Improving the specificity of nucleic acid detection with endonuclease-actuated degradation

Roger S. Zou1,2, Momcilo Gavrilov2,3, Yang Liu4, Dominique Rasoloson4,5, Madison Conte6, Justin Hardick6, Leo Shen1,2, Siqi Chen1,4, Andrew Pekosz7, Geraldine Seydoux4,5, Yukari C. Manabe6 & Taekjip Ha1,2,3,5

Nucleic acid detection is essential for numerous biomedical applications, but often requires complex protocols and/or suffers false-positive readouts. Here, we describe SENTINEL, an approach that combines isothermal amplification with a sequence-specific degradation method to detect nucleic acids with high sensitivity and sequence-specificity. Target single-stranded RNA or double-stranded DNA molecules are amplified by loop-mediated isothermal amplification (LAMP) and subsequently degraded by the combined action of lambda exonuclease and a sequence-specific DNA endonuclease (e.g., Cas9). By combining the sensitivity of LAMP with the precision of DNA endonucleases, the protocol achieves attomolar limits of detection while differentiating between sequences that differ by only one or two base pairs. The protocol requires less than an hour to complete using a 65 °C heat block and fluorometer, and detects SARS-CoV-2 virus particles in human saliva and nasopharyngeal swabs with high sensitivity.
nucleic acid detection has a wide range of applications from epidemiology, disease diagnosis, and pathogen surveillance to laboratory use. While there are many methods for detecting nucleic acids, trade-offs exist between sensitivity, specificity, speed, cost, and convenience. Development of new strategies not only expands the existing toolbox, but also may help resolve bottlenecks in diagnostic testing with significant public health implications, especially for the recent COVID-19 pandemic. Quantitative PCR is the gold standard for nucleic acid detection but requires sophisticated equipment and dedicated laboratory space. Isothermal amplification methods such as loop-mediated isothermal amplification (LAMP), which uses a single heating temperature for target sequence amplification, are an alternate approach that avoids the use of specialized equipment but may suffer from lower specificity and false-positive readouts.

Recently, endonucleases from the clustered regularly interspaced short palindromic repeats (CRISPR) and CRISPR-associated (CRISPR-Cas) adaptive immune systems have been deployed for nucleic acid detection as an additional step on top of initial isothermal amplification to improve the specificity of target sequence recognition. The CRISPR nuclease recognizes the target sequence in the amplified product, which catalyzes the cleavage of quenched fluorescent reporters to generate the detection signal. However, this approach has been limited to Cas12 and Cas13 due to the requirement for non-sequence-specific collateral cleavage activity upon target sequence recognition. We sought to develop a more universal strategy that could be used with any sequence-specific endonuclease.

Sequence-specific endonucleases such as restriction enzymes or RNA-guided endonucleases (i.e., Cris9) are highly efficient and specific. Cleavage generates two new free 5’ ends which can be targeted by lambda exonuclease (λ-exo), a highly processive 5’-3’ exonuclease, for rapid degradation. The initial isothermal amplification (using primers resistant to exonuclease activity) and subsequent degradation after endonuclease cleavage are monitored using convenient fluorescence-based measurements of dsDNA concentration. Samples with the expected target would exhibit an amplification signal after LAMP and a loss of signal after endonuclease and λ-exo digestion. Samples with the incorrect target would either fail to amplify with LAMP, resulting in minimal initial signal, or generate a LAMP amplification product resistant to endonuclease and λ-exo treatment. We term this new rapid nucleic acid detection platform SpCas9 CRISPR Endonuclease Assay (SENTINEL). We demonstrate that this approach combines the attomolar-level sensitivity of isothermal amplification with the nucleotide-level specificity of sequence-specific endonucleases. The SENTINEL assay is highly convenient in requiring minimal laboratory reagents, modular in the choice of sequence-specific endonuclease, fast with under 1-h total time and minimal hands-on time, and versatile in detecting SARS-CoV-2 virus particles in human saliva and nasopharyngeal swabs with high sensitivity.

