Cite as: C. Jiao et al., Science 10.1126/science.abe7106 (2021).

Noncanonical crRNAs derived from host transcripts enable multiplexable RNA detection by Cas9

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CRISPR-Cas systems recognize foreign genetic material using CRISPR RNAs (crRNAs). In Type II systems, a trans-activating crRNA (tracrRNA) hybridizes to crRNAs to drive their processing and utilization by Cas9. While analyzing Cas9-RNA complexes from Campylobacter jejuni, we discovered tracrRNA hybridizing to cellular RNAs, leading to formation of "noncanonical" crRNAs capable of guiding DNA targeting by Cas9. Our discovery inspired the engineering of reprogrammed tracrRNAs that link the presence of any RNA-ofinterest to DNA targeting with different Cas9 orthologs. This capability became the basis for a multiplexable diagnostic platform termed LEOPARD (Leveraging Engineered tracrRNAs and On-target DNAs for PArallel RNA Detection). LEOPARD allowed simultaneous detection of RNAs from different viruses in one test and distinguished SARS-CoV-2 and its D614G variant with single-base resolution in patient samples.

CRISPR-Cas (Clustered Regularly Interspaced Short Palindromic Repeats and CRISPR-associated) immune systems degrade foreign genetic material through the guidance of CRISPR RNAs (crRNAs) (1, 2). crRNAs are encoded as spacerrepeat subunits within a system's CRISPR array (3). Each crRNA typically undergoes processing from a precursor transcribed from the array and then partners with the system's Cas effector nuclease to direct cleavage of target nucleic acids. Within Type II systems, the source of Cas9 nucleases and many CRISPR technologies (4, 5), crRNA processing and subsequent DNA targeting by Cas9 requires a trans-activating crRNA (tracrRNA) (6-8). The tracrRNA hybridizes to the "repeat" portion of each crRNA within the transcribed array. Host-derived RNase III then cleaves the formed RNA stem to generate a processed crRNA:tracrRNA duplex utilized by Cas9 (6). What remains unclear is whether crRNAs are confined to CRISPR-Cas loci or can be derived from elsewhere in the genome. Here, we show that crRNAs can be derived from host RNAs outside the CRISPR-Cas locus, inspiring a novel Cas9-based diagnostic platform that allows scalable detection of multiple biomarkers in a single test.

Cellular RNAs bound to Cas9 from C. jejuni resemble crRNAs

Our prior work interrogating RNAs bound to Cas9 from C. jejuni NCTC11168 (CjeCas9) revealed crRNA-guided RNA

targeting by CjeCas9 (9). To further explore RNA binding partners of CjeCas9, we repeated the immunoprecipitation and RNA sequencing (RIP-seq) approach of epitope-tagged Cas9 using C. jejuni strain CG8421 harboring only two spacers in its endogenous Type II-C CRISPR-Cas system (Fig. 1, A and B, and fig. S1, A and B). RIP-seq identified the CRISPRtracrRNA locus as well as 205 RNA fragments derived from cellular RNAs enriched with Cas9-3xFLAG (Fig. 1C, figs. S1C and table S1). Analyses of the enriched fragments using MEME (10) revealed two significant sequence motifs across all three replicates (Fig. 1D). Motif #1 was complementary to 13 nucleotides (nts) within the guide portion of crRNA2, in line with RNA targeting by crRNAs in NCTC11168 (9). Surprisingly, motif #2 was complementary to 21 nts within the tracrRNA anti-repeat domain. As this domain normally hybridizes to the crRNA repeat as part of crRNA biogenesis (fig. S2, A and B), motif #2 raised the intriguing possibility that these cellular RNAs were hybridizing with the tracrRNA, potentially becoming RNAs that function like crRNAs.

We explored this possibility through two routes. First, for the enriched RNA fragments with motif #2, we aligned the motif within each RNA fragment with a spacer-repeat pair, and we measured the length of each fragment corresponding to the spacer or repeat (Fig. 1E). Most frequently, the spacer part was 15-nt longer than a canonical crRNA spacer while the repeat part was the same size as a canonical crRNA

repeat, similar to slightly extended versions of crRNAs. Second, we predicted how each RNA fragment base pairs with the tracrRNA anti-repeat (*II*). Predicted binding affinities were significantly higher for RNA fragments with motif #2 than for fragments without the motif (p = 3×10^{-7}) (Fig. 1F). However, multiple RNA fragments were predicted to strongly pair with the tracrRNA anti-repeat despite lacking motif #2 (Fig. 1F), likely due to bulges in the RNA duplex creating discontinuities in the motif. For these RNA fragments and those containing motif #2, the predicted interactions between each RNA and the tracrRNA anti-repeat consistently contained imperfect RNA duplexes, with the most extensive pairing near the 3' end of the anti-repeat (Fig. 1G and fig. S2). We made similar observations for motif #1 (fig. S1D).

