Stable Cell Clones Harboring Self-Replicating SARS-CoV-2 RNAs for Drug Screen

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ABSTRACT The development of antivirals against severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) has been hampered by the lack of efficient cell-based replication systems that are amenable to high-throughput screens in biosafety level 2 laboratories. Here we report that stable cell clones harboring autonomously replicating SARS-CoV-2 RNAs without spike (S), membrane (M), and envelope (E) genes can be efficiently derived from the baby hamster kidney (BHK-21) cell line when a pair of mutations were introduced into the non-structural protein 1 (Nsp1) of SARS-CoV-2 to ameliorate cellular toxicity associated with virus replication. In a proof-of-concept experiment we screened a 273-compound library using replicon cells and identified three compounds as novel inhibitors of SARS-CoV-2 replication. Altogether, this work establishes a robust, cell-based system for genetic and functional analyses of SARS-CoV-2 replication and for the development of antiviral drugs.

IMPORTANCE SARS-CoV-2 replicon systems that have been reported up to date were unsuccessful in deriving stable cell lines harboring non-cytopathic replicons. The transient expression of viral sgmRNA or a reporter gene makes it impractical for industry-scale screening of large compound libraries using these systems. Here, for the first time, we derived stable cell clones harboring the SARS-CoV-2 replicon. These clones may now be conveniently cultured in a standard BSL-2 laboratory for high throughput screen of compound libraries. Additionally, our stable replicon cells represent a new model system to study SARS-CoV-2 replication.

KEYWORDS SARS-CoV-2, replicon, stable cell clone, drug screen, replication

The single-stranded, positive-sense SARS-CoV-2 RNA genome is approximately 30 kb in length and comprises a short 5’ untranslated region (UTR), 13 open reading frames (ORFs), a 3’ UTR, and a poly(A) tail. Through discontinuous transcription events, the virus makes at least nine canonical subgenomic RNAs (sgmRNA), which encode structural and accessory proteins (1). The genomic RNA (gRNA) harbors two large ORFs, ORF1a and ORF1ab, which are initially translated into two polyproteins, pp1a and pp1ab, and subsequently processed by viral proteases to produce 16 non-structural proteins (Nsp) that form the viral replication complex and confer immune evasion (2–4).

Viral replication and translation machinery offer some of the most promising targets for antiviral drug development. For example, the main protease (Nsp 5) and the viral RNA-dependent RNA polymerase (Nsp 12) of SARS-CoV-2 are primary targets for antiviral discovery because they are responsible for cleavage of replicase polyproteins 1a/1ab and for virus replication, respectively. A cell-based system that harbors the minimally essential SARS-CoV-2 replication and translation machinery is highly desirable because it enables simultaneous screening of inhibitors of multiple viral proteins in a...
biosafety level 2 setting. To this end, subgenomic viral RNA molecules called replicons are often designed to autonomously replicate in cells without generating infectious virus. Current SARS-CoV-2 replicon systems, however, do not permit persistent replication in cell lines due to intrinsic toxicity (3, 5–8). The limited time window allowable for detection of replication, the inability to generate master and working cell banks for lot consistency, and the challenge to scale up for industrial processes, make it impractical to apply transient replicon systems in high-throughput screening (HTS) of large compound libraries. Here, we describe, for the first time, the derivatization and characterization of stable cell clones harboring autonomously replicating SARS-CoV-2 RNAs. We further demonstrated the applicability of cells harboring SARS-CoV-2 replicon RNA in drug screen.

RESULTS

Initial design of SARS-CoV-2 replicon. To generate stable cell clones harboring replicating SARS-CoV-2 RNAs, we first constructed a replicon termed SARS-CoV-2-Rep-NanoLuc-Neo, in which the spike (S) gene was replaced by a nanoluciferase reporter (NanoLuc), and the envelope (E) and membrane (M) genes were replaced with the neomycin phosphotransferase gene (Neo or NeoR) (Fig. 1A). Electroporation of this replicon RNA along with in vitro transcribed RNA encoding the nucleocapsid protein (NP), which was previously shown to improve launch efficiency (9–11), into Vero E6, Huh7.5.1, A549, and BHK-21 cells resulted in expression of nanoluciferase to varying

