Cas9-mediated targeting of viral RNA in eukaryotic cells

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Clustered, regularly interspaced, short palindromic repeats–CRISPR (CRISPR-Cas) systems are prokaryotic RNA-directed endonuclease machineries that act as an adaptive immune system against foreign genetic elements. Using small CRISPR RNAs that provide specificity, Cas proteins recognize and degrade nucleic acids. Our previous work demonstrated that the Cas9 endonuclease from Francisella novicida (FnCas9) is capable of targeting endogenous bacterial RNA. Here, we show that FnCas9 can be directed by an engineered RNA-targeting guide RNA to target and inhibit a human +ssRNA virus, hepatitis C virus, within eukaryotic cells. This work reveals a versatile and portable RNA-targeting system that can effectively function in eukaryotic cells and be programmed as an antiviral defense.

To determine if FnCas9 was directly associated with HCV RNA, we performed coimmunoprecipitation experiments. We transfected cells with an HA epitope-tagged version of the protein [which maintained its ability to inhibit HCV (Fig. S3 A and B)] as well as the 5′ UTR-targeting rgRNA and subsequently infected the cells with HCV. FnCas9 was immunoprecipitated from cell lysates, associated RNA was purified, and quantitative real-time PCR was performed for the rgRNA and HCV RNA. The rgRNA was present in association with FnCas9 (Fig. 2A), as was HCV RNA (Fig. 2B), suggesting that HCV RNA was directly targeted by the FnCas9:rgRNA complex. A nonspecific rgRNA did not facilitate this interaction (Fig. 2B) but did associate with FnCas9 (Fig. 2C). Thus, FnCas9 can be targeted to associate with viral RNA within eukaryotic cells.

We next sought to determine how FnCas9 inhibited HCV, testing whether its endonucleolytic activity was required and if it

**Significance**

The clustered, regularly interspaced, short palindromic repeats associated endonuclease, Cas9, has quickly become a revolutionary tool in genome engineering. Utilizing small guiding RNAs, Cas9 can be targeted to specific DNA sequences of interest, where it catalyzes DNA cleavage. We now demonstrate that Cas9 from the Gram-negative bacterium Francisella novicida (FnCas9) can be reprogrammed to target a specific RNA substrate, the genome of the +ssRNA virus, hepatitis C virus, in eukaryotic cells. Further, this targeting results in inhibition of viral protein production. Overall, programmable Cas9-mediated viral RNA targeting likely represents one of myriad potential applications of FnCas9 in RNA targeting in eukaryotic cells.
inhibits translation and/or viral replication. As a first control, we found that addition of a nuclear localization signal (NLS) to FnCas9 abrogated its repression of viral protein production (Fig. 3A and Fig. S4), in line with its targeting of cytosolic HCV RNA. Because Cas9 proteins including FnCas9 are known to cleave DNA through two conserved structural domains, the RuvC and HNH endonuclease domains (7), we considered that these regions might be important for inhibiting HCV. We therefore generated alanine point mutations in the conserved RuvC and HNH active sites of FnCas9 (D11A and H969A, respectively). Despite mutation in one or both of these domains, FnCas9 maintained its ability to inhibit HCV (Fig. 3B). However, mutation in the RNA-binding arginine-rich motif (ARM; R59A), which is necessary for the interaction of Cas9 with nucleic acids (8, 14, 15), resulted in diminished HCV inhibition (Fig. 3B). Although these data do not exclude a model whereby accessory cellular RNases are recruited to the target by FnCas9, or that FnCas9 possesses an additional domain with endonucleolytic activity, these data do suggest that endonuclease

**Fig. 1.** FnCas9 can be reprogrammed to inhibit viral protein production in eukaryotic cells. (A) rgRNA schematic with targeting sequences (gray highlight) against the 5′ or 3′ UTR of HCV genomic RNA. (B) Huh-7.5 cells were transfected with the indicated combinations of FnCas9 and rgRNA and infected 48 h later with HCV encoding Renilla luciferase. At 72 h, cells were fixed and stained with anti-E2 antibody and imaged. (C) E2-positive foci from B were quantified and plotted as percent inhibition compared with the vector control. (D) Quantification of viral luciferase production displayed as percent inhibition compared with the vector control (n = 3; bars represent the SEM; data are representative of at least six experiments).
activity of the canonical DNA cleavage motifs are dispensable for FnCas9 inhibition of HCV.

