Simeprevir potently suppresses SARS-CoV-2 replication and synergizes with remdesivir

Authors

Ho Sing Lo^{1,*}, Kenrie Pui Yan Hui^{2,*}, Hei-Ming Lai^{3-5,*}, Khadija Shahed Khan¹, Simranjeet Kaur⁶, Junzhe Huang^{4,5}, Zhongqi Li^{4,5}, Anthony K. N. Chan⁷, Hayley Hei-Yin Cheung⁸, Ka-Chun Ng², John Chi Wang Ho², Yu Wai Chen⁹, Bowen Ma¹, Peter Man-Hin Cheung¹⁰, Donghyuk Shin¹¹, Kaidao Wang¹², Meng-Hsuan Lee¹³, Barbara Selisko¹⁴, Cecilia Eydoux¹⁴, Jean-Claude Guillemot¹⁴, Bruno Canard¹⁴, Kuen-Phon Wu¹³, Po-Huang Liang¹³, Ivan Dikic¹¹, Zhong Zuo¹, Francis K. L. Chan^{4,15}, David S. C. Hui^{4,16}, Vincent C. T. Mok^{4,17}, Kam-Bo Wong⁸, Ho Ko^{3-5,17-19}, Wei Shen Aik⁶, Michael Chi Wai Chan^{2,†}, Wai-Lung Ng^{1,†}

Affiliations

¹School of Pharmacy, Faculty of Medicine, The Chinese University of Hong Kong, Hong Kong ²School of Public Health, Li Ka Shing Faculty of Medicine, The University of Hong Kong, Hong Kong

³Department of Psychiatry, Faculty of Medicine, The Chinese University of Hong Kong, Hong Kong

⁴Department of Medicine and Therapeutics, Faculty of Medicine, The Chinese University of Hong Kong, Hong Kong

⁵Li Ka Shing Institute of Health Sciences, Faculty of Medicine, The Chinese University of Hong Kong, Hong Kong

⁶Department of Chemistry, Faculty of Science, Hong Kong Baptist University, Hong Kong ⁷Department of Systems Biology, Beckman Research Institute, City of Hope, Duarte, CA, USA ⁸School of Life Sciences, Centre for Protein Science and Crystallography, State Key Laboratory of Agrobiotechnology, Faculty of Science, The Chinese University of Hong Kong, Hong Kong ⁹Department of Applied Biology and Chemical Technology and the State Key Laboratory of Chemical Biology and Drug Discovery, The Hong Kong Polytechnic University, Hong Kong ¹⁰School of Public Health, Faculty of Medicine, The Chinese University of Hong Kong, Hong Kong

¹¹Buchmann Institute for Molecular Life Sciences, Goethe University, Frankfurt am Main, Germany

¹²Protein Production Department, GenScript Biotech Corporation, Nanjing, Jiangsu Province, China

¹³Institute of Biological Chemistry, Academia Sinica, Taipei, Taiwan

¹⁴Laboratoire d'Architecture et Fonction des Macromolécules Biologiques, Centre National de la Recherche Scientifique, Université d'Aix-Marseille, Marseille, France

¹⁵Institute of Digestive Disease, Faculty of Medicine, The Chinese University of Hong Kong, Hong Kong

¹⁶Stanley Ho Center for Emerging Infectious Diseases, Faculty of Medicine, The Chinese University of Hong Kong, Hong Kong

 ¹⁷Gerald Choa Neuroscience Centre, Margaret K. L. Cheung Research Centre for Management of Parkinsonism, Faculty of Medicine, The Chinese University of Hong Kong, Hong Kong
 ¹⁸School of Biomedical Sciences, Faculty of Medicine, The Chinese University of Hong Kong, Hong Kong

¹⁹Peter Hung Pain Research Institute, Faculty of Medicine, The Chinese University of Hong Kong, Hong Kong

^{*}These authors contributed equally to this work

[†]To whom correspondence should be addressed: M.C.W.C. (<u>mchan@hku.hk</u>) or W.L.N. (<u>billyng@cuhk.edu.hk</u>)

One Sentence Summary

Discovery of simeprevir as a potent suppressor of SARS-CoV-2 viral replication that synergizes with remdesivir.

Abstract

The outbreak of coronavirus disease 2019 (COVID-19), caused by the severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2), is a global threat to human health. Using a multidisciplinary approach, we identified and validated the hepatitis C virus (HCV) protease inhibitor simeprevir as an especially promising repurposable drug for treating COVID-19. Simeprevir potently reduces SARS-CoV-2 viral load by multiple orders of magnitude and synergizes with remdesivir *in vitro*. Mechanistically, we showed that simeprevir inhibits the main protease (M^{pro}) and unexpectedly the RNA-dependent RNA polymerase (RdRp). Our results thus reveal the viral protein targets of simeprevir, and provide preclinical rationale for the combination of simeprevir and remdesivir for the pharmacological management of COVID-19 patients.

Keywords

Simeprevir, Remdesivir, SARS-CoV-2, COVID-19, Drug repurposing, Drug synergism

Introduction

The recent outbreak of infection by the novel betacoronavirus severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) has spread to almost all countries and claimed more than 760,000 lives worldwide (WHO situation report 209, August 16, 2020). Alarming features of COVID-19 include a high risk of clustered outbreak both in community and nosocomial settings, and up to one-fifth severe/critically ill proportion of symptomatic inpatients reported^{1–4}. Furthermore, a significant proportion of infected individuals are asymptomatic, substantially delaying their diagnoses, hence facilitating the widespread dissemination of COVID-19⁵. With a dire need for effective therapeutics that can reduce both clinical severity and viral shedding, numerous antiviral candidates

have been under clinical trials or in compassionate use for the treatment of SARS-CoV-2 infection⁶.

Several antivirals under study are hypothesized or proven to target the key mediator of a specific step in the SARS-CoV-2 viral replication cycle. For instance, lopinavir/ritonavir (LPV/r) and danoprevir have been proposed to inhibit the SARS-CoV-2 main protease (M^{pro}, also called 3CL^{Pro}) needed for the maturation of multiple viral proteins; chloroquine (CQ) / hydroxychloroquine (HCQ) [alone or combined with azithromycin (AZ)] may abrogate viral replication by inhibiting endosomal acidification crucial for viral entry^{7,8}; nucleoside analogues such as remdesivir, ribavirin, favipiravir and EIDD-2801 likely inhibit the SARS-CoV-2 nsp12 RNA-dependent RNA polymerase (RdRp) and/or induce lethal mutations during viral RNA replication^{9–11}. Unfortunately, on the clinical aspect, LPV/r failed to demonstrate clinical benefits in well-powered randomized controlled trials (RCTs), while HCQ and/or AZ also failed to demonstrate benefits in observational studies¹²⁻¹⁴. Meanwhile, LPV/r, CQ/HCQ and AZ may even increase the incidence of adverse events^{14–16}. Although remdesivir is widely considered as one of the most promising candidates, latest RCTs only revealed marginal shortening of disease duration in patients treated¹⁷. Therefore, further efforts are required to search for more potent, readily repurposable therapeutic agents for SARS-CoV-2 infection, either as sole therapy or in combination with other drugs to enhance their efficacy.