These crRNA-like RNAs raised the question whether these same RNAs were present in our prior RIP-seq analysis with strain NCTC11168 (9). We found that seven of the 96 enriched fragments were predicted to bind the tracrRNA anti-repeat more tightly than at least one crRNA (fig. S3, A and B). Two of these RNAs (derived from *fliF* and *dctA* mRNAs) matched those found in CG8421 (Fig. 1G, and figs. S2D and S3C). The RNA fragment derived from the fliF mRNA could be detected by Northern blot in both total RNA and RIP-seq samples yet disappeared following deletion of cas9 (Fig. 1H and fig. S4, A and B). The dctA RNA fragment was only weakly detected in one strain (fig. S4). While *cas9* deletion did not significantly perturb FliF protein levels in vivo under standard growth conditions (fig. S5), deleting the CRISPR array in NCTC11168 increased levels of the fliF RNA fragment (fig. S4C). Finally, the *fliF* RNA fragment was a processing product as confirmed with Terminator exonuclease treatment (Fig. 1G). These crRNA-like RNAs thus are also present in C. jejuni NCTC11168 and likely exist in other C. jejuni strains based on the shared tracrRNA binding site in *fliF* (fig. S6).

Co-immunoprecipitated RNAs can function as noncanonical crRNAs that direct DNA targeting by Cas9

Cas9 binding, predicted tracrRNA pairing, and the length distribution of many of these enriched RNA fragments suggested the tracrRNA pairs with endogenous RNAs, resulting in "noncanonical" crRNAs (ncrRNAs) (Fig. 2A and fig. S2A). The ncrRNAs therefore would be expected to direct Cas9 to complementary DNA targets flanked by a protospacer-adjacent motif (PAM), similar to a canonical crRNA (*12*). As none of the genes giving rise to the detected ncrRNAs has a correctly-placed PAM, the ncrRNAs are not expected to direct Cas9 to cleave their originating genomic site (table S1).

To evaluate ncrRNA-dependent targeting, we exploited a cell-free transcription-translation (TXTL) assay previously used to characterize CRISPR-Cas systems (13–15). As part of the assay, DNA constructs encoding CjeCas9, an RNA guide, and a GFP reporter harboring a target sequence flanked by a

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recognized PAM are added to the TXTL reaction. GFP fluorescence is then measured over time as a read-out of DNA binding and cleavage by CjeCas9 (Fig. 2B and fig. S7A). We focused on examining the *fliF* ncrRNA given its presence in both *C. jejuni* strains and detection by Northern blotting. Applying this assay to the tracrRNA and mRNA comprising the entire *fliF* coding region (1,683 nts) (Fig. 2C), we found that expressing the mRNA reduced GFP levels 2.5-fold compared to a non-targeting crRNA (p = 5.4×10^{-5}). Expressing the equivalent crRNA reduced GFP levels 15.1-fold compared to the non-targeting control. The reduced GFP silencing for the *fliF* mRNA versus the crRNA potentially reflects not only reduced targeting efficiency but also delayed complex formation. Overall, the TXTL results offer evidence that mRNA-derived ncrRNAs can direct DNA targeting by Cas9.

The reduced performance of the fliF mRNA in TXTL could be due to how an ncrRNA deviates from a standard crRNA. These deviations include the crRNA repeat sequence, the secondary structure of the duplex formed with the tracrRNA anti-repeat, and 5' or 3' extensions to the repeat that do not undergo efficient processing. To evaluate these deviations, we systematically mutated or extended the standard crRNA, either as an sgRNA to ensure duplex formation or as a crRNA/tracrRNA pair, and evaluated GFP silencing in TXTL (Fig. 2D and table S1). CjeCas9 could accommodate some mutations within the region of the repeat:anti-repeat duplex in the sgRNA implicated in nuclease binding (16). The more disruptive mutations spanned more nts, were closer to the 5' end of the repeat, or resulted in a bulge in the tracrRNA (e.g., s3, s6, s7, s9, s12, s18 in Fig. 2D, left panel). Observed differences in GFP silencing do not appear to arise from variable sgRNA levels (fig. S7B). Extending the sgRNA/crRNA ends or mutating the region cleaved by RNase III within the crRNA had minimal impact on GFP silencing (Fig. 2D, right panel). Overall, the tracrRNA can tolerate deviations from a standard crRNA as long as pairing through the 3' end of the tracrRNA anti-repeat is maintained.

