease involved in activating spike and other factors regulating the endocytosis of virion particles.13 In addition, a recent report also discussed that alternative entry points have been identified using neuropilin-1 (NRP1), which was found to significantly enhance the infectivity of SARS-CoV-2 by increasing viral entry into host cells rather than strengthening viral binding.14 Previously, we and others identified the cell surface heparan sulfate proteoglycans (HSPGs) as a critical cofactor that facilitate the entry of SARS-CoV-2 virions.8,9 We further showed that HSPGs, as negatively charged biopolymers, also facilitate the uptake of other positive charge-bearing endocytic cargos such as supercharged green fluorescence protein (GFP) and preformed α-synuclein pathogenic fibrils.15 HSPGs are a family of glycoproteins bearing one or more negatively charged polysaccharide chains consisting of repeated heparan sulfate disaccharide units. Most HSPG family members are anchored to the cell surface either as a single spanning membrane protein (e.g., syndecans) or glycosylphosphatidylinositol (GPI)-anchored protein (e.g., glypicans). Due to the enrichment of negatively charged sulfate groups, HSPGs can effectively serve as an attachment anchor to increase the surface dwell time for endocytic cargos bearing positive charges, facilitating their engagement with a downstream receptor.6,15,16 The internalization of HSPG cargos also requires the cortex actin, a specialized layer of proteins associated with the inner surface of the plasma membrane. A major component of the cortex is actin filament, which associates with myosin motor proteins and other actin-binding proteins. These proteins together maintain plasma membrane dynamics to promote the maturation of clathrin-coated pits.15 The specific mechanism of the cellular entry of SARS-CoV-2 is presented in Figure 1.

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We recently conducted a drug repurposing screen based on our previous study\(^6\) and identified eight drugs that inhibited HSPG-dependent entry of SARS-CoV-2 virions.\(^7\) Intriguingly, despite structural dissimilarity, several of the identified drugs display high potential of binding with heparin, a heparan sulfate analogue, suggesting that they may target the polysaccharide chain on the cell surface of HSPG to inhibit viral entry; however, alternative host protein targets may also exist.\(^8\) In addition to heparin-binding drugs, two structurally unrelated drugs, Sunitinib and BNTX, can both effectively disrupt the actin filaments underlying the plasma membrane (cortex actin) to inhibit HSPG-mediated endocytosis.\(^9\)\(^{12}\) Compared to drugs that target viral proteins, drugs targeting host factors essential for viral entry and replication are less likely to generate drug resistance as a result of viral mutations. While a drug repurposing screen is an effective strategy to rapidly adopt existing drugs for new therapeutic uses, the original target(s) of the approved drugs often reduces their therapeutic specificity, which may cause undesired side effects for treating viral infection. For example, as a heparan sulfate-binding compound, mitoxantrone delivers the most potent antiviral activity in vitro. However, because mitoxantrone was originally approved as anticancer chemotherapy via targeting DNA topoisomerases,\(^17\) cytotoxicity associated with DNA replication inhibition is an obvious concern.

To identify additional inhibitors targeting HSPG-mediated viral entry, we developed a graph convolutional network (GCN)-based classification approach. GCN can efficiently translate 3D structures into molecular graphs composed of nodes and edges and then utilize these graphs to extract spatial information to achieve accurate molecular classification and properties predictions.\(^18\)\(^{21}\) Compared to other traditional computational methods based on molecular dynamics (MD) simulations or density functional theory (DFT), the computational cost of GCN is substantially lower. These features allowed us to rapidly screen 170,000 compounds in several NCATS libraries. From these libraries, we identified and confirmed a set of compounds (256) as inhibitors of HSPG-dependent endocytosis with the most potent IC\(_{50}\) value of 0.95 \(\mu\)M. Further testing with a SARS-CoV-2 pseudotyped particle (PP) entry assay confirmed 16 compounds as entry inhibitors.

