Amilorides inhibit SARS-CoV-2 replication in vitro by targeting RNA structures

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The SARS-CoV-2 pandemic, and the likelihood of future coronavirus pandemics, emphasized the urgent need for development of novel antivirals. Small-molecule chemical probes offer both to reveal aspects of virus replication and to serve as leads for antiviral therapeutic development. Here, we report on the identification of amiloride-based small molecules that potently inhibit OC43 and SARS-CoV-2 replication through targeting of conserved structured elements within the viral 5′-end. Nuclear magnetic resonance–based structural studies revealed specific amiloride interactions with stem loops containing bulge-like structures and were predicted to be strongly bound by the lead amilorides in retrospective docking studies. Amilorides represent the first antiviral small molecules that target RNA structures within the 5′ untranslated regions and proximal region of the CoV genomes. These molecules will serve as chemical probes to further understand CoV RNA biology and can pave the way for the development of specific CoV RNA–targeted antivirals.

INTRODUCTION

Severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) is the etiological agent of the COVID-19 respiratory disease, the largest-scale respiratory virus pandemic the world has witnessed since the 1918 Spanish flu and that has claimed more than 3.9 million lives worldwide as of July 2021 (1–3). Coronaviruses (CoVs) generally cause mild flu-like symptoms in humans but have caused two smaller-scale pandemics in the last two decades: SARS-CoV (2003) and MERS (2012) (4). Recent phylogenetic mapping traced all human CoVs to animal origins (5). While the middle zoonotic carrier of the virus between the animal of origin and humans seems to vary between CoVs, the chronological surfacing of human CoV pandemics seems to follow a dangerous trend of increasing lethality of each pandemic, thereby underscoring the need for a better understanding and targeting of the current and future CoV etiologic agents.

After more than a year since the first cases of SARS-CoV-2 human infection, this virus is expected to remain a global threat until vaccines are available and adopted worldwide. Furthermore, despite the increase in number of approved vaccines, their implementation has been hindered by the scarcity of doses available worldwide. On top of the production rate, pharmaceutical companies are currently facing challenges with the recently reported SARS-CoV-2 variants, against which not all vaccines have proven sustained efficacy, highlighting the need for a synergistic antiviral-based approach (6). While recent treatments have been approved for use within hospital settings, there are no known U.S. Food and Drug Administration (FDA)–approved cures for the infection (7).

Current candidates for treatment have limited approval for emergency use in severe COVID-19 cases. Remdesivir, for example, is an RNA-dependent RNA polymerase inhibitor (RdRp) initially developed during the Ebola outbreak and revisited at the start of the pandemic (8). The compassionate use of the candidate antiviral across many countries reported mixed results, with overall faster recovery time from the virus but no difference in mortality rates (9). While more randomized trials are needed for a final verdict on the efficacy of remdesivir in critical patients, its stereospecific multistep synthetic process highlights the need for new, scalable, and more efficacious antivirals. Baricitinib has been recently approved for emergency use for COVID-19 treatment in conjunction with remdesivir. Also known as Olumiant, this small molecule was approved in 2018 as treatment for moderate to severe rheumatoid arthritis (10). It is proposed that the anti-inflammatory effects of the drug help in decreasing inflammatory cascades associated with COVID-19. While promising, baricitinib has yet to receive approval as a stand-alone treatment and, so far, has been shown to improve recovery time by 1 day when compared with remdesivir-alone treatment (7).

The recent emergence of multiple coronavirus pandemics clearly indicates that SARS-CoV-2 most likely will not be the last CoV pandemic (4). The current limited tools and lack of cures underscore the need for a new approach in developing antivirals that would not only provide novel routes to combat the current pandemic but also provide invaluable information on targetable structures that can aid in the prevention of and fight against future CoV outbreaks. Small molecules are uniquely poised to achieve this goal as their design and development provide both a better understanding of CoV biology and identify druggable targets that could aid in the development of pan-coronavirus antivirals.

Several steps in the coronavirus replication cycle offer potential therapeutic targets for viral inhibition (Fig. 1). CoVs are enveloped...
positive-sense single-stranded RNA genomes of approximately 30 kilobases, making them the largest genomes of RNA viruses (4). SARS-CoV-2 infects human cells by engagement of the ACE2 receptor by the viral spike (S) protein followed by membrane fusion at the plasma membrane or endosomal membranes depending on the availability of host cell proteases that cleave and prime S for entry. Fusion results in the release of its genome and associated proteins in the host cell cytosol. The genome is translated into two large polyproteins, 1a and 1b, which are then processed into individual proteins by the viral protease. Synthesis of full-length negative-strand RNA by products of 1a/1b creates a template to synthesize multiple positive strand copies encapsidated by the viral nucleoprotein into virions (4). The negative RNA strand also serves as a template for the synthesis of shorter subgenomic RNAs that include the essential structural proteins and thus also constitute an attractive therapeutic target (11).

CoV antivirals to date have been developed to target viral proteins, including to prevent endocytosis, assembly of viral protein for export, and condensation of viral genome for packaging (12). While this protein-centric approach has been proven successful in a few cases, the sequence and structural conservation of RNA structural motifs pose an attractive complementary target for small-molecule antiviral development, a strategy that has shown promise against a plethora of viruses (13, 14). Specifically, international efforts that allowed for identification and tracking of SARS-CoV-2 variants highlighted the large number of mutations accumulated in protein-coding regions (15). At the same time, recent global cataloging of mutations in the untranslated regions (UTRs) revealed a lower rate of mutation, highlighting the potential of UTRs as drug targets (16).

Recently published data on genome-wide secondary structure of the virus obtained by in vitro and in vivo SHAPE (selective 2′ hydroxyl acylation analyzed by primer extension) and DMS (dimethyl sulfate) probing of SARS-CoV-2–infected Vero E6 cells, as well as nuclear magnetic resonance (NMR) characterization, recapitulates the computationally predicted stem loops (SLs) at the 5′-region of the genome (Fig. 2) as well other relevant frameshifting and replication-related structures (17–20). Conserved elements at the 5′- and 3′-ends have been identified across many members of the coronavirus family and function as cis-acting elements regulating viral replication (21). Specifically, studies on murine and bovine CoVs showed that phylogenetically conserved SLs in the 5′-UTR are capable of long-range RNA-RNA and protein-RNA interactions responsible for optimal viral replication (22, 23). More recently, studies aimed at uncovering
the pathway that leads to viral protein synthesis via host cell translation machinery revealed that the presence of the full-length 5′-UTR of SARS-CoV-2 leads to a fivefold increase in translation of viral proteins. This preliminary data corroborate the importance of the 5′-UTR region, and the structures within, for efficient viral translation and provide context for the viral hijacking of the host cell translational machinery (24).

