Sequence-specific antimicrobials using efficiently delivered RNA-guided nucleases

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Current antibiotics tend to be broad spectrum, leading to indiscriminate killing of commensal bacteria and accelerated evolution of drug resistance. Here, we use CRISPR-Cas technology to create antimicrobials whose spectrum of activity is chosen by design. RNA-guided nucleases (RGNs) targeting specific DNA sequences are delivered efficiently to microbial populations using bacteriophage or bacteria carrying plasmids transmissible by conjugation. The DNA targets of RGNs can be undesirables such as non-pathogenic enterobacteria and enterohemorrhagic Escherichia coli. Delivery of RGNs significantly improves survival in a Galleria mellonella infection model. We also show that RGNs enable knockdown of targeted strains based on genetic signatures. RGNs constitute a class of highly discriminatory, customizable antimicrobials that can target specific DNA sequences using CRISPR technology and effect cell killing or plasmid loss upon detection of genetic signatures associated with virulence or antibiotic resistance. The specificity of CRISPR-Cas is dictated by short, spacer sequences flanked by direct repeats encoded in the CRISPR locus, which are transcribed into CRISPR RNAs (crRNA). With the aid of a trans-activating small RNA (tracrRNA), crRNAs enable the Cas9 endonuclease to introduce double-stranded breaks in target DNA sequences. Through simple modifications of spacers in the CRISPR locus, an RGN can direct cleavage of almost any DNA sequence, with the only design restriction being a requisite NGG motif immediately 3′ of the target sequence.

There is mounting concern over the emergence and proliferation of multidrug-resistant bacterial pathogens and the dwindling treatment options for these organisms. Recently, carbapenem-resistant Enterobacteriaceae, a group of intestinal gram-negative bacteria known to cause life-threatening opportunistic infections, were highlighted by the US Centers for Disease Control and Prevention as one of three most urgent threats among antibiotic-resistant bacteria. Carbapenems are frequently co-harbored with other resistance determinants on mobile plasmids, we implemented conditional-lethality devices with high specificity, modularity and multiplexability against undesired DNA sequences (Fig. 1a and Supplementary Fig. 1).

Here, we introduce an alternative antimicrobial approach that imposes direct evolutionary pressure at the gene level by using efficiently delivered, programmable RGNs. We engineered the clustered, regularly interspaced, short palindromic repeats (CRISPR–CRISPR-associated (Cas) system, naturally employed in bacteria as a defense strategy against mobile elements, to effect cell death or plasmid loss upon detection of genetic signatures associated with virulence or antibiotic resistance. The Type II CRISPR-Cas system of Streptococcus pyogenes is an effective, programmable tool for genome editing and gene expression in a wide variety of organisms. The specificity of CRISPR-Cas is dictated by short, spacer sequences flanked by direct repeats encoded in the CRISPR locus, which are transcribed and processed into CRISPR RNAs (crRNAs). With the aid of a trans-activating small RNA (tracrRNA), crRNAs enable the Cas9 endonuclease to introduce double-stranded breaks in target DNA sequences. Through simple modifications of spacers in the CRISPR locus, an RGN can direct cleavage of almost any DNA sequence, with the only design restriction being a requisite NGG motif immediately 3′ of the target sequence. By packaging RGNs into bacteriophage particles or harnessing mobilizable plasmids, we implemented conditional-lethality devices with high specificity, modularity and multiplexability against undesired DNA sequences (Fig. 1a and Supplementary Fig. 1).

To establish RGN functionality in mediating sequence-specific cytotoxicity, we designed RGNs to induce double-stranded breaks in blaNDM-1, which encode extended-spectrum and pan-resistance to β-lactam antibiotics, respectively (Supplementary Table 1). Transformation of E. coli by plasmid-borne RGNs (pRGNs) containing a chromosomal copy of these target genes resulted in nearly a 1,000-fold decrease in transformation efficiency as compared to wild-type cells lacking the target (Supplementary Fig. 2a). These results corroborate the mutual exclusivity between a functional crRNA and a cognate target locus. Sequence analysis of 30 escape mutants (cells that receive and maintain an RGN plasmid despite the presence of a target sequence) revealed that tolerance was exclusively due to a defective construct that frequently arose from a spacer deletion within the CRISPR locus (Supplementary Fig. 3). Furthermore, deletion of the tracrRNA as well as inactivation of the RuvC-like nuclease domain of Cas9 (D10A) abrogated the loss of transformation efficiency in cells that harbored a target sequence.

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Thus, a catalytically active endonuclease, tracrRNA and crRNA are necessary and sufficient to mediate sequence-specific cytotoxicity in *E. coli* (Fig. 1a).

