Prioritizing antiviral drugs against SARS-CoV-2 by integrating viral complete genome sequences and drug chemical structures

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The outbreak of a novel febrile respiratory disease called COVID-19, caused by a newfound coronavirus SARS-CoV-2, has brought a worldwide attention. Prioritizing approved drugs is critical for quick clinical trials against COVID-19. In this study, we first manually curated three Virus-Drug Association (VDA) datasets. By incorporating VDAs with the similarity between drugs and that between viruses, we constructed a heterogeneous Virus-Drug network. A novel Random Walk with Restart method (VDA-RWR) was then developed to identify possible VDAs related to SARS-CoV-2. We compared VDA-RWR with three state-of-the-art association prediction models based on fivefold cross-validations (CVs) on viruses, drugs and virus-drug associations on three datasets. VDA-RWR obtained the best AUCs for the three fivefold CVs, significantly outperforming other methods. We found two small molecules coming together on the three datasets, that is, remdesivir and ribavirin. These two chemical agents have higher molecular binding energies of −7.0 kcal/mol and −6.59 kcal/mol with the domain bound structure of the human receptor angiotensin converting enzyme 2 (ACE2) and the SARS-CoV-2 spike protein, respectively. Interestingly, for the first time, experimental results suggested that navitoclax could be potentially applied to stop SARS-CoV-2 and remains to further validation.

In late December, 2019, there was an outbreak of a novel febrile respiratory illness (COVID-19) in Wuhan, Hubei in China1,2. The illness was caused by a novel coronavirus named SARS-CoV-2 by the World Health Organization (WHO) and can transmit from human to human2. As of 10 a.m. Cest time on October, 18, 2020, 40,118,333 cases of SARS-CoV-2 infection and 1,114,749 cases of SARS-CoV-2-caused death have been confirmed around the world3. From February, 2020, WHO is seeking U.S. $675 million for COVID-19 preparedness to prevent human to human transmission4.

SARS-CoV-2 is a new human-infecting single-stranded RNA virus5. It is very similar to two coronaviruses: severe acute respiratory syndrome (SARS) coronavirus (SARS-CoV) and Middle East respiratory syndrome (MERS) coronavirus (MERS-CoV). In November, 2002, SARS first emerged in Guangdong, China, and resulted in 8,098 infection cases with a fatality rate of 9.6%1. In September, 2012, MERS was first found in humans in the Middle East and resulted in 2,465 laboratory-confirmed cases of infection with a fatality rate 34.4%5.

As SARS-CoV-2 is an emerging virus, no specific antiviral treatment has been developed6. Therefore, finding effective drug treatment options is urgently needed for combating SARS-CoV-27. However, it seems unrealistic to test new drugs targeting SARS-CoV-2 within such limited time8. An efficient method is to screen possible drugs from available public data repositories containing FDA-approved compounds7,9. Under such situation, computational methods could be chosen to identify special antiviral drug candidates10–12.

Although little is known about SARS-CoV-2, its complete genome sequence suggests strong homology with SARS-CoV13. To identify possible antiviral drugs, in this study, we investigated the relationship between the complete genome sequences of viruses similar to SARS-CoV-2, the chemical structures of drugs, and Virus-Drug Association (VDA) network topology. We then developed a novel Random Walk with Restart method (VDA-RWR) to find possible VDAs related to SARS-CoV-2 by integrating the genome sequences and the chemical structures into a unified framework. We compared VDA-RWR with NGRHMDA14, SMiR-NBI15 and...
LRLSHMDA. These three methods were applied to biological association prediction in other application areas and obtained better prediction performance. We found that remdesivir and ribavirin come together on three datasets.

Molecular docking is a key bioinformatics modeling tool for drug discovery and used to predict the "best-fit" intermolecular binding between a small molecule and a target or two proteins at the atomic level. It characterizes the behavior of ligands in the binding sites of target proteins as well as elucidates fundamental biochemical processes. The docking process comprises two basic steps: predicting conformation, position, and orientation of ligands within the binding sites and ranking these conformations based on the binding affinity. We used AutoDock, a molecular docking software, to measure the molecular activities of the predicted two compounds (remdesivir and ribavirin) binding to the SARS-CoV-2 spike protein/human receptor angiotensin converting enzyme 2 (ACE2). The docking showed that remdesivir and ribavirin have higher binding energies of $-7.0 \text{ kcal/mol}$ and $-6.59 \text{ kcal/mol}$ with the structure of the spike protein receptor-binding domain bound to the ACE2 receptor, respectively.

