Methods

SEARCH-MaP of E. coli ribosomes

Growth and lysis of *E. coli*

Stellar Chemically Competent Cells (Takara) transformed with an ampicillin resistance plasmid were inoculated into 7 ml of LB Broth (Lennox) containing [CON-CENTRATION] of ampicillin [SUPPLIER]. The culture was incubated at 37°C while shaking at 180 rpm for 18.5 hr. OD_{600} of 1 ml was measured using a NanoDrop One cuvette reader. The remaining 6 ml was spun at 21,000 x g. The pellet was resuspended in 1,158 μ l of 50 mM Tris-HCl pH 7.4 (Invitrogen) and lysed by adding 1.16 μ l of NEBExpress T4 Lysozyme (New England Biolabs) and incubating at room temperature on a tube rotator at 20 rpm for 5 min. The lysate was centrifuged at 16,000 x g at 4°C for 10 min, and the supernatant collected and kept on ice.

DMS treatment of *E. coli* lysate

RNA concentration in the supernatant was estimated by purifying 50 μ l with an RNA Clean & Concentrator-5 kit (Zymo Research) according to the manufacturer's protocol, eluting in 20 μ l of nuclease-free water (Fisher Bioreagents), and measuring with a NanoDrop One (Thermo Fisher Scientific). Supernatant containing an estimated 1.5 pmol (2.2 μ g) of rRNA was mixed with 300 mM Tris-HCl pH 7.4 (Invitrogen), 6 mM magnesium chloride (Invitrogen), and optionally 750 nM ASO (Integrated DNA Technologies) in 98.1 μ l; and incubated at 37°C for 20 min to allow the ASO to bind. 1.9 μ l DMS (Sigma-Aldrich) was added (to 200 mM) and shaken at 500 rpm in a ThermoMixer C (Eppendorf) at 37°C for 5 min. To quench, 20 μ l of beta-mercaptoethanol (Sigma-Aldrich) was added and mixed thoroughly. DMS-modified RNA was purified using an RNA Clean & Concentrator-5 kit (Zymo Research) according to the manufacturer's protocol, eluted in 10 μ l of nuclease-

free water (Fisher Bioreagents), and measured with a NanoDrop One (Thermo Fisher Scientific). ASOs and genomic DNA were removed using $0.5 \,\mu$ l TURBO DNase and 1X TURBO DNase Buffer (Invitrogen) in $25 \,\mu$ l at 37° C for 30 min. RNA was purified again using an RNA Clean & Concentrator-5 kit (Zymo Research) and eluted in $10 \,\mu$ l of nuclease-free water (Fisher Bioreagents).

Library generation *E. coli* RNA

SEARCH-MaP of 2,924 nt SARS-CoV-2 RNA

Design of antisense oligonucleotide groups

Antisense oligonucleotides (ASOs, Supplementary Table ??) and their flanking primers (Supplementary Table ??), with constraints on length and melting temperature, were designed using the Python script https://github.com/rouskinlab/search-map/tree/main/Compute/sars2-2924/olaygo.py.

Synthesis of 2,924 nt SARS-CoV-2 RNA

A DNA template of the 2,924 nt segment of SARS-CoV-2, including a T7 promoter, was amplified from a previously constructed plasmid **?** in 50 μ l using 2X CloneAmp HiFi PCR Premix (Takara Bio) with 250 nM primers TAATACGACT-CACTATAGAATAATGAGCTTAGTCCTGTTGCACTACG and TAAATTGCGGACAT-ACTTATCGGCAATTTTGTTACC (Thermo Fisher Scientific); initial denaturation at 98°C for 60 s; 35 cycles of 98°C for 10 s, 65°C for 10 s, and 72°C for 15 s; and final extension at 72°C for 60 s. The 50 μ l PCR product with 10 μ l of 6X Purple Loading Dye (New England Biolabs) was electrophoresed through a 50 ml gel – 1% SeaKem Agarose (Lonza), 1X tris-acetate-EDTA (Boston BioProducts), and 1X SYBR Safe (Invitrogen) – at 60 V for 60 min. The band at roughly 3 kb was extracted using a Zymoclean Gel DNA Recovery Kit (Zymo Research) according to the manufacturer's protocol, eluted in 10 μ l of nuclease-free water (Fisher Bioreagents), and measured with a NanoDrop (Thermo Fisher Scientific). To in-

crease yield, the gel-extracted DNA was fed into a second round of PCR and gel extraction using the same protocol. Due to remaining contaminants, the DNA was further purified using a DNA Clean & Concentrator-5 kit (Zymo Research) according to the manufacturer's protocol, eluted in 10 μ l of nuclease-free water (Fisher Bioreagents), and measured with a NanoDrop (Thermo Fisher Scientific).