Antibiotic resistance genes are often present on large, multicopy plasmids capable of autonomous transfer in microbial populations, resulting in horizontal dissemination of drug resistance. RGN activity against high-copy plasmids was verified using a GFP-expressing, ColE1-derived vector containing a standard β-lactamase selectable marker (pZ-E-bla~gfp~) or blad~NDM-1~ (pZ-E-bla~NDM-1~). Vectors bearing this ColE1 origin are reported to be present at copy numbers of 50–70 per cell. Transformation of cells containing pZ-E-bla~NDM-1~~gfp~ with pRGN~ndm-1~·1 plasmid-borne RGN targeting blad~NDM-1~, led to a three-log~10~ reduction in transformants retaining resistance to the β-lactam antibiotic carbencillin, whereas transformation of cells containing target-free pZ-E-bla~gfp~ with pRGN~ndm-1~ did not lead to a reduction in resistant transformants (Supplementary Fig. 2b).

The activity of RGNs is therefore sufficient to exclude even high-copy antibiotic resistance plasmids from cells and can reestablish a resistant population to antibiotics. Similarly, transformation of cells possessing pZ-E-bla~NDM-1~~gfp~ with pRGN~ndm-1~ led to an ~1,000-fold decrease in GFP-expressing cells, as measured by flow cytometry, but no decrease was found with transformation of cells possessing pZ-E-bla~gfp~ with pRGN~ndm-1~ (Supplementary Fig. 4).

The usefulness of RGNs for antimicrobial therapy hinges on high-efficiency delivery of genetic constructs to bacterial cells. We explored two mechanisms of horizontal gene transfer that are used by bacteria to acquire foreign genetic elements: plasmid conjugation and viral transduction. Although constrained by requirements for cell-cell contact, conjugal plasmids often have wide host ranges and no recipient factors necessary for DNA uptake have been identified. Efficient transfer of RGNs was achieved using the broad-host-range plasmid R162 mobilized by *E. coli* S17-1, which contains the conjugal machinery of plasmid RP4 integrated into its chromosome. In filter mating experiments, conjugal transfer of RGNs resulted in a 40- to 60-fold reduction in target carbencillin-resistant recipient cells (Supplementary Fig. 1b). Under selection for transconjugants, transfer of RGNs into recipients yielded a 2- to 3-log~10~ reduction in target cells as compared to controls, suggesting that conjugation efficiency, as opposed to RGN activity, limited RGN efficacy in this context (Supplementary Fig. 1c). Future work will be necessary to further optimize the efficiency of conjugation-based delivery vehicles for antimicrobials based on mobilizable RGNs.

Bacteriophages are natural predators of bacteria and are highly adept at injecting DNA into host cells. To adapt phage for RGN delivery, we engineered phagemid vectors by pairing RGN constructs targeting blad~NDM-1~ or blad~SHV-18~ with an f1 origin to facilitate packaging into M13 particles. Phage-packaged RGN~ndm-1~ (ΦRGN~ndm-1~) was capable of comprehensively transducing a population of *E. coli* EMG2 (Supplementary Fig. 5). To test the ΦRGNs, we conjugated native plasmids containing blad~NDM-1~ (pNDM-1) or blad~SHV-18~ (pSHV-18) from clinical isolates into EMG2. Treatment of the EMG2 pNDM-1 or EMG2 pSHV-18 strains with the cognate ΦRGNs resulted in 2- to 3-log~10~ reductions in viable cells even in the absence of any selection (Fig. 1b). Furthermore, ΦRGNs engendered no toxicity against wild-type EMG2 or EMG2 containing noncognate plasmids (Fig. 1b).

In naturally occurring Type II CRISPR-Cas systems, the CRISPR locus may contain multiple spacers, each of which is processed into an independent crRNA molecule that, with tracrRNA, enables Cas9 to cleave cognate DNA sequences. To explore the utility of a single ΦRGN exhibiting activity against more than one genetic signature, we engineered a construct containing two spacers encoding two different crRNAs for targeting the blad~NDM-1~ and blad~SHV-18~ resistance genes (ΦRGN~ndm-1~·1~shv-18~). ΦRGN~ndm-1~·1~shv-18~ generated 2- to 3-log~10~ reductions in viable cells of EMG2 pNDM-1 or EMG2 pSHV-18, but not of wild-type EMG2 (Fig. 1b). Thus, RGNs may be multiplexed against different genetic signatures, enabling simultaneous targeting of a variety of virulence factors and resistance genes that may exist in microbial populations.