FIG 1 Initial design of SARS-CoV-2 replicons. (A) Top, genome organization of SARS-CoV-2. Leader sequence (red), transcriptional regulatory sequence within the leader sequence (TRS-L) and within the body (TRS-B) are highlighted in green. Bottom, the design of SARS-CoV-2-Rep-NanoLuc-Neo. (B, C, D, E) Replication kinetics of SARS-CoV-2-Rep-NanoLuc-Neo in Vero E6 (B), A549 (C), Huh7.5.1 (D), and BHK-21 cells (E). Nano luciferase was measured at indicated time points post-electroporation. Notably, nanoluciferase continued to decrease even under G418 selection. (F) Generation of BHK21 stable cells that express NP in a doxycycline-inducible manner. Cells were induced with 0.5 μg/mL doxycycline and lysed at 48 h postinduction for Western blotting with anti-NP and anti-actin antibodies. Numbers on the left refer to the positions of marker proteins that are given in kilodalton (kDa).
extents (Fig. 1B to E). However, no viable clones could be recovered after 21 days of selection in G418, suggesting that active replication of the replicon RNA is either unsustainable or cytotoxic. Huh7.5.1 and BHK-21 cells supported higher levels of nanoluciferase expression than Vero E6 and A549, although we cannot rule out that the differences are attributed to different electroporation efficiency of the four cell lines. Because electroporating two different RNA species into the same cell is inefficient, we created a BHK-21 stable cell clone (BHK-21-NPDox-ON) in which NP is expressed in a doxycycline-inducible manner (Fig. 1F). Electroporation of SARS-CoV-2-Rep-NanoLuc-Neo RNA into BHK-21-NPDox-ON cells resulted in three neomycin-resistant clones out of four million cells. The resulted clones grew very slowly in the presence of 200 μg/mL G418, and no nanoluciferase activity or viral RNA could be detected, indicating the loss of functional replicon RNA. Altogether, persistent replication of SARS-CoV-2-Rep-NanoLuc-Neo could not be achieved in any of the four mammalian cell lines.

An improved SARS-CoV-2 replicon. Coronaviruses have evolved a variety of mechanisms to shut off host transcription and translation (12–14). Of all SARS-CoV-2 proteins, Nsp1 causes the most severe viability reduction in cells of human lung origin (15). The carboxyl terminus of Nsp1 folds into two helices, which insert into the mRNA entrance channel on the 40S ribosome subunit, preventing both the host mRNA and viral mRNA from gaining access to ribosomes and consequently shutting down translation (16, 17) (Fig. 2A). The first C-terminal helix (residues 153 to 160) makes hydrophobic interactions with the 40S ribosomal protein uS5, and interacts with the 40S ribosomal protein uS3 with salt-bridges (18); the second C-terminal helix (residues 166 to 178) interacts with ribosomal protein eS30 and with the phosphate backbone of h18 of the 18S rRNA via the two conserved arginines R171 and R175 (16). A notably conserved KH dipeptide (K164 and H165) between the two helices forms interactions with h18 through H165 stacking between Uridine 607 (U607) and U630 of 18S rRNA, and through electrostatic interactions between K164 and the rRNA bases G625 and U630 (Fig. 2B). To explore the possibility of disrupting the interaction between Nsp1 and host ribosomes, we performed molecular dynamics simulation followed by free energy perturbation calculation. The results predicted that mutations of residues K164, R171, R175, H165, and S167 of Nsp1 to alanine will reduce the interaction in the order of impact (Fig. 2C and Fig. S1). We hypothesize that a pair of mutations, such as K164A/H165A that weaken the interaction between C-terminus of Nsp1 and ribosome, will lead to a shorter occupation time of Nsp1 on the ribosome and increase the accessibility of ribosomes to host mRNA. As a result, the Nsp1-mediated toxicity to the host should be alleviated. To test this possibility, we created a new replicon construct SARS-CoV-2-Rep-NanoLuc-Neo-Nsp1K164A/H165A in which K164/H165 were mutated to alanine (Fig. 2D). For comparison, we also made two additional replicons, SARS-CoV-2-Rep-NanoLuc-Neo-Nsp1R124S/K125E and SARS-CoV-2-Rep-NanoLuc-Neo-Nsp1N128S/K129E, given that both pairs of mutations (R124S/K125E and N128S/K129E) reportedly reduce Nsp1-mediated cell toxicity in a human lung cell line (15). When electroporating into BHK-21-NPDox-ON cells, all three replicons led to transient expression of nano luciferase (Fig. 2E). However, in the presence of 200 μg/mL G418, only SARS-CoV-2-Rep-NanoLuc-Neo-Nsp1K164A/H165A yielded viable cells (Pool #1), from which 12 stable clones (Clone #2 to #13) were subsequently derived by limiting dilution. Three independent experiments were performed, and each time only electroporation of SARS-CoV-2-Rep-NanoLuc-Neo-Nsp1K164A/H165A resulted in viable clones in BHK-21-NPDox-ON cells (Table 1). It is worth mentioning that SARS-CoV-2-Rep-NanoLuc-Neo-Nsp1K164A/H165A also replicated well in Huh7.5.1 cell line although no viable cells could be recovered after G418 selection (Fig. 2F).