150 ng of DNA template was transcribed using a MEGAscript T7 Transcription Kit (Invitrogen) according to the manufacturer's protocol, incubating at 37°C for 3 hr. DNA template was then degraded by incubating with 1 μ I of TURBO DNase (Invitrogen) at 37°C for 15 min. RNA was purified using an RNA Clean & Concentrator-5 kit (Zymo Research) according to the manufacturer's protocol, eluted in 20 μ I of nuclease-free water (Fisher Bioreagents), and measured with a NanoDrop (Thermo Fisher Scientific).

DMS treatment of 2,924 nt SARS-CoV-2 RNA

ASOs were ordered from Integrated DNA Technologies already resuspended to $10~\mu\mathrm{M}$ in 1X IDTE buffer ($10~\mathrm{mM}$ Tris, $0.1~\mathrm{mM}$ EDTA) in a 96-well PCR plate. Each ASO pool was assembled from 25 pmol of each constituent ASO (Supplementary Table $\ref{Table 1}$); volume was adjusted to $12.5~\mu\mathrm{l}$ by adding TE Buffer $-10~\mathrm{mM}$ Tris (Invitrogen) with $0.1~\mathrm{mM}$ EDTA (Invitrogen). 450 fmol of 2,924 nt SARS-CoV-2 RNA was added to each ASO pool for a total of $13.5~\mu\mathrm{l}$ in a PCR tube. The tube was heated to $95~\mathrm{C}$ for $60~\mathrm{s}$ to denature the RNA, placed on ice for several minutes, and transferred to a $1.5~\mathrm{ml}$ tube. To refold the RNA, $35~\mu\mathrm{l}$ of $1.4\mathrm{X}$ refolding buffer comprising 400 mM sodium cacodylate pH $7.2~\mathrm{(Electron}$ Microscopy Sciences) and $6~\mathrm{mM}$ magnesium chloride (Invitrogen) was added, then incubated at $37~\mathrm{C}$ for $25~\mathrm{min}$. For no-ASO control 1, $12.5~\mu\mathrm{l}$ of TE Buffer was used instead of an ASO pool. For no-ASO control 2, $12.5~\mu\mathrm{l}$ of TE Buffer was added after placing on ice and before refolding to confirm the timing of adding TE Buffer would not alter the RNA structure.

RNA was treated with DMS in 50 μ l containing 1.5 μ l (320 mM) of DMS (Sigma-Aldrich) while shaking at 500 rpm in a ThermoMixer C (Eppendorf) at 37°C for

5 min. To quench, 30 μ I of beta-mercaptoethanol (Sigma-Aldrich) was added and mixed thoroughly. RNA was purified using an RNA Clean & Concentrator-5 kit (Zymo Research) according to the manufacturer's protocol, eluted in 10 μ I of nuclease-free water (Fisher Bioreagents), and measured with a NanoDrop (Thermo Fisher Scientific).

ASOs were removed from 4 μ l of DMS-modified RNA in 10 μ l containing 1 μ l of TURBO DNase (Invitrogen) and 1X TURBO DNase Buffer (Invitrogen), incubated at 37°C for 30 min. To stop the reaction, 2 μ l of DNase Inactivation Reagent was added and incubated at room temperature for 10 min, mixing several times throughout by flicking. DNase Inactivation Reagent was precipitated by spinning on a benchtop PCR tube centrifuge for 10 min and transferring 4 μ l of supernatant to a new tube.

Library generation of 2,924 nt SARS-CoV-2 RNA

4 μ l RNA was reverse transcribed in 20 μ l containing 1X First Strand Buffer (Invitrogen), 500 µM dNTPs (Promega), 5 mM dithiothreitol (Invitrogen), 500 nM FSE primer CTTCGTCCTTTTCTTGGAAGCGACA (Integrated DNA Technologies), 500 nM section-specific reverse primer (Integrated DNA Technologies, Supplementary Table ??), 1 μ l of RNaseOUT (Invitrogen), and 1 μ l of TGIRT-III enzyme (InGex) at 57°C for 90 min, followed by inactivation at 85°C for 15 min. To degrade the RNA, 1 μ l of Hybridase Thermostable RNase H (Lucigen) was added to each tube and incubated at 37°C for 20 min. 1 μ l of unpurified RT product was amplified in 12.5 μ l using the Advantage HF 2 PCR Kit (Takara Bio) with 1X Advantage 2 PCR Buffer, 1X Advantage-HF 2 dNTP Mix, 1X Advantage-HF 2 Polymerase Mix, 250 nM primers (Integrated DNA Technologies) for either the FSE (CCCTGTGGGTTTTA-CACTTAAAAAC and CTTCGTCCTTTTCTTGGAAGCGACA) or specific section (Supplementary Table ??); initial denaturation at 94°C for 60 s; 25 cycles of 94°C for 30 s, 60°C for 30 s, and 68°C for 60 s; and final extension at 68°C for 60 s. $5 \mu l$ of every amplicon from the same RT product was pooled and then purified using a DNA Clean & Concentrator-5 kit (Zymo Research) according to the manufacturer's protocol, eluted in 20 μ l of nuclease-free water (Fisher Bioreagents), and measured with a NanoDrop (Thermo Fisher Scientific).

