Artificial neural network-based study predicts GS-441524 as a potential inhibitor of SARS COV-2 activator protein furin: A polypharmacology approach

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Research Article

Keywords: Furin, Covid-19, Drug repurposing, ANN, SOM, LB-VS, SB-VS, Polypharmacology, GS-441524, Antivirals

Posted Date: November 18th, 2021

DOI: https://doi.org/10.21203/rs.3.rs-932436/v1

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Abstract

Furin, a pro-protein convertase, plays a significant role as a biological scissor in bacterial, viral, and even mammalian substrates which in turn decides the fate of many viral and bacterial infections along with the numerous ailments caused by cancer, diabetes, inflammations, and neurological disorders. In the wake of the current pandemic caused by the virus SARS COV-2, furin has become the center of attraction for researchers as the spike protein contains a polybasic furin cleavage site. In the present work, we have searched for novel inhibitors against this interesting human target from FDA-approved antivirals. To enhance the selection of new inhibitors we employed Kohonen’s artificial neural network-based self-organizing maps for ligand-based virtual screening. Promising results were obtained which can help in drug repurposing and network pharmacology studies addressing the errors generated due to promiscuity/polypharmacology. We found 15 existing FDA antivirals having the potential to inhibit furin. Among these, six compounds have targets on many important human proteins (LDLR, FCGR1A, PCK1, TLR7, DNA, and PNP) also. These 15 drugs inhibiting furin could be studied in patients infected with many viruses including SARS COV-2. We also propose two promising candidate FDA drugs GS-441524 and Grazoprevir (MK-5172) for repurposing as inhibitors of furin. The best results were observed with GS-441524.

1. Introduction

In the recent past even after a huge development in pharma industries new drugs or treatments approved by USFDA have remained constant (1). Repurposing of the drug is always promising as not only it can shorten the time for designing but also be cost-effective with randomized clinical trials (2). Different computational platforms can be used with novel testable hypotheses for systematic drug repositioning (3), (4). Due to the unavailability of 3D-based protein structures in some cases, traditional structure-based methods are not always followed. Furthermore, drug resistance is a big challenge due to the single virus protein target as the viral genome is rapidly evolving (5). Our group recently reported for the first time in India in 2021 about two lineages B.1.112 and B.1.99 and also revealed 1,143 unique single nucleotide variants along with 73 novel variants using COVIDSeq approach for SARS-CoV-2 (6). Most antiviral therapies do not completely eradicate the viral infection. Instead, they limit the immune recovery, and their long-term use results in resistance and toxicity, making space for the search of new antiviral classes with new targets. Specific antiviral agents should essentially target either the viral life cycle or the host immune clearance. There are two categories of targets required for the completion of virus replication and infection; viral molecules and host molecules. In our recent work on molecular docking-based screening, direct antiviral activity by remarkable binding of vitamins as inhibitors to various important proteins of SARS-Cov-2 are suggested (7). Moreover, this work revealed vitamin B12 to be employed as an efficient nutraceutical for almost all the nCoV-2 drug targets including the host protease furin which is a human host molecule and one of the promising targets for therapeutic intervention. It cleaves more than 150 mammalian, viral and bacterial proteins including the disastrous virus coronaviruses (CoVs). Researchers in their recent works have found furin inhibitors as a new avenue
for the treatment of COVID-19 (8). The mechanism for furin-mediated protein processing for health and disease is well-known. Furin is a member of the pro-protein convertase family, PCSK 3 representing furin. The viral and bacterial infection is caused by the production of various toxins by proteolytic cleavage such as Anthrax toxin, Diphtheria toxin, Pseudomonas exotoxin A, Shiga toxin, etc. Furin activation of viral proteins is essential for their entry to the host cell. Furin has basic amino acid motifs as a cleavage site and known as “PACE” paired basic amino acid cleaving enzymes. It is located at chromosome 15 q26.1 and expressed by the FUR (FES upstream region) (9). The hot spot cleavage site of furin is known as R-X-K/R-R↓ (10). Bioinformatic analyses have revealed more than 100 furin cleavage sites in humans such as growth factors, cytokines, collagens, adhesion molecules, MMP (metalloproteinases), receptors, etc. (11). Many recent reports have mentioned peptidic and nonpeptidic inhibitors for therapeutic inhibition of furin resulting in its reduced proteolytic activity. Numerous pharmaceutical companies have come up with solutions for the current global pandemic caused by SARS-CoV-2 with different vaccines and existing medicines. However, vaccination is going on in spite of which many cases are being observed and no effective medications are there due to the appearance of double mutant and triple mutant strains. Hence the need of the hour is a better prognosis of the Covid patients.