Characterization of cells harboring SARS-CoV-2-Rep-NanoLuc-Neo-Nsp1K164A/H165A. To explore the possibility for SARS-CoV-2-Rep-NanoLuc-Neo-Nsp1K164A/H165A to persist in cells without selection pressure, G418 was subsequently withdrawn from Pool #1 cells after the initial selection. Up to 1 week there was no significant loss of nanoluciferase expression, a feature that is compatible with drug screen. The level of
nanoluciferase decreased by one log after 10 days culturing without G418 and then by another two logs after 21 days (Fig. 3A).

Next, we performed quantitative reverse transcription-PCR (RT-qPCR) to profile gRNA and sgRNAs species in cells harboring SARS-CoV-2-Rep-NanoLuc-Neo-Nsp1K164A/H165A. SARS-CoV-2 transcribes multiple canonical sgRNAs, including S, E, M, NP, ORF3a, ORF6, ORF7a, ORF7b, ORF8, and ORF10, although multiple studies have found negligible ORF10
expression and very few ORF7b body–leader junctions (1, 19). In replicon cells, S, M, E ngmRNA are replaced with those encoding NanoLuc and NeoR; ngmRNA encoding ORF6 is lost because its transcriptional regulatory sequence body (TRS-B) resides in the M gene, which is also deleted in the replicon RNA. Hence, cells harboring the replicon would express at least six ngmRNAs, namely, NanoLuc, NeoR, ORF3a, ORF7a, ORF8, and NP. We designed primers/probes to specifically amplify the grRNA of the replicon and ngmRNAs of NanoLuc and NeoR. Shown in Fig. 3B and C, the temporal expression of grRNA and ngmRNA for the NanoLuc and NeoR genes were clearly observed in Pool #1 cells. Western blotting confirmed the presence of Nsp1 and NP in replicon cell lysates (Fig. 3D). To further characterize the replicon grRNA, next-generation sequencing (NGS) was performed on all 12 stable clones. While the replicon grRNA containing Nsp1K164A/H165A was present in all clones, additional synonymous or missense mutations were detected in each clone (Table S1). Among those, Nsp4 R401S substitution was detected in 10 out of 12 clones, and Nsp10 T111I appeared in six out of 12 clones. Ongoing research is studying whether these changes constitute adaptive mutations which had enabled efficient replication in BHK-21 cells. Notably, sequencing the replicon grRNA in clone #9 also revealed a deletion knocking out ORF7a/b, ORF8 and the first 392 amino acids of the NP (Fig. S2), indicating that NP is dispensable for genome replication. The presence of canonical ngmRNA species in each stable clone was also confirmed by NGS, although the method employed in this study could not locate all ngmRNA species due to uneven coverage of sequence reads over different regions (Fig. 3E).

The 12 stable clones were also characterized for cellular morphology and growth kinetics. Shown in Fig. 4A and B, all stable cell clones maintained a BHK-21-like morphology with doubling times ranging from 18 h to 30 h. Finally, every single cell was stained positive when we performed immunostaining on 12 stable clones with an anti-double stranded RNA (dsRNA) antibody (Fig. 4C). Because dsRNA is an intermediate product during the replication of positive sense genome viruses, this finding confirmed the existence of active SARS-CoV-2 replication in these clones.

**Application of replicon cells in drug screen.** To demonstrate the suitability of SARS-CoV-2-Rep-NanoLuc-Neo-Nsp1K164A/H165A cells in drug screen, we tested a library of 273 compounds for inhibitory effects. These compounds were selected to target Nsp5 (3CLpro), Nsp3 (PLpro), Nsp12 (RdRP), Nsp15, Nsp16, and X domain by virtual screen (Table S2). At 5 μM, nine compounds exhibited more than 50% inhibition based on nanoluciferase expression (Fig. 5A). Three compounds, Darapladib, Genz-123346, and JNJ-5207852 (Fig. 5B), were validated in replicon Pool #1 cells and then in three human cell lines A549-hACE2, Calu-3, and Caco-2 cells using live virus. Remdesivir (inhibitor of RdRP) and GC376 (3CL protease inhibitor) were included as positive controls. The IC_{50} of these compounds are provided in Fig. 5C. Interestingly, all three compounds displayed cell type-specific activity against SARS-CoV-2, which others also noticed with repurposed drugs (20). We subsequently sought to evaluate the clonal variability in response to drug treatment. To this end, stable replicon cell clone #3, #5, #7, #9, #11, and #13 were tested for responses to GC376 treatment. Acquired IC_{50} values ranged from 5.9 μM to 13 μM (mean = 9.3 μM, standard deviation = 2.9 μM), suggesting that all clones are potentially suitable for testing drug efficacy (Fig. 6A). Lastly, all 12 clones have now been maintained for more than 20 passages under G418 without losing the expression of nano luciferase (Fig. 6B).