200 ng of pooled PCR product was prepared for sequencing using the NEB-Next Ultra II DNA Library Prep Kit for Illumina (New England Biolabs) according to the manufacturer's protocol with the following modifications. During size selection after adapter ligation, 27.5 μ I and 12.5 μ I of NEBNext Sample Purification Beads (New England Biolabs) were used in the first and second steps, respectively, to select inserts of 280-300 bp. Indexing PCR was run at half volume (25 μ I) for 3 cycles. In lieu of the final bead cleanup, 420 bp inserts were selected using a 2% E-Gel SizeSelect II Agarose Gel (Invitrogen) according to the manufacturer's protocol. DNA concentrations were measured using a Qubit 3.0 Fluorometer (Thermo Fisher Scientific) according to the manufacturer's protocol. Samples were pooled and sequenced using an iSeq 100 Sequencing System (Illumina) with 2 x 150 bp paired-end reads according to the manufacturer's protocol.

Data analysis of 2,924 nt SARS-CoV-2 RNA

Sequencing data were processed with SEISMIC-RNA v0.12 and v0.13 to compute mutation rates, clusters, correlations, and secondary structures. Effects of each ASO group (Figure ??b, Supplementary Figures ?? and ??) were computed with the script https://github.com/rouskinlab/search-map/tree/main/Compute/sars2-2924/run-tile.sh. Clustering and structure modeling (Figure ??c and d, Supplementary Figure ??a and b) were performed with the script https://github.com/rouskinlab/search-map/tree/main/Compute/sars2-2924/run-deep.sh. Because some samples contained amplicons that overlapped each other, sequence alignment map (SAM) files were filtered by amplicon using the script https://github.com/rouskinlab/search-map/tree/main/Compute/sars2-2924/filter-deep.py. The fraction of structures containing long-range stems (Supplementary Figure ??c) was determined using the script https://github.com/rouskinlab/search-map/tree/main/Compute/sars2-2924/fraction folded.py.

SEARCH-MaP of long-range base pairs in multiple coronaviruses

Computational screen for long-range base pairs in coronaviruses

All coronaviruses with reference genomes in the NCBI Reference Sequence Database ? as of December 2021 were searched for using the following query:

The reference sequences (https://github.com/rouskinlab/search-map/ tree/main/Compute/covs-screen/cov refseq.fasta) and table of features(https://github.com/rouskinlab/search-map/tree/main/Compute/ covs-screen/cov features.txt) were downloaded and used to locate the slippery site in each genome using a custom Python script (https://github. com/rouskinlab/search-map/tree/main/Compute/covs-screen/extract long fse.py). For each genome, up to 100 secondary structure models of the 2,000 nt segment from 100 nt upstream to 1,893 nt downstream of the slippery site (excluding genomes with ambiguous nucleotides in this segment) were generated using Fold v6.3 from RNAstructure? via the script https://github.com/ rouskinlab/search-map/tree/main/Compute/covs-screen/fold long fse.py. The fraction of models in which each base paired with any other base between positions 101 and 250 was calculated using the script https://github. com/rouskinlab/search-map/tree/main/Compute/covs-screen/analyze interactions.py. Using these fractions, coronaviruses were clustered via the unweighted pair group method with arithmetic mean (UPGMA) and a euclidean distance metric, implemented in Seaborn v0.11 ? and SciPy v1.7 ? (Supplementary Figure ??). From each cluster with prominent potential long-range interactions involving the FSE, coronaviruses were manually selected for experimental study.

Synthesis of 239 and 1,799 nt coronaviral RNAs

For each selected coronavirus, the 1,799 nt segment from 290 to 1,502 nt downstream of the slippery site was ordered from Twist Bioscience as a gene fragment flanked by the standard 5' and 3' adapters CAATCCGCCCTCACTACAACCG and CTACTCTGGCGTCGATGAGGGA, respectively. Gene fragments were resuspended to 10 ng/ μ l in 10 mM Tris-HCl pH 8 (Invitrogen). Each DNA template for transcription of 1,799 nt RNA segments, including a T7 promoter, was amplified from 0.5 μ l (5 ng) of a gene fragment in 20 μ l using 2X CloneAmp HiFi PCR Premix (Takara Bio) with 250 nM of each primer TAATACGACTCACTATAG-GCAATCCGCCCTCACTACAACCG and TCCCTCATCGACGCCAGAGTAG; initial denaturation at 98°C for 30 s; 30 cycles of 98°C for 10 s, X°C (see Supplementary Table ??) for 10 s, and 72°C for 15 s; and final extension at 72°C for 60 s. DNA templates for transcription of 239 nt RNA segments were amplified using the same procedure but with the forward primers with T7 promoters (F+T7) and reverse primers (R) in Supplementary Table ??. For experiments in which the RNAs were transcribed as a pool of all coronaviruses, all PCR products of the same length (i.e. 239 or 1,799 nt) were pooled, then purified using a DNA Clean & Concentrator-5 kit (Zymo Research) according to the manufacturer's protocol; concentrations were measured with a NanoDrop (Thermo Fisher Scientific). Otherwise, PCR products were purified individually.