In addition to antibiotic-modifying enzymes, such as β-lactamases, alterations in host proteins constitute a major antibiotic resistance
Figure 2 Characterization of ΦRGN-mediated killing of antibiotic-resistant bacteria. (a) Time-course treatment of EMG2 WT or EMG2 pNDM-1 with SM buffer, ΦRGNndm-1 or ΦRGNshv-18 at a multiplicity of infection (MOI) ~20. Data represent the fold change in viable colonies at indicated time points relative to time 0 h. (b) Dose-response curve of EMG2 WT and EMG2 gyrA<sub>ΔD87G</sub> treated with various concentrations of ΦRGN<sub>gyrA<sub>ΔD87G</sub></sub> for 2 h. Data represent fold change in viable colonies relative to samples treated with SM buffer. Error bars (a,b), s.e.m. of three independent biological replicates (n = 3). (c) EMG2 E. coli harboring the natural pNDM-1 plasmid or the <i>blb</i><sub>blbDM-1</sub> gene in a synthetic expression vector (pZA<sub>-ndm1-gfp</sub>) were treated with either ΦRGN<sub>ndm1</sub> or ΦRGN<sub>shv-18</sub> at MOI ~20 and plated onto both nonselective LB and LB + carbenicillin (Cb) to select for <i>blb</i><sub>blbDM-1</sub>-containing cells. ΦRGN<sub>ndm-1</sub> treatment of cells harboring pNDM-1 resulted in a reduction in viability in the absence of selection, whereas ΦRGN<sub>ndm-1</sub> treatment of cells with pZA<sub>-ndm1-gfp</sub> demonstrated similar cytotoxicity only under selective pressure for maintenance of the pZA<sub>-ndm1-gfp</sub> plasmid. (d) EMG2 pSHV-18 complemented with the cognate antitoxin (pZA31<sub>-penII</sub>) for the PemK toxin or a control vector (pZA31<sub>-gfp</sub>) was treated with SM buffer, ΦRGN<sub>ndm-1</sub> or ΦRGN<sub>shv-18</sub>. Cultures were plated on LB and LB + Cb and colonies were enumerated to assess cytotoxicity or plasmid loss.

Mechanism in bacteria. Owing to the specificity of the CRISPR-Cas system, we suspected that RGNs might discriminate between susceptible and resistant strains that differ by a single-nucleotide mutation in DNA gyrase (<i>gyrA</i>), which confers resistance to quinolone antibiotics (Supplementary Table 1) 14. Indeed, ΦRGN<sub>gyrA<sub>ΔD87G</sub></sub> was specifically cytotoxic only for quinolone-resistant E. coli harboring the chromosomal <i>gyrA<sub>ΔD87G</sub></i> mutation and not for otherwise isogenic strains with the wild-type <i>gyrA</i> gene (Fig. 1c).

Killing curves revealed that ΦRGNs mediated rapid killing of target cells, with viable cell counts that decayed exponentially (<i>t</i><sub>1/2</sub> ~ 13 min) and maximal bactericidal effect was achieved by 2–4 h (Fig. 2a). Moreover, ΦRGN antimicrobial activity increased with plasmidic particle concentration (Fig. 2b and Supplementary Fig. 6). To further characterize the cellular response to RGN-mediated targeting, we assessed treatment of cells that harbored a GFP reporter under the SOS response<sup>15</sup>. We observed a 2.6- or 4.0-fold increase in cellular response to RGN-mediated targeting, we assessed treatment of cells that harbored a GFP reporter under the SOS response<sup>15</sup>. We observed a 2.6- or 4.0-fold increase in fluorescence in cells containing the reporter plasmid and a plasmid-borne (p<i>blbDM-1</i>) or chromosomal (gyrA<sub>ΔD87G</sub>) target site, respectively, when treated with the cognate versus noncognate ΦRGNs (Supplementary Fig. 7). These results confirm that RGNs can induce DNA damage in target cells and demonstrate that they can be coupled with SOS-based reporters to detect specific genes or sequences, even at the single-nucleotide level.