Drug-like small molecules offer the ability to develop chemical probes that reveal function and to design bioavailable clinical candidates for treatment. While RNA targeting has lagged behind protein targeting, recent successes in both the laboratory and the clinic support its potential role. The first U.S. FDA–approved small molecule targeting RNA other than the ribosome was approved for treatment of spinal muscular atrophy in August of 2020 (25). Effective small-molecule targeting in the laboratory has also been observed for a plethora of disease-relevant RNAs, including viral RNAs. Specifically, small molecules targeting structures within the 5′-UTR region have shown antiviral activity for a number of positive-sense RNA viruses such as HCV, FMDV, and EV71 (26–28). Recent studies have begun evaluating the potential of small molecules against the frameshifting elements of SARS-CoV-2 RNA in the coding region. These recent reports include evaluation of known SARS-CoV pseudoknot binders and development of a small-molecule binder to the attenuator hairpin preceding the pseudoknot (29–32). Coupling of the latter to the known ribonuclease-targeting chimera technology results in recruitment of cellular ribonucleases leading to viral genomic RNA degradation. In addition, a fragment-based screen against structures in the 5′-UTR of SARS-CoV-2 was successful in identifying potential ligands (31). While the antiviral activity of these molecules has not been published, these preliminary results, combined with known RNA-targeted antivirals for other positive-sense RNA viruses, stand as proof of concept of the targetability of SARS-CoV-2 RNA motifs.

Successful efforts to target viral 5′-UTR structures have often leveraged the synthetic tuning of a known RNA binding scaffold (26, 27, 33). We recently used this approach to develop RNA-targeted antivirals based on dimethylamiloride (DMA) that target the internal ribosomal entry site (IRES) region in the 5′-UTR of EV71. The DMA scaffold had been previously reported to be poised for tuning for specific RNA constructs via a facile three-step synthesis (27). Further investigation and functionalization of the scaffold resulted in a bioactive antiviral analog that formed a repressive ternary complex with IRES SL II RNA and the human AUF1 protein, ultimately inhibiting translation and compromising viral replication. The successes in DMA exploration highlight the scaffold’s potential for tuning and broad applicability as an antiviral scaffold (27).

Here, we report DMA analogs that show promising antiviral properties by reducing SARS-CoV-2 virus titer in a dose-dependent manner in infected cells. In addition, dual-luciferase reporter assays confirmed the antiviral activity of the small molecules to be dependent on the 5′-UTR and proximal region of the SARS-CoV-2 genome. Investigation of possible conserved RNA binding sites of the lead small molecules revealed putative bulge-like binding sites in SL1, SL4, and SL5a, located in the 5′-UTR of the SARS-CoV-2 genome, and in the adjacent SL6 located in ORF1a, supporting both the targetability of 5′-region SLs. Together, these results establish conserved CoV RNA structures as antiviral targets and reveal lead molecules with promising antiviral properties.

**RESULTS**

**Phylogenetic conservation of RNA structural elements**

As the functional significance of the 5′-end SLs is still being elucidated, we examined sequence conservation in the 5′-end region across the *Betacoronavirus* genus as a preliminary approach to assess the
suitability of the known structures in this region as therapeutic targets. Sequence conservation within the contexts of folded RNA domains would imply selective pressures to exert biological function. We constructed a multiple sequence alignment spanning the 5′-UTR and the adjacent region including representatives of all five betacoronavirus subgenera (Fig. 3). Alignments disclosed the highest degree of conservation in the region encoding SL2 and SL3, both of which are relatively short and contain stretches of 5 to 6 nucleotides (nt) that are 100% conserved. By contrast, SL1 and SL4 to SL6 are less conserved, but notably, four of the five SLs that span the 5′-UTR contain stretches of relatively highly conserved nucleotide sequence (i.e., 70 to 100% conservation across the genus). The position of these SL’s within or adjacent to the 5′-UTR (which is relatively conserved in length across all Betacoronaviruses) indicates that these conserved regions are likely to represent homologous nucleotides and suggests that they are, to some degree, functionally equivalent. Our findings are in agreement with recent studies that reported sequence and structural similarity among members of the Coronaviridae family, suggesting that selective pressure plays a central role in conserving RNA secondary structures essential for the viral life cycle.

**DMAs inhibit human coronavirus OC43 virus replication in a dose-dependent manner**

We pursued targeting of the 5′-end SL structures with the DMA scaffold, which has been previously reported as an RNA binding scaffold that can be successfully optimized for selectivity for distinct RNA elements (27). To quickly assess potential CoV antiviral activity, human OC43 betacoronavirus was used because of its lower virulence and thus suitability for use in standard cell culture facilities (3). Vero E6 cells were infected with human coronavirus OC43 at a multiplicity of infection (MOI) = 1. After a 1-hour adsorption, a panel of 23 DMAs at 50 or 100 μM concentration was added to the cells and incubated for 24 hours at 33°C. Virus titers were determined by plaque formation on Vero E6 cells, and DMA-132, DMA-135, and DMA-155 (Fig. 4) reduced virus titer by ~1000-fold at 100 μM concentration (fig. S1). The results further suggest DMA-132, DMA-135, and DMA-155 reduce virus titer in a dose-dependent manner (fig. S2). The parent scaffold, DMA (DMA-1), demonstrated no activity and is used as an inactive control moving forward.

**Antiviral potency of DMAs against SARS-CoV-2**

To determine the antiviral activity and potency of the lead small molecules against SARS-CoV-2, we used a simplified quantitative reverse transcription polymerase chain reaction (qRT-PCR) assay to monitor SARS-CoV-2 viral RNA levels in supernatants of infected Vero E6 cells (34). Similarly to remdesivir, DMA-135 and DMA-155 led to a dose-dependent 10- to 30-fold decrease in cell-free viral RNA levels within 24 hours of infection with an approximate median inhibitory concentration (IC50) of 10 and 16 μM, respectively (fig. S4).

Antiviral activity of the three most active DMA leads (DMA-132, DMA-135, and DMA-155) was confirmed using Vero E6 cells infected with wild-type SARS-CoV-2. Small-molecule treatment was performed at 10 and 50 μM, respectively. DMA-132 and DMA-135 showed dose-dependent reduction in virus titer compared to dimethyl sulfoxide (DMSO), as measured by median tissue culture infectious dose (TCID50) assay, without measurable effect on cellular viability as measured by adenosine 5′-triphosphate (ATP) content (Fig. 5).