50 ng of DNA template was transcribed using a HiScribe T7 High Yield RNA Synthesis Kit (New England Biolabs) according to the manufacturer's protocol but at one-quarter volume (5 μ l), supplemented with 0.25 μ l RNaseOUT (Invitrogen), for 16 hr. DNA template was degraded by incubating with 0.5 μ l of TURBO DNase (Invitrogen) at 37°C for 30 min. RNA was purified using an RNA Clean & Concentrator-5 kit (Zymo Research) according to the manufacturer's protocol,

eluted in 50 μ l of nuclease-free water (Fisher Bioreagents), and measured with a NanoDrop (Thermo Fisher Scientific).

DMS treatment of 239 and 1,799 nt coronaviral RNAs

Antisense oligonucleotides (ASOs) in Supplementary Table **??** were ordered from Integrated DNA Technologies and resuspended to 100 μ M in low-EDTA TE buffer: 10 mM Tris pH 7.4 with 0.1 mM EDTA (Integrated DNA Technologies). For each coronavirus, 5 μ I of each corresponding ASO (Supplementary Table **??**) was pooled; the pool of ASOs was diluted with low-EDTA TE buffer to a final volume of 100 μ I, bringing each ASO to 5 μ M. 1X refolding buffer comprising 300 mM sodium cacodylate pH 7.2 (Electron Microscopy Sciences) and 6 mM magnesium chloride (Invitrogen) was assembled, then pre-warmed to 37°C.

For already-pooled RNA, 300 ng was diluted in 2.5 μ I of nuclease-free water (Fisher Bioreagents) in a PCR tube, heated to 95°C for 1 min to denature, chilled on ice for 3 min, added to 95 μ I of pre-warmed refolding buffer, and incubated at 37°C for 20 min to refold. For individually transcribed RNA, 1 pmol was mixed with 10 μ I of either low-EDTA TE buffer (for probing without ASOs) or the ASO pool for the corresponding coronavirus (for probing with ASOs) in a PCR tube, heated to 95°C for 1 min to denature the RNA, chilled on ice for 3 min, added to pre-warmed refolding buffer for a total volume of 100 μ I, and incubated at 37°C for 20 min to refold the RNA (possibly with ASOs). Subsequently, equimolar amounts of all refolded RNAs were combined into one 97 μ I pool in a 1.5 ml tube.

RNA was treated with DMS (Sigma-Aldrich) – $2.5 \,\mu$ l (260 mM) for RNAs transcribed as pools or 3 μ l (320 mM) for RNAs pooled after transcription – in 100 μ l while shaking at 800 rpm in a ThermoMixer C (Eppendorf) at 37°C for 5 min. To quench, 60 μ l of beta-mercaptoethanol (Sigma-Aldrich) was added and mixed thoroughly. DMS-modified RNA was purified using an RNA Clean & Concentrator-5 kit (Zymo Research) according to the manufacturer's protocol, eluted in 16 μ l of nuclease-free water (Fisher Bioreagents), and measured with a NanoDrop (Thermo Fisher Scientific). If added, ASOs were then degraded in 50 μ l containing 1X

TURBO DNase Buffer (Invitrogen) and 1 μ I of TURBO DNase Enzyme (Invitrogen) at 37°C for 30 min; RNA was purified with an RNA Clean & Concentrator-5 kit (Zymo Research) according to the manufacturer's protocol, eluted in 16 μ I of nuclease-free water (Fisher Bioreagents), and measured with a NanoDrop (Thermo Fisher Scientific).

