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# The novel anti-CRISPR AcrIIA22 relieves DNA torsion in target plasmids and impairs SpyCas9 activity

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#### 11 Abstract

To overcome CRISPR-Cas defense systems, many phages and mobile genetic elements 12 encode CRISPR-Cas inhibitors called anti-CRISPRs (Acrs). Nearly all characterized Acrs 13 14 directly bind Cas proteins to inactivate CRISPR immunity. Here, using functional 15 metagenomic selection, we describe AcrIIA22, an unconventional Acr found in hypervariable genomic regions of clostridial bacteria and their prophages from human gut 16 17 microbiomes. AcrIIA22 does not bind strongly to SpyCas9 but nonetheless potently inhibits its activity against plasmids. To gain insight into its mechanism, we obtained an 18 19 X-ray crystal structure of AcrIIA22, which revealed homology to PC4-like nucleic-acid binding proteins. Based on mutational analyses and functional assays, we deduced that 20 acrIIA22 encodes a DNA nickase that relieves torsional stress in supercoiled plasmids. 21 22 This may render them less susceptible to SpyCas9, which uses free energy from negative supercoils to form stable R-loops. Modifying DNA topology may provide an additional 23 24 route to CRISPR-Cas resistance in phages and mobile genetic elements. (150)

#### 25 Introduction

26 CRISPR-Cas systems in bacteria and archaea confer sequence-specific immunity against invading phages and other mobile genetic elements (MGEs)<sup>1,2</sup>. In response, MGEs can 27 28 circumvent CRISPR-Cas systems by evading CRISPR immunity. In its simplest form, evasion requires only a single mutation within a CRISPR target site, which allows a phage or MGE to 29 30 escape immune recognition<sup>3</sup>. However, CRISPR-Cas systems routinely acquire new spacer sequences corresponding to new sites within phage and MGE genomes<sup>1</sup>. This means that any 31 single-site evasion strategy is likely to be short-lived. Thus, phages also employ forms of CRISPR-32 Cas evasion that are less easily subverted. For instance, some jumbophages assemble a 33 proteinaceous, nucleus-like compartment around their genomes upon infection, allowing them to 34 overcome diverse bacterial defenses, including CRISPR-Cas and restriction-modification (RM) 35 systems<sup>4,5</sup>. Similarly, other phages decorate their DNA genomes with diverse chemical 36 37 modifications such as the glucosylated cytosines used by phage T4 of Escherichia col<sup>6</sup>, which can prevent Cas nucleases from binding their target sequence. 38

MGEs may also overcome CRISPR-Cas systems by inactivating, rather than evading, CRISPR immunity. MGEs encode diverse CRISPR-Cas inhibitors called anti-CRISPRs (Acrs), which allow them to overcome CRISPR-Cas systems and infect otherwise immune hosts<sup>7</sup>. Most known Acrs bind Cas proteins and inhibit Cas activity by restricting access to target DNA, preventing necessary conformational changes, or inactivating critical CRISPR-Cas components<sup>8,9</sup>. The direct inactivation of Cas proteins by Acrs has proven an effective and widespread strategy for overcoming CRISPR immunity<sup>10</sup>.

46 Recent genetic, bioinformatic, and metagenomic strategies have identified many Acrs that independently target the same CRISPR-Cas system<sup>7-10</sup>. Yet, most CRISPR-Cas systems are not 47 inhibited by known Acrs<sup>10</sup>. Thus, many undiscovered strategies to inhibit or evade CRISPR-Cas 48 systems likely exist in nature. Indeed, over half of the genes in an average phage genome have 49 no known function<sup>11</sup>. To uncover new counter-immune strategies, we recently devised a high-50 51 throughput functional metagenomic selection to find genes that protect a target plasmid from 52 Streptococcus pyogenes Cas9 (SpyCas9), the variant used most frequently for genome editing<sup>12</sup>. 53 Our selection strategy was designed to reveal any gene capable of overcoming SpyCas9 activity in this system, regardless of mechanism. With this approach, we previously described a new 54

phage inhibitor of SpyCas9, called AcrIIA11, which exhibits broad-spectrum anti-Cas9 activity and
 is prevalent across human gut microbiomes<sup>12</sup>.