Ideally, the candidate drugs need to be readily available as intravenous and/or oral formulation(s), possess favourable pharmacokinetics properties as anti-infectives, and do not cause adverse events during the treatment of SARS-CoV-2 infection (e.g. non-specific immunosuppression, arrhythmia or respiratory side effects). Two complementary approaches have been adopted to identify novel drugs or compounds that can suppress SARS-CoV-2 replication. One approach relies on *in vitro* profiling of the antiviral efficacy of up to thousands of compounds in early clinical development, or drugs already approved by the U.S. Food and Drug Administration (FDA)^{18–22}. On the other hand, as the crystal structure of the M^{pro 23,24}, papain-like protease (PL^{pro})²⁵ and the cryo-EM structure of the nsp12-nsp7-nsp8 RdRp complex^{11,26} of the SARS-CoV-2 virus became available, the structure-based development of their specific inhibitors becomes feasible. Structure-aided screening will enable the discovery of novel compounds as highly potent inhibitors²⁷ as well as the repurposing of readily available drugs as anti-CoV agents for fast-track clinical trials.

Here we report our results regarding the discovery of FDA-approved drugs potentially active against the SARS-CoV-2. *In vitro* experiments led to the identification of simeprevir, a hepatitis C virus (HCV) NS3A/4 protease inhibitor²⁸, as a potent inhibitor of SARS-CoV-2 replication (**Fig. 1A**). Importantly, simeprevir acts synergistically with remdesivir, whereby the effective dose of remdesivir could be lowered by multiple-fold

by simeprevir at physiologically feasible concentrations. Interestingly, biochemical and molecular characterizations revealed that simeprevir inhibits both the M^{pro} protease and RdRp polymerase activities. This unexpected anti-SARS-CoV-2 mechanism of simeprevir provides hints on novel antiviral strategies.

Results

A prioritized screening identifies simeprevir as a potent suppressor of SARS-CoV-2 replication in a cellular infection model

Given our goal of identifying immediately usable and practical drugs against SARS-CoV-2, we prioritized a list of repurposing drug candidates for *in vitro* testing based on joint considerations on safety, pharmacokinetics, drug formulation availability, and feasibility of rapidly conducting clinical trials (**Table S1**). We focused on FDA-approved antivirals (including simeprevir, saquinavir, daclatasvir, ribavirin, sofosbuvir and zidovudine), and drugs whose primary indication was not antiviral but had reported antiviral activity (including bromocriptine and atovaquone). Remdesivir was also tested for comparison of efficacy and as a positive control.

In a Vero E6 cellular infection model, we found the macrocyclic HCV protease inhibitor simeprevir as the only prioritized drug candidate that showed potent suppression of SARS-CoV-2 replication in the $\leq 10 \ \mu$ M range (**Fig. 1B**). More detailed dose-response characterization found that simeprevir has a potency comparable to remdesivir (**Fig. 1C, D**). The half-maximal effective concentration (EC50) of simeprevir was determined to be 4.08 μ M, while the 50% cytotoxicity concentration (CC50) was 19.33 μ M (**Fig. 1C, D**). In a physiologically relevant human lung epithelial cell model, ACE2-expressing A549 cells (A549-ACE2) infected with SARS-CoV-2, we also observed the strong antiviral effect of simeprevir²⁹ (**Supplementary Fig. 1**). The cytotoxicity data are also in line with the reported *in vitro* and *in vivo* safety pharmacological profiling using human cell lines, genotoxicity assays, and animal models^{28,30}. These data suggest that a desirable therapeutic window exists for the suppression of SARS-CoV-2 replication with simeprevir. bioRxiv preprint doi: https://doi.org/10.1101/2020.05.26.116020; this version posted September 3, 2020. The copyright holder for this preprint (which was not certified by peer review) is the author/funder. All rights reserved. No reuse allowed without permission.



Fig. 1 Repurposing FDA-approved Drugs for SARS-CoV-2 through cellular screening. (A) Summary of methodology used in this paper. (B) Screening for FDA-approved small molecule therapeutics for activities in suppressing SARS-CoV-2 replication in Vero E6 cells. Dose-response curves in the suppression of SARS-CoV-2 replication in Vero E6 cells and cytotoxicity for simeprevir (C) and remdesivir (D) are shown. Data points in all plots represent mean \pm S.E.M.. For all data points, n = 3 replicates.

Simeprevir potentiates the suppression of SARS-CoV-2 replication by remdesivir

While simeprevir is a potential candidate for clinical use alone, we hypothesized that it may also have a synergistic effect with remdesivir, thereby mitigating its reported adverse effects, improving its efficacy, and broadening its applicability¹⁷. Indeed, combining simeprevir and remdesivir at various concentrations apparently provided much greater suppression of SARS-CoV-2 replication than remdesivir alone, while they did not synergize to increase cytotoxicity (**Fig. 2A**). Importantly, such effects were not merely additive, as the excess over Bliss score suggested synergism at 3.3 μ M simeprevir and 1.1 – 10 μ M remdesivir in suppressing SARS-CoV-2 replication (**Fig. 2B**).

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Fig. 2 (A) Viral replication-suppression efficacies of different combinations of simeprevir and remdesivir concentrations. The numbers after S (simeprevir) and R (remdesivir) indicate the respective drug concentrations in μ M. Data points in all plots represent mean ± S.E.M.. For all data points, *n* = 3 replicates. **(B)** Bliss score analyses of synergism. Left panel: Diagram showing 12 combinations of simeprevir and remdesivir and their respective percentage inhibition (% inhibition, color-coded) of SARS-CoV-2 replication in Vero E6 cells compared to DMSO controls. Right panel: Excess over Bliss score (Δ Bliss, color-coded) of different drug combinations. A positive and negative number indicates a likely synergistic and antagonistic effect, respectively, while a zero value indicates independence of action.