**TABLE 1 Vial clone numbers**

| Replicon Name          | 1st transfection | 2nd transfection | 3rd transfection |
|------------------------|------------------|------------------|------------------|
| SARS-CoV-2-Rep-NanoLuc-Neo | 3 \(^{(0)}\) | 0 | 0 |
| SARS-CoV-2-Rep-NanoLuc-Neo-Nsp1K164A/H165A | 132 | 7 | 107 |

\(^{(a)}\)Three clones were initially obtained after G418 selection but found to be devoid of the replicon RNA after deep sequencing. \(^{(b)}\)4 × 10^6 BHK-21-NP\(^{Dox-ON}\) cells were electroporated with 10 μg of each RNA for each transfection.

\(^{(c)}\)The second transfection has lower efficiency, resulting in fewer clones.

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FIG 3 Characterization of cells harboring BHK21-NPDox-ON Rep-NanoLuc-Neo-NspK164A/H165A. (A) Nano luciferase in BHK21-NPDox-ON replicon cells (Pool #1) was measured at given time points following G418 withdrawal. RNA was also extracted at indicate time points and quantified by RT-qPCRs targeting ORF1ab (gRNA in A), or sgmNeoR and sgmNanoLuc (B). (C) RT-PCR analysis of viral RNA from replicon Pool #1 cells. The corresponding primer pairs are

| Lane | Primer name | Sequence | Amplicon Size (bp) |
|------|-------------|----------|-------------------|
| 1    | 32f         | ccaacacactttcagctcttg | 403 |
| 434r | caacctctcaagccacaa | 4 |
| 2    | 1001f       | agagcctgaatcatgcagca | 888 |
| 1892r| caacgacagcctcgatgca | 8 |
| 3    | 3000f       | ttggtgagaattgac | 1095 |
| 4072r| tgctcactaactaaga | 4 |
| 4    | 7000f       | acgctctggctagttgctt | 963 |
| 7965r| tggacataaaaagtctatagtaa | 4 |
| 5    | 8000f       | atgttgttcatgtgccgga | 960 |
| 8932r| gtgtgtacaggtgctcactga | 4 |
| 6    | 17000f      | acctaatctctatcctgag | 1010 |
| 17990r| tttaaagctctctctaatggct | 4 |
| 7    | sgmNanoR    | ccaccaacacactttcagctcttg | 526 |
| Neor | tcaacctctacatgtcttc | 4 |
| 8    | sgeF        | cgatcctcagctctttca | 825 |

(Continued on next page)
DISCUSSION

Two important observations were made in our study regarding establishing stable cell clones harboring SARS-CoV-2 replicon. First, stable cell clones harboring SARS-CoV-2 replicon were obtained when K164A/H165A mutations were introduced to Nsp1. This result is consistent with the structural analysis of Nsp1, which predicts that K164A/H165A reduce the interaction between C-terminus of Nsp1 and the ribosome and hence increase the accessibility of ribosomes to host mRNA. By contrast, R124S/K125E mutations may reduce the binding of the Nsp1 N-terminus to the 5′-UTR of viral mRNA, leaving the C-terminal region of Nsp1 constantly bound to ribosome (15, 18, 21). Consequently, neither viral nor host mRNA can efficiently access the ribosome in the presence of Nsp1R124S/K125E. The structural modeling of N128S/K129E could not be performed due to lack of published structure. N128S/K129E, despite being reported to attenuate SARS-CoV Nsp1-mediated inhibition of interferons (22), failed to derive viable cells in a replicon. Secondly, viable cells were only recoverable from the BHK-21 background, implying that SARS-CoV-2-Rep-NanoLuc-Neo-Nsp1K164A/H165A remains cytopathic in other cell types. In addition to Nsp1, other viral factors have been reported to cause cytopathic effect. For example, SARS-CoV-2 PLpro displayed cytopathogenic effect in Vero E6 cells (23). Notably, as this manuscript was in preparation, a similar but non-identical SARS-CoV-2 replicon bearing Nsp1K164A/H165A reportedly failed to generate stable cell clones in Huh-7.5 and BHK-21 cells (24), raising an interesting question whether the detected additional mutations from our stable cell clones facilitated non-cytopathic replication.
In summary, we have established a robust replicon system for future genetic and functional analyses of SARS-CoV-2 replication in vitro. To the best of our knowledge, this is the first time that stable cell clones harboring a SARS-CoV-2 replicon could be derived. These clones offer significant advantages over existing replicon systems in that cells may now be conveniently cultured, banked, qualified, and used in a standard BSL-2 laboratory for high throughput screen of compound libraries. The stable replicon system also opens a new avenue for molecular characterization of SARS-CoV-2 replication.