**Investigation of small-molecule activity against CoV-2–luciferase reporter gene expression**

To assess the effect of DMAs on reporter gene expression directed by SARS-CoV-2 sequence elements, a reporter plasmid, pCoV-2–5′UTR–FLuc–3′UTR was used as template for in vitro synthesis of CoV-2–5′UTR–FLuc–3′UTR RNA (Fig. 6A). This plasmid contains the 5′-end 805-nt segment from the virus genome and the 3′-UTR. Thus, the 805-nt segment spans the genomic RNA 5′-UTR (SL1 to SL5) and ORF1a encoding a portion of nsp1 (including SL6 to SL8) fused in-frame with the firefly luciferase (FLuc) ORF. Plasmid pRL was used as template for the synthesis of control Renilla luciferase (RLuc) reporter RNA lacking SARS-CoV-2 sequences. The RNAs were cotransfected, and various concentrations of DMAs were added with the transfections. Two days after transfection, RLuc and FLuc activities were measured using a dual-luciferase reporter assay. As shown in Fig. 6A, addition of 10 μM DMA-132 or DMA-135 reduced FLuc activity, which is under the control of SARS-CoV-2 5′-end and 3′-UTR, by approximately 50%. Addition of 10 μM DMA-155 resulted in the largest decrease in FLuc activity; specifically, FLuc signal was reduced by ~90%. A similar level of FLuc signal reduction was achieved in the presence of 10-fold higher concentration of DMA-132 and DMA-135 when compared with DMA-155. Activities of the control RLuc remained relatively constant for all DMAs across all concentrations tested; this control indicates that decreases in FLuc required SARS-CoV-2 sequence elements and were not due to any putative cytotoxic effects of DMAs or to nonspecific effects on translation. To test whether the CoV-2 3′-UTR contributes to DMA-mediated translational repression, we repeated the experiment using a FLuc reporter, CoV-2–5′UTR–FLuc RNA, in which the CoV-2 3′-UTR was replaced with vector-encoded sequence; thus, the CoV-2 5′-ends of both FLuc reporter RNAs are the same (Fig. 6B). The effects of DMAs on translational repression were virtually identical to those in Fig. 6A. We note that select DMAs that inhibited SARS-CoV-2 5′-end–dependent FLuc translation (this work) had little effect on EV71 5′-UTR–dependent FLuc translation (27). Thus, the DMAs act at the level of the SARS-CoV-2 5′-end and not at the level of putative structure present in the FLuc coding region. Together, these results clearly demonstrate that DMA-dependent suppression of SARS-CoV-2 luciferase reporter activity requires only 5′-end sequences of the virus.

**NMR profiling and in vitro affinity of 5′-SL-DMA interactions**

Toward understanding potential mechanisms by which the DMAs inhibit SARS-CoV-2 replication, we carried out single-point 11C,1H TROSY HSQC (transverse relaxation-optimized spectroscopy–heteronuclear single quantum coherence) titrations of DMA-132,
DMA-135, and DMA-155 into SL1 to SL6 of the 5′-region. We selected this region of the 5′-UTR to NMR profile following the reasoning that it contains SLs with assigned functions that can be prepared at sizes amenable to facile NMR screening, shows a degree of selective pressure to maintain sequence and structure, and contains several noncanonical RNA elements that we hypothesize are good targets for the DMAs. An advantage of using 13C-1H TROSY HSQC titrations to profile small molecule–RNA interactions is
that the extent of NMR signal perturbation provides a convenient proxy on the degree of binding specificity, even in the absence of chemical shift assignments. The designs of the individual SLs were based on the reasoning that DMAs likely bind the 5′-region SLs at noncanonical structural elements such as bulge, internal, or apical loops. Therefore, we in vitro transcribed isolated SL domains using 13C, 15N rNTPs (ribonucleoside triphosphate) that would maximize NMR signal detection for the noncanonical elements over the base paired regions (Fig. 7 and fig. S5). For example, SL1 was prepared separately as A(13C, 15N) and C(13C, 15N) selectively labeled constructs because adenosines and cytosines are the most abundant nucleobases within or proximal to its internal and apical loops, respectively. Using this strategy, we are able to efficiently profile each 5′-region SL to determine whether the DMAs bind with reasonable affinity and specificity as determined by qualitative assessment of the DMA-induced NMR signal perturbation. Figure 7 and fig. S5 summarize the effects that the addition of excess (5:1) DMAs has on the NMR spectra of each major SL domain. First, we observed that the DMAs induced differential chemical shift perturbations (CSPs) or line broadening (LB) for each SL, and the largest signal perturbations (CSPs or LB) were observed in spectra recorded for SL1, SL4, SL5a, and SL6 (Fig. 7). The assignments of SL1 and SL5a have been previously reported (35, 36). Each of these 5′ domains contains bulges and/or other internal loops. Second, only a subset of the 13C-1H correlation peaks is perturbed in the spectra, providing initial evidence that the DMAs interact through specific surfaces rather than delocalized binding. Third, the extent of the signal perturbations is also differential between DMAs, with some DMAs inducing shifts of the correlation peaks to new positions within the spectra of a given SL and others inducing severe LB of the NMR signals. These variable signal perturbations suggest that the DMAs interact with different binding affinities and binding modes. None of the DMAs caused substantial changes to spectra recorded on SL2, which contains a 5-nt CUUGU apical loop but no internal loops or bulges. Together, the single-point 13C-1H TROSY HSQC titrations provide compelling evidence that the DMAs make specific interactions with the SARS-CoV-2 5′-region via surfaces composed of noncanonical structural elements.

Fig. 4. Chemical structure of the three lead molecules derived from the focused library screen against OC43-infected Vero E6 cell screening.

Fig. 5. Inhibition of SARS-CoV-2 infection by DMA leads without cellular toxicity. Vero E6 cells were infected with SARS-CoV-2 (MOI = 0.1) in the presence of DMSO or the indicated compounds for 3 days followed by normalized measurement of virus titer in culture supernatant (black bars). Results were generated with two independent experiments, each with two replicates. Cellular viability was assayed by measuring cellular ATP content in uninfected Vero E6 cells after 3 days of treatment with the indicated compounds (gray bars).