Results

Experimental settings. In this section, we conducted extensive experiments to investigate the performance of our proposed VDA-RWR method. For the VDA matrix $Y_{n \times m}$ from $n$ viruses and $m$ drugs, fivefold Cross-Validations (CVs) were performed under the following three different experimental settings.

- **Fivefold Cross Validation 1 (CV1):** CV on viruses, that is, random rows in $Y$ (i.e., viruses) were selected for testing.
- **Fivefold Cross Validation 2 (CV2):** CV on drugs, that is, random columns in $Y$ (i.e., drugs) were selected for testing.
- **Fivefold Cross Validation 3 (CV3):** CV on virus-drug pairs, that is, random entries in $Y$ (i.e., virus-drug pairs) were selected for testing.

Under CV1, in each round, 80% of rows in $Y$ were used as training set and the remaining 20% of rows were used as test set. Under CV2, in each round, 80% of columns in $Y$ were used as training set and the remaining 20% of columns were used as test set. Under CV3, in each round, 80% of entries in $Y$ were used as training set and the remaining 20% of entries were test set. These three settings CV1, CV2, and CV3 specially refer to potential VDA identification for (1) new viruses (especially for SARS-CoV-2), (2) new drugs, and (3) new virus-drug pairs, respectively.

Parameters $r$, $\mu$, and $\alpha$ denote the global restart rate, the transition probability, and the weight between the virus network and the drug network, respectively. For these three parameters, we performed cross validations on the training set to find the optimal values. In addition, the iteration stopped when $\|p_{t+1} - p_t\|_2 \leq 1e^{-11}$. SMiR-NBI need not set the parameters. For the parameters in NGRHMDA and LRLSHMDA, we conducted grid search to find the optimal values. The detailed settings are shown on Table 1.

| Method     | Dataset 1          | Dataset 2          | Dataset 3          |
|------------|--------------------|--------------------|--------------------|
| NGRHMDA    | $\alpha = 0.4, \beta = 0.8$ | $\alpha = 0.6, \beta = 0.9$ | $\alpha = 0.9, \beta = 0.9$ |
| LRLSHMDA   | $\eta_M = 0.9, \eta_D = 0.3$ | $\eta_M = 0.8, \eta_D = 0.1$ | $\eta_M = 0.6, \eta_D = 0.1$ |
| VDA-RWR    | $r = 0.7, \mu = 0.9, \alpha = 0.5$ | $r = 0.5, \mu = 0.9, \alpha = 0.9$ | $r = 0.7, \mu = 0.9, \alpha = 0.9$ |

Table 1. The optimal values of parameters in VDA-RWR, NGRHMDA and LRLSHMDA.

Evaluation metrics. Sensitivity, specificity, F1 score, accuracy and AUC were widely applied to evaluate the proposed methods. Sensitivity denotes the ratio of correctly predicted positive VDAs to all positive VDAs. Specificity is the ratio of correctly predicted negative VDAs to all negative VDAs (all the unknown associations were labeled as negative). F1 score denotes the harmonic mean of recall and precision. Accuracy represents the ratio of correctly predicted positive and negative VDAs to all positive and negative VDAs. We used these five metrics to evaluate the performance of VDA-RWR. They were defined as follows:

\[
\text{Sensitivity} = \frac{TP}{TP + FN} \quad (1) \\
\text{Specificity} = \frac{TN}{TN + FP} \quad (2) \\
\text{Accuracy} = \frac{TP + TN}{TP + TN + FP + FN} \quad (3) \\
\text{F1 score} = \frac{2TP}{2TP + FP + FN} \quad (4)
\]
where $TP$, $FP$, $TN$ and $FN$ represent true positive, false positive, true negative and false negative, respectively.