57 Here, we describe acrIIA22, which was the second most common Acr candidate recovered 58 from our original functional selection. AcrIIA22 encodes a 54 amino acid protein that impairs 59 SpyCas9 activity. We observe that homologs of acrIIA22 are found in hypervariable loci in phage and bacterial genomes. Unlike most other Acrs, AcrIIA22 does not bind strongly to SpyCas9 in 60 61 vitro. Instead, guided by an X-ray crystal structure of AcrIIA22, coupled with mutational and biochemical analyses, we show that AcrIIA22 encodes a DNA nickase. By nicking a supercoiled 62 plasmid substrate and relieving its torsional stress, AcrIIA22 renders the target less susceptible 63 to SpyCas9 activity. AcrIIA22 thus represents a novel mechanism of SpyCas9 evasion, which 64 capitalizes on SpyCas9's preference for negative supercoils to efficiently form R-loops and cleave 65 DNA<sup>13-16</sup>. Such a resistance mechanism could be accessible to diverse MGEs, providing a route 66 to CRISPR-Cas tolerance in many genetic contexts. 67

### 68 Results

### 69 Functional selection reveals a novel anti-CRISPR protein, AcrIIA22

70 We recently carried out a functional selection for SpyCas9 antagonism, recovering clones 71 from metagenomic libraries that could potently inhibit SpyCas9<sup>12</sup>. In this two-plasmid setup, we 72 used an arabinose-inducible SpvCas9 on an expression plasmid to cleave the kanamvcin 73 resistance (Kan<sup>R</sup>) gene of a second 'target' plasmid. We then grew cultures in SpyCas9-inducing conditions and measured the proportion of colony forming units (cfus) that remained kanamycin 74 75 resistant (Figure 1A). This proportion is a measure of how many clones retained their target 76 plasmid and, thus, how effectively that plasmid withstood SpyCas9 attack. In our previously published work, we describe AcrIIA11, a novel anti-CRISPR from a metagenomic clone named 77 F01A 2 (Genbank ID MK637582.1), which was the most abundant clone after functional selection 78 of a human fecal microbiome<sup>12</sup>. This functional selection also revealed another protective clone, 79 F01A 4 (Genbank ID MK637587.1), which was the second most abundant contig following 80 selection. Together, these two contigs (F01A 2 and F01A 4) accounted for >96% of the 81 82 normalized read coverage.

The F01A 4 contig is 685 bp long, encodes three potential open reading frames (ORFs), 83 and confers complete protection against SpyCas9, with plasmid retention equaling that of an 84 uninduced SpyCas9 control (Figure 1B). To determine the genetic basis for SpyCas9 antagonism 85 in this contig, we introduced an early stop codon into each of the three potential ORFs and 86 87 analyzed how these mutations affected the contig's ability to protect a target plasmid from SpyCas9. We found that an early stop codon in orf 1 reduced the proportion of  $Kan^{R}$  cfus by a 88 89 factor of 10<sup>5</sup>, matching the value observed for an empty vector control (Figure 1B). Furthermore, expression of orf 1 alone was sufficient for SpyCas9 antagonism (Figure 1C), protecting a target 90 plasmid from SpyCas9 cleavage as effectively as the potent SpyCas9 inhibitor, AcrIIA4. In this 91 assay, orf 1 was slightly toxic when singly expressed in E. coli, reducing growth rate by 7% 92 (Supplemental Figure 1). Combined, our results indicate that orf\_1 completely accounts for the 93 SpyCas9 protection phenotype of contig F01A 4. 94

One mechanism through which *orf\_1* could apparently antagonize SpyCas9 in our functional assay would be by lowering its expression. To address this possibility, we carried out two experiments. First, we swapped the *spycas9* gene for *gfp* in our expression vector and asked whether *orf\_1* induction impacted fluorescence output. We saw no change in fluorescence upon