Fig. 6. Vero E6 cells were cotransfected with luciferase constructs and cultured with various concentration of small molecule. (A) Vero E6 cells were cotransfected with CoV-2-5′UTR-Fluc-3′UTR. (B) Vero E6 cells were cotransfected with CoV-2–5′UTR-Fluc (left), and no change in inhibition was observed when compared to the presence of 3′-UTR (right, DMA-135). Luciferase activity was measured 2 days later. Mean values and SDs from three independent experiments are shown in the bar graphs. ***P < 0.001; **P < 0.01; N., not significant relative to the DMSO control, except for the 5′UTR-Fluc-3′UTR comparison in (B), which is relative to DMA-135 at 0.1 μM.
Fig. 7. Single-point $^{13}$C-$^1$H TROSY HSQC titrations reveal that DMA-132, DMA-135, and DMA-155 bind with moderate affinity and specificity to SARS-CoV-2 5'-region SLs. The spectra were recorded at 900 MHz in 100% D$_2$O buffer of 25 mM K$_2$HPO$_4$ and 50 mM KCl at pH 6.2. Temperatures (298, 303, or 308 K) were optimized for each RNA construct to maximize the number of observed correlation peaks. The total RNA concentrations were set to 100 µM while titrating fivefold excess DMA.
Notably, SL1 and SL6 experienced the most substantial NMR signal perturbations upon addition of excess DMAs (Fig. 7). NMR signal perturbations are observed at other SLs but to a lesser degree based on our selection criteria. For SL1, addition of DMA-132 and DMA-155 induced the migration of several correlation peaks to new spectral positions (Fig. 7). By comparison, addition of DMA-135 induced severe LB of ~50% of the total observed correlation peaks recorded for the A(13C, 15N) and C(13C, 15N) selectively labeled SL1 constructs (Fig. 7 and fig. S5). In the presence of fivefold excess DMA-135, most of the nuclear Overhauser effect (NOE) stacking patterns disappear for a CG(\textsuperscript{H}), AU(\textsuperscript{H}3,\textsuperscript{g}) selectively labeled SL1 construct, whereas substantially more NOE cross-peaks are still observed for the 5:1 (DMA-155)–SL1 complex (fig. S6). These differences likely reflect binding heterogeneity and collectively suggest that DMA-132, DMA-135, and DMA-155 differentially interact with SL1 despite having similar core scaffolds (Fig. 7).

For SL6, addition of excess DMA-155 induced a combination of CSPs and LB to a greater extent than DMA-132 or DMA-135. To further assess the nature of these signal perturbations on the SL6 structure, a \textsuperscript{1}H–\textsuperscript{1}H NOE spectroscopy (NOESY; \(t_m = 250\) ms) spectrum was recorded with \(\text{U}G(\text{H})\), \(\text{AC}(\text{H}3,\text{g})\) selectively labeled SL6 constructs, which was chosen based on the abundance of A and C residues in the bulge (Fig. 8A). Figure 8 shows the \textsuperscript{1}H–\textsuperscript{1}H NOESY spectra of free SL6 overlaid with its complex to which a 5:1 molar ratio of DMA-155 was added. On the basis of the intra-NOEs observed between A and C residues of SL6, tentative chemical shift assignments were determined (Fig. 8, B and C). The addition of excess small molecule caused CSPs to A339H8, A339H2, A310H8, A310H2, C311H6, and A314H8 spin systems, which are located within or in proximity to the bulge. This supports a model in which DMA-155 preferentially interacts with the bulge region of SL6. Notably, the presence of intra-NOEs in the complex indicates the preservation of base stacking within the (DMA-155)–SL6 complex. In addition, the \textsuperscript{1}H–\textsuperscript{1}H NOESY spectra of \(\text{CUG}(\text{H})\), \(\text{A}(\text{H}3,\text{g})\) selectively labeled SL6 overlaid with its DMA-155–bound form (Fig. 8C) show inter-NOEs between the methyl protons of DMA-155 [2.5 parts per million (ppm)] and both A339H2 (7.94 ppm) and A310H2 (7.88 ppm), although these particular inter-NOEs are weak. As further evidence of the involvement of A339 and A310 in binding, their H8 spin systems also showed CSPs upon the addition of DMA-155. Four additional broad NOE peaks appeared between the methyl protons and protons with chemical shifts between ~6.5 and 7 ppm (Fig. 8C). We reasoned that these might be intra-NOEs between the methyl groups and amino group located on the central six-membered ring system. To further investigate the origin of these NOEs, a \textsuperscript{1}H–\textsuperscript{1}H NOESY spectrum of \(\text{ACGU}(\text{H})\)-labeled SL6 bound to DMA-155 (shown in orange, Fig. 8C) displayed identical broad NOE peaks. This shows that these peaks signify intra-NOEs within DMA-155 itself, which do not appear in the \textsuperscript{1}H spectrum of the free small molecule in D\textsubscript{2}O but only in the environment of the SL6 complex. The possibility of these peaks arising from inter-NOEs between the amino of the RNA and methyl protons was eliminated on the basis of \textsuperscript{15}N-HSQC NMR experiments of DMA-155 titrated into \textsuperscript{15}N fully labeled SL6. In sum, through the support of intra- and inter-NOEs, DMA-155 shows a degree of specificity to the bulge region of SL6.

Moreover, the affinity of the three leads for the SL constructs was investigated via in vitro indicator displacement assays (IDAs) (27, 37, 38). DMA-155 was revealed to be a strong binder with the highest affinity for SL6, in agreement with NOE studies (Fig. 8 and fig. S17). DMA-135 showed weak affinity for SL1, SL4, and SL6; the parent scaffold (DMA-001) and DMA-132 did not show binding affinity against any of the constructs under these conditions. The measured 50% competitive displacement dose (CD\textsubscript{50}) in vitro is about 10-fold weaker than the IC\textsubscript{50} recorded via luciferase assays, which may highlight the differences between these assays. CD\textsubscript{50} measurements depend on both the affinity and binding mode of the indicator used, while in cell assays, these reflect binding in the presence of a larger RNA context and in the presence of proteins. Additional biophysical studies are underway to better understand the relationship between these in vitro and in cellulo assays.

**In silico ligand screening of DMA-focused library against 5′-UTR and adjacent RNA structures**

To generate potential three-dimensional (3D) models of each SL, we used fragment assembly of RNA with full-atom refinement (FARFAR). We chose FARFAR to generate preliminary models as it has consistently been demonstrated to be the most accurate RNA 3D prediction algorithm (39, 40). We generated between 5000 and 100,000 models for each SL and then generated ~15 representative clusters.