**METHODS**

**Computational Details.** GCN Model. GCN-based approaches display considerable robustness for structural elucidations,\(^18\)\(^{19}\)\(^{21}\) because it could fully utilize the molecular graphs for information extraction with substantially reduced computational cost, compared to MD- or DFT-based methods.\(^22\)\(^{20}\) In addition, such an architecture can directly work on graph-based inputs, instead of depending on collected descriptors; thus, it is more suitable for spatial information processing.\(^30\) At the same time, it is still flexible enough to include different chemical knowledge as extra descriptors for specific assignments.\(^22\)\(^{31}\)\(^{40}\) In this study, we employed the self-developed GCN package for activity classifications, and the SchNet architecture was applied.\(^41\) The workflow of the applied GCN is described in Figure 2. For any given drug molecule, its structural information was contained in the simplified molecular-input line-entry system (SMILES) string, and GCN can transform the molecular graph into a set of numerical descriptors for computational processing.

All the collected SMILES strings of drug molecules were first translated into molecular graphs through the TencentAlchemiData set within the Deep Graph Library (DGL) library.\(^42\)\(^43\) Each drug molecule is composed of edges and nodes within 3D space. Within the framework of GCN, the generated nodes represent atomic points within a molecule, while the edges are corresponding to interatomic connections. With these numerically encoded features, structure similarity can be well summarized, and related molecular properties can be mapped correspondingly. In fact, within any molecular or fragmentary graphs, all the connections between every two atoms are fully solved for information extraction; the specific values were recorded in distance tensors at the radial basis function (RBF) layer, guaranteeing there is no omission of important structure information. In addition, within the GCN model, to decently solve molecular graphs at an atomic level, multiple continuous-filter convolutions (cfcconv) layers were employed to optimize and record the interatomic evolution. For instance, at the \(k+1\) layer, the \(i\)th atom’s evolution can be expressed with the following equation

\[
a_i^{k+1} = \sum_{j=0}^{N} a_j^k \omega^k(d_{ij})
\]

in which “o” represents element-wise multiplication, and \(\omega^k\) is the filter generation that can map the atoms’ descriptions to the filter bank. To efficiently control the evolution accuracy via the applied filter values, a Gaussian-type function, \(\text{gauss}_\text{inv}\) was employed, which can be expressed with the following equation
where $\mu_m$ is the preset value of cutoff, and $l_{ij}$ represents the bonding distance among the $i$th atom and $j$th atom. The $\alpha$ is attributed to hyper parameters, and it was set to 0.1 in this study.\(^1\)

For any prediction or classification task, the computed value, $\text{Pro}$, by the GCN model is calibrated with respect to experimental measurement, $\text{Pro}'$, and the accuracy can be well indicated by the squared loss function

$$L(\text{Pro}, \text{Pro}') = (\text{Pro} - \text{Pro}')^2$$

In this study, we applied the developed GCN package for drug activity classification; however, it is worth noting that this promising architecture is also able to include various kinds of chemical and physical knowledge for more challenging structural assignments.

Data Set. We applied the above-described GCN model to a previously reported COVID-19 related drug screening, which identified drugs that block HSPG-dependent entry of $\alpha$-synuclein fibrils. A classification algorithm was based on NCATS’ collected activity values. The model was first trained by the collected data, which consisted of 3832 compounds.\(^6\) Among them, 367 compounds show activities, and 3465 are inactive. These compounds were randomly divided with a ratio of 9:1; 90% was used as the training set and the remaining 10% as the test set. The trained GCN model was validated by the compounds in the test set, which scored an accuracy of 99.5%. The trained model was then used to screen more than 170,000 compounds contained in three independent libraries, Genesis, Sytravon, and NCATS Pharmacologically Active Chemical Toolbox (NPACT), none of which had been experimentally screened by endocytosis or SARS-CoV-2 PP entry assays. NCATS has assembled the Genesis collection with 100,000 compounds to provide a novel modern chemical library that emphasizes high-quality chemical starting points, sp3-enriched.