Cheminformatics incorporating machine learning are breakthrough techniques imperative in the pharmaceutical field today as we live in an era of infectious outbreaks, rising lifestyle diseases, and the age-old time-consuming procedures in drug discovery (12). Artificial intelligence (AI) and machine learning (ML) tools are applied for furin targeting while screening of molecules including higher-dimensional similarity/differences and networking observation has been performed to establish a relationship (interactome) with selected drug compounds and their respective important host factors along with furin.

Kohonen's Self-organizing maps (SOM) are extensively used today to establish the relationship of structure to biological activity and drug discovery in much pharmaceutical research. This methodology is widely applied in drug repurposing and scaffold hopping (13). Self-Organizing Maps (SOM) are mainly an artificial neural network (ANN) that implements virtual screening for assembling thematic molecular libraries. The concept of the SOM technique, case studies, and potential future applications are presented in the report presented by Schneider et al (13). Here the ANN is employed to identify the nonlinear connections of arbitrary dimensions into processable real numbers of one or two-dimensional maps at less computational costs. This mapping technique permits high-dimensional input space to be transformed into lower-dimensional output space (14). Studies with SOM supervised methods were proven more accurate than standard linear QSAR methods (15). ANN is not a single algorithm; instead, it comprises a framework of machine learning algorithms to process complex data. Kohonen's self-organizing map is employed here to screen and predict the active FDA antivirals on Furin basically to transform a pattern of arbitrary dimension into a 2D map adaptively.

Managing large amounts of heterogeneous data in a short period is becoming possible through AI technology. AI/ML, systems biology, including computer-aided drug repurposing are speedily becoming an integral part of the global workflow, to prioritize novel drug targets, new chemical entities, and to
evaluate off-label or drug repositioning proposals (16). All these above-mentioned techniques are utilized in our present work where we focused to present an integrative antiviral drug repurposing methodology for the first time with the combination of systems pharmacology-based network medicine platform that quantifies the interplay between the virus-host interaction and drug targets in the human PPI network. An ANN employed polypharmacology based computational approach has been reported in the present paper with multiple antiviral drugs for furin which uncovers the therapeutic interventions.

2. Materials And Methods

2.1 Datasets:
Human furin protein had 29 bioassays containing 221066 tested compounds in PubChem Repository on 07-07-2020 (https://pubchem.ncbi.nlm.nih.gov/protein/EAX02111). All furin actives and inactives were taken for analysis except inactive of AID 463115 (~ 2.2 lakh) to minimize the bias due to imbalance of diversity of actives and inactives (17). The 386 actives and 93 in-actives in structure format were downloaded from the PubChem repository which was tested on human furin (PACE). These compounds were used for descriptor generation and subsequent model building.

Among the 21 PDB structures available, 5 MIM along with the bound ligand C$_{28}$H$_{37}$N$_{13}$O$_{2}$, (CID − 44593169, IUPAC Name: 2-(4-((1R,2R,4S,5S)-2-(bis (azanyl) methylideneamino)-5-(4-(bis(azanyl)methylideneamino)phenoxy)-5-(4-(bis(azanyl)methylideneamino)phenyl) amino)cyclohexyl)oxyphenyl) guanidine) which is considered as reference molecule in this study, was downloaded from the protein data bank as it has a single chain and zero mutations with a good X-ray Structure Validation Report (https://www.rcsb.org/structure/5MIM). Antiviral drugs which come under approved/investigational category in drugbank (https://go.drugbank.com/) were selected as screening dataset and their SDFs (total 180) were downloaded from PubChem repository.