**Results**

For initial validation, we evaluated wild type SpCas9 as the endonuclease. Using in vitro transcription, we generated synthetic RNA from the N-gene of SARS-CoV-2 as the target of interest. Reverse transcription LAMP (RT-LAMP) was first performed at 65°C for 30 min. The RT-LAMP primers were designed to contain 5’ phosphorothioate modifications, resulting in product DNA that resists λ-exo degradation. Agarose gel electrophoresis visualized the RT-LAMP product, which exhibited multiple bands of increasing molecular weight, as expected. We first verified that Cas9 can efficiently cleave RT-LAMP product by mixing diluted RT-LAMP product with Cas9 in complex with crRNA and tracrRNA (henceforth referred to as gRNA) targeting the SARS-CoV-2 N-gene (Cas9-19n-N2-gRNA) (Supplementary Fig. 1), or a non-targeting gRNA. We observed a band shift in RT-LAMP product on agarose gel with the use of Cas9/gRNA targeting the SARS-CoV-2 sequence (‘19n’) compared to the use of non-targeting Cas9/gRNA (‘neg’), suggesting efficient Cas9 cleavage (Fig. 1b). However, addition of λ-exo to the reaction unexpectedly did not lead to degradation of cleavage product (Fig. 1b). We hypothesized that λ-exo loading was inhibited by Cas9 molecules remaining on the target DNA after cleavage. To overcome this obstacle, we hypothesized that a highly processive helicase, such as the recently engineered Rep-X ‘super’-helicase, could be used to evict Cas9, potentially through loading onto the cleaved nontarget strand. Indeed, we found that addition of Rep-X and ATP to the λ-exo reaction greatly increased digestion (‘19n’) (Fig. 1b, c). Cas9 combined with a non-target gRNA (‘neg’) resulted in no detectable degradation upon addition of Rep-X and/or λ-exo, consistent with Cas9 cleavage being the actual cause for DNA degradation via λ-exo (Fig. 1b, c).

To greatly simplify the diagnostic assay, we created a “master mix” composed of Glycine-KOH buffer with preformed SpCas9-gRNA complex, λ-exo, Rep-X, ATP, and MgCl2. Starting from RNA as the sample of interest, RT-LAMP was first employed for isothermal amplification at 65°C for 30 min on a heat block (alternatively using LAMP for DNA samples of interest) (Fig. 2a). 2 µL of 1:20-diluted RT-LAMP product was incubated with 18 µL of the SENTINEL master mix in two tubes, one composed of on-target (Reaction A) and the other of non-target Cas9/gRNA (Reaction B), for 15–30 min at room temperature. For a negative control reference to account for variation between reagent batches, RT-LAMP was also performed without RNA as the sample of interest, RT-LAMP was first employed for isothermal amplification at 65°C for 30 min on a heat block (alternatively using LAMP for DNA samples of interest) (Fig. 2a). 2 µL of 1:20-diluted RT-LAMP product was incubated with 18 µL of the SENTINEL master mix in two tubes, one composed of on-target (Reaction A) and the other of non-target Cas9/sRNA (Reaction B), for 15–30 min at room temperature. For a negative control reference to account for variation between reagent batches, RT-LAMP was also performed without RNA as the sample of interest, RT-LAMP was first employed for isothermal amplification at 65°C for 30 min on a heat block (alternatively using LAMP for DNA samples of interest) (Fig. 2a). 2 µL of 1:20-diluted RT-LAMP product was incubated with 18 µL of the SENTINEL master mix in two tubes, one composed of on-target (Reaction A) and the other of non-target Cas9/gRNA (Reaction B), for 15–30 min at room temperature. For a negative control reference to account for variation between reagent batches, RT-LAMP was also performed without RNA as the sample of interest, RT-LAMP was first employed for isothermal amplification at 65°C for 30 min on a heat block (alternatively using LAMP for DNA samples of interest) (Fig. 2a). 2 µL of 1:20-diluted RT-LAMP product was incubated with 18 µL of the SENTINEL master mix in two tubes, one composed of on-target (Reaction A) and the other of non-target Cas9/gRNA (Reaction B), for 15–30 min at room temperature. For a negative control reference to account for variation between reagent batches, RT-LAMP was also performed without RNA as the sample of interest, RT-LAMP was first employed for isothermal amplification at 65°C for 30 min on a heat block (alternatively using LAMP for DNA samples of interest) (Fig. 2a).