AUC is the average area under the receiver operating characteristics (ROC) curve. The curve can be plotted by the ratio of True Positive Rate (TPR) to False Positive Rate (FPR) according to different thresholds. TPR and FPR are defined via Eqs. (4–5).

$$TPR = \frac{TP}{TP + FN} = \frac{TP}{F}$$  \hfill (5)

$$FPR = \frac{FP}{FP + TN} = \frac{FP}{F}$$  \hfill (6)

For these five evaluation metrics, higher values represent better performance.

### Performance evaluation under three fivefold CVs.

We compared VDA-RWR with NGRHMDA\textsuperscript{14}, SMiR-NBI\textsuperscript{15} and LRLSHMDA\textsuperscript{16}. NGRHMDA was presented to find potential microbe-disease associations by integrating neighbor-based collaborative filtering and graph-based scoring\textsuperscript{14}. SMiR-NBI can comprehensively identify new pharmacogenomic biomarkers by constructing a heterogeneous network connecting genes, drugs, and miRNAs\textsuperscript{15}. LRLSHMDA was applied to predict human microbe-disease associations based on Laplacian regularized least squares\textsuperscript{16}. These three state-of-the-art approaches obtained good performance in their corresponding applications. We performed these four methods for 100 times on three different fivefold CV settings on three datasets. The final performance was averaged over the five rounds for 100 times. The results are shown in Tables 2, 3, and 4. The best results were shown in bold in each column.

On dataset 1 and dataset 3, VDA-RWR outperformed other three methods in terms of specificity, accuracy, F1 score and AUC under three CVs. On dataset 2, although the sensitivity of VDA-RWR was slightly lower, VDA-RWR computed better specificity, accuracy, F1 score and AUC under majority of conditions. The slight difference can be produced by different data structures. AUC is one more important evaluation metric compared to other four measurements. AUC = 0.5 represents random performance and AUC = 1 shows perfect performance.

| Datasets | Methods     | Sensitivity | Specificity | F1 score | Accuracy | AUC  |
|----------|-------------|-------------|-------------|----------|----------|------|
| Dataset 1 | NGRHMDA     | 0.6435      | 0.6719      | 0.0850   | 0.6713   | 0.8329 |
|          | SMiR-NBI    | 0.8510      | 0.1917      | 0.0393   | 0.2064   | 0.6021 |
|          | LRLSHMDA    | 0.7938      | 0.5773      | 0.1122   | 0.5820   | 0.8249 |
|          | VDA-RWR     | 0.5070      | 0.8932      | 0.1294   | 0.8846   | 0.9182 |
| Dataset 2 | NGRHMDA     | 0.4867      | 0.8027      | 0.0719   | 0.7967   | 0.8017 |
|          | SMiR-NBI    | 0.9971      | 0.0929      | 0.0404   | 0.1098   | 0.7205 |
|          | LRLSHMDA    | 0.7720      | 0.4166      | 0.0639   | 0.4232   | 0.7334 |
|          | VDA-RWR     | 0.5045      | 0.7981      | 0.0814   | 0.7926   | 0.8025 |
| Dataset 3 | NGRHMDA     | 0.4579      | 0.6785      | 0.0279   | 0.6758   | 0.6772 |
|          | SMiR-NBI    | 0.9751      | 0.0434      | 0.0243   | 0.0549   | 0.5665 |
|          | LRLSHMDA    | 0.7420      | 0.5264      | 0.0493   | 0.5290   | 0.7468 |
|          | VDA-RWR     | 0.5054      | 0.8098      | 0.0628   | 0.8061   | 0.8168 |

Table 2. The performance comparison of four methods on three datasets under CV1. Bold values indicates the best values for the different methods under the same evaluation.