We were intrigued to observe that targeted cleavage of <i>blbDM-1</i> with ΦRGN<sub>ndm-1</sub> in the context of the native plasmid was lethal to host cells, whereas targeted cleavage of the same gene in a standard cloning vector was not (pNDM-1 versus pZA<sub>-ndm1-gfp</sub>), respectively, in Fig. 2c). Therefore, we hypothesized that ΦRGN-induced plasmid loss in itself does not elicit lethality, but rather results in cytotoxicity by means of other co-harborred plasmid-borne functions. Toxin-antitoxin systems are components of natural plasmids that ensure their persistence in bacterial populations by inhibiting the growth of daughter cells that do not inherit the plasmid. Toxin-antitoxin systems traditionally consist of a labile antitoxin that quenches the activity of a stable toxin. Owing to the differential stability of these two components, cessation of gene expression in bacteria<sup>14</sup>.
expression upon plasmid loss leads to depletion of the antitoxin pool faster than the toxin pool, resulting in de-repression of toxin activity and, ultimately, stasis or programmed cell death. Analysis of the sequenced pSHV-18 plasmid revealed the presence of a single toxin-antitoxin module, pemIK, which is commonly found among isolates harboring extended-spectrum β-lactamases. When complemented with the Peml antitoxin expressed constitutively in trans (pZA31-peml), ΦRGNshv-18 treatment of EMG2 pSHV-18 abrogated cytotoxicity and instead resulted in resensitization of this multidrug-resistant strain to carbenicillin (Fig. 3d). We attributed this effect to Peml inactivation of the pemK toxin, thus enabling loss of the pSHV-18 plasmid without concomitant bacterial killing. Toxin-antitoxin systems can therefore dictate the outcome of ΦRGN activity on episomal targets, as their presence leads to cytotoxicity and their absence or neutralization to plasmid loss.

To further demonstrate the versatility of RGNs for specifically combating pathogens, we designed a ΦRGN to target intimin, a chromosomally encoded virulence factor of enterohemorrhagic O157:H7 (EHEC) necessary for intestinal colonization and pathology. Encoded by the eae gene, intimin is a cell-surface adhesive that mediates intimate attachment to the host epithelium, permitting subsequent disruption of intestinal tight junctions and effacement of microvilli. Treatment of EHEC with ΦRGNeae resulted in a 20-fold reduction in viable cell counts; this cytotoxicity was increased an additional 100-fold under kanamy cin selection for ΦRGNeae transductants (Fig. 3a). The increase in cytotoxicity with selection for cells receiving the construct suggests that the efficacy of ΦRGN treatment was limited by delivery in this strain. Furthermore, ΦRGN treatment was assessed in G. mellonella larvae, an infection model that yields virulence data often predictive for higher-order mammals. This model has also been used to evaluate the efficacy of antimicrobials or phage therapy against various Gram-negative, Gram-positive and fungal pathogens. Administration of ΦRGNeae to HEC-infected G. mellonella larvae significantly improved survival over no treatment or an off-target ΦRGN control (log-rank test, \( P < 0.001 \)) (Fig. 3b). Moreover, ΦRGNeae was significantly more effective than chloramphenicol treatment, to which the HEC strain was resistant (log-rank test, \( P < 0.05 \)) (Supplementary Fig. 8 and Supplementary Table 1). Although ΦRGNeae treatment was inferior to carbenicillin, to which the bacteria were susceptible (Supplementary Fig. 8 and Supplementary Table 1), these data support RGNs as viable alternatives for cases where bacteria are highly resistant to existing antibiotics. Improvements in delivery efficiency with ΦRGNeae would be expected to improve treatment efficacy and outcome.

In addition to implementing targeted antimicrobial therapies, RGNs can be used to modulate the composition of complex bacterial populations (Fig. 4). Current therapies that use a prebiotic, probiotic or drug to modify the human microbiota have demonstrated potential for alleviating various disease states, but such therapies remain poorly characterized in terms of off-target effects and the specific mechanisms by which they act. In concert with the host range of the delivery vehicle, RGN activity can selectively remove bacteria with specific genomic content. This could reduce the prevalence of unwanted genes, including antibiotic resistance and virulence loci, or metabolic pathways from bacterial communities without affecting bystanders. As a proof-of-principle experiment for ‘bacterial knockdowns’ using RGNs, we constructed a synthetic consortium comprising three plasmid-susceptible E. coli strains with differential antibiotic resistance profiles. We used β-lactam–resistant E. coli EMG2 pNDM-1, quinolone-resistant RFS289 (gyrA<sub>D87G</sub>) and chloramphenicol-resistant CJ236. Application of ΦRGNndm-1 elicited >400-fold greater killing of EMG2 pNDM-1, compared to control, while leaving RFS289 and CJ236 cell populations intact. Treatment with ΦRGNgyr<sub>A<sub>D87G</sub></sub> resulted in >20,000-fold greater killing of RFS289, compared to control, without a concomitant reduction in EMG2 pNDM-1 or CJ236 (Fig. 4). These results demonstrate that RGNs can selectively knockdown bacteria that contain target DNA sequences, thereby allowing the remaining nontarget bacteria to dominate the population. Adapting this approach for tuning endogenous microbiota could be accomplished by delivering RGNs in vivo by means of broad-host-range phages or phage cocktails, or with conjugative plasmids. An appropriately targeted bacterial knockdown approach could be used in functional studies of complex microbiota and to complement additive therapies, such as probiotics, for microbiome-associated diseases by clearing specific niches or removing defined genes from bacterial populations.