We aimed to summarize the three fluorescence measurements (A–C) as a single numerical score, where a high score would indicate detection of the target nucleic acid, i.e., a positive test. First, we calculated the fractional increase in LAMP product due to the presence of target sample (B/C), since samples can only have a positive test if LAMP successfully amplified. Second, we determined the fractional reduction in dsDNA fluorescence of the LAMP product with use of target versus non-targeting Cas9/gRNA (1–A/B). If the target product was amplified by LAMP, target Cas9/gRNA should efficiently cleave to allow λ-exo degradation, resulting in a larger reduction. In contrast, if undesired, off-target product was amplified by LAMP, then use of target Cas9 would have minimal effect relative to a non-targeting Cas9/gRNA, resulting in no reduction. Both quantities, (1–A/B) and (B/C), should have large values to have a positive test, motivating the final SENTINEL score to be computed as (1–A/B)×(B/C). The SENTINEL score is high if LAMP results in fluorescence increase upon dsDNA intercalation.

With the assay and its performance metric at hand, we evaluated the compatibility of the assay for different input types, as well as its limit of detection. Using SARS-CoV-2 in vitro transcribed single-stranded RNA (ssRNA) or a double-stranded DNA
(dsDNA) plasmid encoding the N-gene, we performed (RT)-LAMP using 5′ phosphorothiolated LAMP primers, followed by incubation with SENTINEL master mix containing either on-target gRNA targeting the SARS-CoV-2 template sequence or a non-targeting gRNA. We evaluated the capability of the assay to detect varying concentrations of template through sequential serial dilutions. First, we determined that the master mix was functional—input concentrations above 1 copy per microliter resulted in average SENTINEL scores of over 4, versus a score that approaches 0 when no template was supplied or template quantity under the limit of detection (Fig. 2b, c). Furthermore, we found that this strategy could reliably detect down to approximately 1 ssRNA or dsDNA molecules per microliter of input sample, which is in the attomolar concentration range. The sensitivity of this assay is bounded by the sensitivity of the LAMP step, which has been shown to be comparable to gold standard quantitative PCR (qPCR)23. Together, these results demonstrate that our assay exhibits attomolar levels of sensitivity with clear discrimination between positive and negative test results.

Next, we evaluated the assay’s ability to distinguish between closely related target sequences, the stability of its reagents, and its required reaction duration. We performed the assay on in vitro
transcribed synthetic RNA encoding either SARS-CoV-2 (19nCoV) or the closely related bat-SL-CoVZC45 (batCoV)24 N-gene. For the SENTINEL reaction, they were paired with Cas9 gRNAs targeting either 19nCoV (Cas9-19n-N2-gRNA#1) or batCoV (Cas9-bat-N2-gRNA#1), whose sequences differed by 5 nucleotides (Supplementary Fig. 1). Notably, use of RT-LAMP alone for 19nCoV detection resulted in a false positive readout when batCoV ssRNA was in the sample solution (Fig. 3a). In comparison, our method did not have a false positive readout–correct pairings of RT-LAMP product with gRNA resulted in a score of over 6, while incorrect pairings (i.e., 19nCoV gRNA with batCoV RT-LAMP product) resulted in a SENTINEL score of under 2 (Fig. 3b). We verified that this difference was due to large, expected reductions in DNA concentrations with the use of the on-target versus non-targeting gRNA, as measured by PicoGreen fluorescence (Fig. 3c). In addition, the master mix was stable for multiple –80 °C to room temperature freeze–thaw cycles (Fig. 3b, c), as well as long-term storage up to 1 month in –80, −20, and 4 °C, but not room temperature (Fig. 3d). Next, by performing the endonuclease reaction as a function of time, we found that even 5 min was sufficient to achieve clear discrimination between expected positive and negative samples (Fig. 3e).

Because SENTINEL relies on targeted endonuclease cleavage to actuate exonuclease-driven target discrimination, we evaluated whether other CRISPR endonucleases such as enhanced-specificity Cas9s and Cas12a25 would be compatible. To better evaluate the specificity advantages of engineered Cas9s in the SENTINEL system, we designed new gRNAs targeting the N-gene of SARS-CoV-2 (Cas9-19n-N2-gRNA#2) and bat-SL-CoVZC45 (Cas9-19n-N2-gRNA#2) that differed by only two nucleotides in the protospacer (Supplementary Fig. 1). Use of wild type Cas9 for 19nCoV detection was notable for a false positive readout when batCoV ssRNA was in the sample solution (Fig. 4a). In contrast, use of enhanced specificity HiFi Cas9 and eSpCas926,27 resulted in a much lower SENTINEL score for incorrect pairings of input nucleic acids and gRNA, consistent with better discrimination of two-nucleotide mismatches using enhanced specificity Cas9 systems (Fig. 4b–c). We also determined that use of Cas12a (AsCpf1) with gRNAs targeting either the 19nCoV (Cpf1-19n-N2-gRNA#1) or batCoV (Cpf1-bat-N2-gRNA#1) N-gene, which differed by 5 nucleotides in the protospacer (Supplementary Fig. 1), resulted in assay performance comparable to that of Cas9 (Fig. 4d). Notably, Rep-X and ATP were dispensable for the Cas12a reaction, consistent with automatic Cas12a departure from target DNA after cleavage28, which would facilitate λ-exo loading and target DNA degradation without need for Rep-X.