2.2 Model building by SOM clustering:
The SDF files of furin actives and in-actives downloaded from PubChem bioassays were used to build a model using self-organizing map, the unsupervised machine learning clustering algorithm in Canvas of the Schrodinger suite (Schrodinger, Inc., LLC, New York, USA) and to screen the activity of the FDA approved/investigational antiviral drugs. The physicochemical, topological, ligfilter, and the qikprop descriptors generated were fed as input into the neural network which in turn had undergone artificial neural network-based competitive learning to develop the model cluster. The SOM is generated by the property-based clustering using all available descriptors of categories mentioned above generated by the software.

2.3 Ligand docking:
The protein was pre-processed in the protein preparation wizard of maestro 9.3 (Schrodinger, Inc., LLC, New York, USA) and the states were generated at a pH 7.0 ± 2.0. The metals and the water molecules (Beyond 5Å from the hit groups) were removed and minimized in the OPLS-3e force field. The ligand is
also prepared using the ligprep wizard of maestro taking all stable conformations up to 32. The grid for docking is prepared by taking the centroid of the PDB inbuilt inhibitor in Glide. Ligand docking is done with the prepared protein on the generated grid with the prepared ligand (the ligands from best clusters). The glide score along with the glide e-model are tabulated and analyzed for binding affinity.

2.4 Network interaction analysis

The interactome was built for (i) FDA drug and human target network and (ii) FDA drug and viral target network in connection with human target furin. All the target proteins examined in this study are of antiviral FDA drugs (supplementary table S6 and S7) which are considered active on furin and obtained from the drug bank. The information of targets for each drug is collected from UniProt id and the interactions downloaded from bio grid interactions. The network is built and visualized in Cytoscape 3.8.2 (https://cytoscape.org/).

2.5 Molecular Dynamics simulation studies

Molecular dynamic simulations on the furin-reference ligand, furin-GS-441524, and furin-Grazoprevir complexes were carried out using the GROMACS 5.1 software and the regular CHARMM36 force field. The ligand topology was optimized in a high-throughput way using CGenFF. The CHARMM General Force Field (CGenFF) tool combines atom typing and analogy assignment of parameters and charges. Atom typing is done by a testable configurable decision tree (18). The role of a protein or enzyme is inextricably related to its internal and external dynamics. Since there are no experimental options, the only way to test protein dynamics is by quantitative analytical techniques. A customized GROMOS966 was used to run a 50-ns MD simulation to investigate protein-ligand binding and unbinding in terms of energy and geometry shifts, as well as the mechanism and functional behavior of the furin protein after binding to both ligands subsequently. Furthermore, the energy of systems was reduced by using the steepest descent approaches. To keep the systems running for 100 ps, the canonical ensemble of NPT and NVT was used. For 50 ns, molecular dynamics simulation was done. The protein-ligand complexes main chain backbone atoms’ root-mean-square divergence (RMSD), root-mean-square fluctuation (RMSF) the radius of gyration (Rg), and solvent-accessible surface area (SASA) values were calculated.

3. Results

3.1 Building of model for antiviral drugs via SOM clustering

The Furin actives and in-actives are manually curated from the PubChem database (supplementary table S1 and S2) for model building for the ligand-based virtual screening (LB-VS). ANN-based self-organizing maps were generated using unsupervised machine learning clustering algorithms in the canvas Schrodinger suite and using the above-mentioned model the activity of the FDA-approved/investigational antiviral drugs were screened (supplementary table S3). The descriptors generated in canvas were applied for the ANN modeling inputs and the SOM was generated by the property-based clustering. However, SOM is considered as a supervised training method in this study and we employed the actives
as input data without labeling while submitting the job; Most of the time the output-data analysis became more straightforward with known labels (19). The number of molecules from each inadequate dataset is presented in Table 1.