| Datasets | Methods     | Sensitivity | Specificity | F1 score | Accuracy | AUC  |
|----------|-------------|-------------|-------------|----------|----------|------|
| Dataset 1 | NGRHMDA     | 0.7278      | 0.3991      | 0.0643   | 0.4092   | 0.7026 |
|          | SMiR-NBI    | 0.8086      | 0.2164      | 0.0366   | 0.2296   | 0.5806 |
|          | LRLSHMDA    | 0.1299      | 0.6171      | 0.0084   | 0.6057   | 0.1844 |
|          | VDA-RWR     | 0.4977      | 0.7959      | 0.1055   | 0.7905   | 0.8157 |
| Dataset 2 | NGRHMDA     | 0.3987      | 0.5521      | 0.0329   | 0.5495   | 0.4301 |
|          | SMiR-NBI    | 0.8238      | 0.0949      | 0.0332   | 0.1087   | 0.4003 |
|          | LRLSHMDA    | 0.3507      | 0.4667      | 0.0179   | 0.4643   | 0.3173 |
|          | VDA-RWR     | 0.5106      | 0.6832      | 0.0844   | 0.6801   | 0.6932 |
| Dataset 3 | NGRHMDA     | 0.4435      | 0.4560      | 0.0232   | 0.4563   | 0.4058 |
|          | SMiR-NBI    | 0.9124      | 0.0459      | 0.0227   | 0.0567   | 0.4092 |
|          | LRLSHMDA    | 0.1801      | 0.5817      | 0.0074   | 0.5766   | 0.2920 |
|          | VDA-RWR     | 0.5270      | 0.7025      | 0.0812   | 0.7006   | 0.7276 |

Table 3. The performance comparison of four methods on three datasets under CV2. Bold values indicates the best values for the different methods under the same evaluation.
VDA-RWR obtained the best AUCs under majority of conditions. In general, VDA-RWR is proper to discover potential VDAs.

In addition, under CV1, VDA-RWR computed better specificity, accuracy, F1 score and AUC on the three datasets. This result showed that VDA-RWR can effectively find possible antiviral drugs for new viruses (for example, SARS-CoV-2). Under CV2, VDA-RWR outperformed other three methods in terms of specificity, accuracy, F1 score and AUC on dataset 1 and dataset 3. Although the sensitivity, specificity and accuracy values of VDA-RWR were slightly lower than other individual methods on dataset 2, it obtained the best F1 score and AUC. Thus AUC can identify potential viruses associated with new drugs. Under CV3, VDA-RWR calculated the best specificity, F1 score and accuracy. Figure 1 showed the AUC values of four methods under CV1, CV2, and CV3. The results demonstrated that VDA-RWR obtained relatively higher AUCs under three different CVs. It suggested that VDA-RWR could be used to infer potential VDAs.

Case study. In this section, we want to find possible drugs for SARS-CoV-2 after verifying the performance of our proposed VDA-RWR method. We predicted the top 10 drugs with the highest association scores with SARS-CoV-2 on three datasets. The results were shown in Tables 5, 6, and 7, respectively. Among the predicted top 10 small molecules associated with SARS-CoV-2 on dataset 1, all drugs were supported by recent works. Among the predicted top 10 chemical agents related to SARS-CoV-2 on dataset 2, there were 9 VDAs validated by current literatures, that is, 90% chemical agents were reported. Among the predicted top 10 antiviral drugs against SARS-CoV-2 on dataset 3, all compounds were validated by recent publications.

The results on Tables 5, 6, and 7 showed that there were two FDA-approved drugs coming together on three datasets, that is, remdesivir and ribavirin. Remdesivir is a small molecular compound undergoing a clinical trial and shows superior antiviral activity against many RNA viruses including orthocoronavirinae, filoviridae, paramyxoviridae, and pneumoviridae. Sheahan et al. presented that it can improve pulmonary function and reduce severe lung pathology in mice. Similar to SARS-CoV-2, both Ebola virus (EBOV) and MERS-CoV may result in severe acute respiratory diseases. And remdesivir has been used as inhibitors of EBOV and MERS-CoV. More importantly, an array of works have reported that remdesivir is highly effective in controlling SARS-CoV-2 infection and has been directly applied to the treatment of COVID-19. Interestingly, for the first time, experimental results suggested that navitoclax could be potentially applied to stop SARS-COV-2. Navitoclax has been applied to boost the treatment and basic science of chronic lymphoid leukemia, hematological malignancies, non-Hodgkin’s lymphoma, solid tumors, and EGFR activating mutation.