We also tested the capability of restriction enzymes29 to act as the sequence-specific endonuclease to actuate λ-exo cleavage. We identified the 6-cutter restriction enzyme Afel to have one target site in the 19nCoV RT-LAMP product, but not in the batCoV product due to a single mismatch. We replaced the previous Cas9/gRNA and Rep-X components with Afel, then performed SENTINEL (30 min at room temperature) on the RT-LAMP products of samples containing either 19nCoV or batCoV ssRNA. Because the majority of restriction enzymes are known to be functional in the acetate-based CutSmart buffer, we evaluated the reaction using that buffer as well as the previous Glycine–KOH-based λ-exo buffer. For both CutSmart and λ-exo buffer, we verified positive signal only for the 19nCoV sample, but not for the batCoV sample (Fig. 4e). The SENTINEL score was higher using Afel compared to CRISPR enzymes, due to a modest increase in degradation efficiency (1–A/B) and a greater increase in the ratio of B/C attributed to reduced background fluorescence (Supplementary Fig. 3a, b). Together, these results demonstrate that the assay has a high degree of flexibility for endonuclease selection and high specificity.

Next, we determined the ability of this assay to detect SARS-CoV-2 virus particles in human saliva. We obtained heat-inactivated SARS-CoV-2 virus originating from the culture medium of SARS-CoV-2-infected VeroTMRSS2 cells30. To mimic detection of viral particles in patient samples, we diluted virus titer in viral transport media (VTM), followed by 1:4
addition into mixed human saliva. To lyse the virus and extract its RNA, we adapted a previous protocol by adding 1:1 of QuickExtract lysis buffer, followed by heating to 95 °C for 5 min. 1 µL of lysed viral particles was used for RT-LAMP and subsequent steps of the SENTINEL assay (Fig. 4f). By evaluating serial dilutions of the virus titer in saliva, we observed high sensitivity, consistently detecting down to 10 particles per microliter of the original input (Fig. 4g).

Finally, we evaluated the ability for SENTINEL to detect SARS-CoV-2 virus from blinded nasopharyngeal (NP) swabs of 50 patients, 25 of which had detectable quantities of SARS-CoV-2 virus from quantitative PCR (qPCR). We directly mixed 2.5 µL of NP swabs with qPCR-detectable SARS-CoV-2, higher SENTINEL scores corresponded to greater quantity of target sequence, i.e., lower qPCR quantification cycle (Cq) (Fig. 5b). SENTINEL exhibited a sensitivity of 0.72 for this blinded set of patient samples, sensitivity of 1.00 (no false negatives) for the subset of samples with Cq under 30 (p < 0.001), and specificity of 1.00 (no false positives) (Fig. 5b, c). Receiver operating characteristic (ROC) analysis compared to qPCR Cq showed superior performance of SENTINEL over RT-LAMP alone (Fig. 5d), due to potential for non-specific amplification from RT-LAMP (Supp. Fig. 3c, d). Together these results demonstrate the applicability of SENTINEL for rapid detection of SARS-CoV-2 virus directly from patient-derived samples, and improvement over isothermal amplification (RT-LAMP) alone by eliminating potential false positives derived from non-specific amplification.

Discussion
SENTINEL is a first-in-class method for nucleic acid detection that utilizes the target cleavage properties of sequence-specific endonucleases to verify sequence identity. Our approach is generalizable, in that any sequence-specific endonuclease can be utilized, and orthogonal to existing technologies by using reagents, such as lambda exonuclease and Rep-X, not previously used for nucleic acid detection. These features also allow SENTINEL to be rapidly deployed to meet the demands of diagnostic nucleic acid testing, which, as demonstrated by the recent COVID-19 pandemic, is greatly expedited by the availability of diverse assay methodologies that can overcome supply chain bottlenecks. Furthermore, the limited number of liquid-handling steps and use of plate readers for readout allow SENTINEL to be amenable to 96/384-well plate formats and automation using liquid-handling robots for high-throughput sample processing. Incorporation of rapid RNA cleanup methods, such as using magnetic beads, may further improve detection sensitivity.