We applied insights from our mutational analyses to prioritize putative ncrRNAs from *C. jejuni* CG8421 for functional tests in TXTL. In total, we identified eight RNA fragments predicted to base pair extensively with the 3' end of the tracrRNA anti-repeat (Fig. 2E and fig. S8). We then assessed GFP silencing by expressing up to 350 nts upstream and downstream of each associated ncrRNA-encoding gene with tracrRNA, CjeCas9, and the GFP reporter harboring each cognate DNA target. Of the eight tested RNAs, three (from *rseP*, *nuoL*, and *dctA*) yielded a >2-fold reduction in GFP reporter levels compared to a non-targeting crRNA control (p < 0.001). Furthermore, targeting was specifically through the predicted ncrRNA, as mutating the "seed" region of the putative ncrRNA (*17*), scrambling the tracrRNA anti-repeat, or replacing CjeCas9 with the orthogonal FnCas12a nuclease fully

relieved GFP repression (Fig. 2E). Multiple factors such as mRNA folding or accessibility during translation may explain why the other five ncrRNAs did not exhibit targeting activity in TXTL, as a linear-regression model built around the sgRNA mutants had limited ability to predict the targeting activity of these ncrRNAs (text S1). The current lack of predictability parallels guide design for RNA-targeting Cas13a nucleases, which only became predictable with extensive datasets and machine learning (18).

Beyond TXTL, we assessed ncrRNA function as part of DNA targeting in C. jejuni CG8421 and in E. coli. For CG8421, transformation interference assays did not yield any significant DNA targeting directed by ncrRNAs derived from the rseP, dctA, and nuoL mRNAs (fig. S9A), likely due to low ncrRNA abundance compared to the strain's crRNAs under the examined growth conditions. For E. coli, overexpressing the dctA mRNA, CjeCas9, and the tracrRNA led to moderate (15.5-fold) clearance of a transformed plasmid with the putative dctA ncrRNA target (p = 0.0036), but not when the tracrRNA anti-repeat was scrambled (1.6-fold) (p = 0.068) (fig. S9B). We therefore conclude that ncrRNAs derived from mRNAs can elicit DNA targeting in both in vivo and cell-free systems.

The tracrRNA can be reprogrammed to direct Cas9 activity by an RNA-of-interest

The conversion of a cellular RNA into an ncrRNA was based on sequences bearing complementarity to the tracrRNA antirepeat, analogous to natural crRNA biogenesis (fig. S2A). What if the tracrRNA anti-repeat sequence could be changed to hybridize to other RNAs while maintaining the appropriate structure for Cas9 recognition? If so, then the resulting reprogrammed tracrRNA (Rptr, pronounced "raptor") could specifically derive an ncrRNA from a cellular RNA. The resulting ncrRNAs can then guide Cas9 to matching DNA targets (Fig. 3A). While tracrRNA engineering has rarely been explored outside of sgRNAs or crRNA:tracrRNA duplexes (19), multiple studies have shown that the repeat:anti-repeat duplex of the sgRNA for the *Streptococcus pyogenes* Cas9 (SpyCas9) can be extensively modified as long as the secondary structure is maintained (20, 21).

CjeCas9 recognizes a perfect RNA duplex formed between the crRNA repeat and tracrRNA anti-repeat (fig. S2B) (16). Based on our mutational analysis of the crRNA repeat, we already observed that Cas9 can accommodate several mutations within the crRNA repeat:tracrRNA anti-repeat duplex (Fig. 2D). Therefore, we evaluated GFP silencing in TXTL after mutating both sides of the duplex in the *fliF* sgRNA while preserving the secondary structure (s26-s31, Fig. 3B). GFP silencing was maintained even when exchanging the sequence of the entire duplex (s31, Fig. 3B). Next, we reprogrammed the tracrRNA anti-repeat to form perfect 25-bp duplexes with

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three putative ncrRNAs (derived from fliF, CJ8421_04245, CJ8421_04975) exhibiting at most modest GFP silencing in TXTL (Fig. 2, C and E, and fig. S8). In all three cases, GFP repression was significantly enhanced with Rptrs than with WT tracrRNAs (p = 1×10^{-6} - 0.0011), even if repression was not as strong as with the canonical crRNA:tracrRNA pair (Fig. 3C). Finally, we reprogrammed the tracrRNA anti-repeat to base pair with entirely new regions of an mRNA (Fig. 3D). Starting with the CJ8421_04975 mRNA, we designed five different CjeCas9 Rptrs hybridizing to different locations (n1n5) in the mRNA (Fig. 3D and fig. S10A). Of these Rptrs, four yielded significantly reduced GFP levels compared to non-targeting crRNA controls (p = 6×10^{-7} - 0.002). Importantly, mutating the predicted seed region, scrambling the tracrRNA anti-repeat, or replacing CieCas9 with FnCas12a restored GFP expression (Fig. 3D). Northern blot analysis from TXTLextracted RNAs further revealed no detectable processed RNAs with a size resembling mature ncrRNA. Complete ncrRNA processing to a size similar to canonical crRNAs therefore may not be necessary for DNA targeting by CieCas9 (fig. S11), in line with the dispensability of RNase III for crRNA-mediated DNA targeting through the II-C CRISPR-Cas system in Neisseria meningitidis (22).