In light of the rising tide of antibiotic resistance, interest in engineered cellular and viral therapeutics as potential biological solutions to infectious disease has resurged. By repurposing parts developed by nature, synthetic biologists have designed artificial gene circuits for antimarialar production and engineered probiotics and phage therapeutics to eradicate biofilms or potentiate antibiotic activity. Here, we demonstrate that transmissible CRISPR-Cas systems can act as a platform for programmable antimicrobials that harness site-specific cleavage to induce cytotoxicity, activate toxin-antitoxin systems, resensitize bacterial populations to antibiotics and modulate bacterial consortia. This work complements the recent finding that...
the Vibrio cholerae phage, ICP1, encodes its own CRISPR-Cas system to counteract a host-encoded phage defense locus, and that CRISPR-Cas constructs transformed into electrocompetent cell populations are incompatible with cells that contain cognate target sequences. In contrast to these other studies, we show that CRISPR-Cas technology can be applied both in situ for the removal of undesired genes from microbial populations and in vivo to treat infection in the absence of artificial selection. Moreover, we demonstrate that RNAs can be used to activate plasmid-borne toxin-antitoxin systems that have recently become an attractive antimicrobial strategy. In addition to validating antimicrobial activity, we further demonstrate potential applications of CRISPRs in the deletion of plasmids from cells or the detection of DNA elements with up to single-nucleotide resolution using a DNA-damage-responsive reporter.

Because CRISPR-Cas systems are widely conserved in bacteria and archaea, the isolation, development, and optimization of delivery vehicles will be required for the creation of CRISPRs capable of targeting additional strains, including multidrug-resistant pathogens as well as key members of endogenous microbiota. These next-generation phage systems should be designed with the goal of avoiding anticipated shortcomings, including host range limitations, phage resistance and concerns around immune reactions. Delivery systems that function in higher organisms could also enable CRISPRs to modulate the prevalence of specific genes in wild-type populations.

Owing to the modularity and simplicity of CRISPR-Cas engineering, libraries of multiplexedΦRNAs could be rapidly constructed to simultaneously target a plethora of antibiotic resistance and virulence determinants and to modulate the composition of complex microbial communities. The addition of facile, sequence-informed, rational design to a field that has been dominated by time- and cost-intensive screening for broad-spectrum, small-molecule antibiotics has the potential to reinvigorate the pipeline for new antimicrobials.

METHODS

Methods and any associated references are available in the online version of the paper.

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AUTHOR CONTRIBUTIONS

R.J.C. and M.M. designed and performed experiments. R.J.C., M.M. and T.K.L. conceived this study, analyzed the data, discussed results and wrote the manuscript.

COMPETING FINANCIAL INTERESTS

The authors declare competing financial interests: details are available in the online version of the paper.

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ONLINE METHODS

Strains and culture conditions. Unless otherwise noted, bacterial cultures were grown at 37 °C with Luria-Bertani (LB) medium (BD Difco). Bacterial strains used in this study are listed in Supplementary Table 2. Where indicated, antibiotics were added to the growth medium to the following final concentrations: 100 µg/ml carbenicillin (Cb), 30 µg/ml kanamycin (Km), 25 µg/ml chloramphenicol (Cm), 100 µg/ml streptomycin (Sm) and 150 ng/ml ofloxacin (Ox)

Strain construction. E. coli EMG2 SmR was generated by plating an overnight culture of E. coli EMG2 onto LB+Sm. Spontaneous resistant mutants were re-streaked onto LB+S+Sm and an isolated colony was picked and used as the recipient for conjugation of the multidrug resistance plasmids. Overnight cultures of EMG2 SmR (recipient), E. coli CDC1001728 (donor for pNDM-1) and K. pneumoniae K6 (donor for pSHV-18) were washed in sterile PBS and 100 µl of donor and recipient were spotted onto LB agar plates and incubated at 37 °C overnight. Transconjugants were harvested by scraping the cells in 1 ml of sterile PBS and plating onto LB+Sm+Cb. The chromosomal integrations of the blsNDM-1 and blsSHV-18 β-lactamase genes and generation of EMG2 gyrAΔPGN, which were performed by λ-red recombination using the pSI9M system31. Templates for integration at the nonessential lacZTA locus were generated by amplifying the blsNDM-1 and blsSHV-18 genes from lysates of CDC1001728 and K6 using the primers rcD77/78 and rcD73/74, respectively. Templates for construction of EMG2 gyrAΔPGN were obtained by amplifying gyrA from RFS289 using primers mmD155/161.