Future directions include extension to other formats such as portable diagnostic devices and further optimization of the protocol for patient-derived samples to achieve clinical-grade diagnostic testing for COVID-19 using SENTINEL. The emergence of new COVID-19 variants may present new challenges to nucleic acid detection with respect to increased false negatives, but SENTINEL is poised to address these challenges by leveraging capacity for single nucleotide discrimination to distinguish between different virus variants.

Other CRISPR-based methods using Cas12 (DETECTR) and Cas13 (SHERLOCK) exhibited comparable features to our strategy. The original SHERLOCK and DETECTR assays also require at least one heating temperature (37 or 65 °C) for isothermal amplification before CRISPR-based detection, and require ~1 h to complete the entire protocol. Recently, both strategies have been adapted for SARS-CoV-2 detection, STOPCovid.v2 simplified the protocol by designing a one-pot reaction that only requires one heating temperature. Compared to
to grow overnight (37 °C, 220 rpm) and then transferred to 2 L of LB supplemented
with detectable virus from qPCR—Cq below 40, 25 negative controls—Cq above 40). qPCR Cqs were initially measured from 300 µL of patient sample, and adjusted here to be consistent with the 2.5 µL of patient samples used for SENTINEL by adding log$_{10}$(300/2.5) = 6.9. The four red labels divide the graph into True Positives (‘TP’), False Positives (‘FP’), True Negatives (‘TN’), and False Negatives (‘FN’). e True positives with SENTINEL (+‘) had significantly lower qPCR Cq compared to false negatives (‘–’) (p < 0.001) from Student’s t-test. d Receiver operating characteristic (ROC) curves using SENTINEL versus RT-LAMP alone.

DETECTR for SARS-CoV-2, our strategy requires only 1 versus 2 different heating temperatures (because our second step is performed at room temperature whereas DETECTR’s second step is performed at 37 °C). Because DETECTR requires Cas12, only sequence regions adjacent to a Cas12-specific protospacer adjacent motif (PAM) can be evaluated, whereas the flexible choice of endonuclease with SENTINEL allows evaluation of regions next to PAMs of other CRISPR enzymes, or potentially even without PAM restrictions using near-PAMless Cas9 variants 38. Our design is also related to others that combine target cleavage with isothermal amplification 39,40. SENTINEL and related methods all have the risk of post-amplification contamination from downstream manipulation of amplification product, which can potentially be resolved using validated strategies such as UV-light irradiation, enzymatic inactivation with uracil-N-glycosylase 41, and newer enzymatic “eraser” strategies 42.

In conclusion, SENTINEL expands the scope of nucleic acid detection by combining the convenience of isothermal amplification with improved specificity to discriminate between one or two base pair differences. It extends nucleic acid detection by utilizing the on-target cleavage properties, rather than bystander collateral cleavage, of CRISPR enzymes to enable use by diverse CRISPR systems from enhanced-specificity Cas9 to Cas12a, as well as other endonucleases such as restriction enzymes. The entire protocol can be completed in under 1 h and only requires a 65 °C heat block and fluorometer on top of basic laboratory materials. The assay master mix is also stable to multiple freeze–thaw cycles and long-term storage.