Given the functionality of CjeCas9 Rptrs, we asked whether tracrRNAs for other Cas9 homologs can be similarly reprogrammed. We selected the well-characterized Streptococcus pyogenes Cas9 (SpyCas9) and the Streptococcus thermophilus CRISPR1 Cas9 (Sth1Cas9) as examples. In both cases, we devised design rules for Rptrs based on the known secondary structure of the crRNA:tracrRNA duplex and the preference of RNase III to cleave double-stranded RNA with AT-rich sequences (fig. S10A) (20, 23). All ten designed Rptrs significantly reduced GFP levels compared to the non-targeting crRNA control (p = 1×10^{-7} - 1×10^{-4}) (Fig. 3D). As before, GFP expression was restored by disrupting the seed sequence in the ncrRNA guide, scrambling the tracrRNA anti-repeat, or swapping either of the Cas9's for FnCas12a. In many cases, the extent of GFP silencing approached that of the targeting crRNA control. We also evaluated plasmid clearance with Rptrs in E. coli for all three Cas9 orthologs, finding that each could elicit efficient plasmid clearance for at least one tested Rptr (fig. S10B). The targeted plasmid was efficiently cleared even when expressing the sensed mRNA at low levels (fig. S12A) or when deleting RNase III (fig. S12B). Overall, the tracrRNA for different Cas9 orthologs can be converted into Rptrs to elicit DNA targeting based on the presence of a selected cellular RNA.

Reprogrammed tracrRNAs enable sequence-specific detection by Cas9

By linking DNA targeting to an RNA-of-interest, Rptrs offer a unique opportunity for RNA detection and a different

paradigm for CRISPR diagnostics. Current CRISPR diagnostics principally rely on Cas12a or Cas13 searching for double-stranded DNA or RNA targets in a sample, where target recognition elicits non-specific single-stranded DNA or RNA cleavage of a fluorescent reporter (24–26). The non-specific readout practically limits one test to one target sequence. In contrast, Rptrs convert sensed RNAs into ncrRNAs, which would direct Cas9 to matching DNA. Cas9 binding or cleavage of a DNA sequence would then indicate the presence of the sensed RNA in the sample. Because the sequence of each DNA target is unique, large numbers of target sequences could be monitored in parallel in one test. We call the resulting diagnostic platform LEOPARD, for Leveraging Engineered tracrRNAs and On-target DNAs for PArallel RNA Detection (Fig. 4A).

To begin assessing LEOPARD, we performed a simplified in vitro reaction using T7-transcribed RNAs, commercially available SpyCas9 protein, and linear DNA targets (fig. S13A). We began with RNA corresponding to one of the synthetic ncrRNA loci within CJ8421 04975 (n4 under SpvCas9, Fig. 3D). Introducing an annealing step to hybridize the Rptr to the T7-transcribed ncrRNA vielded DNA target cleavage without adding RNase III or RNase A for ncrRNA: Rptr processing (fig. S13, B and C). The cleavage efficiency was also similar to the equivalent crRNA:tracrRNA pair, even when the ncrRNA sequence was extended on either end (fig. S13D). The timescale of the annealing step could also be minimized by rapid cooling of the samples (fig. S13E). With the annealing step, efficient cleavage occurred with a 100-fold excess of yeast total RNA, but only when the ncrRNA was present (Fig. 4B). LEOPARD therefore can report the presence of a specific RNA-of-interest based on cleavage of a DNA target and can be streamlined through further optimization.

LEOPARD allows for multiplexed RNA detection by Cas9 with single-base resolution

Realizing the full multiplexing potential of LEOPARD requires monitoring many DNA targets at once. To initially demonstrate this multiplex capability, we devised a readout scheme based on resolving distinct cleavage products from pooled DNA targets by gel electrophoresis (Fig. 4C). Each target is labeled with a fluorophore on one end, producing only two visualizable products-cleaved and uncleaved. We then applied this scheme to specifically detect nine ~150-nt RNA fragments associated with respiratory viruses, including two from SARS-CoV-2 coronavirus (the causative agent of COVID-19), six from other coronaviruses, and one from influenza H1N1 (Fig. 4D, and fig. S14). Each DNA target was cleaved by Cas9 only in the presence of the corresponding RNA, even when detecting three or five specific RNA fragments in the same reaction (Fig. 4D and fig. S14).

As the viral RNA sequences were selected to minimize

homology, we asked if LEOPARD could detect even a singlenucleotide difference. The D614G mutation in the spike protein of SARS-CoV-2 served as an example, as it comprises a single base change (A23403G) that increased infectivity and drove global spread (27). By placing this nt change within the seed region of the target, we could detect the WT or D614G RNA using one Rptr combined with either the WT or D614G target (Fig. 4E). The matching DNA target was preferentially cleaved when testing each target individually, although some cleavage of the non-matching target was observed. However, combining the two targets in a single reaction yielded discernable cleavage only for the matching target, presumably through preferential binding and cleavage of the perfect target by Cas9 (28). LEOPARD therefore can confer multiplexed RNA detection in a single reaction with single-base resolution.