Simeprevir weakly inhibits the *M*^{pro} and RdRp but does not inhibit PL^{pro} at physiologically feasible concentrations

The desirable anti-SARS-CoV-2 effect of simeprevir prompted us to determine its mechanism of action. Given that simeprevir is an HCV NS3/4A protease inhibitor, we first investigated its inhibitory activity against SARS-CoV-2 M^{pro} and PL^{pro} using biochemical assays^{31,32} (**Fig. 3**). We found inhibition of M^{pro} by simeprevir with half-maximal inhibitory concentration (IC₅₀) of 9.6 ± 2.3 µM (**Fig. 3A**), two times higher than the EC₅₀ determined from our cell-based assay. The substrate cleavage was further verified with SDS-PAGE (**Supplementary Fig. 3**). Docking simeprevir against the apo

protein crystal structure of SARS-CoV-2 M^{pro} suggested a putative binding mode with a score of -9.9 kcal mol⁻¹ (**Supplementary Fig. 4**). This binding mode is consistent with a recent docking study using a homology model of SARS-CoV-2 M^{pro 33}. On the other hand, no inhibition of PL^{pro} activity was observed at physiologically feasible concentrations of simeprevir, with either ISG15 or ubiquitin as substrate (**Fig. 3B**, **Supplementary Fig. 5**, **6**).

We speculated that the weak inhibition of M^{pro} protease activity by simeprevir could not fully account for its antiviral effect towards SARS-CoV-2. To identify additional target(s), we next docked simeprevir alongside several nucleoside analogues (remdesivir, ribavirin, and favipiravir) against the motif F active site of the cryo-EM structure of the SARS-CoV-2 nsp12 RdRp (**Supplementary Fig. 7A**). Interestingly, the docking results revealed that simeprevir had a higher binding score than the nucleoside analogues (**Supplementary Fig. 7B**). To test this experimentally, we established and performed RdRp primer extension assays using recombinant nsp12, nsp7, and nsp8 of SARS-CoV³⁴. Intriguingly, simeprevir showed low micromolar-range inhibition towards SARS-CoV RdRp as validated by both a gel-based assay (**Supplementary Fig. 3C**). Collectively, the assay data suggested that simeprevir inhibits the enzymatic activities of both M^{pro} and RdRp but not PL^{pro}.



Fig. 3 Simeprevir weakly inhibits M^{pro} **and RdRp.** Assay scheme and enzyme activity of main protease (M^{pro}), papain-like protease (PL^{pro}) and RNA-dependent RNA polymerase (RdRp) under various concentrations of simeprevir. **(A)** For M^{pro} , a CFP-YFP conjugate with a M^{pro} cleavage site linker is utilized, where relative activity is determined by the residual FRET efficiency after cleavage. **(B)** For PL^{pro} , rhodamine-conjugated ISG15 is used as a substrate for the enzyme, whose relative activity is determined by release of the fluorophore. **(C)** For RdRp, an extension assay based on the dsRNA-binding property of the intercalating agent Picogreen was established. Data points in all plots represent mean ± S.E.M.. For all data points, n = 3 replicates.

RNA Sequencing identifies significant downregulation of viral defense responses upon the treatment of simeprevir

While inhibition of viral targets seems to be a primary mechanism of action of simeprevir, the weak inhibitory effects observed in biochemical assays as well as its cytotoxicity suggest the possibility of additional host-mediated antiviral response. To further elucidate the antiviral mechanism of simeprevir, we next performed RNA sequencing on Vero E6 cells to reveal the transcriptomic changes upon drug treatment (Fig. 4A). In line with the literature, SARS-CoV-2 infection induced type I interferon and chemokine response (Supplementary Fig. 9)^{35,36}. In mock-infected cells, simeprevir treatment (at 1.1 µM or 3.3 µM) did not induce any significant changes of differentially expressed genes (DEGs); while in SARS-CoV-2-infected cells, a small number of DEGs was observed (Fig. 4B). Gene set enrichment analyses (GSEA) of infected cells using Reactome gene sets revealed significant positive enrichment of 93 gene sets in the simeprevir-treated samples, including histone lysine/arginine methylation, histone demethylation, and cell cycle control (Fig. 4C, Supplementary table 2). On the other hand, gene sets with the gene ontology (GO) terms "defense response to virus" and "response to type I interferon" were negatively enriched, suggesting overall downregulation of these gene sets (Fig. 4D). In the latter set, green monkey orthologs of crucial human innate immune-related genes (e.g. IFIT1-3, USP18) and interferonstimulated genes (e.g. ISG15) were downregulated with simeprevir treatment in a dosedependent manner (Fig. 4E). Similarly, the downregulation of some of these genes as well as the proinflammatory cytokines IL-6 and interferon IFNL1 were also observed with remdesivir treatment (Fig. 4F).



Fig. 4 RNA-seq analysis of simeprevir-mediated host response and antiviral activity. (A) Schematic representation of RNA-seq sample preparation. Treatment sequence and incubation time of simeprevir and SARS-CoV-2 was indicated with arrows and legends. **(B)** Venn diagrams showing differentially expressed genes (DEGs) comparing simeprevir-treated (1.1µM or 3.3μ M), infected and mock-infected samples. **(C)** Bubble plot of top 20 hits of positively enriched reactome gene sets under simeprevir treatment using gene set analysis (GSEA). Enriched gene sets were filtered with criteria false discovery rate (FDR) q-value < 0.25 and nominal p-value < 0.05 before ranked with their normalized enrichment scores (NES). **(D)** Enrichment plots of GSEA results using gene ontology (GO) gene sets. **(E)** Clustered heatmap showing the row-normalized expression level of genes belonging to GO term "response to type I interferon". **(F)** Bar charts showing normalized counts from RNA-seq data of five genes involved in antiviral response. Data points in this plot represent mean ± S.E.M. For all data points, *n* = 3 replicates.

Discussion

The novel coronavirus SARS-CoV-2 has gone from an emerging infection to a global pandemic with its high transmissibility. As human activities are becoming more aggressive and damaging to nature, future coronavirus pandemics are bound to happen. It is therefore essential to reduce the casualties by effective pharmacological

management. Our study has successfully identified the readily repurposable, clinically practical antiviral simeprevir that could target SARS-CoV-2. Specifically, we found up to fourth-order suppression of viral genome copies by simeprevir at $\leq 10 \ \mu$ M in cell-based viral replication assays - a concentration that is expected to be attainable in human lung tissues with ≥ 150 mg daily dosing based on available pharmacokinetic data^{37,38}.