MATERIALS AND METHODS

Cell culture and reagents. The human kidney epithelial cell line Lenti-X 293T was purchased from TaKaRa. The human liver cell line Huh7.5.1 was provided by Dr. Francis Chisari (Scripps Research Institute). The baby hamster kidney fibroblast cell line BHK21 (CCL-10), African green monkey kidney epithelial cells (Vero E6; HTB-37), Calu-3 (HTB-55), and A549 (CCL-185) were purchased from the American Type Culture Collection. A549-hACE2 (NR-53821) cells were obtained from BEI Resources. All cell lines were maintained in Dulbecco’s Modified Eagle Medium supplemented with 5% penicillin and streptomycin, and 10% fetal bovine serum (FBS) at 37°C with 5% CO2. The SARS-CoV-2 Nucleocapsid antibody (40143-MM05) was purchased from Sino Biological. The SARS-CoV-2 Nsp1 antibody (PA5-116941) was purchased from Thermo Fisher Scientific. The β-actin antibody (GTX109639) was purchased from Gentex. Secondary antibodies were purchased from LI-COR Bioscience. GC376 Sodium was purchased from Aobious (AOB36447). Remdesivir was purchased from MedChemExpress (HY-104077).

Plasmid construction. Doxycycline-inducible expression of SARS-CoV-2 NP was established in Vero E6, HuH7.5.1, and BHK-21 using TripZ-NP plasmid. NP cDNA was subcloned into pTripZ (Agel/MluI) using the following primers: TripZ-NPf: 5’-ATATAGACCGGTCCACCATGTCTGATAATGGACCCCA-3’, TripZ-NPr: 5’-ATATAGACGCGTTTAGGCCTGAGTTGAGTCAG-3’. Production of SARS-CoV-2 reporter viruses. SARS-CoV-2 recombinant virus could be generated using a 7-plasmid reverse genetic system which was based on the virus strain (2019-nCoV/USA_WA1/2020) isolated from the first reported SARS-CoV-2 case in the United States. (10). The initial seven plasmids were generous gifts from Dr. P-Y Shi (UTMB). Upon receipt, fragment 4 was subsequently subcloned into a low-copy-number plasmid pSMart LCmP (Lucigen) to increase stability. Standard molecular biology technique was employed to create the SARS-CoV-2 nanoluciferase and firefly reporter...
viruses. In vitro transcription and electroporation were carried following procedures that were detailed elsewhere (25).

**SARS-CoV-2 replicon.** The SARS-CoV-2-Rep-NanoLuc-Neo replicon was constructed based on the full-length SARS-CoV-2 cDNA infectious clone by replacing the S gene with a nano luciferase gene, and by replacing M and E genes with a neomycin phosphotransferase (Neo) gene. To introduce Nsp1 R124S/K125E, N128S/K129E, and K164A/H165A mutations into the SARS-CoV-2-Rep-NanoLuc-Neo replicon, puc57-CoV2-F1 plasmids containing mutated Nsp1 were first created by using overlap PCR method with the following primers: M13F: 5'-GTAAAACGACGGCCAGT-3', R124S/K125E-F: 5'-CAAGGTTCTTCTTTCGGAGAACGGTAGTGATGATG-3', R124S/K125E-R: 5'-TTATTACCGTTCTCCGAAAGAAGAACCTTGCGGTAAG-3', N128S/K129E-F: 5'-TAAGAACGGTAGTGAGGGAGCTGGTGGCCATAGTTA-3', N128S/K129E-R: 5'-CACCAGCTCCCTACCTACC GTTCTTACGAAAGAAGAA-3', K164A/H165A-F: 5'-AAAACTGGAAACCTGGCGACGACGATGTGTTACCCGTAAG-3', K164A/H165A-R: 5'-GGGTAACACCACTGCTGGCGGCAGTGTTCCAGTTTTCTTGAA-3', NheI-R: 5'-CACGAGCAGCCTCTGATGCA-3'. PCR fragments were digested by Bgl II/Nhe I and ligated into Bgl II/Nhe I digested F1 plasmid. The resulted plasmids were validated by restriction enzyme digestion and Sanger sequencing. Assembly of the seven plasmids into the replicon and in vitro transcription were performed following a published protocol (25).