99 orf 1 induction, indicating that orf 1 neither suppressed transcription from our expression vector nor altered its copy number (Supplemental Figure 2). Second, we used Western blots to test 100 whether orf 1 expression impacted SpyCas9 protein levels through the course of a plasmid 101 102 protection assay. We used a crRNA that did not target our plasmid backbone to ensure that orf\_1 expression remained high to maximize its potential impact on SpyCas9 expression levels. We 103 observed that orf 1 expression had no meaningful effect on SpyCas9 expression at any timepoint 104 (Supplemental Figure 2). Thus, we conclude that orf 1 does not impact SpyCas9's translation or 105 degradation rate. Therefore, orf 1 must act via an alternative mechanism to inhibit SpyCas9 106 107 activity. Based on these findings, we conclude that orf 1 encodes a bona fide anti-CRISPR protein and hereafter refer to it as acrIIA22. 108

Next, we investigated whether acrIIA22 could also allow phages to escape from SpyCas9 109 immunity (Supplemental Figure 3). We measured SpyCas9's ability to protect E. coli from infection 110 by phage Mu, in the presence or absence of acrIIA22. As a control, we carried out similar phage 111 infections in the presence or absence of the well-established SpyCas9 inhibitor, acrIIA4. As 112 anticipated, SpyCas9 significantly impaired Mu when targeted to the phage's genome but not if a 113 non-targeting CRISPR RNA (crRNA) was used. Consistent with previous findings<sup>12</sup>, phage Mu 114 could infect targeting strains equally well as non-targeting strains when acrIIA4 was expressed, 115 116 indicating that SpyCas9 immunity was completely abolished by this acr. In comparison, acrIIA22 improved the infectivity of phage Mu by a factor of 100 to 1,000 across multiple experimental 117 conditions (Supplemental Figure 3). We therefore conclude that acrIIA22 only partially protects 118 119 phage Mu from SpyCas9 whereas it completely protects plasmids against SpyCas9 cleavage.

### 120 AcrIIA22 homologs are found in hypervariable regions of bacterial and prophage genomes

AcrIIA22 is 54 amino acids in length and has no sequence homology to any protein of known 121 function, including all previously described Acrs. We examined the distribution of acrIIA22 122 homologs in NCBI's NR and WGS databases but found just seven hits, limiting our ability to make 123 evolutionary inferences about its origins or prevalence. We therefore expanded our search to 124 125 include IMG/VR, a curated database of cultured and uncultured DNA viruses<sup>17</sup>, and assembly data from a meta-analysis of 9.428 diverse human microbiome samples<sup>18</sup>. With an additional 23 126 unique homologs from these databases, we found that the majority of acrIIA22 homologs exist in 127 128 either of two genomic contexts: prophage genomes (Figure 2A, Supplemental Figure 4A) or small, hypervariable regions of bacterial genomes, which we refer to hereafter as 'genomic islands' 129 (Figure 2B, Supplemental Figure 4B). The original metagenomic DNA fragment from our 130 selection, F01A\_4, shared perfect nucleotide identity with one of these genomic islands (Figure 131 132 2B).

Because most acrs are found in phage genomes, we first examined the prophages that 133 encoded AcrIIA22 homologs. These prophages were clearly related, based on many homologous 134 genes and a similar genome organization (Supplemental Figure 4A). We found that these 135 prophages had inserted into several different bacterial loci, including one site between the 136 bacterial genes purF and radC (locus #3, Supplemental Figure 4A). This insertion site is nearly 137 identical to the highly conserved sequences that flanked acrIIA22-encoding bacterial genomic 138 islands (Supplemental Figure 4B). Based on this common insertion site, we hypothesize that the 139 140 apparently bacterial genomic islands with acrIIA22 homologs originated from a common prophage insertion at this locus. We speculate that the original acrIIA22-encoding bacterial genomic island 141 142 resulted from the incomplete excision of an ancestral, acrIIA22-encoding prophage. Supporting

this hypothesis, *acrIIA22* homologs are always found at the end of prophage genomes, near their
 junction with a host bacterial genome (Figure 2A, Supplemental Figure 4A).