| Cluster no | Clusters | Actives | Inactives | FDA Considered for analysis | Reason of consideration of molecules |
|------------|----------|---------|-----------|-----------------------------|--------------------------------------|
| 1          | 69       | 69      |           | Discarded                   | No Actives                           |
| 2          | 33       | 20      | 13        | Selected                    | FDA + Active & No Inactive           |
| 3          | 29       | 29      |           | Discarded                   | No FDA Drugs                         |
| 4          | 27       | 25      | 2         | Selected                    | FDA + Active & No Inactive           |
| 5          | 23a      | 19      | 4         | Selected                    | FDA + Active & No Inactive           |
| 6          | 23b      | 14      | 9         | Selected                    | FDA + Active & No Inactive           |
| 7          | 21       | 21      |           | Discarded                   | No FDA Drugs                         |
| 8          | 20a      | 19      | 1         | Selected                    | FDA + Active & No Inactive           |
| 9          | 20b      | 20      |           | Discarded                   | No Actives                           |
| 10         | 20c      | 19      | 1         | Selected                    | FDA + Active & No Inactive           |

The SOM results are presented in distinct rows and columns in specific color codes that help us to scrutinize the high dimensional data of complex datasets into lower-dimensional space. In this study, the properties of Furin actives and in-actives accompanying with FDA drugs had undergone a mapping to place each molecule into individual clusters (10*10 cells) possessing varied populations, a maximum of showing in red color (69 molecules) and most clusters with least members, null values are shown in blue color (see Fig. 1a). Figure 1b illustrates the Euclidean distances in the hyperspace between the molecules. When the difference between Euclidean distances of data points was large and was depicted as red, whereas small distances were in blue. The variations among the population and the distance were interpreted by various colors between blue and red. The bioassay actives are presented in blue. The slightly variant molecules are depicted in green for a very small distance (in the) analysis and they are resembling in property space as well. The greener section owned higher numbers of FDA drugs.

Each cluster was manually examined to confirm the most desirable molecules comprising only actives along with FDA drugs and restrict in-actives molecules. The clusters in the combination of all three active/inactive/FDA were discarded. Out of 15 clusters, the clusters holding more than 10 molecules were selected for further analysis. Cluster number 2 holding 33 molecules has the maximum number of FDA.
antivirals and actives together (20 actives + 13 FDA). Cluster number 4 carrying 27 molecules (25 actives + 2 FDA) which is the second-largest membered cluster yet possessing just two FDA drugs (GS-441524 and Grazoprevir). Cluster 4 possessing active ligand C_{28}H_{37}N_{13}O_{2} (PubChem CID: 44593169) which was in interaction with furin from the protein structure database (PDB Id 5MIM) acknowledged as the reference active molecule in this study (20). Cluster numbers 5, 6, and 8 also own the active and FDA drugs screened out and granted for additional analysis.

### 3.1.1 Pharmacokinetics profile of selected antiviral drugs from SOM clustering

The physicochemical descriptors generated in canvas, the important ones are tabulated in Table 2. The higher values of molecular weight, AlogP, rotatable bonds, ring count, a higher number of N atoms, and a lower number of O atoms, all can make a compound promiscuous (21). The Grazoprevir is a larger molecule compared to the reference ligand though showing a similar structure with properties of higher values. GS-441524 is smaller and all physicochemical properties are promising compared to the reference ligand. The list of both clusters 2 and 4 with their structures are listed in supplementary tables S4 and S5.
3.2 Screening for FDA antivirals against furin using molecules elected from SOM clustering via molecular docking

Molecular docking analysis is a vital tool for structure-based virtual screening (SB-VS) to predict the promising structural conformations amongst ligand and active sites of an enzyme/receptor. The SOM examination unveiled five clusters 2, 4, 5, 6, and 8 are notable for additional knowledge hence all active ligands along with FDA drugs were docked against the furin. The docking result has shown the conformation and the compound with the least docking score was taken as the most reliable docked antiviral. The best 15 FDA antiviral drugs fall in two clusters 2 and 4 with an immeasurable docking score.
(Fig. 2c) with furin. Figure 2a shows that all 60 (33 + 27) molecules of clusters 2 and 4 have docking scores close to each other and in the model and it is evident that most of their energy falls in the −40 to −90 range in Fig. 2b.

The best docked FDA drug was an outlier while comparing with furin active molecules with reference to Pharmacokinetic and topological descriptors as well. (Table 2) These drugs and their former targets were further considered for the network analysis.