Plasmid construction. To generate the RN G plasmids (Supplementary Fig. 2), we created an intermediate vector pZA-RGN0, which lacks a CRISPR locus. The tracrRNA and P Lump-1 promoter were synthesized (Genewiz) and amplified using primers mmD98/99, cas9 was amplified from pMJ806 (ref. 7) using primers mmD74/75, and the vector backbone was amplified from pZA11G using primers mmD82/83. Each PCR product was purified and ligated by Gibson assembly32. To create the final backbone vector for the RN G plasmids, the pBBR1 origin, chloramphenicol resistance marker, tL17 terminator, and CRISPR locus cloning site were amplified from an intermediate vector pBBR1-MCS1-tL17 using mmD151/154, digested with Nhel and SacI-HF, and ligated with pZA-RGN0 digested with SacI-HF and AvrII to create pZB-RGN0. Digestion of this vector with PstI-HF and XbaI allowed for the insertion of assembled CRISPR loci. The tracrRNA pRGNndm-1 plasmid was created by amplification of pRGNndm-1 with mmD162/163, ClaI digestion, and self-ligation. The Cas9Δ10Δ40 mutant plasmid was constructed through site-directed mutagenesis of pRGNndm-1 with primers mmD108/109 and the KAPA HiFi PCR kit (KAPA Biosystems).

The CRISPR loci were constructed through isothermal annealing and ligation of short, single-stranded oligonucleotides (Integrated DNA Technologies). Each spacer and repeat piece was built by a corresponding oligo duplex connected to adjacent pieces by 6-bp overhangs. In addition, the terminal repeats were designed to contain a 17-bp extension comprised of a Bsal restriction site to generate an overhang that allowed insertion into the terminal repeats were designed to contain a 17-bp extension comprised of a Bsal restriction site to generate an overhang that allowed insertion into the target vector using PstI-HF and XbaI and ligated into pZB-RGN0 digested with the same enzymes to create the final RN G plasmids. Phagemid vector pZEE-Δgfp was created previously by adding the λ origin amplified from the yeast shuttle pRS series13 into pZEE-Δgfp12. The RN G constructs consisting of the genes encoding the tracrRNA, Cas9 and a sequence-targeting crRNA were amplified as a single product from the respective pRGN vectors using KAPA HiFi polymerase (Kapa Biosystems) with primers rcdD169/183 and digested with AvrII and Xmal (New England Biolabs). These inserts were ligated with a backbone derived from amplifying the kanamycin resistance cassette, ColE1 replication origin and the λ origin required for packaging into M13 particles off of pZEE-Δgfp with primers rcdD184/185 and digesting with the same enzymes. E. coli DH5αPro was transformed with ligated plasmids for sequence verification and plasmid purification. The pZEE-bl3NDM-1-Δgfp and pZA-bl3SHV-18-Δgfp vectors were constructed by swapping the antibiotic resistance cassette of the Lutz-Bujard vectors pZE12G and pZA12G32. The bl3NDM-1 gene was amplified from a lysate of CDC1001728 using primers mmD8/9, and the PCR product was digested with SacI-HF and XhoI. The digested product was ligated into the Lutz-Bujard vectors digested with the same enzymes. The PemI antitoxin complementation plasmid pZA31-peml was created by first amplifying the pemI coding sequence from pSHV-18 with mmD253/254. The PCR product was digested with BamHI and KpnI and ligated with the large fragment of a pZA31G digest with the same enzymes. The SOS-responsive pZASHL reporter plasmid was derived from pZEL1G34 by swapping the origin of replication and antibiotic resistance marker with pZASHL using AatII and AvrII as restriction enzymes.

Mobile RN Gs were created by first amplifying the R162 replication origin and orf7 using mmD266/267. The chloramphenicol selection marker and RN G locus were amplified from pRGNndm-1 and pRGNshv-18 with mmD247/248. PCR products were digested with SpeI and Xmal, ligated and used to transform E. coli S17-1 λpir to create the donor cells used in matings.

Minimum inhibitory concentration (MIC) determination. MICs were determined by broth microdilution using LB broth according to the CLSI guidelines35.