To extend LEOPARD beyond this proof-of-principle demonstration, we made two additions. First, we added target-specific RT-PCR and in vitro transcription similar to Cas13-based diagnostics (29) to improve assay sensitivity beyond the threshold set by detection of a cleaved DNA product. Second, we resolved DNA targets using a Bioanalyzer as a more practical readout (Fig. 5A). Applying this modified workflow to sense the in vitro-transcribed WT SARS-CoV-2 RNA fragment, we could detect as little as approximately one copy, or 1.7 aM in the original dilution, of this RNA (Fig. 5B) and fig. S15A) compared to 3×10^8 copies, or 0.6 nM in the original dilution, without pre-amplification (fig. S15B). As this sensitivity would be sufficient for detecting SARS-CoV-2 RNA in patient samples, we applied LEOPARD to evaluate samples confirmed positive or negative for SARS-CoV-2 by RT-qPCR (Fig. 5C and table S1). The positive samples reflected a range of SARS-CoV-2 RNA levels down to ~2 aM. We then probed for both SARS-CoV-2 and the D614G variant as well as Influenza H1N1, the C. jejuni CJ8421_04975 mRNA (n4 under SpyCas9, Fig. 3D) as a non-human negative control, and the mRNA encoding human RNase P to confirm correct administration of the nasal swab using four Rptrs (one for both WT and D614G SARS-CoV-2 (Fig. 4E)) and five DNA targets. Of the four SARS-CoV-2-positive and five SARS-CoV-2-negative samples tested, RNase P mRNA but not the CJ8421 04975 mRNA or H1N1 RNA was detected in all nine samples (Fig. 5C and fig. S16). Strikingly, we detected the D614G variant of SARS-CoV-2 in all four positive samples, which was confirmed by sequencing pre-amplified cDNA (Fig. 5D). Although the sample size is small, detection of this variant suggests that it was spreading in Germany when the samples were collected. WT or D614G SARS-CoV-2 RNA was not detected in any of the negative samples, paralleling the RT-qPCR results (Fig. 5C). Importantly, each reaction allowed for parallel testing for five different RNAs, including controls, that would require separate reactions for other diagnostic platforms. These findings demonstrate the practical utility of LEOPARD for multiplexed RNA detection.

Discussion

Starting from the characterization of a native CRISPR-Cas9 system in the bacterial pathogen *C. jejuni*, we discovered that cellular transcripts can be the source of noncanonical crRNAs through hybridization with the tracrRNA. This discovery adds ncrRNAs to the list of RNA guides found in nature, including crRNAs, scaRNAs that similarly pair with the tracrRNA anti-repeat, and "natural" sgRNAs formed through upstream transcription of the tracrRNA (30, 31). Importantly, these prior examples are all encoded within CRISPR-Cas loci. So far, it remains unclear if ncrRNAs serve a physiological role in *C. jejuni*. Future studies therefore could help clarify whether ncrRNAs derived from outside CRISPR arrays are spurious off-target products that are tolerated by the host or if they confer yet-to-be-discovered functions extending beyond adaptive immunity.

We further demonstrated that Rptrs can link the presence of an RNA-of-interest to sequence-specific DNA targeting by Cas9. This capability could enable in vivo applications with Cas9 such as multiplexed transcriptional recording or transcription-dependent editing. The most immediate application involved multiplexed RNA detection in vitro through LEOPARD. LEOPARD adds to the existing CRISPR diagnostic platforms principally based on Cas12a or Cas13 (24-26, 29) while offering scalable multiplexing in a single reaction. Our reliance on gel electrophoresis or a Bioanalyzer provided a proof-of-principle demonstration of multiplexed detection, although either are difficult to implement beyond a dozen targets. Instead, incorporating microarrays or next-generation sequencing (32) can potentially monitor up to millions of targets by linking the presence of a specific RNA to binding of labeled Cas9 or cleavage of labeled DNA target at a specific location on a chip. Either approach could also enhance assay sensitivity due to the limited number of DNA molecules in a given cluster, potentially circumventing the need for RNA pre-amplification. Simpler setups involving lateral flow assays could also be developed, paralleling other CRISPR diagnostics (26, 33). With further development, LEOPARD could become a powerful diagnostic tool not only for the detection of viral variants distinguished by individual nts; it could also be used for other applications such as screening for cancer mutations, identifying pathogens and antibiotic-resistance markers, or determining gene expression profiles for drug susceptibility. Moreover, extending Rptrs to tracrRNA-dependent nucleases within Type V systems could help incorporate their unique attributes, such as signal amplification or programmable transposition (24, 34).