In addition, we discovered that simeprevir can synergize with remdesivir in inhibiting SARS-CoV-2 replication in a cellular model, potentially allowing lower doses of both drugs to be used to treat COVID-19. In a global pandemic with patients having diverse clinical characteristics, providing additional therapeutic options to remdesivir will be important to treat those who are intolerant or not responding to the drug ¹⁷, which can easily amount to tens of thousands of patients. As there is only one confirmed and approved therapy for COVID-19, a potentially repurposable drug can be rapidly tested in animal models before clinical trials to prepare for supply shortages or when remdesivir-resistant mutations arise. Combination treatment, such as simeprevir-remdesivir, may also help to reduce the dose required to alleviate side effects.

We note, however, there are also several limitations of simeprevir and the proposed simeprevir-remdesivir combination. Simeprevir requires dose adjustments in patients with Child-Pugh Class B or C cirrhosis, as well as in patients with East Asian ancestry³⁷. In addition, simeprevir has been taken off the market since 2018 due to the emergence of next-generation HCV protease inhibitors, hence its supply may not be ramped up easily. Noteworthily, simeprevir is metabolized by the CYP3A4 enzyme with saturable kinetics³⁷ while remdesivir itself is not only a substrate of CYP3A4 but also a CYP3A4 inhibitor. Whether such theoretical pharmacokinetic interaction will exacerbate liver toxicity or provide additional pharmacokinetic synergy (in addition to pharmacodynamic synergy) *in vivo* remains to be tested.

Mechanistically, we found that simeprevir suppresses SARS-CoV-2 replication by targeting at least two viral proteins – it weakly inhibits M^{pro} at ~10 µM and unexpectedly inhibits RdRp at ~5 µM. The potency towards M^{pro} is consistent with the IC₅₀ of ~13.7 µM as determined in a parallel study³⁹. Our gel-based assay (**Supplementary Fig. 9**) suggested that simeprevir interferes with RNA-binding of RdRp because less probe was extended but to full length. This is also supported by the *in silico* docking results, in which simeprevir is docked to a highly conserved RNA binding site showing no amino acid polymorphism between SARS-CoV and SARS-CoV-2 (**Supplementary Fig. 8A**). This putative binding mode hints that simeprevir might block the RNA binding site while remdesivir might target the nucleoside entry site, potentially resulting in a synergistic effect. Importantly, the high similarity in sequence (96% identity) and structure between SARS-CoV-2 and SARS-CoV-2 and SARS-CoV-2 and sa a broad-spectrum antiviral in the *Coronaviridae* family.

Furthermore, the discrepancy between RdRp and M^{pro} inhibitory potency versus *in vitro* inhibitory potency of SARS-CoV-2 replication suggested additional mechanism(s) of action of simeprevir. Our RNA-seq and GSEA analysis revealed several molecular pathways that warrant future investigation. A possible direction is immune modulation *via* epigenetic regulations, which could mediate viral infection (*via* SWI/SNF chromatin remodeling complex and histone H3.3 complex)⁴⁰, interferon-induced antiviral response (*via* H3K79 methylation)⁴¹, and host immune evasion (*via* alteration of DNA methylome)⁴². Whether the downregulation of type I IFN-related genes stems directly from simeprevir's action or a reduction in viral load remains an open question. Collectively, simeprevir targets two viral proteins but may also act on the host proteins to suppress SARS-CoV-2 replication.

Given that simeprevir is originally a non-nucleoside antiviral targeting HCV protease, its inhibition towards RdRp is largely unexpected and represents a novel mechanism of action. Simeprevir thus holds promise to be a lead compound for the future development of dual inhibitors of M^{pro} and RdRp. It should be noted that the potencies of M^{pro} and RdRp inhibition may not entirely account for the strong suppression of SARS-CoV-2 viral replication. Therefore, further investigation of the mechanism of action of simeprevir may uncover new druggable targets for inhibiting SARS-CoV-2 replication.

Materials and Methods

Chemicals and reagents

Bromocriptine mesylate (BD118791), saquinavir (BD150839), bictegravir (BD767657), atovaquone (BD114807) and asunaprevir (BD626409) were purchased from BLD Pharmatech (Shanghai, China). Entecavir (HY-13623), zidovudine (HY-17413), sofosbuvir (HY-15005), daclatasvir (HY-10466), simeprevir (HY-10241), remdesivir (HY-104077) and remdesivir triphosphate sodium (HY-126303A) were purchased from MedChemExpress (Monmouth Junction, NJ). Drug stocks were made with DMSO.

In vitro SARS-CoV-2 antiviral tests

SARS-CoV-2 virus (BetaCoV/Hong Kong/VM20001061/2020, SCoV2) was isolated from the nasopharyngeal aspirate and throat swab of a COVID-19 patient in Hong Kong using Vero E6 cells (ATCC CRL-1586). Vero E6 or A549-ACE2 cells were infected with SCoV2 at a multiplicity of infection (MOI) of 0.05 or 0.5, respectively, in the presence of varying concentrations and/or combinations of the test drugs. DMSO as the vehicle was used as a negative control. Antiviral activities were evaluated by quantification of SARS-CoV-2 ORF1b copy number in the culture supernatant by using

quantitative real-time RT-PCR (qPCR) at 48 h post-infection with specific primers targeting the SARS-CoV-2 ORF1b⁴³.

In vitro drug cytotoxicity assays

In vitro cytotoxicity of the tested drugs was evaluated using thiazolyl blue tetrazolium bromide (MTT, Sigma-Aldrich)-based cell viability assays. Vero E6 cells were seeded onto 48-well plates and treated with indicated concentrations of simeprevir and/or remdesivir for 48 h. Treated cells were incubated with DMEM supplemented with 0.15 mg/mL MTT for 2 h, and formazan crystal products were dissolved with DMSO. Cell viability was quantified with colorimetric absorbance at 590 nm.

Molecular docking simulations

Three-dimensional representations of chemical structures were extracted from the ZINC15 database (http://zinc15.docking.org)⁴⁴, with the application of three selection filters — Protomers, Anodyne, and Ref. ZINC15 subset DrugBank FDA (http://zinc15.docking.org./catalogs/dbfda/) were downloaded as the mol2 file format. The molecular structures were then converted to the pdbqt format (the input file format for AutoDock Vina) using MGLTools2-1.1 RC1 (sub-program "prepare_ligand") (http://adfr.scripps.edu/versions/1.1/downloads.html). AutoDock Vina v1.1.2 was employed to perform docking experiments⁴⁵. Docking of simeprevir on SARS-CoV-2 M^{pro} was performed with the target structure based on an apo protein crystal structure (PDB ID: 6YB7); the A:B dimer was generated by crystallographic symmetry. Docking was run with the substrate-binding residues set to be flexible. Docking of simeprevir and other active triphosphate forms of nucleotide analogues was performed against the nsp12 portion of the SARS-CoV-2 nsp12-nsp7-nsp8 complex cryo-EM structure (PDB ID: 6M71).