The lowest energy conformation from each cluster was used to generate an ensemble for each SL that was submitted to ICM pocket finder to find and characterize possible binding pockets (table S1). Notably, SL2, which showed no change in \textsuperscript{13}C–\textsuperscript{15}N HSQC NMR chemical shifts upon small-molecule addition, did not have any identifiable binding pocket. SL1, SL3, and SL5b presented binding pockets with low to intermediate scores in terms of volume, area, hydrophobicity, buriedness, and druggability score (DLID) parameters often used to describe a binding pocket’s fitness. Notably, SL3 and SL5b presented minor CSPs upon small-molecule addition. \(\text{\textsuperscript{13}}\text{C}\textsuperscript{-15}\text{H}\) TROSY HSQC NMR experiments showed notable changes upon small-molecule binding to SL1, SL5a, and SL6 (Fig. 7), all of which had the highest scores in all parameters (Fig. 9). The structure that presents the highest CSPs upon binding to the small-molecule leads, SL6, is the only SL to have a binding pocket with positive DLID score (0.45; table S1). DLID score has been found as a good predictor of druggability of protein binding pockets but often presents highly negative scores in RNA due to the charged backbone of RNA (41, 42). Overall, this binding pocket analysis strongly correlates with NMR experimental data.

We then docked our published 55-member DMA library against the clustered FARFAR-generated SL structures (figs. S7 to S12). In line with the results from the pocket analysis and NMR experimental data, the overall number of hits across constructs is highest for SL1, SL4, SL5a, and SL6, corroborating their potential as therapeutic targets. Hit ligands DMA-132, DMA-135, and DMA-155 were among the top predicted binding ligands. Furthermore, if the docked structures are limited to the sublibrary of DMAs tested against OC43 and SARS-CoV-2, the three hit ligands score in the top 5 of every SL. Notably, when analyzing the binding location that yielded the best docking scores of the three hit molecules against SL6, we note that while DMA-135 and DMA-155 bound best in one binding pocket (Fig. 9F, blue), DMA-132 docks best in the adjacent binding pocket (Fig. 9F, red). When comparing the NMR profiles of the three small molecules, we note that DMA-135 and DMA-155 have a similar perturbation pattern, while DMA-132 has a slightly different profile, suggesting a possible different binding mode. To further investigate this hypothesis, FARFAR 3D models were generated for every individual bulge and apical loop motif to increase the conformational

diversity at these sites within the SL clusters, which are otherwise largely dictated by the respective positions of the double stranded regions. The DMA library was then rescreened against each of the clustered motifs. The three small-molecule leads DMA-132, DMA-135, and DMA-155 consistently scored among the top hits (figs. S8 to S12). Furthermore, each of the molecules reported the best docking score for SL1, SL5a, SL6, and, more specifically, for motifs created for internal bulge motifs (figs. S13 to S15). In the case of DMA-135, for example, the best docking score was found for the internal loop motif of SL1, in agreement with HSQC data that reported DMA-135 as the best binder for SL1 (fig. S16). Analogously, DMA-155 had the best docking score for the internal loop pocket of SL6, in line with NOE data that identified DMA-155 as the best and most specific binder for SL6 (fig. S17). While preliminary, this docking analysis corroborates the NMR data and the fitness of these ligands for SARS-CoV-2 5′-SL targeting while also supporting its utility as a tool in the identification of new SARS-CoV-2 RNA-targeting ligands.

Fig. 8. Detailed interaction of DMA-155 with SL6. (A) Representation of SL6 secondary structure and labeled nucleotides (red). (B) $^{1}H-^{1}H$ NOESY spectra (900 MHz, $t_m = 250$ ms) of free UG(2H), AC(2H3′-5″) selectively labeled SL6 (blue) and its DMA-155 complex (red), which were collected in 25 mM K$_2$HPO$_4$ and 50 mM KCl (pH 6.2) at 308 K in 100% D$_2$O, show that DMA-155 has a degree of binding specificity. (C) $^{1}H-^{1}H$ NOESY spectra (900 MHz and $t_m = 250$ ms) of free CUG(2H), A(2H3′-5″) selectively labeled SL6 (red), its DMA-155 complex (black), and the fully deuterated RNA complexed with DMA-155 (orange) were collected in 25 mM K$_2$HPO$_4$ and 50 mM KCl (pH 6.2) at 308 K in 100% D$_2$O. On the secondary structure of SL6, the nucleotides highlighted in red represent the labeling scheme of the RNA.
probing. An initial DMA-focused library screen against OC43-adjacent SL6 ideal therapeutically relevant targets for small-molecule probes have been reported to bind most effectively, making the 5′-UTR and the internal loops, the secondary structural elements that amilorides have been synthetically tuned for a range of RNA secondary structures (43). Amiloride, a known RNA binding scaffold, has been found to bind preferentially to SLs that contain large internal or bulge loops. SL1’s NMR data revealed CSPs and LB with DMA-135, which signifies its binding onto a specific surface with a disruption in stacking interactions, unlike DMA-155. Notably, SL6, which contains a moderately conserved and weakly paired ~16-nt bulge loop, showed one of the most notable CSPs when titrated with DMAs. Specifically, NOE experiments revealed that DMA-155 binds specifically to SL6, a finding that was corroborated by in vitro IDAs, which identified SL6 as the preferred target for DMA-155. The 5′-side of the SL6 bulge loop has been proposed to interact with the SARS-CoV-2 nucleocapsid (N) protein under phase separation conditions and may also affect genome packaging (44). We will investigate the modulation of the N protein:SL6 interaction as a potential antiviral mechanism in future work.

In silico analysis corroborated the experimental trend observed with NMR experiments, identifying some of the SLs with predicted bulges (SL1, SL4, SL5a, and SL6) as those with binding pockets with highest druggability. The small molecules that showed the highest antiviral activity, namely, DMA-132, DMA-135, and DMA-155, scored highest in the SLs that reported substantial CSPs upon small-molecule binding. In addition, refined docking of the three small-molecule leads against motif-specific clusters revealed SL1 to be the preferred target for DMA-135 and SL6 as the favored target for DMA-155, in line with NMR and IDA results.