**RNA electroporation.** Forty-eight hours post-doxycycline treatment, BHK-21-NPDox-ON cells were washed with phosphate-buffered saline (PBS), trypsinized, and resuspended in complete growth medium. Cells were pelleted by centrifugation (1,000 × g for 5 min at 4°C), washed twice with ice-cold DMEM, and resuspended in ice-cold Gene Pulser Electroporation Buffer (Bio-Rad) at 1 × 10^7 cells/mL. Cells (0.4 mL) were then mixed with 10 μg of replicon RNA and 2 μg NP RNA, placed into 4 mm gap electroporation cuvettes, and electroporated at 270 V, 100 μF, and 950 μF in a Gene Pulser Xcell Total System (Bio-Rad). To establish stable replicon cells, 200 μg/mL of G418 was added to the media (without doxycycline) between 24 h and 48 h following electroporation, after which culture medium was changed every 2 to 3 days. Three weeks after G418 selection, the resultant foci were counted. All cells were trypsinized and pooled together in a T-75 flask for expansion. Limiting dilution was subsequently performed to derive single cell clones.

**Determination of cell doubling time.** Stable replicon clones and the parent BHK-21-NPDox-ON cells (without doxycycline) were seeded in multiple 48-well plates in triplicates at a density of 1,000 cells per well. One plate was removed every 24 h for fixation in 4% PFA followed by Hoechst 33342 staining. Cell numbers were quantified on a BioTek Cytation 7 cell imaging multimode reader. Doubling time was determined by GraphPad Prism 9.

**Quantification of viral RNA.** Viral RNA was quantified by reverse-transcription quantitative PCR (RT-qPCR) on a StepOnePlus real-time PCR system (Applied Biosystems) using Luna Universal Probe One-Step RT-qPCR Kit (New England Biolabs) with an in-house developed protocol. Primers and probes for qPCR were as follows: ORF1ab forward: 5'-CCCTGTCGGTTTACACTTAA-3', reverse: 5'-ACGATTTGCGATCGCTGA-3',
probe: FAM-CCGTCTGCGGTATGTGGAAAGGTTATGG-BHQ1; NanoLuc gene subgenomic mRNA forward: 5’-CCAACCAACTTTCGATCTCTTG-3’, reverse: 5’-GGACTTGGTCCAGGTTGTAG-3’, probe: FAM-ACGAACAATGGTC –BHQ1; Neomycin phosphotransferase gene subgenomic mRNA forward: 5’-CGATCTCTTGTAGATCTGTTCT-3’, reverse: 5’-GCCCAGTCATAGCCGAATAG-3’, probe: FAM-ACAAGATGGATTGCACGCAGGTTC-BHQ1. To generate standard plasmids, the cDNAs of SARS-CoV-2 ORF1ab gene, NanoLuc gene sgmRNA, and neomycin phosphotransferase gene sgmRNA were cloned into a pCR2.1-TOPO plasmid, respectively. The copy number of replicon RNA was calculated by comparing to a standard curve obtained with serial dilutions of the standard plasmid.

**Immunoblotting.** Cells were grown in 24-well plates and lysates were prepared with Radioimmunoprecipitation assay buffer (50 mM Tris-HCl [pH 7.4]; 1% NP-40; 0.25% sodium deoxycholate; 150 mM NaCl; 1 mM EDTA; protease inhibitor cocktail [Sigma]; 1 mM sodium orthovanadate), and insoluble material was precipitated by brief centrifugation. Lysates were loaded onto 4% to 20% SDS-PAGE gels and transferred to a nitrocellulose membrane (LI-COR, Lincoln, NE), blocked with Intercept (TBS) Blocking Buffer (LI-COR, Lincoln, NE) for 1 h, and incubated with 1:1,000 primary antibody (SARS-CoV-2 Nsp1 Antibody, Thermo Fisher PA5-116941; SARS-CoV-2 NP Antibody, Sino Biological 40143-MM05; Anti-β-Actin antibody, Sigma clone AC-74) overnight at 4°C. Membranes were washed three times with 1/20 TBS containing 0.05% Tween 20 (vol/vol), incubated with IRDye secondary antibodies (LI-COR, Lincoln, NE) for 1 h, and washed again to remove unbound antibody. Odyssey CLx (LI-COR Biosystems, Lincoln, NE) was used to detect bound antibody complexes.

**Compound screen.** Two-hundred and 73 compounds (assembled by TargetMol, not previously approved drugs) were diluted in culture media to a final concentration of 5 uM for initial screen. Approximately 1.5 × 10^4 replicon cells/well were seeded in 96-well plates in the absence of G418. Twenty-four hours later, cell culture media (without G418) was replaced with media containing 5 uM compounds or the same volume of diluent DMSO. After incubation at 37°C for specified periods, cells were assayed for NanoLuc activity using Nano-Glo Luciferase Assay System (N1130, Promega) or cell viability using CellTiter-Glo (G7571, Promega). For validation, A549-hACE2, Caco-2, or Calu-3 cells were seeded in 96-well plates at a density of 10^4 cells/well. Twenty-four hours later, cells were infected with SARS2-NanoLuc reporter virus in triplicates at an MOI of 0.05 in culture medium containing compounds. After 24 h at 37°C, cells were assayed for NanoLuc activity using Nano-Glo Luciferase Assay System (N1130, Promega) or cell viability using CellTiter-Glo (G7571, Promega).