### 3.3 Unraveling furin inhibitor by network-based polypharmacology study

The pharmacokinetic descriptors are high for most of the drugs listed in Table 2 and can have promiscuity (21). It is necessary to analyze all possible polypharmacology before we practice FDA-approved drug candidates on any other targets. For this purpose, we utilized the network-based interaction study of the drugs selected to repurpose. This analysis aims to check the interaction network of present targets before we move further, whether it is in host or pathogen and how costly it becomes while repurposing it on furin. Concerning FDA drugs with multiple targets to be active against furin, its target proteins should be within or near the corresponding subnetwork in the host-pathogen interactome.

The structure-based virtual screening outcomes uncovered 15 promising antiviral FDA drugs for repurposing against the furin even though one was an outlier from the reference ligand values. The molecular targets of FDA antivirals are arranged in Table 3.
The drug-target network exploration represents (Fig. 3) that nodes (red with label) at the center are human targets and white with label are their interactions. The plotted networks manifest six molecules in the interaction network shown in red as Pormer Sodium (LDLR, FCGR1A), Isatoribine (TLR7), GppCp (PCK1), Didanosine (PNP), Valganciclovir and Entecavir. A drug with multiple targets can be efficient for a disease or may cause off-target side effects. Hereabouts, the targets are associated with multiple significant biological pathways (Fig. 3) therefore FDA drugs associated with these targets are eliminated in the further analysis. The proteins that act as a drug target for one disease may also be appropriate drug
targets for possible antiviral infections due to common protein-protein interactions (PPI) and functional pathways explained by the human interactome (22). From network analysis, we found that human protein furin is directly associated with viral drug targets in the PPIs. The most abundant group related to furin is SARS CoV-2 related interactome. We identified that furin is exclusively found in the GS-441524 drug’s subnetwork. Conversely, Valaciclovir, Penciclovir, Acyclovir, and Ganciclovir presently acting on Human Herpesvirus (HHV) protein UL30 (DNA polymerase), and no interaction is observed with human proteins. Grazoprevir is associated with the NS3/4A protein of the Hepatitis C virus but no interactions to analyze. The network-based drug-disease vicinity sheds light on the connection between drug targets and molecular factors in disease pathobiology modules within the PPIs and can serve as a useful tool for efficient screening of potentially new suggestions for FDA-approved drugs. In brief, the up-to-date interactome examination disclosed that GS-441524 can be a repurposed drug on furin.

3.4 Insight into structural perturbation and interaction profile of Furin bound with GS-441524, Grazoprevir, and reference ligand

The molecular dynamics simulation approach has been systematically employed to inspect features of conformational changes leading to modifications in protein-ligand interactions, molecular dynamics, and protein folding (23). In this study, according to SOM clustering and network-based polypharmacology analysis, Grazoprevir and GS-441524 can be repurposed on human furin without many complications as per the available pieces of information today. Taking into consideration the parameters such as root means square deviation (RMSD), root means square fluctuation (RMSF), the radius of gyration (Rg), and solvent-accessible surface area (SASA), MD simulation was performed for 50 ns to observe the change (if any) in the conformation of furin with drugs, as compared to that of the active reference ligand \( \text{C}_{28}\text{H}_{37}\text{N}_{13}\text{O}_{2} \).