While systematically perturbing the standard crRNA:tracrRNA duplex, we found that many deviations-particularly

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outside of the 5' end of the repeat-were tolerated by Cas9 and still led to targeting of designed DNA targets. However, despite this promiscuity, targeting is still determined by the upstream guide sequence and the requirement for a flanking PAM. Thus, both anti-repeat hybridization and guide-dependent DNA targeting may limit off-targeting activity. While we did not observe any detectable off-targeting in vitro or in vivo, future work could devise design rules for Rptrs that account for potential off-targeting as well as on-target activity, similar to existing sgRNA design algorithms (35). In turn, these rules would help advance the utilization of any RNA into a sequence-specific guide for CRISPR technologies.

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ACKNOWLEDGMENTS

We thank Tatjana Achmedov for assistance with Northern blotting; Jiaqi Yu and Elena Vialetto for help with cloning; Daphne Collias for helping devise the acronym LEOPARD; Daphne Collias, Peter Fineran, Ioannis Mougiakos, and Jörg Vogel for critical feedback on the manuscript; Patricia Guerry for sharing the C. jejuni CG8421 strain; Rogier Louwen for serum against CjeCas9, Christina Homberger for technical support with the Bioanalyzer; Neva Caliskan for providing human DNA; Victor Corman for providing SARS-CoV-2 RNA for test validation; and Theresa Höfert and Lena Zubrod for conducting RT-gPCR with COVID-19 patient samples. Funding: Deutsche Forschungsgemeinschaft SH 580/9-1 (CMS), BE 6703/1-1 (CLB) Interdisciplinary Center for Clinical Research Würzburg (IZKF) project Z-6 Free State of Bavaria for COVID-19 research at the IHM. Author contributions: Conceptualization: CJ, SS, GD, CMS, CLB; Discovery of ncrRNAs: GD, CMS; Methodology: CJ, SS, GD, CMS, CLB; NGS Software: TB; Formal Analysis: CJ, SS, GD, TB, CMS, CLB; Investigation: CJ, SS, GD, NLP, TB, FW; Linear-regression model: YY, LB; Obtaining and assessing patient samples: CS, OK; Writing - Original Draft: CJ, SS, CLB; Writing - Review and Editing: CJ, SS, GD, NLP, TB, FW, CS, CMS, CLB; Visualization: CJ, SS, CLB; Supervision: CMS, CLB; Funding Acquisition: CMS, CLB. Competing interests: A provisional patent application has been filed on the related concept by CJ, GD, CMS, and CLB. The other authors declare that they have no competing interests. Data and materials availability: The NGS data from RIP-seq and differential RNA-seq were deposited into NCBI GEO under accession number GSE156266. Code for automating NUPACK folding predictions and calling enriched RNA peaks are available through Zenodo (36, 37). All other data in the main text or the supplementary materials are available upon reasonable request. This work is licensed under a Creative Commons Attribution 4.0 International (CC BY 4.0) license, which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited. To view a copy of this license, visit https://creativecommons.org/licenses/bv/4.0/. This license does not apply to figures/photos/artwork or other content included in the article that is credited to a third party; obtain authorization from the rights holder before using such material.

SUPPLEMENTARY MATERIALS

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Materials and Methods Supplementary Text Figs. S1 to S16 Table S1 References (38–51) MDAR Reproducibility Checklist