In summary, we here identified drug-like small molecules that reduce SARS-CoV-2 replication and are the first antivirals to target the conserved RNA SLs in the 5′-end region of SARS-CoV-2. Work is underway to further characterize the mode of action of these ligands, particularly putative impacts on RNA:protein interactions and specific steps in the viral replication cycle. These leads are uniquely poised to further elucidate the relationship between in vitro preferential binding and small molecule–mediated SL–specific alterations of virus:host interactions. These small molecules offer the opportunity to understand the contribution of individual 5′-end SLs to viral proliferation in a system where mutational studies are difficult because of genome size. Once characterized, we expect these amiloride-based ligands to serve as chemical biology tools to help understand CoV RNA molecular biology, such as N-dependent genome packaging and other cellular stages of the viral RNA replication process. We have established an efficient framework to identify novel RNA-targeted CoV antivirals that will serve not only the SARS-CoV-2 pandemic but future coronavirus pandemics.

### DISCUSSION

Screening of synthetic RNA-focused libraries in recent years has provided the field with many RNA binding bioactive small molecules and some of the highest hit rates among small-molecule screening approaches (43). Amiloride, a known RNA binding scaffold, has been synthetically tuned for a range of RNA secondary structures and recently yielded a novel antiviral lead for the treatment of enterovirus 71. In this case, amiloride inhibited viral translation by binding the viral 5′-UTR and modulating RNA:host protein interactions. SARS-CoV-2 also contains a highly conserved 5′-end that is reported to play a crucial role in viral replication and hijacking of host cell translational machinery. The presence of multiple bulge or internal loops, the secondary structural elements that amilorides have been reported to bind most effectively, makes the 5′-UTR and the adjacent SL6 ideal therapeutically relevant targets for small-molecule probing. An initial DMA-focused library screen against OC43-infected Vero E6 cells allowed for the identification of three lead compounds, namely, DMA-132, DMA-135, and DMA-155, that significantly reduced virus titer. Initial structure-activity relationships could also be resolved, highlighting the critical substitution of the dimethylamine group at the C5 position and rigid aromatic substituents at the C6 position. These molecules were also found to be active against SARS-CoV-2 in both screening-format qRT-PCR and infectious virus titer assays. Luciferase assays revealed the presence of the 5′-UTR and proximal region as necessary and sufficient for translation inhibition upon small-molecule treatment. Markedly, all in vitro assays identified DMA-155 as the strongest SARS-CoV-2 inhibitor, with the largest reduction in luciferase signal and highest decrease in virus titer at 10 μM.

NMR profiling of leads DMA-132, DMA-135, and DMA-155 against each of the major 5′-UTR and adjacent SL domains revealed that the DMAs bind preferentially to SLs that contain large internal or bulge loops. SL1’s NMR data revealed CSPs and LB with DMA-135, which signifies its binding onto a specific surface with a disruption in stacking interactions, unlike DMA-155. Notably, SL6, which contains a moderately conserved and weakly paired ~16-nt bulge loop, showed one of the most notable CSPs when titrated with DMAs. Specifically, NOE experiments revealed that DMA-155 binds specifically to SL6, a finding that was corroborated by in vitro IDAs, which identified SL6 as the preferred target for DMA-155. The 5′-side of the SL6 bulge loop has been proposed to interact with the SARS-CoV-2 nucleocapsid (N) protein under phase separation conditions and may also affect genome packaging (44). We will investigate the modulation of the N protein:SL6 interaction as a potential antiviral mechanism in future work.

In silico analysis corroborated the experimental trend observed with NMR experiments, identifying some of the SLs with predicted bulges (SL1, SL4, SL5a, and SL6) as those with binding pockets with highest druggability. The small molecules that showed the highest antiviral activity, namely, DMA-132, DMA-135, and DMA-155, scored highest in the SLs that reported substantial CSPs upon small-molecule binding. In addition, refined docking of the three small-molecule leads against motif-specific clusters revealed SL1 to be the preferred target for DMA-135 and SL6 as the favored target for DMA-155, in line with NMR and IDA results.

In summary, we here identified drug-like small molecules that reduce SARS-CoV-2 replication and are the first antivirals to target the conserved RNA SLs in the 5′-end region of SARS-CoV-2. Work is underway to further characterize the mode of action of these ligands, particularly putative impacts on RNA:protein interactions and specific steps in the viral replication cycle. These leads are uniquely poised to further elucidate the relationship between in vitro preferential binding and small molecule–mediated SL–specific alterations of virus:host interactions. These small molecules offer the opportunity to understand the contribution of individual 5′-end SLs to viral proliferation in a system where mutational studies are difficult because of genome size. Once characterized, we expect these amiloride-based ligands to serve as chemical biology tools to help understand CoV RNA molecular biology, such as N-dependent genome packaging and other cellular stages of the viral RNA replication process. We have established an efficient framework to identify novel RNA-targeted CoV antivirals that will serve not only the SARS-CoV-2 pandemic but future coronavirus pandemics.

### MATERIALS AND METHODS

#### Analysis of sequence conservation

Representative betacoronavirus sequences were selected according to official taxonomy as represented by the International Committee for the Taxonomy of Viruses (45). Multiple sequence alignments of
coronavirus sequences were constructed using BLAST and MAFFT as implemented in GLUE (46–48). Alignments were manually inspected and adjusted using Se-Al. Position coverage and percentage identity were calculated and visualized using JalView (49).

**Cells and virus**

Vero E6 (African green monkey kidney; ATCC CRL-1586) cells were cultured in minimum essential medium (MEM) supplemented with 10% fetal bovine serum (FBS) (Thermo Fisher Scientific) and maintained at 33°C. Cells were infected with human coronavirus OC43 cultured in minimum essential medium (MEM) supplemented with Vero E6 (African green monkey kidney; ATCC CRL-1586) cells were inspected and adjusted using Se-Al. Position coverage and percentage identity were determined by plaque assay on Vero E6 cells. SARS-CoV-2 fresh medium with various concentrations of DMAs. Media from (ATCC VR-1558) at indicated MOI and incubated 1 hour at 33°C for adsorption. Unbound virus was removed, and cells were refed fresh medium with various concentrations of DMAs. Media from infected cells were harvested 24 hours postinfection, and virus titers were determined by plaque assay on Vero E6 cells. SARS-CoV-2 diluted in Dulbecco’s modified Eagle’s medium (DMEM)/2% FBS for a solution of 20,000 plaque-forming units (pfu)/ml. Growth media was aspirated from 24-well plates and replaced with 495 μl of DMEM/2% FBS containing SARS-CoV-2 (20,000 pfu/ml) for an MOI of 0.1. Five microliters of DMSO or compound was then immediately diluted into each well for final concentrations of 50 and 10 μM. Plates were incubated at 37°C for 72 hours. Media were then harvested, centrifuged at room temperature for 10 minutes at 1500g, and then used for TCID₅₀ assay. Serial dilutions of supernatant from the treated cells were added to Vero E6 cells in 96-well plates, and cells were monitored for cytopathic effect. Virus titer was calculated from the numbers of positive wells using a modified Reed and Muench method.