Sequencing library generation of 239 and 1,799 nt coronaviral RNAs

100 ng of DMS-modified RNA was prepared for sequencing using the xGen Broad-Range RNA Library Preparation Kit (Integrated DNA Technologies) according to the manufacturer's protocol, with the following modifications. During fragmentation, 8 μ l of RNA was combined with 1 μ l of Reagent F1, 4 μ l of Reagent F3, and $2 \mu l$ of Reagent F2. For reverse transcription, $1 \mu l$ of Enzyme R1, $2 \mu l$ of TGIRT-III enzyme (InGex), and 1 μ I of 100 mM dithiothreitol (Invitrogen) was used instead of the reaction mix, then incubated at room temperature for 30 minutes before adding 2 μ l of Reagent F2. Reverse transcription was stopped by adding 1 μ l of 4 M sodium hydroxide (Fluka), heating to 95°C for 3 min, chilling at 4°C, then neutralizing with 1μ l of 4 M hydrochloric acid. Instead of a bead cleanup after the final PCR, unpurified PCR products with 6X DNA loading dye (Invitrogen) were elecrophoresed through an 8% polyacrylamide Tris-borate-EDTA (TBE) gel (Invitrogen) at 180 V for 55 min. The gel was stained with SYBR Gold (Invitrogen); the section between 250 and 500 bp was excised and placed in a 0.5 ml tube with a hole punctured in the bottom by an 18-gauge needle (BD Biosciences), which was nested inside a 1.5 ml tube and centrifuged at 21,300 x g for 1 min to crush the gel slice into the larger tube. Crushed gel pieces were suspended in 500 μ l of 300 mM sodium chloride (Boston Bioproducts), shaken at 1,500 rpm in a ThermoMixer C (Eppendorf) at 70°C for 20 min, and centrifuged at 21,300 x g through a 0.22 µm Costar Spin-X filter column to remove the gel pieces. Filtrate was mixed with 600 μ l isopropanol (Sigma-Aldrich) and 3 μ l GlycoBlue Coprecipitant (Invitrogen), vortexed briefly, and stored at -20°C overnight. DNA was then pelleted by centrifugation at 4° C at $18,200 \times g$ for 45 min. The supernatant was aspirated, and the pellet was washed with 1 ml of ice-cold 70% ethanol (Sigma-Aldritch), resuspended in $15 \mu l$ nuclease-free water (Fisher Bioreagents), and quantified using the 1X dsDNA High Sensitivity Assay Kit for the Qubit 3.0 Fluorometer (Thermo Fisher Scientific) according to the manufacturer's protocol. Samples were pooled and sequenced using an iSeq 100 Sequencing System (Illumina) with $2 \times 150 \text{ bp}$ paired-end reads according to the manufacturer's protocol.

Data analysis of 239 and 1,799 nt coronaviral RNAs

Sequencing data were processed with SEISMIC-RNA v0.11 and v0.12 to compute mutation rates, correlations between samples, and secondary structure models using the commands in the shell script https://github.com/rouskinlab/ search-map/tree/main/Compute/covs-1799/run.sh. For the 239 and 1,799 nt RNAs that had been pooled during transcription, the two replicates for each coronavirus for each length were confirmed to give similar results, then merged before comparing the 239 and 1,799 nt RNAs to each other. For the comparison of RNAs with and without ASOs, the no-ASO samples that had been transcribed individually were confirmed to give similar results to those transcribed as a pool; then, all no-ASO samples were pooled before comparing to samples with ASOs. For each coronavirus, the DMS reactivities of the combined no-ASO samples were used to model up to 20 secondary structures of the 1,799 nt segment using Fold from RNAstructure v6.3 ?. Structure models were checked manually for correspondence with the rolling correlation between the +ASO and no-ASO conditions; the minimum free energy structure was chosen for every coronavirus except for transmissible gastroenteritis virus, in which the first sub-optimal structure - but not the minimum free energy structure - contained long-range base pairs supported by the rolling correlation. Rolling correlations between +ASO and no-ASO conditions superimposed on secondary structure models (Figure ??) were graphed using the Python script https://github.com/rouskinlab/ search-map/tree/main/Compute/util/pairs_vs_correl.py.

SEARCH-MaP of 1,799 nt SARS-CoV-2 RNA

RNA synthesis of 1,799 nt SARS-CoV-2 RNA

A DNA template for transcription, including a T7 promoter, was amplified from the 1,799 bp gene fragment of SARS-CoV-2 (Twist Bioscience) as described above but with primers TAATACGACTCACTATAGGTACTGGTCAGGCAATAACAGTTA-CAC and GACCCCATTTATTAAATGGAAAACCAGCTG (Integrated DNA Technologies), an annealing temperature of 65°C, and an extension time of 10 s; eluted in 18 μ I of 10 mM Tris-HCl pH 8 (Invitrogen); and measured with a NanoDrop One (Thermo Fisher Scientific). 100 ng of DNA template was transcribed using a HiScribe T7 High Yield RNA Synthesis Kit (New England Biolabs) according to the manufacturer's protocol for 11 hr. DNA template was degraded by incubating with 1 μ I of TURBO DNase (Invitrogen) at 37°C for 30 min. RNA was purified using an RNA Clean & Concentrator-25 kit (Zymo Research) according to the manufacturer's protocol, eluted in 50 μ I of nuclease-free water (Fisher Bioreagents), and measured with a NanoDrop One (Thermo Fisher Scientific).