9 September 2020; accepted 9 April 2021 Published online 27 April 2021 10.1126/science.abe7106

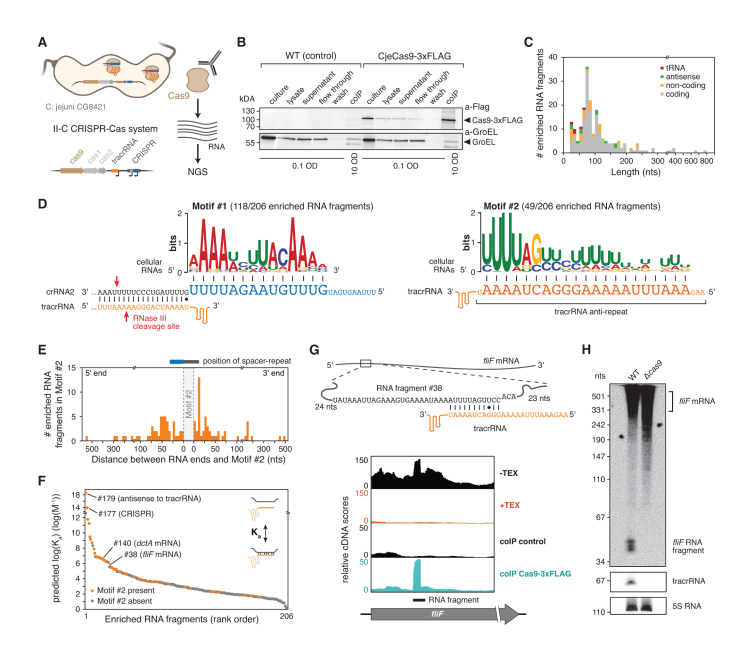


Fig. 1. Fragments of cellular RNAs bound by Cas9 in *Campylobacter jejuni* resemble crRNAs. (A) CoIP and sequencing RNAs bound to Cas9-3xFLAG from *C. jejuni* CG8421 using RIP-seq. (B) Western blot analysis of samples from *C. jejuni* strains with Cas9-3xFLAG or untagged WT control before and after immunoprecipitation. (C) Size range of the cellular RNA fragments identified through RIP-seq. Colors indicate the class of RNA. (D) Two motifs extracted by MEME from the enriched RNA fragments and their predicted interaction with crRNA2 or the tracrRNA. $E = 8.8 \times 10^{-22}$ (motif 1), 8.9×10^{-14} (motif 2). (E) Distribution of RNA lengths centered around motif #2. (F) Predicted binding affinity (K_a) between the tracrRNA anti-repeat and each enriched RNA fragment. Orange indicates the presence of motif #2. (G) Predicted *fliF* mRNA:tracrRNA duplex and mapped reads from differential RNA-seq (top) or RIP-seq (bottom) performed in CG8421. (H) Northern blot analysis for *fliF* RNAs from CG8421 WT and *cas9*-deletion strains.

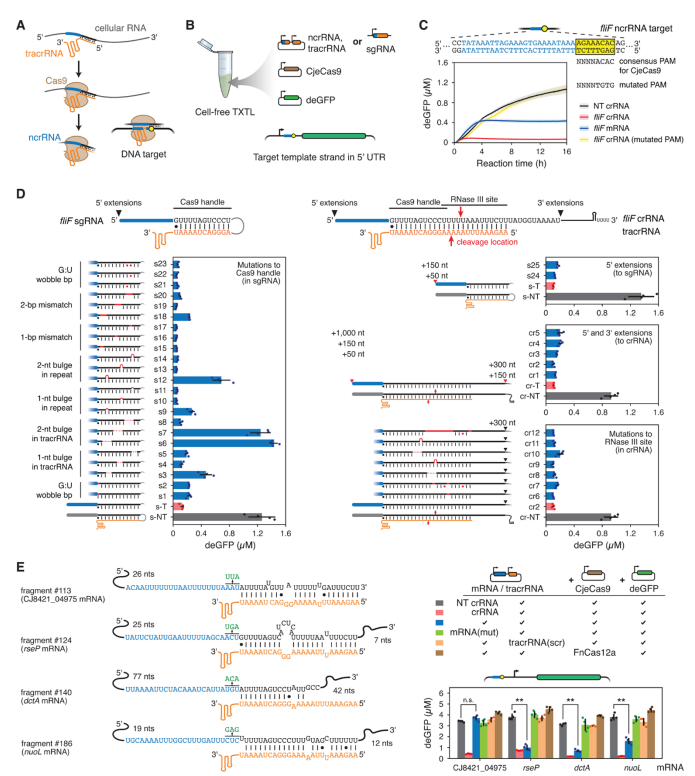


Fig. 2. Noncanonical crRNAs can direct DNA cleavage by CjeCas9. (A) General process for ncrRNA generation. (B) Applying the TXTL assay to characterize putative ncrRNAs. (C) DNA targeting through the *fliF* ncrRNA in TXTL. Lines and shaded regions indicate the mean and standard deviation from four separately-mixed replicates. NT: non-targeting. (D) Systematic evaluation of mutating the repeat:anti-repeat duplex for CjeCas9 with TXTL. Endpoint GFP levels are shown. Mutations and extensions to the *fliF* sgRNA/crRNA repeat are indicated in red. See table S1 for sequences. (E) DNA targeting by selected ncrRNAs predicted in TXTL. Check marks indicate use of the construct above the line. mRNA(mut): mRNA encoding the ncrRNA with point mutations in the predicted "seed" region of the guide. tracrRNA(scr): tracrRNA with the anti-repeat sequence scrambled. Values in D, E represent the mean and standard deviation from four separately-mixed replicates. **: p < 0.001. n.s.: not significant.