**Effects of SARS-CoV-2 5′- and 3′-end sequence elements on luciferase reporter activity**

The reporter plasmid pCoV-2–5′UTR–FLuc–3′UTR contains the SARS-CoV-2 5′-UTR and adjacent coding sequences in ORF1a fused in-frame with the FLuc open reading frame, followed by the SARS-CoV-2 3′UTR. For plasmid pCoV-2–5′UTR–FLuc, the CoV-2 3′UTR was replaced with vector-encoded sequence. The plasmids were provided by S.-R. Shih (Chang-Gung University, Taiwan). Plasmid pRL, the Renilla luciferase control reporter vector, was purchased from Promega. CoV-2–5′UTR–FLuc–3′UTR, CoV-2–5′UTR–FLuc, and RLuc RNAs were in vitro synthesized from these plasmid templates using the MEGAscript T7 Transcription Kit (Thermo Fisher Scientific).

Vero E6 cells were seeded in 24-well plates. Two hundred nanograms of reporter RNA, 5 μl of SuperFect (Qiagen), and 400 μl of MEM with 10% FBS were combined and added to one well of cells. Cells were incubated at 33°C for 4 hours, media were changed, and various concentrations of DMAs were added. Two days after transfection, IRES activity was determined by measuring RLuc and FLuc activities using the Dual-Luciferase Reporter Assay System (Promega).

**Cytotoxicity assays**

Various concentrations of DMAs were added to Vero E6 cells in culture. The cells were incubated at 33°C for 96 hours. Cell viability was determined by MTT assay and measured at 570 nm according to the manufacturer’s instructions (EMD Millipore). All experiments were performed in triplicate. The concentration of DMAs required to reduce cell viability to 50% of the control cells was expressed as CC₅₀.

**SARS-CoV-2 antiviral assays**

Analysis of SARS-CoV-2 growth in cell culture supernatants was performed using a simplified qRT-PCR assay as explained before (34). Briefly, Vero E6 cells were infected with SARS-CoV-2 at an MOI of 0.1 IU per cell in 96-well plates, and virus inoculum was removed 1 hour postadsorption and replaced with media containing serial dilutions of the DMA compounds. Five microliters of cell culture media containing released virions was collected at 24 hours post infection and processed as detailed in previous studies (34). Viral RNA levels were quantitated by qRT-PCR using primers specific to SARS-CoV-2 N gene and a standard curve derived from in vitro synthesized RNA encoding N.

In addition, antiviral activity was tested against SARS-CoV-2 using a TCID₅₀ assay, and Vero E6 cells were seeded at 1 x 10⁵ per well in a 24-well plate at 37°C for 24 hours. Cells and samples were then transferred to a Biosafety Level 3 facility. Stocks of SARS-CoV-2 were diluted in Dulbecco’s modified Eagle’s medium (DMEM)/2% FBS with a final volume of 200 μl. The plates were incubated at 37°C for 72 hours. Media were then harvested, centrifuged at room temperature for 10 minutes at 1500g, and then used for TCID₅₀ assay. Serial dilutions of supernatant from the treated cells were added to Vero E6 cells in 96-well plates, and cells were monitored for cytopathic effect. Virus titer was calculated from the numbers of positive wells using a modified Reed and Muench method.

**Synthesis and purification of RNA SL constructs present at 5′-end**

SL1 to SL6 of the 5′-end were in vitro transcribed using a standard protocol from synthetic DNA templates from Integrated DNA Technologies (Coralville, IA) (50, 51). The 3- to 6-ml reactions involved the use of purified recombinant T7 RNA polymerase expressed in BL21 (DE3) cells. Depending on the labeling scheme of each RNA, double-labeled, 13C and 15N, rNTPs and unlabeled rNTPs were used in the reaction. Nucleotide labeling of the SLs was based on the abundance of nucleotides in bulges and loops. The labeling pattern for the 1H-13C HSQC experiments was as follows: SL1 CG(1H, 15N)-labeled and A(13C, 15N)-labeled, SL2 U(13C, 15N)-labeled, SL3 AU(13C, 15N)-labeled, SL4 A(13C, 15N)-labeled, SL5a AU(13C, 15N)-labeled, SL5b AU(13C, 15N)-labeled, and SL6 AC(13C, 15N)-labeled. For the 1H-1H NOESY experiments, the labeling scheme was SL6 UG(2H) AC(2H, 3′-5″), CUG(2H) A(2H, 3′-5″), and ACUG(2H) selectively labeled constructs, along with a CG(2H), AU(2H, 3′-5″) selectively labeled SL1 construct.

Next, the SLs were purified on denaturing gels, ranging from 8 to 16%, and extracted using electroelution. After desalting with a Millipore Amicon Ultra-4 centrifugal filter, the RNAs were annealed by heating for 2 min at 95°C and flash cooled on ice. The samples were thoroughly washed and concentrated down with a 100% D₂O buffer of 25 mM K₂HPO₄ and 50 mM KCl at pH 6.2. Using the NanoDrop 2000 software (Thermo Fisher Scientific), the theoretical extinction coefficients of the SLs were calculated to determine RNA concentrations. Samples for NMR titrations contained 100 μM RNA in D₂O buffer with a final volume of 200 μl.

**NMR profiling of DMA interactions with 5′-end structures**

A 900-MHz spectrometer was used to record all NMR data. The 1H-13C HSQC titrations were recorded with 100 μM of selectively labeled SL1 to SL6 in a 100% D₂O buffer of 25 mM K₂HPO₄ and 50 mM KCl at pH 6.2 with a 200-μl sample volume. Titrations of the DMA molecules into the different SLs were collected at a molar ratio of 5:1. DMA to RNA, at a temperature of either 298, 303, or 308 K. For each construct, temperature optimization experiments were determined to do the optimum temperature to conduct the titrations with. Also, based on the optimized temperatures, 1H-1H
NOESY spectra (τm = 250 ms) of selectively labeled SL6 samples and their DMA-155 complexes were collected with a 200 μM sample concentration in 25 mM K2HPO4, 50 mM KCl (pH 6.2) in 100% D2O at a 200-μl sample volume. At similar conditions, H-1-H NOESY spectra (τm = 250 ms) of selectively labeled SL1 were collected in the absence and presence of DMA-135 and DMA-155 at a 5:1 molar ratio. The NMR spectra were processed using NMRPipe/NMRDraw and analyzed with NMRView J or Sparky (52–54).