DMS treatment of 1,799 nt SARS-CoV-2 RNA

1.15X refolding buffer comprising 345 mM sodium cacodylate pH 7.2 (Electron Microscopy Sciences) and 7 mM magnesium chloride (Invitrogen) was assembled and pre-warmed to 37°C. 1 pmol of RNA was mixed with 100 pmol of each ASO (Integrated DNA Technologies, Supplementary Table $\ref{thm:pmodel}$) in 10 μ l total, heated to 95°C for 60 s to denature, chilled on ice for 5-10 min, and added to 87.1 μ l of pre-warmed refolding buffer. If no ASO would be added during refolding, then 1 μ l of nuclease-free water (Fisher Bioreagents) was added. RNA was incubated at 37°C for 15-20 min to refold. If an ASO would be added during refolding, then 100 pmol (1 μ l) of ASO was added. RNA was incubated for another 15 min to allow any newly added ASOs to bind.

RNA was probed in 100 μ l containing 1.9 μ l (300 mM) DMS (Sigma-Aldrich) while shaking at 500 rpm in a ThermoMixer C (Eppendorf) at 37°C for 5 min.

To quench, $20 \mu l$ of beta-mercaptoethanol (Sigma-Aldrich) was added and mixed thoroughly. DMS-modified RNA was purified using an RNA Clean & Concentrator-5 kit (Zymo Research) according to the manufacturer's protocol, eluted in $15 \mu l$ of nuclease-free water (Fisher Bioreagents), and measured with a NanoDrop One (Thermo Fisher Scientific).

Library generation 1,799 nt SARS-CoV-2 RNA

1 μ I of DMS-modified RNA was reverse transcribed in 20 μ I using Induro Reverse Transcriptase (New England Biolabs) according to the manufacturer's protocol with 500 nM of primer CTTCGTCCTTTTCTTGGAAGCGACA (Integrated DNA Technologies) at 57°C for 30 min, followed by inactivation at 95°C for 1 min. 1 μ I of unpurified RT product was amplified in 20 μ I using Q5 High-Fidelity 2X Master Mix (New England Biolabs) with 500 nM of each primer CCCTGTGGGTTT-TACACTTAAAAAC and CTTCGTCCTTTTCTTGGAAGCGACA (Integrated DNA Technologies); initial denaturation at 98°C for 30 s; 30 cycles of 98°C for 10 s, 65°C for 20 s, and 72°C for 20 s; and final extension at 72°C for 120 s. The PCR product was purified using a DNA Clean & Concentrator-5 kit (Zymo Research) according to the manufacturer's protocol, eluted in 20 μ I of 10 mM Tris-HCI pH 8 (Invitrogen), and measured with a NanoDrop One (Thermo Fisher Scientific).

50-100 ng of purified PCR product was prepared for sequencing using the NEBNext Ultra II DNA Library Prep Kit for Illumina (New England Biolabs) according to the manufacturer's protocol with the following modifications. All steps were performed at half of the volume specified in the protocol, including reactions, bead cleanups, and washes. During size selection after adapter ligation, $14 \mu l$ and $7 \mu l$ of SPRIselect Beads (Beckman Coulter) were used in the first and second steps, respectively, to select inserts of 283 bp. Indexing PCR was run with 400 nM of each primer for 4 cycles. After indexing, PCR products were pooled in pairs; in lieu of the final bead cleanup, 405 bp products were selected using a 2% E-Gel SizeSelect II Agarose Gel (Invitrogen) according to the manufacturer's protocol. DNA concentrations were measured using a Qubit 4 Fluorometer (Thermo Fisher

Scientific) according to the manufacturer's protocol. Samples were pooled and sequenced using a NextSeq 1000 Sequencing System (Illumina) with 2 x 150 bp paired-end reads according to the manufacturer's protocol.

Data analysis of 1,799 nt SARS-CoV-2 RNA

Sequencing data were processed with SEISMIC-RNA v0.11 and v0.12 to compute mutation rates, clusters, and correlations between samples using the commands in the shell script https://github.com/rouskinlab/search-map/ tree/main/Compute/sars2-1799/run.sh. Heatmaps of the reproducibility of clustering between replicates (Supplementary Figure ??) were generated using the Python script https://github.com/rouskinlab/search-map/tree/ main/Compute/sars2-1799/compare-clusters.py. After the two replicates were confirmed to give similar clusters, they were pooled for subsequent analyses. Secondary structures with rolling correlations (Figure ??b) were drawn using the Python script https://github.com/rouskinlab/search-map/ tree/main/Compute/sars2-1799/draw-structure.py. Alternative structure models (Figure ??c) were selected and created with the help of the Python scripts https://github.com/rouskinlab/search-map/tree/main/Compute/ sars2-1799/choose-model-parts.py and https://github.com/rouskinlab/ search-map/tree/main/Compute/sars2-1799/make-models.py. Heatmaps of areas under the curve (Figure ??d) were generated using the Python script https://github.com/rouskinlab/search-map/tree/main/Compute/sars2-1799/ atlas-plot.py.