We subsequently tested whether FnCas9 could inhibit translation of HCV RNA. We performed an in vitro translation reaction using immunoprecipitated FnCas9 from transfected Huh-7.5 cells, purified RNA from cells transfected with either the 5′ UTR-targeting rgRNA or the nontargeting RNA, as well as an in vitro transcribed HCV genomic RNA. Addition of both FnCas9 and the 5′ UTR-targeting rgRNA resulted in decreased translation of the HCV genome, as measured by viral luciferase production (Fig. 3C). Furthermore, a catalytically inactive FnCas9 (D11A/H969A) maintained its ability to inhibit translation of HCV, whereas the ARM (R59A) mutant displayed less translational inhibition (Fig. 3C). Taken together, these data suggest that FnCas9 is capable of directly inhibiting translation of target RNA. To determine whether FnCas9 also inhibits replication of HCV RNA, we targeted the negative-sense strand of the 5′ UTR (generated during replication and which is untranslated). Such targeting resulted in inhibition of HCV (Fig. S5). Together with the targeting of the 3′ UTR (which is involved in viral replication, but not translation), this suggests that FnCas9 is capable of inhibiting viral replication as well. Overall, these data strongly support a model whereby FnCas9 binds HCV RNA and inhibits the function of both translational and replication machineries.

We subsequently tested the sequence requirements for RNA targeting. Cas9 proteins require a short sequence adjacent to the targeted region, called a proto-spacer adjacent motif (PAM) (14). We sought to determine if a similar conserved adjacent region was necessary for HCV inhibition. A 5′ UTR-targeting rgRNA shifted to lack similar adjacent sequences still inhibited HCV (Fig. 4A). In fact, no common features are observed in the sequences adjacent to the targets of rgRNAs used in this study (Fig. S6A). In contrast, DNA targeting by FnCas9 endogenously within F. novicida absolutely requires a PAM (Fig. S6 B and C). Thus, these data demonstrate that FnCas9-mediated inhibition of HCV is independent of sequences adjacent to the targeted region.

DNA-targeting RNAs used by Cas9 require a 3′ seed sequence within the targeting region, and even a single mismatch in this region can abrogate function (16, 17). To test if there was a
similar requirement for RNA targeting, we generated a panel of rgRNA mutants containing mismatches within the targeting sequence. We found that mutation of up to six bases within the 3′ targeting region of the rgRNA was tolerated without loss of HCV inhibition (Fig. 4B). Longer regions of mismatched bases at the 3′ end resulted in a loss of activity (Fig. 4B). Specific mismatches in the 5′ region of rgRNA were also nonfunctional (Fig. 4C). A single mismatch in either the first or second base was sufficient to abrogate viral inhibition (Fig. 4C). However, a mismatch in the third base alone did not lead to a loss of activity (Fig. 4C). These data strongly suggest that unlike DNA targeting by other Cas9 proteins, FnCas9 inhibition of HCV requires a critical sequence in the 5′ end rather than the 3′ end of the targeting region of the guiding RNA (16, 17).

The previous experiments demonstrated that FnCas9 could target an RNA and facilitate resistance to HCV infection. We next tested whether FnCas9 could be introduced following an established viral infection and inhibit the virus (Fig. 5A). Transfection of HCV-infected Huh-7.5 cells with FnCas9 and 5′ or 3′ UTR-targeting rgRNAs resulted in decreased viral protein production (Fig. 5B), whereas expression of either FnCas9 or rgRNAs alone was not sufficient to inhibit HCV infection (Fig. 5B). Thus, the FnCas9:rgRNA machinery can combat both new and established viral infections.

**Fig. 4.** RNA sequence requirements for FnCas9 inhibition of HCV. (A) Huh-7.5 cells were transfected with FnCas9 using the rgRNA mutants in the indicated shifted alignments. At 72 h, viral luciferase was quantified and the percent inhibition compared with the nontargeting rgRNA is displayed (n = 12; data are compiled from three independent experiments). (B) Experiments were performed as above, using the mutants in the 3′ region indicated in the alignment below the figure (n = 12; bars represent the SEM; data are compiled from three independent experiments). (C) Experiments were performed as above, using the mutants in the 5′ region indicated in the alignment below the figure (n = 12; bars represent the SEM; data are compiled from three independent experiments).
Discussion
These data demonstrate the successful adaptation of the CRISPR-Cas prokaryotic immune system as an intracellular eukaryotic antiviral defense. Although other CRISPR-Cas systems can target RNA in archaea (18–20) and bacteria (21), and recently Streptococcus pyogenes Cas9 (SpCas9) has been shown to cleave RNAs in vitro (22), this work demonstrates the reprogramming of a Cas protein (FnCas9) to target RNA within a eukaryotic cell. Intriguingly, we find that orthologous Cas9 proteins from diverse type II CRISPR-Cas families, including S. pyogenes, Streptococcus thermophilus, and Neisseria meningitidis, are also capable of inhibiting HCV during cellular infection (Figs. S7 and S8). This suggests a broader capability of diverse Cas9 proteins to target and associate with RNAs of interest. Our results further demonstrate that FnCas9 inhibition of HCV is PAM-independent, unlike the in vitro RNA-targeting ability of SpCas9, which requires exogenous PAM-encoding oligomers (22). Thus, this method of RNA inhibition may be more flexible in its targeting. Importantly, expression of FnCas9 and either the HCV targeting or control rgRNA did not lead to significant changes in host cell gene expression compared with cells expressing a vector control, demonstrating the specificity of these complexes (Fig. S9 and Dataset S2). Furthermore, the presence of FnCas9 in the cytosol is necessary to inhibit HCV, in contrast to DNA targeting by Cas9 that relies on nuclear localization (23). Cytosolic RNA targeting would potentially limit FnCas9 off-target effects on host DNA.