145 To better understand acrIIA22's gene neighborhood, we again searched the assemblies of 146 over 9,400 human microbiomes for more examples of these genomic islands<sup>18</sup>. We did not include acrIIA22 as a query. Instead, we only considered contigs with ≥98% nucleotide identity to purF 147 and radC, the conserved genes that flanked the genomic islands. This search yielded 258 contigs. 148 149 Aligning these sequences revealed that each contig encoded a short, hypervariable region of small ORFs which was flanked by conserved genomic sequences (Figure 2B, Supplemental 150 Figure 4B). In total, we observed 128 unique examples of these hypervariable loci, which 151 displayed considerable gene turnover, resulting in 54 distinct gene arrangements among the 128 152 153 unique loci. Despite not being included in our search criteria, acrIIA22 homologs were universally 154 conserved in all 128 unique genomic islands. In contrast, no other gene was present in more than two-thirds of the 54 distinct gene arrangements (Figure 2B, Supplemental Figure 4C). Based on 155 156 this finding, we infer that the arrival of acrIIA22 preceded the diversification seen at this locus and 157 has been retained despite the considerable gene turnover that has occurred subsequently.

Though most ORFs in these islands were of unknown function, many had close homologs in 158 159 the genomes of nine representative acrIIA22-encoding phage (Supplemental Figures 4A, 4B, 4C). 160 This suggests that phages continue to supply the genetic diversity seen at these hypervariable genomic loci. These rapid gene gains and losses likely occur as they do in other genomic islands, 161 via recombination between this locus and related MGEs that infect the same host bacterium. 162 163 without the MGE necessarily integrating into the locus<sup>19</sup>. Taken together, our data suggest that an incomplete prophage excision event left acrIIA22 behind in a bacterial genomic locus, which 164 165 then diversified via gene exchange with additional phage genomes (Figure 2C, Supplemental 166 Figure 4D).

Like in the genomic islands (Figure 2B), we found acrI/A22 homologs in hypervariable regions 167 of prophage genomes, where they were consistently near the junction with a host bacterial 168 169 genome (Supplemental Figure 4A). Thus, nearly all acrIIA22-encoding loci show signatures of 170 frequent recombination. Despite this, we could find no gene consistently present within or outside of acrIIA22-encoding genomic islands that could account for their hypervariable nature (e.g. an 171 172 integrase, transposase, recombinase, or similar function that is typically associated with locusspecific recombination<sup>20</sup>). Instead, acrIIA22 was the only gene conserved at this locus. This 173 conservation led us to speculate that acrIIA22 might promote recombination, either alone or with 174 other factors. If this were true, it could explain the high rates of gene exchange observed adjacent 175 176 to the *acrIIA22* gene in phage and bacterial genomes (Figure 2, Supplemental Figure 4).

In total, we identified 30 unique acrIIA22 homologs, 25 of which were predicted to originate 177 from the unnamed clostridial genus, CAG-217 (Figure 3A). Because acrs are only beneficial to 178 phages if they inhibit CRISPR-Cas activity, they are typically found only in taxa with a high 179 prevalence of susceptible Cas proteins9. If AcrIIA22 functions naturally as an Acr, we would 180 predict that Cas9-encoding, type II-A CRISPR-Cas systems like SpyCas9 would be common in 181 CAG-217 bacteria. To test this idea, we examined 779 draft assemblies of CAG-217 genomes 182 and found that 179 of the 181 predicted CRISPR-Cas systems were type II-A systems (the 183 remaining two loci were Cas12-encoding, type V-A systems). This enrichment for Cas9 is 184 particularly striking as *Clostridia* typically encode other CRISPR-Cas defenses and only rarely 185 186 encode Cas9-based systems<sup>21</sup>. Moreover, prophages from CAG-217 encode 78 type II-A Acrs (homologs of AcrIIA7, AcrIIA17, and AcrIIA21), suggesting they are actively engaged in an arms 187

race with Cas9-based defenses in these bacteria. In one case, we found *acrIIA17* and *acrIIA22* homologs within one kilobase of each other in a prophage genome (Supplemental Figure 5)<sup>22</sup>.
 Phages often collect *acr* genes in the same genomic locus<sup>23</sup>, commonly pairing narrow-spectrum
 *acr*s that act during lytic infection alongside broad-spectrum *acr*s that operate during lysogeny<sup>24</sup>.
 Together, these observations support our hypothesis that prophages encode *acrIIA22* homologs

to inhibit type II-A CRISPR-Cas (Cas9) systems in CAG-217 genomes.