Molecular docking. The molecular docking between the above two antiviral drugs (remdesivir and ribavirin) and the spike protein and ACE2 are described in Table 8. The results showed that remdesivir and ribavirin have higher binding energies of $-7.0$ kcal/mol and $-6.59$ kcal/mol with the structure of the spike protein receptor-binding domain bound to the ACE2 receptor, respectively. The subfigure in each circle denotes the residues at the binding site of the spike protein/ACE2 and their corresponding orientations. For example, the amino acids K68 and Q493 were predicted to be the key residues for remdesivir binding to the SARS-CoV-2 spike protein/ACE2 while K353, R403, Q493 and G496 were predicted as the key residues for ribavirin binding to these two target proteins.

In Table 8, green denotes the structure of ACE2 and cyan denotes the SARS-CoV-2 spike protein in the figures of molecular docking.
Discussion
Finding possible antiviral drugs against SARS-CoV-2 is extremely urgent with the rapid spread of COVID-19. However, it seems very difficult to design a novel drug for COVID-19 within a very short time. One of efficient ways is to identify new clues of the treatment from FDA-approved drugs.

In our proposed VDA-RWR method, we computed the association scores for each virus-drug pair to predict potential antiviral drugs against SARS-CoV-2 based on random walk with restart and biological information.

Figure 1. The AUC values of VDA-RWR under different CVs on three datasets.

Table 5. The predicted top 10 drugs associated with SARS-CoV-2.

| Rank | Drug         | Evidence                                                                 |
|------|--------------|---------------------------------------------------------------------------|
| 1    | Remdesivir   | PMID: 32020029, 31996494, 32022370, 31971553, 32035018, 32035533, 32036774, 32194944, 32275812, 32145386, 32838064 |
|      |              | https://doi.org/10.1101/2020.01.28.922922                                  |
| 2    | Oseltamivir  | PMID: 32034637, 32127666                                                  |
| 3    | Ribavirin    | PMID: 32034637, 32127666, 32227493, 26492219, 32771797                     |
| 4    | Zanamivir    | PMID: 32511320                                                            |
| 5    | Presatovir   | PMID: 32147628                                                            |
| 6    | Elvitegravir | PMID: 32147628                                                            |
| 7    | Zidovudine   | PMID: 32568013                                                            |
| 8    | Emtricitabine| PMID: 32488835                                                            |
| 9    | Mycophenolic acid | PMID: 32579258                  |
| 10   | Chloroquine  | PMID: 32020029, 32145363, 32074550, 32236562                        |
of viruses and drugs. The originality of our proposed VDA-RWR method remains, constructing three small datasets and inferring possible antiviral chemical agents against SARS-CoV-2 from FDA-approved drugs. The comparative experiments showed better performance of the VDA-RWR method. Higher AUC values under three fivefold CVs on three datasets and molecular binding energies indicated that the selected small molecules are likely to be used to stop the transmission of COVID-19.

VDA-RWR can obtain superior performance under the three fivefold CVs on three datasets. This observation may be attributed to random walk with restart, a state-of-the-art model that can randomly walk on the heterogeneous virus-drug network and effectively compute association score for each virus-drug pair. More importantly, VDA-RWR integrated various biological information including the complete genome sequences of viruses and chemical structures of chemical agents.

The proposed VDA-RWR method is also helpful in design and interpretation of pharmacological experiment related to COVID-19. More importantly, VDA-RWR can be further applied to predict antiviral drugs against novel viruses without any associated chemical agents.

Methods

Virus-drug association data. **Dataset 1.** Virus data. We considered 11 viruses similar to SARS-CoV-2. These viruses include influenza A viruses including A-H1N132, A-H5N133, and A-H7N934, chronic hepatitis C virus (HCV)35, human immunodeficiency virus type 1 (HIV-1)36, human immunodeficiency virus type 2 (HIV-2)37, hendra virus38, human cytomegalovirus39, MERS-CoV40, respiratory syncytial virus41 and SARS-CoV42. The complete genome sequences of these viruses are downloaded from the NCBI database43. We used MAFFT44 (https://mafft.cbrc.jp/alignment/software/, version 7, open source license: GPL or BSD), a multiple sequence alignment tool, to compute virus-virus sequence similarity matrix $S_v$.