Because the lifecycles of viruses with both RNA and DNA genomes require an RNA stage (generated during transcription, replication, or both), and FnCas9 can target both negative- and positive-sense strands of RNA and inhibit by blocking both viral translation and replication machineries, it is likely that the FnCas9:rgRNA machinery can be used to target diverse viruses, including both +ssRNA viruses (such as flavivirus, poliovirus, and rhinovirus) and –ssRNA viruses (such as filovirus, paramyxovirus, and orthomyxovirus). Furthermore, some eukaryotic viruses have mechanisms to circumvent eukaryotic RNA-targeting antiviral defenses, such as classical RNAi systems (24–26); however, these viruses have not evolved in the presence of Cas9, so it is unlikely that they have Cas9 evasion strategies. Thus, the FnCas9:rgRNA machinery could facilitate the targeting of viruses as soon as their genome sequences are available, without knowledge of the virus lifecycle or host receptors.

Given the vast success of Cas9 as a mediator of genome engineering in a multitude of species (7, 9, 10, 16, 27–33), our data suggest that FnCas9 could be used in a broad range of systems, representing an innovative paradigm in Cas9-mediated genetic engineering. This work demonstrates a portable, interdomain machinery capable of viral inhibition, likely just one of myriad potential biotechnological and medical applications of Cas9-mediated RNA targeting.

Materials and Methods

Plasmid Construction. FnCas9 was amplified using the primers found in Table S1 and cloned into the Xbal and Pmel sites of pdCNA3.3 (Invitrogen). FnCas9 point mutants were amplified from strains published previously (8). SpCas9, StCas9, and NmCas9 were amplified from Addgene vectors 48669, 41815, and 47872, respectively. To create rgRNA vectors, F. novicida CRISPR repeat sequences and the indicated targeting sequence were cloned into the gRNA-encoding plasmid from the Church Lab (Addgene 41824) (9), within the pCR-Blunt-II (Invitrogen) backbone, using overlapping PCR and the primers indicated in Table S1.

Cell Lines and Culture Conditions. Huh-7.5 cells were maintained in Dulbecco’s modified Eagle medium (DMEM; BioWhittaker) containing 10% (vol/vol) FBS (HyClone) and 100 μg/mL of penicillin/streptomycin (Cellgro) at 37° in 5% CO₂.

Plasmid Transfection. Huh-7.5 cells were seeded to 80% confluence in 24-well plates in DMEM without antibiotics the day before transfection. We transfected 800 ng of total plasmid DNA using Lipofectamine 3000 transfection reagent (Invitrogen) according to the manufacturer’s instructions. Six hours following transfection, cell supernatants were aspirated and replaced with DMEM supplemented with FBS and antibiotics.

Virus Transcription and Transfection. Rluc virus used for luciferase assays encodes the Renilla luciferase gene between the p7 and NS2 coding sequences of the previously described JFH/JFH genotype 2a infectious clone Cp7 (13). Rluc and Cp7 viral RNA were prepared as previously described (13). Purified plasmid encoding the CDNA copy of the full-length viral genome was linearized, and 5’ overhangs were digested with Mung Bean Nuclease (New England Biolabs). Linearized DNA was purified with phenol–chloroform extraction and ethanol precipitation. Transcription of the linearized DNA template was performed using a MEGAscript T7 kit, and the linear template was treated with Dnase I (Ambion). RNA was purified with phenol–chloroform extraction and isopropanol precipitation, and concentration and purity were determined using a Nanodrop spectrophotometer and standard agarose gel electrophoresis.