**Methods**

**SpCas9 purification.** BL21-CodonPlus (DE3)-RIL competent cells (Agilent Technologies 230245) were transformed with Cas9 plasmid (Addgene #67881) and inoculated in 25 mL of LB-ampicillin media. The bacteria culture was first allowed to grow overnight (37 °C, 220 rpm) and then transferred to 2 L of LB supplemented with ampicillin and 0.1% glucose until OD$_{600}$ of ~0.5. Subsequently, the cells were induced with IPTG at a final concentration of 0.2 mM and maintained overnight at 18 °C. The bacteria cells were pelleted at 4500 × g, 4 °C for 15 min and resuspended in 40 mL of lysis buffer containing 20 mM Tris pH 8.0, 250 mM KCl, 20 mM imidazole, 10% glycerol, 1 mM TCEP, 1 mM PMSF, and cOmplete™ EDTA-free protease inhibitor tablet (Sigma-Aldrich 1183617001). This cell suspension was lysed using a microfluidizer and the supernatant containing Cas9 protein was clarified by spinning down cell debris at 16,000 × g, 4 °C for 40 min and filtering with 0.2 µm syringe filters (Thermo Scientific™ F25006). 4 mL Ni-NTA agarose bead slurry (Qiagen 30210) was pre-equilibrated with lysis buffer. The clarified supernatant was then loaded at 4 °C. The protein-bound Ni-NTA beads were washed with 40 mL wash buffer containing 20 mM Tris pH 8.0, 800 mM KCl, 20 mM imidazole, 10% glycerol, and 1 mM TCEP. Gradient elution was performed with buffer containing 20 mM to 400 mM HEPEs pH 8.0, 500 mM KCl, 10% glycerol, and varying concentrations of imidazole (100, 150, 200, and 250 mM) at 7 mL collection volume per fraction. The eluted fractions were tested on a SDS–PAGE gel and imaged by Coomasie blue (Bio-Rad 1610400) staining. To remove any DNA contamination, 5 mL Hitrap Q HP (Cytiva 17115401) was charged with 1 M KCl and then equilibrated with elution buffer containing 250 mM imidazole. The purified protein solution was then passed over the Q column at 4 °C. The flow-through was collected and dialyzed in a 10 kDa SnakeSkin™ dialysis tubing (Thermo Fisher 68100) against 1 L of dialysis buffer (20 mM HEPES pH 7.5, 500 mM KCl, 20% glycerol) at 4 °C overnight. Next day, the protein was dialyzed for an additional 3 h. in fresh 1 L of dialysis buffer. The final Cas9 protein was concentrated to 10 µg/µL using Amicon Ultra-15 Centrifugal Filter Unit, Ultracel-10 (Millipore Sigma UFC901008), aliquotted, and flash-frozen and stored at −80 °C.

eSpCas9 purification. The purification protocol was adapted from the manuscript associated with the eSpCas9 plasmid (Addgene #126769). Briefly, the eSpCas9 plasmid was transformed into BL21 Rosetta 2 (DE3) cells (Millipore EMD 71397) then grown in LB media with 10 µg/mL Kanamycin overnight at 37 °C. 10 mL of this culture was inoculated into 1 L of LB media with 10 µg/mL Kanamycin and grown to a final cell density of 0.6 OD$_{600}$ then chilled at 18 °C. The protein was expressed at 18 °C for 16 h. following induction with 0.2 mM IPTG. The cells were centrifuged at 6000 × g for 15 min at 4 °C, resuspended in 30 mL of lysis buffer (40 mM Tris pH 8.0, 500 mM NaCl, 20 mM imidazole, 1 mM TCEP) supplemented with cOmplete™ EDTA-free protease inhibitor tablet (1 tablet/30 mL; Sigma-Aldrich 1183617001), and then sonicated on ice. The lysate was cleared by centrifugation at 48,000 × g for 40 min at 4 °C, which was then bound to a 5 mL Mini Nuvia IMAC Ni-Charged column (Bio-Rad 7800812). The resin was washed extensively with a solution of 40 mM Tris pH 8.0, 500 mM NaCl, 20 mM imidazole, and the bound protein was eluted by a solution of 40 mM Tris pH 8.0, 250 mM imidazole, 150 mM NaCl, 1 mM TCEP, 10% glycerol was added to the eluted sample and the His6-MBP fusion protein was cleaved by TEV protease (Addgene plK793) (3 h at 25 °C). The volume of the protein solution was made up to 100 mL with buffer (20 mM HEPES pH 7.5, 100 mM KCl, 1 mM DTT). The cleaved protein was purified on a 5 mL HiTrap SP HP cation exchange column (GE Healthcare 17115201) and eluted with 1 M KCl, 20 mM HEPES pH 7.5, 1 mM DTT. The protein was further purified by size exclusion chromatography on a HiPrep 26/60 Sephacryl S-200 HR column (GE 17-1195-01) in 20 mM HEPES pH 7.5, 200 mM
KCI, 1 mM DTT, and 10% glycerol. The eluted protein was confirmed by SDS–PAGE and SimplyBlue™ SafeStain (Invitrogen LC6060). The protein was stored at −20 °C.