194 We next tested whether the ability to inhibit type II-A CRISPR-Cas systems was shared among acrIIA22 homologs from CAG-217 bacteria. To do so, we selected acrIIA22 homologs that 195 spanned the phylogenetic diversity present among CAG-217 genomes (Figure 3A) and tested 196 their ability to protect a target plasmid from SpyCas9 elimination. These analyses revealed that 197 198 diverse acrIIA22 homologs from CAG-217 bacteria (for example, sharing only 56.9% identity) 199 could antagonize SpyCas9 activity at least partially (Figure 3B), reminiscent of the broad inhibition that has been previously observed for some other type II-A Acrs<sup>12</sup>. To determine if this anti-200 201 CRISPR activity extended beyond SpyCas9, we used a slightly modified plasmid protection assay (see methods) to test whether acrIIA22 could inhibit other type II and type V CRISPR-Cas 202 systems, as these were the only two CRISPR-Cas types present in CAG-217 genomes. Though 203 204 acrIIA22 could not inhibit any of the type V (Cas12-encoding) systems we tested, it did protect a 205 target plasmid from two substantially diverged type II CRISPR-Cas systems, consistent with the high prevalence of Cas9-based systems among CAG-217 bacteria (Figure 3C). Such broad-206 spectrum inhibition can occur either by targeting a conserved feature of Cas9 or by inhibiting Cas9 207 208 via an indirect mechanism that it cannot easily evade.

### 209 AcrIIA22 functions via a non-canonical mechanism

210 Almost all characterized Acrs inhibit their cognate Cas proteins via direct binding without the involvement of additional co-factors. As a result, they exhibit strong inhibitory activity when tested 211 in vitro (Supplemental Table 1). To determine if this was the case for AcrIIA22, we purified it from 212 213 E. coli and asked whether it could bind and inhibit SpyCas9. To test for binding, we asked whether 214 twin-strep-tagged AcrIIA22 co-precipitated with untagged SpyCas9 when mixed as purified proteins. Unlike with AcrIIA4, which binds strongly to SpyCas9 and inhibits its activity in vitro, we 215 detected little to no binding between AcrIIA22 and SpyCas9, regardless of whether a single-guide 216 217 RNA (sqRNA) was included or not (Supplemental Figure 6). We also observed that AcrIIA22 had no impact on SpyCas9's ability to cleave linear, double-stranded DNA (dsDNA), even when 218 219 AcrIIA22 was included at substantial molar excess over SpyCas9 (Supplemental Figure 7). These results suggest that AcrIIA22 cannot bind and inhibit SpyCas9, at least in isolation. Thus, AcrIIA22 220 221 lacks the predominant biochemical activities exhibited by previous Acrs that have been 222 mechanistically characterized.

We therefore considered the possibility that AcrIIA22 encodes an unconventional anti-223 CRISPR that acts via a non-canonical mechanism. However, the only AcrIIA22 homologs we 224 could detect using BLAST were proteins of unknown function, which provided few clues about 225 226 AcrIIA22 activity or biochemical mechanisms. Anticipating that structural homology might provide 227 better insight into its mechanism of inhibition, we solved AcrIIA22's structure using X-ray crystallography. We first built a *de novo* model from AcrIIA22's primary sequence with Robetta<sup>25</sup>. 228 We then used this model as a molecular replacement probe to solve its structure at 2.80Å 229 230 resolution (PDB:7JTA). The asymmetric unit in AcrIIA22's crystal comprises two monomers 231 stacked end-to-end, with each monomer folding into a four-stranded  $\beta$ -sheet (Figure 4A, Table 232 1). A DALI structure-structure search revealed that the AcrIIA22 monomer is similar to members

of the newly recognized PC4-like structural fold (Figure 4B, Supplemental Table 2). PC4-like proteins have independently evolved in all domains of life, typically adopt a  $\beta$ - $\beta$ - $\beta$ - $\beta$ - $\alpha$  topology, and often homodimerize to bind diverse RNA and DNA species using variably positioned  $\beta$ sheets<sup>26</sup>.