Virtual ligand screening against SARS-CoV-2 5′-end RNA structures

**FARFAR model generation**

FARFAR is part of the Rosetta 3 software package and can be obtained for free academic usage (https://www.rosettacommons.org/software/license-and-download). Each SL was generated through the following protocol.

* RNA helix.py* is a python wrapper for the Rosetta executable *rna_helix*. *RNA helix.py* and is available in $ROSETTA/tools/rna_tools/bin, where $ROSETTA is the Rosetta installation path. Below are the commands to generate SL1.

```bash
RNA helix.py -seq gguu aacc -resnum 1-4 24-27 -o helix_1.pdb
RNA helix.py -seq acc ggu -resnum 8-10 17-19 -o helix_2.pdb
```

For each helix in an SL other than the nucleotides that flank bulges and loops, we prebuilt as idealized A-form helices with the above commands. Modeling helical residues as idealized A-form substantially reduce computational time, allowing for more models to be built.

FARFAR modeling is performed through the *rna_denovo* executable

```
rna_denovo -nstruct 1000 -s helix_1.pdb helix_2.pdb -fasta input.fasta -secstruct input.secstruct -minimize_rna true -out:file:silent farfar.out
```

where *-nstruct* is the maximum number of models requested, *-fasta* is the path of a fasta file containing the RNA sequence, *-secstruct* is the path of a file containing the RNA secondary structure in dot-bracket notation, *-minimize_rna true* minimizes the RNA after fragment assembly, and *-out:file:silent farfar.out* is the path of the file containing all generated models. Each SL was run on 100 cores for 24 hours; the number of generated models is reported in table FARFAR (table S1).

To reduce the number of models to dock against, we performed fixed-width clustering using the *rna_cluster* executable

```
rna_cluster -in:file:silent farfar.out -nstruct 15 -cluster:radius RADIUS
```

where *-nstruct* is the maximum number of clusters requested, and *-cluster:radius* is max distance in heavy-atom root mean square deviation between members of the same cluster.

**Small-molecule docking**

Virtual docking simulations were performed using ICM (55) (Molsoft LLC, La Jolla, CA) using the SARS-CoV-2 5′-UTR structures obtained from FARFAR modeling and clustering. RNA structural elements’ binding pockets were defined using ICM Pocket Finder module, and all the small-molecule protonation states were adjusted to pH 7.0 using ChemAxon (www.chemaxon.com). RNA ensembles were then combined into conformational stacks using ICM’s “impose conformations.” Then, to create “flexible receptors” to dock against that would reflect all of the conformations of each structure, ICM’s “create 4D grid” function was used for each docking project. Each of the structures was then docked against a library of 55 DMA molecules. The DMA library was saved in .sdf file format, which was indexed for virtual ligand screening using ICM-Pro. The virtual screening simulation was implemented with a conformational search and optimization with a limit of 10 conformers per molecule. The thoroughness was left at level 10.

**Indicator displacement assay**

A serial dilution of the seven SL RNAs (SL1, SL2, SL3, SL4, SL5a, SL5b, and SL6) was performed in tris buffer (50 mM tris-HCl and 50 mM KCl, at pH 7.4) in a 96-well plate in triplicate. Eight microliters of each dilution was transferred to a 384-well plate followed by 8 μl of a 500 nM solution of RiboGreen dye in the same buffer (Invitrogen). The plates were excited at 487 nm (8-nm slit), and emission was recorded at 525 nm (8-nm slit; focal height, 11.3 mm) using a CLARIOstar plate reader (BMG Labtech). The affinity of the dye for the RNA construct was determined by fitting the raw fluorescence in GraphPad Prism version 8.3.1 for Macintosh [GraphPad Software, La Jolla, California, USA (www.graphpad.com)] by fitting to the [Agonist] versus response – variable slope model that uses Eq. 1

$$\frac{Y}{X} = \frac{\text{Top} - \text{Bottom}}{\text{HillSlope} + \text{EC}_{50}^{\text{HillSlope}}}$$

where $Y$ is normalized percent change in fluorescence intensity, $X$ is RNA concentration, Bottom is lowest fluorescence percent change, and Top is highest fluorescence percent change. Affinity of the dye for the RNA construct was used as the ideal RNA concentration for small-molecule tiritations.

The two most promising small-molecule leads (DMA-135 and DMA-155) were then screened against SL1, SL4, SL5a, SL5b, and SL6, but not against the short SL2 and SL3 for which the RiboGreen dye showed weak affinity.

A serial dilution of DMA-135 and DMA-155 (0, 0.5, 1.0, 2.5, 5, 10.0, 12.5, 15, 20, 25, 40, 50, 75, 100, 225, 400, and 450 μM) was performed in phosphate buffer tris buffer (50 mM tris-HCl and 50 mM KCl, at pH 7.4) in a 96-well plate in triplicate. Eight microliters of each dilution was transferred to a 384-well plate, followed by 8 μl of a solution of SL RNA and 0.5 μM RiboGreen (Invitrogen). The 384-well plates were at 4000 rpm for 1 min and incubated in the dark for 30 min. The plates were excited at 487 nm (8-nm slit), and emission was read at 525 nm (8-nm slit; focal height, 11.3 nm) using a CLARIOstar plate reader (BMG Labtech). Percent fluorescence indicator displacement (%FID) was calculated by subtracting and, subsequently, dividing by the blank wells with RNA-dye complex and no small molecule as shown in Eq. 2

$$\%\text{FID} = \left(\frac{F_0 - F}{F_0}\right) \times 100$$

where $F_0$ is the fluorescence of the blank well with RNA + dye and no small molecule, and $F$ is the fluorescence of the well with all three components (RNA + dye + small molecule).

Each technical triplicate was averaged, and the resulting FID values were averaged between three independent experiments. The binding curve and median effective concentration (EC50) value were obtained by using a nonlinear fit curve agonist versus response with variable slope at four parameters [GraphPad Prism Software version for Macintosh 8.3.1, La Jolla, California, USA (www.graphpadprism.com)]
as shown in Eq. 1. Reported values are averages of three independent experiments ± SD.

**SUPPLEMENTARY MATERIALS**

Supplementary material for this article is available at https://science.org/doi/10.1126/sciadv.abl6096

**REFERENCES AND NOTES**


Amilorides inhibit SARS-CoV-2 replication in vitro by targeting RNA structures

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