Figure 3. (a–c) Distribution of the chemical properties, molecular weight (MW), SlogP, and logS for Genesis library. (d–f) Distribution of the chemical properties, molecular weight, SlogP, and logS for Sytravon library. (g–i) Distribution of the chemical properties, molecular weight, SlogP, and logS for NPACT library. (k) PCA analysis for the three independent libraries: Genesis, Sytravon, and NPACT.
chemotypes, and core scaffolds that enable rapid purchase and derivatization via medicinal chemistry. The Sytravon library is a retired Pharma screening collection that contains 44,000 diverse and novel small molecules, with an emphasis on medicinal chemistry-tractable scaffolds. The NPACT is a library of about 5,000 annotated compounds that inform on novel phenotypes, biological pathways, and cellular processes. There are more than 7,000 mechanisms and phenotypes identified in the literature and worldwide patents that cover biological interactions within mammalian, microbial, plant, and other model systems. The physicochemical properties and the chemical diversity coverage of these three libraries are shown in Figure 3.

α-Synuclein Fibrils Uptake Assay and Drug Verification. Fluorescence-labeled α-synuclein fibrils were generated as previously described.15 HEK293T cells were dispensed into black, clear-bottomed 1536-well microplates (Greiner BioOne) at 5000 cells/well in 5 L media with 200 nM pHrodo red-labeled α-syn fibrils and incubated at 37 °C, 5% CO2 and 85% humidity overnight (∼16 h). Compounds picked from the virtual screen were titrated 1:3 with 11 points in DMSO and dispensed into the assay plate at 23 nL/well via pintool. Cells were incubated with compounds for 1 h at 37 °C with 5% CO2 before 15 μL/well of PPs were added. The plates were then spinoculated by centrifugation at 1500 rpm (453g) for 45 min and incubated for 48 h at 37 °C, 5% CO2 to allow cell entry of PPs and the expression of luciferase. After the incubation, the supernatant was removed with gentle centrifugation using a Blue Washer (BlueCat Bio). Then 20 μL/well of Bright-Glo luciferase detection reagent (Promega) was added to assay plates and incubated for 5 min at room temperature. The luminescence signal was measured using a PHERAStar plate reader (BMG Labtech). Data were normalized with wells containing PPs as 100% and wells containing control DEnv PP as 0%.

ATP Content Cytotoxicity Assay. HEK293T-ACE2-GFP cells were seeded in white, solid-bottomed 384-well microplates (Greiner BioOne) at 6000 cells/well in 15 μL medium and incubated at 37 °C with 5% CO2 overnight (∼16 h). Compounds were titrated 1:3 in DMSO and dispensed into the assay plate at 23 nL/well via pintool. Cells were incubated with compounds for 1 h at 37 °C with 5% CO2 before 15 μL/well of media was added. The plates were then incubated at 37 °C for 48 h at 37 °C, 5% CO2. After incubation, 30 μL/well of ATPLite (PerkinElmer) was added to assay plates and incubated for 15 min at room temperature. The luminescence signal was measured using a Viewlux plate reader (PerkinElmer). Data were normalized with wells containing cells as 100% and wells containing media only as 0%.

RESULTS AND DISCUSSION

Overall performance of GCN Model. Unlike traditional computational drug discovery methods such as structural homology-based drug search, the GCN classification model utilizes molecular graphs to extract spatial information. The modeling process computes in a bonding environment at an atomic or interatomic level within a fully connected graph as opposed to utilizing simple descriptors. As a result, the structural features of drug molecules can be well captured and built from low-level logic,36,41 making no omission of important possibilities. This method results in a robust performance with the classification accuracy as high as 99.5% for the training set (workflow is described in Figure 4).

We also did a benchmarking analysis by comparing the classification accuracy of GCN with several popular machine learning algorithms. From the calculated AUC values, GCN performs comparably better than other methods (Table 1), especially for the case in which the ratio of active/inactive
compounds is small; in another aspect, from the confusion matrix of validation set, it is notable that GCN is advantageous to exclude inactive compounds with a high accuracy (more details can be found in Table S1, Supporting Information). At the same time, based on a Y-randomization test using a random counter training set (the activity was randomly shuffled in the training data set), we noticed that the AUC value of GCN is largely reduced, indicating its dependence on highly solved structure similarity for active compounds detection. Additionally, the identified new compounds from our in-house libraries also show some structural novelty with respect to the active compounds in our training set, further highlighting its high applicability in real practice of decent drug screenings, compared to other structural assignment-based approaches.