DMS-MaPseq of transmissible gastroenteritis virus in ST cells

Cells and Viruses

Transmissible gastroenteritis virus (TGEV, TC-adapted Miller strain, ATCC VR-1740) and ST cells (ATCC CRL-1746) were ordered from American Type Culture Collection (ATCC). ST cells were maintained in Eagle's Minimum Essential Medium (EMEM, Gibco) supplemented with 10% fetal bovine serum (Gibco), 1% sodium pyruvate (Gibco), and 1% Pen Strep (Gibco) at 37°C with 5% carbon dioxide. For TGEV, the infection medium (IM) comprised EMEM (Gibgo) supplemented with 10% fetal bovine serum (Gibco), 1% sodium pyruvate, and 1 μ g/ml of TPCK trypsin (Thermo Fisher Scientific).

Production and titering of TGEV

A 150 mm dish was seeded with 1 x 10^7 ST cells, grown overnight, and washed twice with phosphate-buffered saline (PBS, Gibco). Cells were inoculated with 8 ml of TGEV in IM at a multiplicity of infection (MOI) of 0.1, which was kept on for 60 min with rocking every 15 min. The inoculum was removed, cells were washed twice with PBS, and 26 ml of IM was added. Cells were checked daily for cytopathic effects (CPE); after 5 days, upon development of 80% CPE, the supernatant was centrifuged at 200 x g for 5 min and then filtered through a 0.45 μ m filter to harvest TGEV, which was frozen at -80°C in 1 ml aliquots.

Harvested TGEV was titered via tissue culture infectious dose (TCID₅₀). Briefly, ST cells were seeded in a poly-L-lysine coated 96-well plate at 4 x 10⁴ cells per well and grown overnight. TGEV was thawed on ice and serially diluted from 10^{-1} to 10^{-10} in IM. Cells were washed once with PBS, and each well was inoculated with 100 μ I of either one serial dilution of TGEV (8 replicates per dilution level) or sterile IM (for negative controls). The plate was wrapped in parafilm and incubated at 37°C until CPE appeared. Then, media was aspirated and cells were fixed with 4% paraformaldehyde for 30 min and decanted. 0.5% crystal violet was

then added to each well; the plate was rocked for 10 min, submerged in water to remove excess crystal violet, and dried. Wells with CPE were counted and the titer determined using the Spearman-Kärber method.

TGEV infection and DMS treatment

Four 150 mm dishes were each seeded with 1 x 10⁷ ST cells, grown overnight, and washed twice with phosphate-buffered saline. Cells were inoculated with 8 ml of TGEV in IM at a multiplicity of infection (MOI) of 2, which was kept on for 60 min with rocking every 15 min. The inoculum was removed, cells were washed twice with PBS, and 26 ml of IM was added.

After 48 hr, media was aspirated. $250 \,\mu\text{l}$ of DMS was mixed with 10 ml of IM and immediately added to two plates; the other two received 10 ml IM without DMS. Plates were incubated at 37°C for 5 min. The media was aspirated and replaced with stop solution (30% beta-mercaptoethanol in 1X PBS). Cells were scraped off using a cell scraper and spun down at 3,000 x g for 3 min. The pellet was washed with stop solution, spun down again, washed with 10 ml PBS, dissolved in 3 ml of TRIzol (Invitrogen), and split into 1 ml techincal replicates.

RNA purification

200 μ l of chloroform was added to each 1 ml techincal replicate, vortexed for 20 s, and rested until the phases separated. Samples were then spun at 18,200 x g for 15 min at 4°C; the aqueous phase transferred to a new tube and mixed with an equal volume of 100% ethanol (Koptec). RNA was purified using a 50 μ g Monarch RNA Cleanup Column (New England Biolabs), eluted in 20 μ l of nuclease-free water, and quantified with a NanoDrop.

To remove rRNA, 10 μ g of total RNA was diluted in 6 μ l of nuclease-free water and mixed with 1 μ l of anti-rRNA ASOs (Integrated DNA Technologies) and 3 μ l of HYBE buffer (200 mM sodium chloride, 100 mM Tris-HCl pH 7.5). The mixture was incubated at 95°C for 2 min and cooled by 0.1°C/s until reaching 45°C. A

preheated mixture of 10 μ l of RNase H (New England Biolabs) and 2 μ l of RNase H Buffer (New England Biolabs) was added and incubated at 45°C for 30 min. RNA was purified using a 10 μ g Monarch RNA Cleanup Column (New England Biolabs) and eluted in 42 μ l of nuclease-free water.