Despite crystallizing as a homodimer, AcrIIA22 migrated from a size exclusion 237 chromatography (SEC) column at an elution volume corresponding to a calculated mass 238 239 approximately four times larger than its expected monomeric molecular weight (Figure 4B). This 240 suggested that AcrIIA22 may oligomerize in vivo. Indeed, AcrIIA22 was predicted to form a stable 241 tetramer when analyzed with PISA, a tool for inferring macromolecular assembles from crystal structures<sup>27</sup> (Figure 4C, Supplemental Figures 8A, 8B). This putative tetramer has a molecular 242 mass consistent with that observed by SEC and comprises pairs of outward-facing, concave β-243 244 sheets. A series of hydrophobic interactions likely stabilize this configuration of β-sheets instead of the typical  $\alpha$ -helical interactions seen in other PC4-like proteins, potentially explaining the 245 246 absence of an α-helix in AcrIIA22 (Supplemental Figures 8C, 8D). Interestingly, many PC4-like 247 proteins bind nucleic acids using similar concave β-sheets, and in some instances form higherorder oligomers as an obligate step for binding and/or unwinding DNA or RNA<sup>26</sup>. Consistent with 248 this possibility, AcrIIA22's  $\beta$ -sheets orient along each outward face of the putative tetramer, 249 250 resemble those in PC4-like proteins, and form a groove that could potentially accommodate a nucleic acid substrate (Figure 4D, Supplemental Figures 8A, 8B, 8E). Thus, AcrIIA22's structural 251 and functional attributes led us to suspect that it could also interact with nucleic acids and 252 253 potentially affect their topology.

Our tetramer model predicts that an interface at the C-terminus of AcrIIA22 is required for 254 255 adjacent  $\beta$ -sheets to bind one another and form a grooved, oligomeric structure (Figures 4C, 4F). 256 We reasoned that a two-residue. C-terminal truncation of AcrIIA22 would disrupt this interface (Figure 4F, Supplemental Figure 8G). To test this hypothesis, we examined the oligomeric state 257 of this 2-aa AcrIIA22 deletion mutant by SEC. Consistent with our hypothesis, we found that the 258 259 mutant AcrIIA22 complexes migrated at half the size of the wild-type complexes, corresponding to approximately twice AcrIIA22's molecular weight (Figure 4B). This suggests that the C-terminal 260 interface is required to progress from a two to four-membered oligomer, consistent with our model. 261 Moreover, we found that the 2-aa deletion mutant was also impaired for SpyCas9 antagonism in 262 our plasmid protection assay (Supplemental Figure 9A). Thus, this C-terminal motif is necessary 263 264 for protection from SpyCas9 and for higher-order oligomerization, suggesting that oligomerization may be necessary for AcrIIA22's anti-SpyCas9 activity. 265

### AcrilA22 is a DNA nickase that relieves torsion of supercoiled plasmids

Our structural analyses indicated that AcrIIA22 is a PC4-like nucleic acid-interacting protein. 267 Like AcrIIA22, many of the known PC4-like proteins are encoded in phage genomes. Among 268 these is AcrIIA22's closest structural relative in the PC4-like family: a predicted single-stranded 269 270 binding (SSB) protein from phage T5 (Figure 4E)<sup>28</sup>. This putative SSB protein has been predicted to directly stimulate recombination during the recombination-dependent replication of phage T5's 271 272 genome<sup>29</sup>. This prediction, together with our inference from genomic analyses (Figure 2, Supplemental Figure 4), led us to hypothesize that AcrIIA22 may have similar recombination-273 274 stimulating activity. Indeed, other PC4-like proteins have been observed experimentally to unwind duplex DNA, a function consistent with their proposed roles in transcription and recombination<sup>26,30</sup>. 275 276 Therefore, we investigated whether AcrIIA22 might also similarly interact with duplexed DNA to 277 affect its topology.