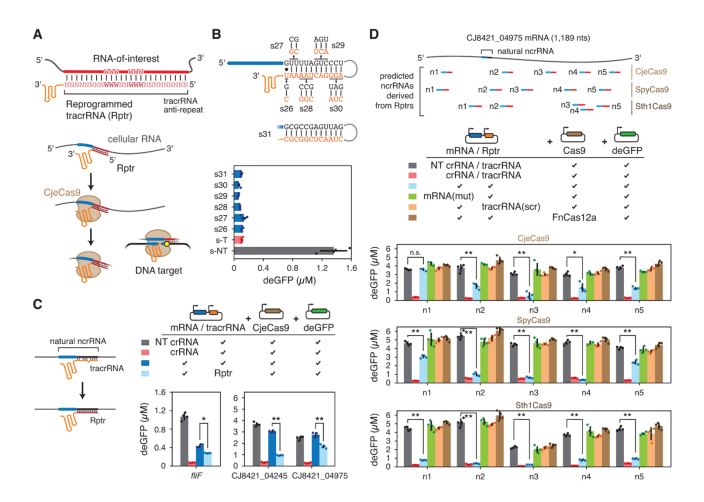


Fig. 3. Reprogrammed tracrRNAs co-opt RNA transcripts to guide different Cas9 orthologs. (A) Design of reprogrammed tracrRNAs (Rptrs) utilized by CjeCas9. W = A or T. PAM: yellow circle. (B) Toleration of mutations to the RNA duplex of an sgRNA that preserve secondary structure in TXTL. (C) Enhancing DNA targeting by less functional or non-functional ncrRNAs by converting the tracrRNA into a Rptr in TXTL. (D) Sequence-specific DNA targeting in TXTL using Rptrs compatible with three different Cas9 nucleases. Values in B - D represent the mean and standard deviation from four replicates in TXTL. *: p < 0.01. **: p < 0.001. n.s.: not significant.

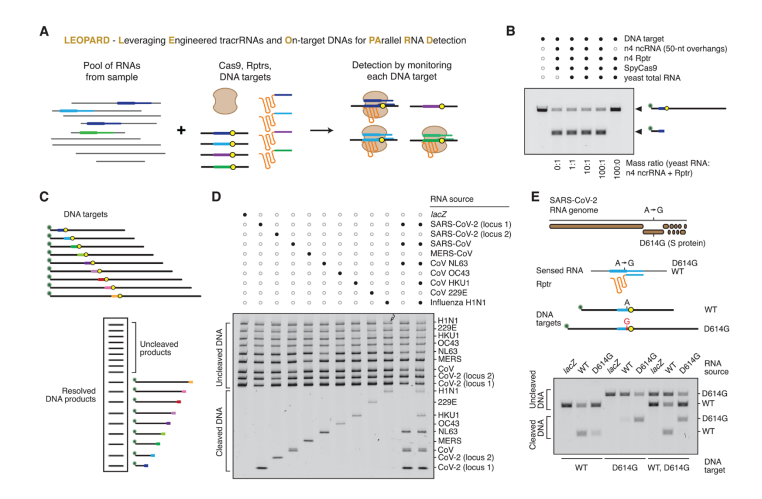


Fig. 4. Reprogrammed tracrRNAs enable multiplexable RNA detection with single-base resolution in vitro. (A) Overview of the multiplexed diagnostic platform LEOPARD. (B) Highly specific cleavage of target DNA by SpyCas9 in vitro. Targeting was directed by the ncrRNA associated with the n4 locus in CJ8421 04975. (C) Multiplexed monitoring of DNA targets. (D) Parallel detection of nine different RNA fragments associated with respiratory viruses by denaturing PAGE. Filled dots in D and E indicate the presence of the component listed on the right. (E) Specific detection of the D614G point mutation within the spike protein of SARS-CoV-2. Each detected T7-transcribed RNA in B, D and E comprised the intended ncrRNA sequence with 50-nt extensions on either end.

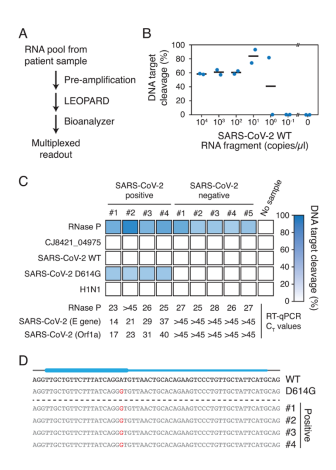


Fig. 5. LEOPARD with RNA pre-amplification and Bioanalyzer readout allows for multiplexed detection of SARS-CoV-2 in patient samples. (A) General workflow for LEOPARD with target-specific pre-amplification and DNA target resolution on a Bioanalyzer. (B) Sensitivity of the workflow for detecting dilutions of an in vitro-transcribed WT SARS-CoV-2 RNA fragment. One microliter was added for each test. Bars represent the average of independent duplicates. (C) Multiplexed detection of five RNAs in patient samples confirmed positive or negative for SARS-CoV-2 by RT-qPCR. (D) Sanger sequencing results of the detected region in SARS-CoV-2 cDNA from the positive patient samples. Blue bar: Position of the ncrRNA with the thick part indicating the resulting ncrRNA portion.



Noncanonical crRNAs derived from host transcripts enable multiplexable RNA detection by Cas9

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published online April 27, 2021

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