| Rank | Drug          | Evidence                              |
|------|---------------|---------------------------------------|
| 1    | Favipiravir   | PMID: 32346491, 32967849, 32972430    |
| 2    | Remdesivir    | PMID: 32020029, 31996494, 32022370, 31971553, 32035018, 32035533, 32036774, 32194944, 32275812, 32145386, 32838064 |
|      |               | [https://doi.org/10.1101/2020.01.28.922922](https://doi.org/10.1101/2020.01.28.922922) |
| 3    | Cidofovir     | PMID: 32546018                         |
| 4    | Galidesivir   | PMID: 32711596                         |
| 5    | Niclosamide   | PMID: 32125140, 32221153              |
| 6    | Mycophenolic acid | PMID: 3257258                 |
| 7    | Iracanazole   | [https://doi.org/10.22541/au.159467021.16927198](https://doi.org/10.22541/au.159467021.16927198) |
| 8    | Brequinar     | PMID: 32426387                        |
| 9    | Navitoclax    | Unconfirmed                           |
| 10   | Ribavirin     | PMID: 32034637, 32127666, 32227493, 26492219, 32771797 |

| Rank | Drug          | Evidence                              |
|------|---------------|---------------------------------------|
| 1    | Nitazoxanide  | PMID: 32127666, 32568620, 32448490    |
| 2    | Ribavirin     | PMID: 3203637, 32127666, 32227493, 26492219, 32771797 |
| 3    | Chloroquine   | PMID: 32020029, 32145363, 32074550, 32236562 |
| 4    | Hexachlorophene | PMID: 15950190                          |
| 5    | Camostat      | PMID: 32347443                         |
| 6    | Favipiravir   | PMID: 32346834                        |
| 7    | Umifenovin    | PMID: 32941741                        |
| 8    | Remdesivir    | PMID: 32020029, 31996494, 32022370, 31971553, 32035018, 32035533, 32036774, 32194944, 32275812, 32145386, 32838064 |
|      |               | [https://doi.org/10.1101/2020.01.28.922922](https://doi.org/10.1101/2020.01.28.922922) |
| 9    | Amantadine    | PMID: 32361028                        |
| 10   | Niclosamide   | PMID: 32125140, 32221153             |

**Table 6.** The predicted top 10 drugs associated with SARS-CoV-2 on dataset 2.

**Table 7.** The predicted top 10 drugs associated with SARS-CoV-2 on dataset 3.

Drug data. We manually curated drugs associated with these 11 viruses from the DrugBank45 and NCBI46 databases and published literatures reported by the PubMed database46 and collected 78 small molecules after removing macromolecules. Based on the assumption that two drugs are more similar if they share more chemi-
cal substructures, drug-drug similarity can be computed. Extended connectivity fingerprints (ECFPs) are circular fingerprints and developed for structure–activity modeling and molecular feature description. We used RDKit (https://github.com/rdkit/rdkit, releases 131, open source license: BSD), an open-source cheminformatics software, to compute ECFPs of drugs with a radius of 2. Drug-drug chemical structure similarity matrix $S_d$ can be computed by the ECFPs of drugs.

VDAs. We searched the publicly available repositories including the DrugBank45 and NCBI43 databases and publications reported by the PubMed database46. At the time of writing, we obtained 96 virus-drug associations (VDAs) between 11 viruses and 78 drugs. We described A-H1N132, A-H5N133, and A-H7N934 as three viruses although they belong to influenza A for the sake of description.

Dataset 2. The DrugVirus.info database49 (https://drugvirus.info/) provided various VDA-related resources. We obtained 770 VDAs from 69 viruses and 128 drugs after removing the viruses whose complete genome sequences are unknown from the database. The chemical structure of drugs and the complete genome sequences of viruses were downloaded from the DrugBank database and the NCBI database, respectively. Similar to dataset 1, we used RDKit and MAFFT to calculate drug similarity and virus similarity.

Dataset 3. We retrieved 407 VDAs from 34 viruses and 203 drugs by searching documents related to viruses and drugs based on text mining techniques. Similar to dataset 1, we computed drug similarity matrix and virus similarity matrix. The details of three datasets are shown in Table 9.