Expression and purification of M^{pro} and its substrate for FRET assay

The sequence of SARS-CoV-2 M^{pro} was obtained from GenBank (accession number: YP_009725301), codon-optimized, and ordered from GenScript. A C-terminal hexahistidine-maltose binding protein (His₆-MBP) tag with two in-between Factor Xa digestion sites were inserted. Expression and purification of SARS-CoV-2 M^{pro} was then performed as described for SARS-CoV M^{pro 31}. The protein substrate, where the cleavage sequence "TSAVLQSGFRKM" of M^{pro} was inserted between a cyan fluorescent protein and a yellow fluorescent protein, was expressed and purified as described³¹.

In vitro M^{pro} inhibition assay

The inhibition assay was based on fluorescence resonance energy transfer (FRET) using a fluorescent protein-based substrate previously developed for SARS-CoV M^{pro 31,46}. 0.1 µM of purified SARS-CoV-2 M^{pro} was pre-incubated with 0 - 250 µM simeprevir in 20 mM HEPES pH 6.5, 120 mM NaCl, 0.4 mM EDTA, 4 mM DTT for 30 min before the reaction was initiated by addition of 10 µM protein substrate³². Protease activity was followed at 25 °C by FRET with excitation and emission wavelengths of 430 nm and 530 nm, respectively, using a multi-plate reader as previously described^{31,46}. Reduction of fluorescence at 530 nm was fitted to a single exponential decay to obtain the observed rate constant (k_{obs}). Relative activity of M^{pro} was defined as the ratio of k_{obs} with inhibitors to that without. The relative IC₅₀ value of simeprevir was determined by fitting the relative activity at different inhibitor concentration to a four-parameter logistics equation.

Expression and purification of SARS-CoV nsp7L8, nsp12

The fusion protein nsp7-nsp8 (nsp7L8) was generated by inserting a GSGSGS linker sequence between the nsp7 and nsp8 coding sequences³⁴. The nsp7L8, nsp8 and nsp12 were produced and purified independently as described⁴⁷. The complex, referred to as the replication/transcription complex (RTC), was reconstituted with a 1:3:3 ratio of nsp12:nsp7L8:nsp8.

In vitro RdRp inhibition assay - fluorescence-based

The assay was performed as previously described⁴⁸. The compound concentration leading to a 50% inhibition of RTC-mediated RNA synthesis was determined as previously described. Briefly, poly(A) template and the SARS-CoV RTC was incubated 5 min at room temperature and then added to increasing concentration of compound. Reaction was started by adding UTP and incubated 20 min at 30°C. Reaction assays were stopped by the addition of 20 µl EDTA 100 mM. Positive and negative controls consisted of a reaction mix with 5% DMSO (final concentration) or EDTA (100 mM) instead of compounds, respectively. Picogreen® fluorescent reagent was diluted to 1/800 final in TE buffer according to the data manufacturer and aliquots were distributed into each well of the plate. The plate was incubated for 5 min in the dark at room temperature and the fluorescence signal was then read at 480 nm (excitation) and 530 nm (emission) using a TecanSafire2 microplate reader. IC_{50} was determined using the following equation: % of active enzyme = $100/(1+(1)^2/IC_{50})$, where I is the concentration of inhibitor and 100% of activity is the fluorescence intensity without inhibitor. IC_{50} was determined from curve-fitting using the GraphPad Prism 8.

In vitro RdRp inhibition assay - gel-based

Enzyme mix (10 μ M nsp12, 30 μ M nsp7L8, 30 μ M nsp8) in complex buffer (25 mM HEPES pH 7.5, 150 NaCl, 5 mM TCEP, 5 mM MgCl₂) was incubated for 10 min on

ice and then diluted with reaction buffer (20 mM HEPES pH 7.5, 50 mM NaCl, 5 mM MgCl₂) to 2 μ M nsp12 (6 μ M nsp7L8 and nsp8) to a final volume of 10 μ l. The resulting enzyme complex was mixed with the 10 μ l of 0.8 μ M primer/ template (P/T) carrying a Cy5 fluorescent label at the 5' end (P:Cy5-GUC AUU CUC C, T: UAG CUU CUU AGG AGA AUG AC) in reaction buffer, and incubated at 30°C for 10 min. Inhibitor was added in 2 μ l to the elongation complex and reactions were immediately started with 18 μ l of NTP mix in the reaction buffer. Final concentrations in the reactions were 0.5 μ M nsp12 (1.5 μ M nsp7L8 and nsp8), 0.2 μ M P/T, 50 μ M NTPs and the given concentrations of inhibitors. Samples of 8 μ l were taken at given time points and mixed with 40 μ l of formamide containing 10 mM EDTA. Ten- μ l samples were analyzed by denaturing PAGE (20 % acrylamide, 7 M urea, TBE buffer); and product profiles visualized by a fluorescence imager (Amersham Typhoon). Quantification of product bands and analysis were performed using ImageQuant and Excel.

In vitro PL^{pro} inhibition assay - fluorescence-based

The purification and assay of PL^{pro} activity was adapted from as previously described²⁵. Briefly, a ISG15-C-term protein tagged with rhodamine was used as a substrate for the enzymatic assay. Because of the solubility of Simeprevir, we used an optimized reaction buffer (5% DMSO, 15% PEG300, 50 mM Tris-HCI (pH 7.5), 50 mM NaCl, 5 mM DTT). 5 uL of solution containing 0 - 2 mM of Simeprevir and 2 μ M of ISG15c-rhodamine were aliquoted into a 384-well plate. Reaction was initiated by addition of 5 μ L of 40 nM PL^{pro} to the well. Initial velocities of rhodamine release (36 - 240 seconds) were normalized against DMSO control. Reactions were conducted for 12 minutes with monitoring of fluorescence intensity at 485/520 nm using a microplate reader (PHERAstar FSX, BMG Labtech). The same experiment was repeated with ubiquitin-rhodamine substrate.