**Rep-X purification.** E. coli Rep helicase was purified and crosslinked as previously described (Arslan et al.7). Briefly, the pET28+ vector containing Rep-DM4 sequence was transformed into E. coli BL21(DE3) cells and grown in TB medium. When optical density reached OD600 = 0.6 the protein overexpression was induced with 0.5 mM IPTG. The cells were incubated overnight at 18 °C and harvested by centrifugation at 10,000 × g. Our Rep-DM4 contains 6x-His tag on its N-terminus for Ni-NTA affinity-column-based purification. The cell pellet was resuspended in a lysis buffer and cells were lysed with a sonicator. After binding Rep to Ni-NTA column and several washes, Rep was eluted with 150 mM imidazole-containing buffer. Rep concentration was kept below 4 mg/mL (~50 µM) to avoid aggregation.

Our Rep-DM4 mutant has four native cysteines removed (C18L, C43S, C167V, C400) and SSO mutagenesis is introduced for crosslinking. C178 and C400 are linked with bismaleimidoethane crosslinker (BMOE) to lock Rep in a closed conformation and form Rep-X. Optimal crosslinking is achieved at Rep concentration between 20 and 25 µM, with Rep to BMOE ratio of 1:5. Excess imidazole and crosslinker were removed by overnight dialysis in the storage buffer (50% glycerol, 600 mM NaCl, 50 mM Tris, pH 7.6). The samples are stored at −80 °C. This method achieves nearly 95% crosslinking efficiency.

**Generating synthetic ssRNA via in vitro transcription.** For SARS-CoV-2 N-gene, perform PCR using 0.5 µl each of 10 µM forward and reverse primers [IVT_19n_N2_FWD, IVT_19n_N2_REV; primer sequences in Table S1]. Mix with 4 µl of water, 0.5 µl of 2019-nCoV_N, Positive Control template plasmid (Integrated DNA technologies 100006625), and 10 µl of Q5 2× master mix (New England BioLabs M0494). Thermocycle at 98 °C for 30 s for an initial denaturation, followed by 35 cycles of 98 °C for 0.9 s, 72 °C for 30 s, then final extension of 72 °C for 2 min and 4 °C hold. Use QIAquick PCR purification kit (Qiagen 28104) to clean up PCR elution and elute in 35 µl EB supplied with kit—a agarose gel of PCR product should visualize band at 931 base pairs. For in vitro transcription, use the HiScribe T7 kit (New England BioLabs E2050) by mixing 17 µl of purified PCR product with 3 µl of 10× T7 transcription buffer, 8 µl of NTP mix, 2.5 µl of 50 mM DTT, and 2 µl of T7 RNA polymerase mix for a 30 µl final volume. iTermate at 37 °C at least 2 h. Then, add 20 µl of water and 1 µl of NTP mix, 2.5 µl of 50 mM DTT, and 2 µl of T7 RNA polymerase mix for a 30 µl final solution (HBSS), 10 ml heat-inactivated FBS (Corning), and 2 ml of filter-sterilized Gentamicin/Ampicillin B mixture. Supernatant from SARS-CoV-2-infected Vero cell culture with 1057 viral particles per µl was heated at 65 °C for 1 h for virus inactivation. Serial dilutions were performed using VTM. 1 ml of the dilution was mixed with 4 ml pooled human saliva (Innovative Research IRHUSLM5L) to simulate virus presence in human saliva. This was mixed 1:1 with QuickExtract™ DNA Extraction Solution (Lucigen QEO9050), then immediately heated to 95 °C for 5 min. 1 µl of this final solution was used in a 20 µl RT-LAMP reaction, with the remainder of the protocol identical to the section “CRISPR-based SENTINEL assay on synthetic ssRNA or dsDNA”.

**Crispr-based SENTINEL assay on heat-inactivated viral particles.** Gentamicin/Ampicillin B mixture was first made by mixing equal volumes of 50 mg/ml Gentamicin (Sigma-Aldrich G1397) with 250 µg/ml Ampicillin B (Sigma-Aldrich A2942), then filter-sterilized using a 0.22 µm pore size filter unit. Viral transport media (VTM) was prepared by mixing 500 ml of Hanks balanced salt solution (HBSS), 10 ml heat-inactivated FBS (Corning), and 2 ml of filter-sterilized Gentamicin/Ampicillin B mixture.

**Patient consent statement and study cohort.** From April 21, 2020 to July 16, 2020, non-hospitalized adults who were self-isolating after receiving a positive NP SARS-CoV-2 real-time reverse-transcription polymerase chain reaction (RT-PCR) result from the Johns Hopkins Medical Microbiology laboratory were approached for participation by telephone using a verbal consent script. Obtaining signed informed consent form for subjects enrolled in this study was not initially feasible for study staff due to the contagious nature of COVID-19 being studied under this protocol. Instead, the study staff used a consent waiver and obtained verbal consent. Inclusion criteria were age ≥18 years, able to receive study materials while remaining in isolation, and able and willing to perform self-collection of specimens. This protocol and verbal consent were approved by the Johns Hopkins University School of Medicine Institutional Review Board (IRB), and are identical to that of Manabe et al. (2020) and Antar et al. (2021). All procedures were in accordance with the ethical standards of the Helsinki Declaration of the World Medical Association.