Huh-7.5 cells were grown to 70% confluence, trypsinized, and washed twice in PBS. We mixed 10 μg of purified RNA with 0.4 mL Huh-7.5 cells suspended at a concentration of 2 × 10⁵ cells/mL. Samples were electroporated in BTX 2-mm gap cuvettes (Harvard Apparatus, Inc.) using an ECM 830 apparatus (BTX Genetronics) with five pulses of 99 μs at 820 V over 1.1 s. Cells were suspended in 25 mL complete DMEM, and virus was harvested and stored at ~80 °C 3 d following transfection.

Fig. 5. FnCas9 can inhibit an established viral infection. (A) Experimental outline. HCV-transfected Huh-7.5 cells were transfected with FnCas9 and the indicated targeting RNAs, after 72 h postinfection. (B) Quantification of viral luciferase production, displayed as percent inhibition compared with the vector control (n = 3; bars represent the SEM; data are representative of at least 12 experiments).
Immunoprecipitation. Anti-HA IP was performed according to the manufacturer’s instructions from lysates of Huh-7.5 cells infected with HCVcc and transfected with FncCas9 and rrRNA expression vectors as indicated (Sigga Aldrich). Total RNA was extracted from the precipitated fraction using an RNeasy Mini Kit (Qiagen) according to the manufacturer’s instructions. Translated viral luciferase was measured as described below.

In Vitro Translation Assay. Immunoprecipitated FncCas9 was incubated with 1 μg HCV RNA, and 500 ng RNA from Huh-7.5 cells transfected with either the 5’ UTR-targeting rrRNA or the control rrRNA, in conjunction with the rabbit reticulocyte lysate in vitro translation kit (Life Technologies) according to the manufacturer’s instructions. Translated viral luciferase was measured as described below.

Quantitative RT-PCR. Quantitative reverse transcription reactions were performed using Taqman One Step RT-PCR Master Mix Reagent (Applied Biosystems) and primers specific for the 5’ UTR of HCV (Table S1). Sample analysis was performed on an Applied Biosystems 7500 apparatus. rrRNAs were quantified via Syber Green One Step RT-PCR, with primers found in Table S1.

Luciferase Assays. Huh-7.5 cells in a 96-well plate format were lysed and analyzed for relative light activity using the Renilla Luciferase Assay System (Promega) according to the manufacturer’s instructions. Briefly, 50 μL of Renilla substrate in assay buffer was added to 50 μL of cell lysate, and relative light units were quantitated on a Clarity 4.0 microplate luminescent (Biotek).

Immunohistochemistry. Six thousand Huh-7.5 cells per well were plated in collagen-coated 96-well plates. The following day, cells were infected with the J6/JFH genotype 2a virus C7. Following 3 d of incubation, cells were fixed with methanol, washed twice with PBS, and permeabilized with PBS containing 0.1% Tween-20 (PBS-T). Fixed cells were incubated in blocking buffer containing 1% BSA and 0.2% skim milk for 30 min. Endogenous peroxidase was blocked using 3% (vol/vol) H2O2, and then cells were washed twice with PBS and once with PBS-T. Cells were then incubated with the 2C1 monoclonal antibody to HCV E2 glycoprotein for 1 h at room temperature. After two washes with PBS and one with PBS-T, cells were incubated with ImmPRESS anti-mouse HRP (Vector Laboratories), washed, and developed using DAB substrate (Vector Laboratories).

Plasmid Inhibition Transformation Assay. The spacer sequence from the F. novicida crRNA1 (nucleotides 818163–818196 in the F. novicida U112 genome) was cloned into the pBav plasmid (KanR) (Addgene 26702) using the overlapping PCR primers found in Table S1 to create a dsDNA target for FncCas9. The pBav plasmids were transfected individually (250 ng plasmid/ transformation) into chemically competent U112 (300 μL) as described previously (4). Transformants were selected for on TSA containing 0.2% cysteine and 30 μg/mL kanamycin.

RNA-Seq Preparation and Analysis. Two sets of Huh-7.5 cells transfected with Cas9 + 5’ UTR-targeting rrRNA, Cas9 + Control rrRNA, or vector only were sorted by GFP on a FACs Aria. Total RNA was prepared using the QIAGEN RNeasy Micro Kit. Libraries were generated from 500 pg of total RNA using the CLONTECH SMARTer Ultralow V3 kit, and barcoding and sequencing primers were added using the Illumina NexeraXT DNA kit. Libraries were assessed by Bioanalyzer capillary electrophoresis, quantified, pooled, and sequenced on an Illumina MiSeq system as 150-base single-read reactions. Average read depth was 5.8 M, and all samples had Q30 scores >93.9%. Transcripts were aligned to the hg19 reference genome using the STAR v1.4.0g1 software (34). Transcript quantitation and differential expression were performed with the Cushlinks suite of tools and Cuffdiff v2.1.1 (35). Sequenced reads were deposited into the Sequence Read Archive database.

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