**Identification of Inhibitors for HSPG-Mediated Endocytosis.** We used the GCN-based model to screen 170,000 compounds. Approximately 2000 compounds were shortlisted by the virtual screening, which generated a small library that could be rapidly processed by a conventional quantitative high-throughput screen (qHTS) (Figure 5a). We

| Method | AUC score |
|--------|-----------|
| 1 Random forest | 0.585 |
| 2 AdaBoost | 0.589 |
| 3 GaussianNB | 0.537 |
| 4 LogisticRegression | 0.604 |
| 5 GradientBoosting | 0.575 |
| 6 SVM | 0.594 |
| 7 GCN (SchNet) | 0.683 |

"The technical details can be found on our GitHub page: https://github.com/tcsfrank0177/Graph-convolutional-network-DrugScreening.

![Figure 5](https://example.com/figure5.png)

**Figure 5.** Identification of inhibitors for HSPG-mediated endocytosis. (a) Workflow of α-synuclein fibrils uptake assay for confirmation of hits from virtual compound screen. (b) Summary of activities of the top 10 compounds; IC_{50} was determined by titration experiments. (c) Dose−response curves of compound’s inhibitory effect on α-synuclein fibrils uptake. (d) Measured fluorescence intensity of internalized α-synuclein fibril-Alexa596 by U2OS cells treated with compound at 2-fold of its IC_{50}. The experimental repeat number is 3.

**Table 1. Calculated AUC Scores of Machine Learning-Based Classification Methods upon the Applied Data Set**
then employed pHrodo red-labeled α-synuclein fibrils as a HSPG cargo in a combination screen because α-synuclein fibrils share a similar entry mechanism as SARS-CoV-2. Importantly, the fluorescence intensity of cells treated with pHrodo-labeled α-synuclein fibrils is only dependent on the amount of internalized cargo and the endolysosomal pH. By comparison, the luciferase-based pseudoviral entry assay can be influenced not only by the level of viral entry but also by other factors that impact mRNA expression, translation, and luciferase stability. The screen identified 256 active compounds with the most potent IC₅₀ value of 0.95 μM. We picked the 10 top compounds based on their potency and structural novelty (Figure 5b) and measured their cytotoxicity by an ATP content assay. The results showed that for four out of the 10 compounds, the IC₅₀ for cytotoxicity was at least 10-fold larger than that for the inhibition of α-synuclein fibril uptake (Figure 5b and c), suggesting a safety window for the usage of these drugs as endocytosis inhibitors. These newly identified compounds displayed significant structural novelty when compared to drugs in the training set. This can be verified by the Tanimoto similarity analysis (see the Support Information Table S2 for more details). Interestingly, among the 10 confirmed drugs, six of them share some common structural characteristics (NCGC00411718-01, NCGC00411727-01, NCGC00411705-01, NCGC00411733-01, NCGC00411718-01, NCGC00411588-01), suggesting a possible common mechanism for inhibiting HSPG-mediated endocytosis.

To rule out false-positive hits due to compound-induced changes in lysosomal pH, which could reduce the fluorescence of internalized α-synuclein fibrils, we measured the uptakes of α-synuclein fibrils labeled with a pH-insensitive dye (Alexa596) in U2OS cells. When cells were treated with the top 10 inhibitors at concentrations 2-fold higher than their respective
IC₅₀ values, we found that all compounds tested could significantly inhibit the uptake of α-synuclein fibrils compared to control treated cells (Figure 5d). These results suggest that these chemicals are indeed endocytosis inhibitors that block HSPG-mediated entry of α-synuclein fibrils. We then treated cells with increased concentrations of NCGC00411718 and NCGC00159478, which showed the highest inhibition on the entry of pHrodo-labeled α-synuclein fibrils. Drug-treated cells were incubated with Alexa596-labeled α-synuclein fibrils in the presence of the inhibitor for 2 h and imaged by a confocal microscope. The results suggest that both compounds inhibit α-synuclein fibril uptake in a dose-dependent manner with IC₅₀ comparable to that measured by pHrodo-labeled α-synuclein fibrils (Figure 6a−d).