The RMSD value of the backbone atoms of the ligand GS-441524-furin complexes was observed at about 0.15Å which is found to be as good as the RMSD of Grazoprevir-furin (0.2Å). However, in comparison to the reference furin inhibitor complex, different types of deviation were observed throughout the simulation. Overall, RMSD ranges of ~0.1–0.15, ~0.1–0.25 nm, and ~0.1–0.15 perceived for the GS-441524, Grazoprevir, and the reference complexes respectively (Fig. 5a.). There is retreating of RMSD in the ligand orientation near ~30 ns and ~40 ns in the reference ligand complex. The GS-441524 complex scale of fluctuation and the dissimilarity between the average RMSD values acclaimed that simulation produced stable trajectories. Therefore, these outcomes established that FDA drug GS-441524 is more enduring complex with furin in correlation to the other two molecules. The backbone RMS fluctuations were also calculated as they can postulate the clue about backbone mobility. The RMSF study correlates the functional analysis (Fig. 5b.), the highest fluctuation ~0.3 nm observed between 245–265 residues in the reference complex, furthermore 120–130 and 340–350 residues in Grazoprevir. In the furin-GS-441524 complex, the highest fluctuations ~0.2 nm and ~0.15 were recorded near 345–370 and 120–130 residues respectively. These described major reductions in scores of particular residues propose their direct contribution in enhancing the overall stability of the FDA drug-bound furin complex in divergence to reference complex. In Fig. 6d, the radius of gyration (Rg) graph illustrates that the circulation of atoms of
a protein around its axis in all the complexes is stable. The Rg of furin- GS-441524 complex is ~ 2.2 to ~ 2.22 nm which qualifies one to assess the compactness changes of a complex. Similarly, all complexes exhibited a virtuous solvent accessible area graph (Fig. 5c.). Overall analysis spectacles furin in complex with GS-441524 was observed to maintain equilibrium throughout the simulation. Afterward MD simulation, the protein-ligand complex is proximal to the in-vitro environment, hence further wet-lab experiments may qualify the GS-441524 as a furin inhibitor.

4. Discussion

In the present article, we focused on our particular interest in drug repurposing, since already approved drugs can go through shortened clinical trials. As our knowledge is exponentially expanding with constantly emerging “omics” (e.g., transcriptomic, proteomic, metabolomic), data on drug/disease-associated mechanisms could easily be incorporated and analyzed. Our group working in this area has reported through immunoinformatics approach for immunogenic protein from the proteome of Leptospira and a codon-based analytical study for susceptible hosts of Nipah Virus (24), (25). Our work will add to a broad arena towards the analysis of quantitative structure-activity relationships, pharmacophores, homology models, and other molecular modeling approaches like deciding conformational stability (26) as well as network analysis of biological functions, machine learning, and any other analysis tools that include using a computer.

In this study, each molecule is considered as a data point, no labeling was done during submission of data for analysis, but generated descriptors propose the input into the neuron vectors which in turn make the input nodes. The data processing can be categorized into three broad interconnected networks in ANN, the input layer, the hidden layer, and the output layer. The connections in the network were assigned with weighted values which help the learning process and create the nodes in the hidden layers and finally the optimized signal after all the iterations are passed on to the output layer. According to the input given through descriptors of each data point, the molecules, within the different neurons in the network, there will be competitive and cooperative learning iterations by the end of which the winning neuron among the neuron vectors was determined and will be the closest to the molecule or the data point. After the determination of the winning neuron, update it with the weight of the neighboring neurons to place each one in possible closest positions, this process is continued until the best optimization is achieved and the clustered molecules have fallen into the cells of the map (27) (Fig. 1a). Each cluster was manually analyzed to confirm the best ones containing actives together with FDA drugs and have no in-actives. Out of 180 FDA antivirals, 15 predicted by clusters 2 and 4 demonstrated good results in molecular docking studies. However, it is necessary to analyze polypharmacology before we take a possible drug onto any other targets and not that binding affinity and matching properties alone. In recent years researchers are becoming dependent on a computational approach for their study on the prediction of targets, off-targets, and poly-pharmacology of hit compounds. Thus the network pharmacology comprises components functioning in stepwise order “multi drugs to multi-targets and multi-diseases (28).
There are many important host proteins like FCGRIA, LDLR, PNP, PCK1, and TLR 7 that have well-established interactions with the selected drugs. All of them are having more or less connections with various diseases (viral, tumor, etc) positively or negatively. Purine-nucleoside phosphorylase (PNP) is a very important enzyme that catalyzes nucleotide synthesis in the salvage pathway. Inherited deficiency of PNP causes T cell deficiency and abnormal B cell function in some patients as adenosine deaminase activity is affected. The enzyme deficiency was observed in HIV patients (29). FCGRIA is a leucocyte membrane receptor protein capable of binding the Fc domain of IgG with high affinity and signaling for a variety of inflammatory and proinflammatory responses. In human immunodeficiency virus (HIV) infection, it has a role in sepsis and rheumatoid arthritis. FCGRIA has been proposed as an attractive target for immunotherapy by various workers (30). Phosphoenolpyruvate carboxykinase (PEPCK also known as PCK, EC number 4.1.1.32) a cytoplasmic protein catalyzes gluconeogenesis implicating its important role in cancer biology and virology. Anti-tumor treatment based on gluconeogenesis has become an advanced concept nowadays. According to a recent report, gluconeogenic pathway inhibition of PCK1 enhanced HBV replication (31). Several repeated amino acid sequences are present in the ligand-binding domain for LDL in LDLR (low-density lipoprotein receptor). Lipoprotein receptor-mediated HCV entry into the host cell is also well documented.