**Specimen collection.** Participants were mailed a sample collection kit that included an international air transport association (IATA)-approved biologic specimen containment unit as well as sample collection materials and written instructions for sample collection. Participants self-collected mid-turbinate nasal and oropharyngeal (nasal-OP) swabs; both swabs were placed in 3 ml viral transport medium. All samples were immediately placed in the IATA container and stored in the participant’s freezer before shipping. Participants self-collected samples on the day they received the collection materials (day 0) and subsequently on study days 3, 7, 14, and on day 14, the participant shipped the collected samples on ice-cold packs to Johns Hopkins University for analysis using an overnight courier service. The procedure was identical to that of Manabe et al. (2020) and Antar et al. (2021)

**Blinding schema.** Individual 1 has information on ground-truth quantitative PCR readout for each patient sample. 25 SARS-CoV-2 positive samples and 25 SARS-CoV-2 negative samples were randomly placed in a freezer box by Individual 1. When Individual 1 was not present, Individual 2 removed the samples from the freezer box and performed the SENTINEL assay. Individual 1 sends the SENTINEL results to Individual 2 for verification.
SENTINEL assay on patient nasopharyngeal swabs. 35 µL of patient nasopharyngeal swabs was mixed 1:1 with QuickExtract™ DNA extraction solution (Lucigen Q50950), then immediately heated to 95 °C for 8 min. 2 µL of this final solution was used in a 50 µL SENTINEL reaction. 45 min was used for the RT-LAMP step, and detection was performed on the SpectraMax i3x plate reader (Molecular Devices). Everything else was identical to the section “CRISPR-based SENTINEL assay on synthetic ssRNA or dsDNA.”

Computation of SENTINEL score. Reaction A corresponds to the on-target SENTINEL reaction (on-target Cas9/gRNA or on-target restriction enzyme). Reaction B corresponds to the non-target SENTINEL reaction (non-target Cas9/gRNA or water in the place of restriction enzyme). Reaction C corresponds to the negative control—the sample expected to have no target nucleic acid, exposed to the non-target SENTINEL reaction. A–C will be the results of Reactions A–C, respectively, as fluorescence measurements (in arbitrary units) on a plate reader. The first part of the SENTINEL score is fractional reduction in fluorescence with on-target versus non-target Cas9/gRNA, i.e., 1–A/B. Next, this value is scaled by the fractional increase in DNA quantity due to LAMP amplification (B/C). Together, the SENTINEL score is computed using the following formula:

\[ (1−A/B) \times (B/C) \]

Statistics and reproducibility. All statistical analyses were conducted on either Microsoft Excel or GraphPad Prism 9. All data for statistical analysis were numerical, and a sample size and/or replicates of at least 3 were used for hypothesis testing using Student’s t-test. Replicates were all ‘biological’, in that each sample of the replicate was independently conducted and in separate test tubes.

Reporting summary. Further information on research design is available in the Nature Research Reporting Summary linked to this article.

Data availability. Source data underlying main figures are presented in Supplementary Data 1. Uncropped versions of gels and blots are presented in Supplementary Fig. 4.

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Author contributions
R.S.Z. and T.H. initially conceived the project. R.S.Z., L.S., and S.C. designed and performed experiments. M.G. purified Rep-X, Y.L. purified SpCas9, D.R. purified eSpCas9 using protocol adapted by D.R. and G.S. A.P. generated heat-inactivated SARS-CoV-2 virus. M.C., J.H., and Y.C.M. obtained patient samples and performed clinical qPCR. All authors contributed to the writing of the manuscript. T.H. supervised the project.

Competing interests
The authors declare the following competing interests: R.S.Z. and T.H. are listed as authors of a patent application submitted by Johns Hopkins University on the method of nucleic acid detection presented in this manuscript. G.S. serves on the Scientific Advisory Board of Dewpoint Therapeutics, Inc. The remaining authors declare no competing interests.

Additional information
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Correspondence and requests for materials should be addressed to Taekjip Ha.