In vitro PL^{pro} inhibition assay - Gel-based

All proteins used for ubiquitination and PL^{pro} biochemical assays were expressed in *E. coli* RIL cells with 0.6 mM IPTG over-induction at 16 °C. Rsp5 WW3-HECT with 2 mutations (Q808M, E809L) at the C-terminus was expressed using GST-fusion affinity tag followed by TEV protease digestion and purification by size exclusion chromatography. His-tagged PL^{pro} , UBA1, UBCH7, ubiquitin K63R with extra cysteine at the N-terminus were also expressed and purified similarly, except hexahistidine tag used. Ubiquitin was further cross-linked with fluorescein-5-maleimide (Anaspec, Fremont, CA, US) and the poly-ubiquitination sample was generated following previous protocol ⁴⁹. Deubiquitination assays using PL^{pro} (SARS-CoV-2) were carried out at 37 °C for 10 minutes using 1-100 μ M Simeprevir and 1 μ M PL^{pro} . Final DMSO concentration of each reaction is 2%. The reaction was quenched by SDS sample buffer and analyzed by 4-20% SDS-gel (GenScript, Piscataway, NJ, US). Fluorescent ubiquitin signals were imaged using Thermo iBright exposed for 750 ms.

Sample preparation for RNA-seq

Approximately 4 x 10⁵ Vero E6 cells were seeded onto each well of 12-well plates, in DMEM supplemented with 2% FBS, 4.5 g/L D-glucose, 4 mM L glutamine, 25 mM HEPES and 1% penicillin/streptomycin. Infections of SARS-CoV-2 were performed at a MOI of 1 for 24 hours, followed by drug treatment or 2% DMSO in triplicates. Uninfected cells were also treated with the same concentrations of drug or DMSO in triplicates. After 24 hours of drug treatment, total RNA from infected cells and uninfected cells were extracted using Qiagen RNeasy mini kit (Qiagen) following the manufacturer's instructions. Then, we performed pair-end sequencing on a NovaSeq 6000 PE150 platform and generated 20 million reads per sample at Novogene Bioinformatics Institute (Novogene, Beijing, China).

Bioinformatic Analyses

The raw reads quality was checked by the FastQC (0.11.7) and aligned to ChlSab1.1 (Chlorocebus sabaeus) reference genome by the <u>STAR</u> (2.5.0a) with default parameters. The count matrix was generated by the <u>featureCounts</u> (as a component of Subread package 2.0.1) program. Differentially expressed genes (DEGs) were calculated by <u>DESeq2</u> package (1.26.0) under R environment (3.6.1) and characterized for each sample (|L2FC| > 1, p-adjusted-value < 0.05). Gene set enrichment analysis (GSEA) was performed as previously described using normalized counts with orthology gene converting to human gene by <u>biomaRt</u> package (2.42.1)⁵⁰. Bubble plots and heatmaps were generated using *ggplot* package and *heatmap()* function in R respectively.

Statistical Analyses

For in vitro experiments, four-parameter dose response curve fitting was performed with constraints: Top = 1, $IC_{50} > 0$ using GraphPad Prism 8.

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Competing interests: CUHK and HKU have filed a US provisional patent application based on the finding of this manuscript. W.L.N., M.C.W.C., K.P.Y.H., H.S.L., K.S.K., H.K. and H.M.L. are inventors of the patent. F.K.L.C. has served as a consultant to Eisai, Pfizer, Takeda and Otsuka, and has been paid lecture fees by Eisai, Pfizer, AstraZeneca and Takeda.

Data and materials availability: The raw data and data analysis codes used in this project are available from the authors upon reasonable request.

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Supplementary Fig. 1 Dose-response curves in the suppression of SARS-CoV-2 replication in A549-ACE2 cells and cytotoxicity for simeprevir (Left) and remdesivir (Right) are shown. Data points in all plots represent mean \pm S.E.M.. For all data points, n = 3 replicates.

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Supplementary Fig. 2 SDS-PAGE analysis of recombinant SARS-CoV-2 M^{pro}.

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Supplementary Fig. 3 (A) The addition of SARS-CoV-2 M^{pro} led to the cleavage of the substrate, causing detectable decline in FRET signal with 430-nm excitation and 530-nm emission. **(B)** Confirmation of substrate cleavage by M^{pro} using SDS-PAGE. **(C)** The addition of simeprevir at varying concentrations attenuated the rate of FRET substrate cleavage by M^{pro}.



Supplementary Fig. 4 Docking of simeprevir on SARS-CoV-2 M^{pro} (performed with AutoDock Vina version 1.1.2). The M^{pro} structure was based on an apo protein crystal structure (PDB ID: 6YB7); the A:B dimer was generated by crystallographic symmetry. Docking was run with the substrate-binding residues set to be flexible; and a 30 x 30 x 30 Å³ search box centered near the side-chain Nɛ2 atom of His163. The top docking mode shown here scored -9.9 kcal mol⁻¹. The protein is shown as a semi-transparent molecular surface encasing its stick model with the catalytic residues His41 and Cys145 in green. The ligand is shown as a stick model with oxygen atoms in red; nitrogen atoms in blue, sulfur atoms in yellow and carbon atoms in white.



Supplementary Fig. 5 Simeprevir did not inhibit PL^{pro} activity in a repeated PL^{pro} cleavage assay using a different substrate ubiquitin-rhodamine.



Supplementary Fig. 6 Inhibitor deubiquitination assay of PL^{pro} shows PL^{pro} (SARS-CoV-2) actively cleaves poly-ubiquitinated substrate into di-ubiquitin or longer ubiquitin chain. While simeprevir was added in reaction solution, the activity of PL^{pro} was not profoundly inhibited compared to control experiments (no inhibitor and no inhibitor with 2% DMSO).

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Supplementary Fig. 7 (A) Binding mode of simeprevir and other inhibitors against SARS-CoV-2 nsp12 (PDB ID: 6M71). Motif F is highlighted in yellow. **(B)** Docking scores of drug candidates against nsp12.



Supplementary Fig. 8 (A) Gel-based assay scheme for SARS-CoV-RdRp. Two partially complementary RNA probes are extended by the enzyme, and the extent of extension is visualized by detecting the Cy5 labeling of the shorter probe P. **(B)** Time-dependent elongation of probe P. Fraction of extension is determined by densitometry of extended product versus total RNA. **(C)** RNA-PAGE imaged with Cy5 mode.

DMSO infected vs. DMSO uninfected

Gene Set	NES	FDR q-val
RESPONSE_TO_TYPE_I_INTERFERON	1.88	0.101
DEFENSE_RESPONSE_TO_VIRUS	1.51	0.101
RESPONSE_TO_CHEMOKINE	1.36	0.133



Supplementary Fig. 9 Gene set enrichment analysis of SARS-CoV-2 infection. Relevant statistics and enrichment plots for three gene sets in response to virus infection are shown.