**Next-generation sequencing.** To prepare sequencing libraries, 100 µL of total RNAs were extracted from 5 × 10^7 replicon cells using RNeasy minikit (Qiagen, Gaithersburg, MD). Then, 2 µL of each sample was used to assess RNA quality using Agilent 2100 Bioanalyzer (Agilent Technologies, Santa Clara, CA), and the RNA integration numbers (RIN) were all greater than 9. A total of 300 ng of total RNA was used to prepare the sequencing library using Illumina Stranded Total RNA Prep, Ligation with Ribo-Zero Plus. After rRNA removal, adaptor ligation, and cDNA library concentration and normalization, prepared libraries were loaded onto a NextSeq sequencer (Illumina, San Diego, CA) for deep sequencing of paired-end reads of 2 × 74 cycles. The numbers of reads mapped to the constructed virus genome range between 27,000 and 544,000 for individual samples. Variant calling was performed using Qiagen CLC Genomics Workbench V20 low-frequency variant detection with the requirement of significance of ≥ 5% and minimum frequency of ≥ 20%. For canonical sgRNA identification, a set of six sequences were constructed based on the replicon genome, each consisting of 49 nucleotides upstream of the 6-bp transcription regulatory sequence (TRS) motif (ACGAAC) found in the Leader sequence, the TRS-B sequence and 50 nucleotides downstream of each TRS-B, which extend into the coding region of each ORF. The specific sequences covering the junctions are shown in Table 2.

### Table 2: Canonical sgRNA sequences

| Name of subgenomic mRNA | Sequence of subgenomic mRNA |
|------------------------|-----------------------------|
| sgmNanoLuc             | CAGTGAACAAACCCCACTTGTGAGCTTTGTCTTCTAAACGACAAATGGTCTTTCACTCGAAGATTTCGTTGGGAGCTTGCGGACAGACACCGGG (Green, leader sequence; red, TRS-B; orange, NanoLuc) |
| sgmORF3a               | CAGTGAACAAACCCCACTTGTGAGCTTTGTCTTCTAAACGACAAATGGTCTTTCACTCGAAGATTTCGTTGGGAGCTTGCGGACAGACACCGGG (Green, leader sequence; red, TRS-B; orange, ORF3a) |
| sgmNeoR                | CAGTGAACAAACCCCACTTGTGAGCTTTGTCTTCTAAACGACAAATGGTCTTTCACTCGAAGATTTCGTTGGGAGCTTGCGGACAGACACCGGG (Green, leader sequence; red, TRS-B; orange, ORF3a) |
| sgmORF7                | CAGTGAACAAACCCCACTTGTGAGCTTTGTCTTCTAAACGACAAATGGTCTTTCACTCGAAGATTTCGTTGGGAGCTTGCGGACAGACACCGGG (Green, leader sequence; red, TRS-B; orange, ORF7) |
| sgmORF8                | CAGTGAACAAACCCCACTTGTGAGCTTTGTCTTCTAAACGACAAATGGTCTTTCACTCGAAGATTTCGTTGGGAGCTTGCGGACAGACACCGGG (Green, leader sequence; red, TRS-B; orange, ORF8) |
| sgmNP                  | CAGTGAACAAACCCCACTTGTGAGCTTTGTCTTCTAAACGACAAATGGTCTTTCACTCGAAGATTTCGTTGGGAGCTTGCGGACAGACACCGGG (Green, leader sequence; red, TRS-B; orange, NP) |
Short, paired-end reads of RNA samples from 12 stable replicon cell clones were uploaded and analyzed on the NGS platform high performance integrated virtual environment (HIVE) (26). The reads were indexed, deduplicated, and quality metrics were collected upon data ingestion, which verified the high quality of the reads (Fig. S3). Alignment of the reads was performed with HIVE’s native (27) against seven reference sequences. HIVE Hexagon default parameters, tuned for viral analysis allowing for small indels and mutations, were used and a threshold of 65 bases or longer of the aligned query was applied. Given the fact that the length of the reads was 70 bp and the way the subject sequences were constructed, the alignments returned only split reads with patterns from both sides of the investigated junctions.