We target furin as part of the treatment of infectious diseases by making sure that the other targets plotted here will not get disturbed. But if we take the drugs presently acting on the 5 targets of human (FCGRIA, LDLR, PNP, PCK1, and TLR 7) and repurpose them on furin, their existing network of interactions will be disrupted; All the four drugs which are known to interact with these targets namely Porfimer Sodium, Isatorbine, 5’-Guanylyl methylene bisphosphonate, and Didanosine, are not to be carried further to repurpose without detailed studies. We strongly recommend collecting detailed data regarding the effect on furin from the patients who take these medicines and the repercussions due to this while infected by viral diseases like SARS-CoV-2. The drugs Entecavir and Valganciclovir also need proper studies before attempting to repurpose as their known current target is human DNA. Although no interactions are available, no inference can be made from this study at present. We collected information from drug bank details about all the drugs under discussion the two best-docked molecules Porfimer Sodium and Isatoribine are presently used, former for esophageal cancer treatment and the latter a compound that elevates levels of interferon-alpha.

After the poly-pharmacology analysis followed by molecular dynamics simulation, we found GS-441524 and Grazoprevir to be the most promising molecules to repurpose against the furin. GS-441524 is the nucleoside, a metabolite of the phosphoramidate prodrug Remdesivir. GS-441524 was initially developed by Gilead Sciences, Inc for the treatment of filovirus infections. It is a 1’-cyano substituted C-nucleoside derived from 4-aza-7, 9-dideazaadenosine. Remdesivir (GS-5734), a prodrug of GS-441524 used for the treatment of Ebola infections (32). Current investigations demonstrate that this molecule has been designed to target the RNA-dependent RNA polymerase (RdRp) of several viruses, including the Ebola virus, MERS-CoV, SARS-CoV, and potentially SARS-CoV-2. The latest study has further confirmed that the direct use of GS-441524 can potently inhibit the replication of SARS-CoV-2 in a mouse model (33). In the present study, another prominent antiviral molecule Grazoprevir screened out as a furin inhibitor.
Grazoprevir is a hepatitis C protease inhibitor given in combination with Elbasvir (under the brand name Zepatier) for the treatment of chronic hepatitis C virus (HCV) infection in adults. Up-to-date examinations unveiled that it has striking anti-COVID activity for the essential proteins involved in viral entry into host cells ACE2 and TMPRSS2\(^{(34)}\). The HCV drugs Simeprevir, Vaniprevir, Paritaprevir, and Grazoprevir in combination with Remdesivir gained the effectiveness of Remdesivir alone by 10-fold \(^{(35)}\). Nguyenla et al (2020), published that Remdesivir has a synergistic effect with Velpatasvir and Elbasvir the HCV drugs \(^{(36)}\). Additionally in combination with HCV partner drugs Sofosbuvir and Grazoprevir promote efficacy, synergistically with Remdesivir in a human lung epithelial cell line infected with a clinical isolate of SARS-CoV-2. Herewith we strongly recommend that Remdesivir analog GS-441524 alone or in combination with Grazoprevir may have a possible inhibitory effect on furin. Nonetheless further wet-lab experiments may shed light on this investigation.