In this study, the set of known VDAs was considered as the ‘gold standard’ dataset and was applied to evaluate the performance of our proposed VDA-RWR method. We described the known VDAs as a matrix $Y$:

| Ligand | Molecular formula | Molecular docking | Binding energy (kcal/mol) | Binding sites | Distance(Å) |
|--------|-------------------|-------------------|---------------------------|--------------|-------------|
| Remdesivir | C_{27}H_{35}N_{6}O_{8}P | ![Image](remdesivir.png) | 7.0 | K68 | 2.0 |
| Ribavirin | C_{18}H_{26}CIN_{3} | ![Image](ribavirin.png) | 6.59 | K353 | 2.2 |

Table 8. Molecular docking between remdesivir and ribavirin and the SARS-CoV-2 spike (S) protein/ACE2.

| Datasets | Viruses | Drugs | VDAs |
|----------|---------|-------|------|
| Dataset 1 | 12      | 78    | 96   |
| Dataset 2 | 69      | 128   | 770  |
| Dataset 3 | 34      | 203   | 407  |

Table 9. Statistics for the virus-drug association networks.
where $v_i$ and $d_j$ represent the $i$th virus and $j$th drug, respectively.

The VDA-RWR method. Inspired by the method provided by Valdeolivas et al., we developed a VDA prediction method based on Random Walk with Restart on the heterogeneous network (VDA-RWR). The proposed VDA-RWR method comprised two steps. First, a random walk-based model integrating various biological data was learned to explain the constructed ‘gold standard’ dataset. Second, this model was used to find potential VDAs for viruses and drugs absent from the ‘gold standard’ dataset. The details are shown in Fig. 2.

We first considered virus-virus similarity graph $G_v$, drug-drug similarity graph $G_d$, and VDA graph $G_a$, which formed a heterogeneous network. We defined $S_v(n \times n)$, $S_d(m \times m)$, and $Y(n \times m)$ as their corresponding adjacency matrices. The adjacency matrix of the heterogeneous network can be denoted as:

$$W = \begin{bmatrix} S_v & Y \\ Y^T & S_d \end{bmatrix},$$

where $Y^T$ denoted the transpose of the VDA matrix $Y$.

We then calculated the transition probabilities of random walk with restart on the heterogeneous network. Suppose $W = \begin{bmatrix} W_{vv} & W_{vd} \\ W_{dv} & W_{dd} \end{bmatrix}$ represented the matrix of transitions on the heterogeneous network, where $W_{vd}/W_{dd}$ denoted the walk within the virus/drug network, $W_{vd}/W_{dd}$ described the jump from the virus/drug network to the drug/virus network. Given the probability $\mu$ of jumping from the virus/drug network to the drug/virus network, the transition probability from virus $v_i$ to virus $v_j$ was defined as

$$W_{vv}(i,j) = \begin{cases} \frac{S_v(i,j)}{\sum_{k=1}^{n} S_v(i,k)} & \text{if } \sum_{k=1}^{n} Y(i,k) = 0 \\ \frac{(1-\mu)S_v(i,j)}{\sum_{k=1}^{n} S_v(i,k)} & \text{otherwise} \end{cases}$$

The transition probability from virus $v_i$ to drug $d_j$ was defined as

$$Y_{ij} = \begin{cases} 1 & \text{if } v_i \text{ associates with } d_j \\ 0 & \text{otherwise} \end{cases}$$

Figure 2. Flowchart of the VDA-RWR method based on the genome sequences of viruses, the chemical structures of drugs, and random walk with restart on the heterogeneous network.
The transition probability from drug $d_i$ to drug $d_j$ was defined as

$$W_{di}(i,j) = \begin{cases} \frac{\mu Y(i,j)}{\sum_{k=1}^{m} Y(i,k)} & \text{if } \sum_{k=1}^{m} Y(i,k) \neq 0 \\ 0 & \text{otherwise} \end{cases}$$

The transition probability from drug $d_i$ to virus $v_j$ was defined as

$$W_{dv}(i,j) = \begin{cases} \frac{\mu Y(i,j)}{\sum_{k=1}^{n} Y(k,i)} & \text{if } \sum_{k=1}^{n} Y(k,i) \neq 0 \\ 0 & \text{otherwise} \end{cases}$$