Transformation assays. Overnight cultures were diluted 1:100 in fresh LB and grown to an optical density (OD600) of ~0.3~0.5. Following 15 min of incubation on ice, cultured cells were centrifuged at 3,200, and pellets were resuspended in one tenth volume of TSS buffer (LB, 10% polyethylene glycol, 5% dimethyl sulfoxide, 50 mM Mg2+ at pH 6.5)36. A 100 µl aliquot of cells was inoculated with 10 ng of RN G plasmid DNA. Plasmids were purified from the DH5αPro cloning host using a Qiagen QIAprep Spin Miniprep Kit and the concentration was determined using a Quant-iT PicoGreen dsDNA Assay Kit (Invitrogen). Following 30 min of incubation on ice, cells were heat shocked at 42 °C for 30 s, returned to ice for 2 min and recovered for 1.5 h at 37 °C in 1 ml of SOC broth (HiMedia). For the chromosomal target assay, serial dilutions of cells were plated on LB+Cm to select for transformants. Plates were incubated overnight at 37 °C, and the number of colony-forming units (CFU) were enumerated the following day. Transformation efficiency was used to assess whether the given RN G plasmid was toxic to cells and was calculated as the CFU/ml per µg of DNA transformed (Supplementary Fig. 2a).

For the episomal target assay (Supplementary Fig. 2b), following recovery, cultures were washed in fresh LB, diluted 1:100 in LB supplemented with chloramphenicol to select for transformants and incubated for 16 h at 37 °C. Samples were washed in sterile PBS, serially diluted and plated on LB+Cm and LB+Cm+Cb or analyzed by flow cytometry (Supplementary Fig. 4). Colonies were enumerated the following day and plasmid loss was inferred by calculating the ratio of Cb6+CmR CFUs to CmR CFUs.

Overnight cultures of RN G transformants were also diluted 1:100 in sterile PBS, aliquoted in duplicates in a 96-well plate and immediately assayed using a BD LSFRFortessa cell analyzer. Cells were consistently gated by side scatter and forward scatter across independent biological replicates. Fluorescence measurements were performed using a 488-nm argon excitation laser. The GEP gate and laser voltages were initially determined using untreated pZEE-bl3NDM-1 and EMG2 cells as positive and negative controls, respectively, and implemented across biological replicates. BD FACSDIVA software was used for data acquisition and analysis.

Sequence analysis. Escape mutants from transformation assays were re-isolated by passaging surviving colonies onto LB+Cm+Cb. DNA isolation for escape sequencing was done by either extracting plasmid DNA from isolated
escape mutants using the Qiagen QIAprep Spin Miniprep Kit or by amplifying the integrated target locus using primers mmD9/234 or mmD3/4 for blsNDM-1 and blsNDM-1 respectively. Sequencing was performed by Genewiz using the primers mmD112–115/153 and mC111 for analysis of the RGN plasmids and mmD3 or mmD234 for examination of the integrated resistance genes.

Phagemid purification. Phagemids encoding the RGNs were purified using the Qiagen QIAprep Spin Miniprep Kit (Qiagen) and used to transform E. coli DH5α Pro along with the m3cp helper plasmid for generation of phagemid-loaded M13 particles.37 Strains were inoculated and grown overnight in 250 ml LB + Km to maintain M13cp and the phagemid, respectively. Cells were pelleted and the supernatant fluid containing the phagemid particles was passed through a 0.2-µm filter. For all purifications except the PhRGNgRfAMAG purification for the dose-response curve, M13 phagemid particles were precipitated by the addition of 5% polyethylene glycol (PEG-6000) and 0.5M NaCl and incubation overnight at 4 °C (ref. 38) and pelleted at 12,000g for 1 h. Purified phagemid pellets were resuspended gently in 1/100th volume of SM buffer (50 mM Tris-HCl [pH 7.5]), 100 mM NaCl, 10 mM MgSO4 and stored at 4 °C. For the PhRGNgRfAMAG purification for the dose-response curve (Fig. 2b), M13 phagemid particles were precipitated39 through the addition of concentrated HCl to pH 4.2 and subsequently pelleted at 13,000g for 15 min. The phagemid pellet was resuspended in 1/100th volume of water and concentrated NaOH was added to pH 7.0 to solubilize phagemid particles. Tris-HCl [pH 7.5], NaCl and MgSO4 were added to reconstitute the composition of SM buffer.

Titers were measured by incubating sample dilutions with E. coli EMG2 for 30 min and enumerating transductants by plating on LB and LB + Km. Titers were defined in TFU µl/ml, which is the concentration of phagemid at which ~100% of a recipient population of an equivalent cell concentration would be transduced.