To remove DNA (including anti-rRNA ASOs), 5 μ I of 10X Turbo DNase Buffer (Thermo Fisher) and 3 μ I of TURBO RNase (Thermo Fisher) were added and incubated at 37°C for 20 min. RNA was purified using a 10 μ g Monarch RNA Cleanup Column (New England Biolabs) and eluted in 10 μ I of nuclease-free water.

Library generation for the full TGEV genome

RNA was prepared for sequencing using the xGen Broad-Range RNA Library Preparation Kit (Integrated DNA Technologies) according to the manufacturer's protocol, with the same modifications as described above (1.3.4), notably the substitution of TGIRT-III (InGex) for the kit's reverse transcriptase. Samples were pooled and sequenced using a NextSeq 1000 Sequencing System (Illumina) with 2 x 150 bp paired-end reads according to the manufacturer's protocol.

Library generation for amplicons

1 μl of rRNA-depleted, DNased RNA was reverse transcribed in 20 μl using Induro Reverse Transcriptase (New England Biolabs) according to the manufacturer's protocol with 500 nM of primer ACAATTCGTCTTAAGGAATTTACCAATACACG-CAA (Integrated DNA Technologies) at 57°C for 30 min, followed by inactivation at 95°C for 1 min. 1 μl of unpurified RT product was amplified in 10 μl using Q5 High-Fidelity 2X Master Mix (New England Biolabs) with 1 μM of each primer, either GC-CGCTACAAAGGTAAGTTCGTGCAAATACCAACT and ACAATTCGTCTTAAGGAATTTACCAATACACGCAA or GTGAAAAGTGACATCTATGGTTCTGATTATAAGCAGTA and CTATACCAAGTTGTTTGAAATGGTAACCTGCAGTAACA (Integrated DNA Technologies); initial denaturation at 98°C for 30 s; 30 cycles of 98°C for 5 s, 69°C for 20 s, and 72°C for 15 s; and final extension at 72°C for 120 s. Amplification was

confirmed by electrophoresing 1 μ I of each PCR product. PCR products for both pairs of primers were pooled and then purified using a DNA Clean & Concentrator-5 kit (Zymo Research) according to the manufacturer's protocol, eluted in 18 μ I of 10 mM Tris-HCl pH 8 (Invitrogen), and measured with a NanoDrop (Thermo Fisher Scientific).

175-225 ng of purified PCR product was prepared for sequencing using the NEBNext Ultra II DNA Library Prep Kit for Illumina (New England Biolabs) according to the manufacturer's protocol with the following modifications. All steps were performed at half of the volume specified in the protocol, including reactions, bead cleanups, and washes. During size selection after adapter ligation, $14~\mu l$ and $7~\mu l$ of SPRIselect Beads (Beckman Coulter) were used in the first and second steps, respectively, to select inserts of 295 bp. Indexing PCR was run with 400 nM of each primer for 4 cycles. In lieu of the final bead cleanup, 415 bp products were selected using a 2% E-Gel SizeSelect II Agarose Gel (Invitrogen) according to the manufacturer's protocol. DNA concentrations were measured using a Qubit 4 Fluorometer (Thermo Fisher Scientific) according to the manufacturer's protocol. Samples were pooled and sequenced using a NextSeq 1000 Sequencing System (Illumina) with 2 x 150 bp paired-end reads according to the manufacturer's protocol.

Data analysis of transmissible gastroenteritis virus in ST cells

The genomic sequence of this TGEV strain was determined using the script https:
//github.com/rouskinlab/search-map/tree/main/Compute/tgev-virus/
consensus.sh: reads from the untreated sample were aligned to the TGEV reference genome (NC_038861.1) using Bowtie 2 ? and the consensus sequence
was determined using Samtools ?. All reads were processed with SEISMIC-RNA
v0.15 to compute mutation rates, correlations between samples, and secondary
structure models using the commands in the shell script https://github.com/
rouskinlab/search-map/tree/main/Compute/tgev-virus/run.sh. Positions in the untreated sample with mutation rates greater than 0.01 were masked.

Replicates were checked for reproducibility and pooled for clustering and structure modeling. A model of short-range base pairs (maximum distance 300 nt) in the TGEV genome was generated from the DMS reactivities using Fold-smp from RNAstructure? in five overlapping 10 kb segments, which were merged using the script https://github.com/rouskinlab/search-map/tree/main/Compute/tgev-virus/assemble-tgev-ss.py. Rolling area under the curve superimposed on secondary structure models in Figure??d was graphed using the script https://github.com/rouskinlab/search-map/tree/main/Compute/tgev-virus/make-figure-6d.py, and in Supplementary Figure?? using the script https://github.com/rouskinlab/search-map/tree/main/Compute/tgev-virus/plot genome.py.