Drug candidate	Approved indication(s)	Common/notable adverse effects	Drug formulation(s) available	Contraindications (except known hypersensitivity to the drug)
Simeprevir	Chronic HCV infection	Photosensitivity, rash, fatigue, myalgia, dyspnea	Oral	None
Daclatasvir	Chronic HCV infection	Fatigue, nausea, anemia, diarrhea	Oral	None
Saquinavir	HIV infection	Nausea, diarrhea, QT prolongation	Oral	Prolonged QT interval, Patients at risk/having complete AV block
Bromocriptine	Hyperprolactinemia, pituitary prolactinoma	Constipation, dizziness, nausea, fatigue, orthostatic hypotension, vasospasm, abdominal pain	Oral	Uncontrolled hypertension, psychosis, syncopal migraine
Asunaprevir	HIV infection	Fatigue, rash, nausea, neutropenia, anemia, deranged liver function tests	Oral	Moderate/severe hepatic impairment (Child B or C)
Bictegravir	HIV infection	Increased serum creatine kinase, deranged liver function tests, neutropenia	Oral	None
Entecavir	HBV infection	Deranged liver function tests	Oral	Moderate/severe hepatic impairment (Child B or C)
Zidovudine	HIV infection	Headache, malaise, rash, nausea, neutropenia, anemia	Oral	None
Sofosbuvir	Chronic HCV infection	Fatigue, headache, insomnia, nausea, diarrhea, anemia, myalgia, rash	Oral	None
Atovaquone	Protozoal infection or prophylaxis (Pneumocystic jirovecii, Plasmodium spp. etc)	Headache, insomnia, rash, diarrhea, myalgia, drug fever, hyponatremia, neutropenia	Oral	None
Remdesivir	COVID-19	Deranged liver function tests	Intravenous	Serum alanine transaminase ≥5x the upper-limit of normal

Supplementary Table 1 FDA-approved repurposable drug candidates tested in this study.

NAME	SIZE	NES	FDR q-val
REACTOME_PKMTS_METHYLATE_HISTONE_LYSINES	46	2.035874	0.044
REACTOME_MITOTIC_PROPHASE	109	1.795425	0.213909
REACTOME_RMTS_METHYLATE_HISTONE_ARGININES	49	1.748761	0.228738
REACTOME_NRIF_SIGNALS_CELL_DEATH_FROM_THE_N UCLEUS	15	1.739992	0.193812
REACTOME_SCF_SKP2_MEDIATED_DEGRADATION_OF_ P27_P21	56	1.705256	0.24902
REACTOME_RHO_GTPASES_ACTIVATE_PKNS	58	1.70523	0.21485
REACTOME_CELLULAR_SENESCENCE	154	1.700467	0.203287
REACTOME_MEIOTIC_RECOMBINATION	51	1.70042	0.183376
REACTOME_MEIOSIS	77	1.698298	0.17955
REACTOME_AUF1_HNRNP_D0_BINDS_AND_DESTABILIZ ES_MRNA	51	1.689022	0.170498
REACTOME_GENE_SILENCING_BY_RNA	98	1.680767	0.185941
REACTOME_ASYMMETRIC_LOCALIZATION_OF_PCP_PR OTEINS	58	1.662479	0.215862
REACTOME_CONDENSATION_OF_PROPHASE_CHROMO SOMES	43	1.661445	0.209014
REACTOME_MICRORNA_MIRNA_BIOGENESIS	24	1.645813	0.21082
REACTOME_INTERLEUKIN_7_SIGNALING	24	1.643195	0.199699
REACTOME_TRANSCRIPTIONAL_REGULATION_BY_SMALL_RNAS	75	1.642892	0.189968
REACTOME_RUNX1_REGULATES_TRANSCRIPTION_OF_ GENES_INVOLVED_IN_DIFFERENTIATION_OF_HSCS	93	1.639989	0.183625
REACTOME_HDMS_DEMETHYLATE_HISTONES	30	1.624097	0.189556
REACTOME_ESTROGEN_DEPENDENT_GENE_EXPRESSI ON	114	1.623586	0.181895
REACTOME_G1_S_DNA_DAMAGE_CHECKPOINTS	63	1.622633	0.177329
REACTOME_REGULATION_OF_MRNA_STABILITY_BY_PR OTEINS_THAT_BIND_AU_RICH_ELEMENTS	81	1.619633	0.179618
REACTOME_DEFECTIVE_CFTR_CAUSES_CYSTIC_FIBRO	56	1.616455	0.179718

REACTOME_APC_C_CDH1_MEDIATED_DEGRADATION_O F_CDC20_AND_OTHER_APC_C_CDH1_TARGETED_PROT EINS_IN_LATE_MITOSIS_EARLY_G1	67	1.613433	0.177683
REACTOME_PRC2_METHYLATES_HISTONES_AND_DNA	41	1.606214	0.17865
REACTOME_EUKARYOTIC_TRANSLATION_INITIATION	81	1.603774	0.174798
REACTOME_HDACS_DEACETYLATE_HISTONES	55	1.59932	0.175764
REACTOME_CELLULAR_RESPONSE_TO_HYPOXIA	67	1.596512	0.170884
REACTOME_RNA_POLYMERASE_I_TRANSCRIPTION	78	1.595998	0.166353
REACTOME_HATS_ACETYLATE_HISTONES	103	1.594675	0.169724
REACTOME_RNA_POLYMERASE_I_PROMOTER_ESCAPE	59	1.593383	0.167005
REACTOME_FORMATION_OF_THE_EARLY_ELONGATION _COMPLEX	32	1.593123	0.163037
REACTOME_SWITCHING_OF_ORIGINS_TO_A_POST_REP LICATIVE_STATE	84	1.590331	0.165325
REACTOME_FANCONI_ANEMIA_PATHWAY	35	1.587149	0.16675
REACTOME_APOPTOTIC_FACTOR_MEDIATED_RESPONS E	16	1.585396	0.16314
REACTOME_ACTIVATION_OF_THE_MRNA_UPON_BINDIN G_OF_THE_CAP_BINDING_COMPLEX_AND_EIFS_AND_S UBSEQUENT_BINDING_TO_43S	42	1.585282	0.159736
REACTOME_ERCC6_CSB_AND_EHMT2_G9A_POSITIVELY _REGULATE_RRNA_EXPRESSION	44	1.577036	0.163907
REACTOME_ACTIVATION_OF_ANTERIOR_HOX_GENES_I N_HINDBRAIN_DEVELOPMENT_DURING_EARLY_EMBRY OGENESIS	87	1.5695	0.168932
REACTOME_HCMV_LATE_EVENTS	77	1.569343	0.165644
REACTOME_STABILIZATION_OF_P53	52	1.5621	0.176659
REACTOME_HIV_TRANSCRIPTION_ELONGATION	40	1.56044	0.175868
REACTOME_NEGATIVE_EPIGENETIC_REGULATION_OF_ RRNA_EXPRESSION	76	1.560153	0.172651
REACTOME_FORMATION_OF_TC_NER_PRE_INCISION_C OMPLEX	52	1.556269	0.171755
REACTOME_P75NTR_SIGNALS_VIA_NF_KB	15	1.55442	0.170508
REACTOME_ACTIVATED_PKN1_STIMULATES_TRANSCRI PTION_OF_AR_ANDROGEN_RECEPTOR_REGULATED_G ENES_KLK2_AND_KLK3	33	1.550302	0.172005