Identification of SARS-CoV-2 Entry Inhibitors. To test whether the newly identified endocytosis inhibitors could inhibit the entry of SARS-CoV-2, we used a previously established pseudotyped particle entry assay (Figure 7a). As shown previously,6 the entry of the pseudoviral particles into cells results in the expression of the luciferase reporter. To control the impact of ACE2-GFP expression levels on viral entry under drug-treated conditions, we normalized the luciferase signals by the ACE2-GFP level. We also measured the cytotoxicity of these chemicals in ACE2-GFP-expressing cells using an ATP-based cell viability assay. We analyzed the top 27 compounds from the 256 inhibitors identified from the α-synuclein fibril uptake screen. Among them, 16 in total showed an inhibitory activity against the viral entry with the most potent IC₅₀ value of 0.76 μM. It is notable that some toxicity was observed for these compounds in HEK293T-ACE2-GFP cells after 48 h treatment. The viral inhibition and cytotoxicity curves of the top six compounds are shown in Figure 7b.

NCGC00115755 Inhibited SARS-CoV-2 Pseudotyped Particle Entry by Disrupting Actin Filaments. We previously showed that the actin network under the plasma membrane is critical for the entry of HSPG-dependent endocytosis cargos including SARS-CoV-2.6,15 We therefore asked whether any of the newly identified endocytosis could inhibit the actin cytoskeleton. To this end, we stained U2OS cells with Alexa488-labeled phalloidin, an actin binding dye. In control-treated cells, actin filaments were readily detected, which often run in parallel (Figure 8a). When cells treated with the top 10 endocytosis inhibitors were stained by Alexa488-labeled phalloidin, we observed dose-dependent disruption of cortex actin filaments (Supporting Information: Video 1, Video 2, Video 3). By contrast, in drug treated cells, the number of actin stress fibers were significantly reduced, and actin comet tails were barely detectable (Supporting Information: Video 1, Video 2, Video 3). Altogether, these findings suggest that NCGC00115755-02 disrupts actin filament assembly, resulting in an endocytosis defect.

**CONCLUSION**

Artificial intelligence-based virtual screening technologies have the potential to efficiently select drug candidates for specific targets with high accuracy at an affordable cost, and, therefore, are important complementaries to conventional high-throughput small molecule screening (HTS). SARS-CoV-2 viruses co-
opt a cellular endocytosis pathway to enter human airway epithelial cells. This key viral entry step has been subjected to conventional drug repurposing screens and computer docking-based screens, yielding several viral entry inhibitors.6,14 In this study, we developed and trained a GCN model using the structural information from previously identified SARS-CoV-2 entry inhibitors. When this model was applied to untested chemical libraries, it efficiently selected compounds with high probability of showing anti-SARS-CoV-2 activity. This model, when combined with conventional drug screening assays, generates a powerful strategy that allows rapid identification of new SARS-CoV-2 entry inhibitors. In principle, this strategy can be applied to any drug targets, which can quickly expand the existing inhibitor repertoire of any class. The findings shown in this study have revealed a promising venue for accelerated drug development.

#### DATA AND SOFTWARE AVAILABILITY

Technical details of the developed package can be found on our GitHub page: https://github.com/tcsnfrank0177/Graph-convolutional-network-DrugScreening. Programming environment: Python 3.6 or higher is recommended. Supporting videos are provided in the Supporting Information.

#### ASSOCIATED CONTENT

**Supporting Information**
The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.jcim.2c00222.

- Detailed similarity analysis (PDF)
- Video 1 (AVI)
- Video 2 (AVI)
- Video 3 (AVI)

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**Figure 8.** NCGC00115755-02 targets cellular actin cytoskeleton. (a) Cells treated with NCGC00115755-02 at the indicated concentrations were incubated with Alexa594-labeled α-synuclein fibrils for 2 h. Cells were stained with Phalloidin-Alexa488 in green to detect actin filaments and DAPI in blue to reveal the nuclei and then imaged. Note that cells treated with the drug have reduced levels of internalized α-synuclein fibrils. NCGC00115755-02 treatment also causes the disassembly of actin stress fiber and generates large actin aggregates. (b) Quantification of Alexa594-labeled α-synuclein fluorescence intensity in panel a. Error bars indicate SEM. The experimental repeat number is 2.
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Notes

The authors declare no competing financial interest.

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