Phagemid kill assays. Cultures were inoculated and grown overnight in LB with appropriate antibiotics at 37 °C with shaking. The following day, overnight cultures were subcultured 1:100 into 3 ml LB (no antibiotics) and grown at 37 °C with shaking until the OD600 reached ~0.8. Cultures were diluted into LB to -108 CFU/ml for pNDM-1 and pSHV-18 assays (Fig. 1b, 2a) or -109 CFU/ml for gyrAΔAMAG (Fig. 1c) and EHEC assays (Fig. 3a) and 245 µl of the suspension was added to 5 µl of purified phagemid stock in a 96-well plate and incubated static at 37 °C. The number of viable cells in samples at each interval during the time-course or at 2 h for endpoint assays was determined by serial dilution and spotting onto LB, LB + Chp, and LB + Km to analyze cytotoxicity, plasmid loss and phagemid delivery, respectively. Initial suspensions were also diluted and plated onto LB to quantify the initial bacterial inocula. Colonies were enumerated after 8–9 h incubation at 37 °C to calculate cell viability (CFU/ml) and averaged over three independent experiments. Nonlinear curve fitting of the time-course to an exponential decay curve was performed using GraphPad Prism.

G. mellonella model. Larvae of the model organism G. mellonella were purchased from Vanderhorst Wholesale, Inc. (St. Marys, OH, USA) and received in the final larval instar for survival assays. Larvae were removed from food source, allowed to acclimate for at least 24 h at room temperature in the dark, and used within 4 d of receipt. For all injections, a KDS100 (KD Scientific) or Pump 11 Elite (Harvard Apparatus) automated syringe pump was set to dispense a 10-µl dose of receipt. For all injections, a KDS100 (KD Scientific) or Pump 11 Elite (Harvard Apparatus) automated syringe pump was set to dispense a 10-µl dose of SM buffer, antibiotic or PhRGNgRfAMAG treatment was administered behind the final right proleg (Fig. 3c, Supplementary Fig. 8). Larvae were inoculated at 37 °C and survival was monitored at 12 h intervals for 72 h, with death indicated by lack of movement and unresponsiveness to touch.34 Kaplan-Meier survival curves were generated and analyzed with the log-rank test using GraphPad Prism.

LexA reporter assay. Overnight cultures of EMG2 WT, EMG2 pNDM-1 and EMG2 gyrAΔAMAG containing the SOS-responsive reporter plasmid pZ3A3LG were diluted 1:50 in LB and incubated with either SM buffer, PhRGNgRfAMAG or PhRGNgRfAMAG at MOI ~5 for 2 h at 37 °C (Supplementary Fig. 7). Cultures were diluted 1:5 in 250 µl of sterile PBS and analyzed using a BD LSR Fortessa cell analyzer, as above. BD FACSDIVA software was used for data acquisition and analysis was performed using FlowJo software.

Bacterial conjugation. Donor and recipient strains grown overnight in LB with appropriate antibiotics were diluted 1:100 in fresh media and grown to an OD600 ~1. Cells were pelleted and resuspended in sterile PBS, and mating pairs were mixed at a donor to recipient ratio of 340 ± 66:1. Mating mixtures were pelleted, resuspended in 20 µl of PBS and spotted onto nitrocellulose filters placed on LB agar plates. Initial bacterial suspensions were serially diluted and plated on LB agar plates to quantify the initial inocula. Matings proceeded at 37 °C for 3 h with a single mixing step. At 90 min, mating mixtures were collected by vigorously vortexing the filters in 1 ml sterile PBS. Cells were pelleted, resuspended in 20 µl PBS and re-seeded onto filters and incubated as above for the remaining 90 min. At the end of the 3 h mating, cells were again recovered by vigorously vortexing the filters in 1 ml sterile PBS. Mating mixtures were serially diluted in PBS and plated onto LB+Chp to select for total number of Chp-resistant recipient cells and LB + Chp + Km to select for transconjugants. Colonies were enumerated following overnight incubation at 37 °C to determine viable cell counts and were averaged over nine independent biological replicates (Supplementary Fig. 1).

Synthetic consortia remodeling. E. coli CJ236, EMG2 pNDM-1 and RFS289 strains grown overnight in LB with appropriate antibiotics were diluted 1:100 into fresh LB (no antibiotics) and grown to OD600 ~0.8. Cultures were seeded into fresh LB such that the initial mixture contained ~1 x 106 CFU/ml of each strain and 245 µl of the suspension was added to 5 µl of SM buffer or purified PhRGNgRfAMAG in triplicate in a 96-well plate and spotted onto LB + Chp, + Sm and + Oxf to quantify the initial concentration of CJ236, EMG2 pNDM-1 and RFS289, respectively. Samples were then incubated, plated and enumerated as in phagemid kill assays. The composition of the synthetic ecosystem under each treatment condition was determined by counting viable colonies on plates selective for each strain as above and data were calculated as viable cell concentration (CFU/ml) averaged over three biological replicates (Fig. 4).

Data analysis and statistics. All data were analyzed using GraphPad Prism version 6.0 (GraphPad Software, San Diego, CA, USA, http://www.graphpad.com/).

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