**MD simulations.** We carried out all-atom MD simulations for the complex of nsp1 and the fragment of rRNA (Charcoal gray in Fig. 2) using the NAMD2.13 package (28) running on the IBM Power Cluster. The atomic coordinates for the complex (a bound state) were obtained from the crystal structure (PDB code: 7K5I) (18). The complex was further solvated in a cubic water box that measures about 78 × 78 × 78 Å³. Na⁺ and Cl⁻ were added to neutralize the entire simulation system and set the ion concentration to be 0.15 M (Fig. S1A). The final simulation system comprises 47,971 atoms. The built system was first minimized for 10 ps and then equilibrated for 1,000 ps in the NPT ensemble (P = 1 bar and T = 300 K), with atoms in the backbones (of both nsp1 and RNA) harmonically restrained (spring constant k = 1 kcal/mol/Å²). The production run (~200 ns) was performed in the NPT ensemble, when only constraining the terminals of nsp1 (both N- and C-terminals) and rRNA (both 5' and 3'-terminals). The same approach was applied in the production run for nsp1 in a 0.15 M NaCl electrolyte (Fig. S1B), a free state required in the free energy perturbation (FEP) calculations (see below). The water box for the nsp1-only simulation also measures about 78 × 78 × 78 Å² (29). Note that the similar system size for the bound and free states are required for free energy perturbation calculations for mutations with a net charge change.

We used the CHARMM36m force field (29) for proteins and rRNA, the TIP3P model for water (30, 31), the standard force field (32) for Na⁺ and Cl⁻. The periodic boundary conditions (PBC) were applied in all three dimensions. Long-range Coulomb interactions were computed using particle-mesh Ewald (PME) full electrostatics with the grid size of about 1 Å in each dimension. The pairwise van der Waals (vdW) energies were calculated using a smooth (10 to 12 Å) cutoff. The temperature T was kept at 300 K by applying the Langevin thermostat (33), while the pressure was maintained constant at 1 bar using the Nosé-Hoover method (34). With the SETTLE algorithm (35) enabled to keep all bonds rigid, the simulation time-step was 2 fs for bonded and non-bonded (including vdw, angle, improper and dihedral) interactions, and the time-step for Coulomb interactions was 4 fs, with the multiple time-step algorithm (36).

**Free energy perturbation calculations.** After equilibrating the structures in bound and free states, we performed FEP calculations (37). In the perturbation method, many intermediate stages (denoted by λ) whose Hamiltonian \( H(\lambda) = \lambda H_I + (1-\lambda) H_F \) are inserted between the initial (\( H_I \)) and final (\( H_F \)) states to yield a high accuracy. With the softcore potential enabled, \( \lambda \) in each FEP calculation for the bound or free state varies from 0 to 1 in 20 perturbation windows (lasting 300 ps in each window), yielding gradual and progressive annihilation and exhillination processes for mutations at residue 164 (K to A), 165 (H to K), 171 (R to A), 167 (S to A), 171 (R to P), and 175 (R to A), respectively. In FEP runs for the K164A mutation, the net charge of the MD system changed from 0 to −1 e (where e is the elementary charge). It is important to have similar sizes of the simulation systems for the free and bound states (38, 39) so that the energy shifts from the Ewald summation (due to the net charge in the final simulation system) approximately cancel out when calculating \( \Delta G \). The same approaches were applied to investigate mutations of R171A and R175A. More detailed procedures can be found in our previous work (40, 41).

**In vitro cytotoxicity assay and CC₅₀ determination.** Cytotoxicity was determined by cell viability assay as previously described. In brief, the cell viability was measured using Cell-Titer Glo (Promega) according to the manufacturers’ instructions, and luminescence signals were measured by GloMax luminometer. CC₅₀ values were calculated using a nonlinear regression curve fit in Prism Software version 9 (GraphPad). The reported CC₅₀ values were the results of at least three biological or technical replicates.

**Immunofluorescence.** Stable cells were plated on collagen-coated glass coverslips into a 24-well plate at 1 × 10⁴ cells/well 2 days before fixation. The cells were washed for 5 min three times with 1X phosphate-buffered saline. After wash, cells were fixed in 4% paraformaldehyde for 15 min and then permeabilized in 0.2% Triton X-100 at room temperature. An anti-dsRNA antibody (Kerafast, rJ2) was added at 1:40 for overnight at 4°C followed by incubation with 1:5,000 Alexa 568 conjugated Goat anti-mouse antibody. Images were captured by a Leica STELLARIS laser scanning confocal microscope.

**SUPPLEMENTAL MATERIAL**
Supplemental material is available online only.

**SUPPLEMENTAL FILE 1**, PDF file, 2 MB.
**SUPPLEMENTAL FILE 2**, XLSX file, 0.1 MB.

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