For a given virus/drug, the particle can either jump between graphs or stay in the current graph with a defined probability $r \in (0, 1)$. Therefore, we finally defined the random walk with a restart probability $r$ as:

$$p_{t+1} = rWp_t + (1-r)p_0$$

where $p_t$ represented the computed association probability at the $t$-th step random walk. We defined the initial probability as: $p_0 = \begin{bmatrix} \alpha u_0 \\ (1-\alpha) t_0 \end{bmatrix}$, where $u_0$ and $t_0$ denoted the initial probability on the drug network and the virus network, respectively. If we tend to identify possible drugs for a given virus $v_j$, it is considered as a seed node in the virus network. Here, $v_j$ was assigned as 1 and other nodes as 0, constructing the initial probability of the virus network $t_0$. All nodes in the drug network $u_0$ were assigned as equal probabilities with the sum of 1. For example, to find potential antiviral drugs against SARS-CoV-2, we set SARS-CoV-2 as a seed node, and all drugs in the drug network were assigned as the same probabilities with the values of 1/m. The parameter $\alpha$ was used to control the weight of the virus network and the drug network. In addition, a virus is new if it does not associate with any drugs, and a drug is new if it is not applied to any viruses.

Molecular docking. Molecular docking technique was applied to compute the intermolecular binding abilities between the predicted anti-SARS-CoV-2 drugs and the SARS-CoV-2 spike protein/human ACE2. The chemical structures of drugs were downloaded in the form of the PDB format files from the DrugBank database. We used AutoDockTools to convert these PDB files into pdbqt files needed by AutoDock4. The structures of SARS-CoV-2 spike receptor-binding domain bound with ACE2 (PDB ID: 6M0I) were downloaded from the RCSB Protein Data Bank. The spike protein and ACE2 were used as receptors, and the predicted anti-SARS-CoV-2 drugs were used as ligands for the molecular docking.

We first removed solvent and organic compounds and preprocessed the receptor proteins based on PyMOL. We used AutoDockTools to convert these PDB files into pdbqt files needed by AutoDock4. The spike protein and ACE2 were used as receptors, and the predicted anti-SARS-CoV-2 drugs were used as ligands for the molecular docking.

Conclusion
To find potential antiviral drugs, in this study, we integrated the complete genome sequences of viruses, the chemical structures of drugs, and the VDA network. We then developed a VDA prediction method based on random walk with restart on the heterogeneous network. The results suggested that remdesivir and ribavirin may be applied to the treatment of COVID-19. In the emergency situation, this study focused more on finding antiviral drugs. In the future, we will further integrate more biological data and design more powerful models to improve the accuracy of VDA identification. We hope that our proposed VDA-RWR method could help the screening of drugs for preventing COVID-19.

Data availability
Source codes and datasets are freely available for download at https://github.com/plhhn/VIAD-RWR/.

Received: 18 July 2020; Accepted: 18 December 2020
Published online: 18 March 2021

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Acknowledgements
We are thankful for help from Guangyi Liu, Ming Kuang, and Longjie Liao from Hunan University of Technology, and Ruyi Dong, Lebin Liang, Qinqin Lu, and Jidong Lang from Geneis (Beijing) Co. Ltd. We would like to thank all authors of the cited references.

Author contributions
L.-H.P. and L.S. contributed equally to this work. L.-H.P., J.-L.X., J.-L.Y., and L.-Q.Z. conceived the study, designed the schedule, and analyzed the data. F.-X.L. screened the viruses similar to SARS-CoV-2, X.-F.T. downloaded the genome sequences of viruses and computed virus similarity matrix, L.S. computed drug similarity matrix, L.S., X.-F.T., F.-X.L., and J.-J.W. constructed VDA network, L.S. run random walk algorithm, L.-H.P., G.T., and L.-Q.Z. wrote the paper, J.-L.Y. revised the original draft. All authors read and approved the final manuscript.

Funding
This research was funded by National Natural Science Foundation of China (Grant 61803151) and Natural Science Foundation of Hunan province (Grant 2018JJ3570, 2018JJ2461).

Competing interests
Authors Geng Tian and Jialiang Yang were employed by the company Geneis (Beijing) Co. Ltd. The remaining authors declare that the research was conducted in the absence of any commercial or financial relationships that could be constructed as a potential conflict of interest.

Additional information
Supplementary Information The online version contains supplementary material available at https://doi.org/10.1038/s41598-021-83737-5.

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