A number of very recent researches are evidencing the importance of GS-441524 as a potent antiviral drug due to its strong binding properties with NSP3 macrodomain and NSP12-NSP8-NSP7 of SARS Cov 2 \(^{(37)}\), \(^{(38)}\). It has also been proven that this molecule has very less retention value inside the host body as most of the residue is eliminated with urine suspecting no side effect \(^{(39)}\). Amazingly, out of the majority of Remdesivir in clinical trial experiments, recovered in urine was found to be in the form of GS-441524; whereas only 10% recovered as remdesivir \(^{(40)}\), \(^{(41)}\). Moreover, no report has shown any host protein interaction except some metabolizing enzymes like CPY450. Our work has achieved the interesting interaction of GS-441524 with furin, though indirectly. Hence results of our work could be a very powerful support to the scientific community to take a final decision towards repurposing GS-441524 as SARS Cov 2 antiviral drug.

5. Conclusion

The present study evidenced that all the 15 FDA-approved antiviral drugs have the potential to inhibit the pro-protein convertase furin. These drugs are presently being successfully used to prevent many diseases (viral, tumor, etc). However, furin inhibiting properties of these drugs in these patients can be investigated to support our results. With proper studies digging deep into the biological/interactome/pharmacokinetics studies all these 15 FDA approved drugs can be repurposed according to the pros and cons of each disease, but we can propose with existing data that Grazoprevir and GS-441524 be repurposed on human furin without developing many complications as per the available information today. These results can open up more options to reduce death tolls in comorbidity conditions during the SARS COV-2 infections.

Declarations

Acknowledgment

The authors are grateful for the support given by the ministry of education and innovation cell for software facilities as part of the drug discovery Hackathon 2020, Govt of India. All authors are grateful to
the “Covid Omics Research Consortium” (CORC) initiated by Dr. Jayashankar Das, especially all members of the CORC repurposing wing for instilling enthusiasm igniting the passion in these tough times.

**Ethical Approval**

This manuscript does not contain any studies with human or animal participants performed by any of the authors.

**Consent to participate:** Not applicable

**Consent to publish:** All authors hereby agree to give their consent for publication

**Authors’ contributions**

DM, MP, and JD conceptualized the project and methodology. DM performed the data collection, data analysis, validation data curation, and prepared the draft of the manuscript. MP, SS, SD, and KD supported the analysis, writing of the manuscript, and scientific discussions. AG supported during the validation, and preparation of the final manuscript. All authors contributed to the manuscript writing. All authors read and approved the final manuscript.

**Funding**

This research did not receive any specific grant from funding agencies in the public, commercial, or not-for-profit sectors.

**Competing interests**

The authors declare that they have no known competing financial interests that could have appeared to influence the work reported in this paper.

**Availability of data and material**

All data are available either in the manuscript and supplementary files or in the public repositories. The code is available in Schrodinger Canvas.

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Figures

(a) Cell Population Analysis of antiviral FDA drugs and bioassay actives of Furin (b) Euclidean distance between the antiviral FDA drugs and bioassay actives of Furin in the property based SOM. (SOM Colour Schema: If the difference is high, then that ensemble is represented in red, and if the distance is low, it is represented in blue. The various colors between blue and red are used to represent the variations among the properties, population, and distance)
Figure 2

(a) Molecular docking score of all molecules from cluster 2 and cluster 4 (b) Glide e-model of all molecules from cluster 2 and cluster 4 (c) Molecular docking score of FDA drugs from cluster 2 and cluster 4 (d) Glide e-model of FDA drugs from cluster 2 and 4
Figure 3

Drug target network analysis for drugs acting on human targets, the red polygons are drugs, red lines are edges connecting their targets, white with labels are their interacting proteins.
Figure 4

Drug target network analysis for viral host interactome: furin with drugs acting on viral targets, GS-441524 having maximum interactions with SARS-COV2 related proteins, the red circles are drugs red lines are edges connecting their targets and it reaches to furin through common targets.
Figure 5

Molecular dynamics simulation analysis of protein-ligand complex of furin-Grazoprevir, furin-reference ligand, and furin-GS-441524: (a) Root means square deviation (RMSD), (b) Root means square fluctuation (RMSF), (c) solvent-accessible surface area (SASA), (d) Radius of gyration (Rg).

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