REACTOME_TRANSCRIPTIONAL_REGULATION_OF_GRA	56	1.549487	0.169161
REACTOME_FORMATION_OF_THE_BETA_CATENIN_TCF _TRANSACTIVATING_COMPLEX	61	1.548597	0.167521
REACTOME_EPIGENETIC_REGULATION_OF_GENE_EXP RESSION	113	1.545807	0.169608
REACTOME_MRNA_CAPPING	28	1.539692	0.171467
REACTOME_B_WICH_COMPLEX_POSITIVELY_REGULAT ES_RRNA_EXPRESSION	58	1.535715	0.172508
REACTOME_CYCLIN_A_CDK2_ASSOCIATED_EVENTS_AT _S_PHASE_ENTRY	80	1.533588	0.171664
REACTOME_OXIDATIVE_STRESS_INDUCED_SENESCEN CE	88	1.532286	0.170114
REACTOME_AMYLOID_FIBER_FORMATION	64	1.531269	0.168856
REACTOME_INFLUENZA_INFECTION	117	1.524853	0.177163
REACTOME_RRNA_PROCESSING	161	1.523799	0.175456
REACTOME_ABORTIVE_ELONGATION_OF_HIV_1_TRANS CRIPT_IN_THE_ABSENCE_OF_TAT	22	1.523257	0.175333
REACTOME_POSITIVE_EPIGENETIC_REGULATION_OF_R RNA_EXPRESSION	72	1.519612	0.179256
REACTOME_METABOLISM_OF_POLYAMINES	54	1.513546	0.18058
REACTOME_NONSENSE_MEDIATED_DECAY_NMD	79	1.510758	0.182073
REACTOME_DEGRADATION_OF_DVL	53	1.50713	0.1833
REACTOME_SIRT1_NEGATIVELY_REGULATES_RRNA_EX PRESSION	37	1.502177	0.186786
REACTOME_PRE_NOTCH_EXPRESSION_AND_PROCESSI NG	80	1.502052	0.184445
REACTOME_DNA_METHYLATION	34	1.49901	0.186451
REACTOME_REPRODUCTION	91	1.497232	0.188127
REACTOME_TRANSCRIPTION_COUPLED_NUCLEOTIDE_ EXCISION_REPAIR_TC_NER	76	1.493491	0.188147
REACTOME_REGULATION_OF_EXPRESSION_OF_SLITS_ AND_ROBOS	129	1.488774	0.191595
REACTOME_SRP_DEPENDENT_COTRANSLATIONAL_PR OTEIN_TARGETING_TO_MEMBRANE	75	1.481835	0.201191

REACTOME_CELLULAR_RESPONSES_TO_EXTERNAL_ST IMULI	486	1.480829	0.198844
REACTOME_EUKARYOTIC_TRANSLATION_ELONGATION	58	1.474574	0.207194
REACTOME_ORC1_REMOVAL_FROM_CHROMATIN	66	1.471954	0.208534
REACTOME_ASSEMBLY_OF_THE_PRE_REPLICATIVE_CO MPLEX	63	1.470746	0.208734
REACTOME_KERATAN_SULFATE_BIOSYNTHESIS	21	1.464303	0.215357
REACTOME_SENESCENCE_ASSOCIATED_SECRETORY_ PHENOTYPE_SASP	75	1.464137	0.213509
REACTOME_TRANSLATION	240	1.45829	0.223907
REACTOME_VOLTAGE_GATED_POTASSIUM_CHANNELS	20	1.454349	0.226557
REACTOME_DNA_DAMAGE_RECOGNITION_IN_GG_NER	37	1.447582	0.233274
REACTOME_INTERCONVERSION_OF_NUCLEOTIDE_DI_A ND_TRIPHOSPHATES	25	1.447349	0.231582
REACTOME_APC_C_MEDIATED_DEGRADATION_OF_CEL L_CYCLE_PROTEINS	81	1.443777	0.23438
REACTOME_POLYMERASE_SWITCHING_ON_THE_C_STR AND_OF_THE_TELOMERE	16	1.439511	0.23541
REACTOME_TNFR1_INDUCED_NFKAPPAB_SIGNALING_P ATHWAY	27	1.437981	0.233472
REACTOME_DUAL_INCISION_IN_TC_NER	63	1.437643	0.231104
REACTOME_REGULATION_OF_RUNX3_EXPRESSION_AN D_ACTIVITY	51	1.432937	0.234545
REACTOME_KERATAN_SULFATE_KERATIN_METABOLIS M	27	1.430326	0.236033
REACTOME_PCP_CE_PATHWAY	85	1.427146	0.237307
REACTOME_NEGATIVE_REGULATION_OF_NOTCH4_SIG NALING	51	1.42502	0.237688
REACTOME_SIGNALING_BY_NOTCH4	78	1.420591	0.246818
REACTOME_KSRP_KHSRP_BINDS_AND_DESTABILIZES_ MRNA	16	1.420209	0.244459
REACTOME_TRANSCRIPTIONAL_REGULATION_BY_RUN X2	108	1.419713	0.243666
REACTOME_DECTIN_1_MEDIATED_NONCANONICAL_NF_ KB_SIGNALING	58	1.419439	0.241926

REACTOME_RESPONSE_OF_EIF2AK4_GCN2_TO_AMINO _ACID_DEFICIENCY	66	1.418276	0.240912
REACTOME_RUNX1_REGULATES_GENES_INVOLVED_IN _MEGAKARYOCYTE_DIFFERENTIATION_AND_PLATELET _FUNCTION	64	1.417405	0.239215
REACTOME_ESR_MEDIATED_SIGNALING	178	1.414636	0.242566
REACTOME_DNA_REPLICATION	119	1.407263	0.249869
REACTOME_NUCLEOTIDE_SALVAGE	20	1.406257	0.249112

Supplementary Table 2. Gene sets enriched in Sim 1.1 vs. DMSO in infected samples according to Gene set enrichment analysis (GSEA).