278 We investigated whether we could detect any biochemical effect of acrIIA22 on a double-279 stranded DNA (dsDNA) plasmid *in vivo*. In this experiment, we compared two *acrIIA22* genotypes: 280 the wild-type sequence and a null mutant with a single base pair change to create an early stop 281 codon. We grew overnight cultures of plasmids expressing each genotype, purified plasmid DNA, 282 and analyzed its topology using gel electrophoresis (Figure 5A). As is typical for plasmid purifications from *E. coli*, the plasmid encoding the null mutant was predominantly recovered in a 283 supercoiled form. In contrast, AcrIIA22 expression shifted much of the target plasmid to a slowly 284 migrating form, consistent with an open-circle conformation. These findings suggest that AcrIIA22 285 286 expression could relieve plasmid supercoiling, potentially via DNA nicking activity.

Though acrIIA22 expression appeared to alter plasmid topology in vivo. DNA topology is a 287 dynamic process, regulated by many competing factors and dependent on cellular physiology<sup>31</sup>. 288 289 Thus, we could not attribute the observed change in plasmid conformation solely to AcrIIA22. To more directly investigate AcrIIA22's effect on plasmid topology, we purified an N-terminal, His6-290 291 tagged AcrIIA22 protein and examined its impact on a plasmid DNA substrate in vitro. By gel 292 electrophoresis, we observed that AcrIIA22 shifted a supercoiled plasmid to a slowly migrating form in a time and concentration-dependent manner (Figure 5B, Supplemental Figure 10D). For 293 294 comparison, we also treated a plasmid with the nickase Nb.BssSI, yielding a band that migrated 295 at the same position as the putatively open-circle product generated via AcrIIA22 activity (Figure 5B). High concentrations of AcrIIA22 resulted in conversion of plasmids to a linearized DNA 296 product, consistent with a nickase-like nuclease activity acting on both strands of DNA (Figure 297 5B). This nicking activity was strongly stimulated in the presence of Mn<sup>2+</sup>, Co<sup>2+</sup>, and Mg<sup>2+</sup>, weakly 298 299 with Ni<sup>2+</sup> and Zn<sup>2+</sup>, and not at all with Ca<sup>2+</sup> (Supplemental Figure 11). To confirm that the observed gel-shift was the result of nicking activity and not protein-bound DNA, we purified an AcrIIA22-300 treated plasmid with phenol-chloroform and re-examined it by gel electrophoresis. We observed 301 that the open-circle form of the plasmid persisted through purification, establishing it as the 302 303 product of a bona-fide nickase (Supplemental Figure 11). Consistent with our in vivo observations (Supplemental Figure 9A), we found that the 2-aa deletion mutant was impaired for nicking activity 304 relative to wildtype AcrIIA22 (Supplemental Figure 9B). These data suggested that acrIIA22 may 305 306 encode for a protein that nicks DNA.

307 No known nuclease has been previously characterized among the PC4-family proteins<sup>26</sup>. Therefore, to further test our hypothesis that AcrIIA22 nicks supercoiled plasmids, we performed 308 309 several additional experiments. First, we re-purified an N-terminal, His6-tagged AcrIIA22 protein, but this time examined individual fractions for nicking activity. Consistent with AcrIIA22's 310 hypothesized function, nicking activity correlated with AcrIIA22 concentration across these 311 312 fractions (Supplemental Figures 10B, 10C); no co-purifying contaminant was detected via Coomassie stain (Supplemental Figure 10A). This nicking activity, however, was low enough that 313 we could not eliminate the possibility that another protein, undetectable via Coomassie stain, 314 might have co-purified with AcrIIA22 and could explained this behavior. Reasoning that different 315 contaminating proteins would result from different purification strategies, we generated a new 316 version of the AcrIIA22 protein and purified it via a C-terminal, twin-strep-tag. A more sensitive, 317 silver-stained gel indicated that this AcrIIA22 preparation was also very pure (Supplemental 318 Figure 10E). We subsequently confirmed that it nicked supercoiled plasmids with a specific 319 activity of 5.1x10<sup>-7</sup> nmol/min/mg (Figures 5C, Supplemental Figures 10G, 10H). This activity is 320 comparable to other nickases involved in phage-bacterial conflicts (including SspB, which nicks 321 at a rate of 8.9x10<sup>-7</sup> nmol/min/mg)<sup>32</sup>. Notably, this specific activity is significantly higher than we 322 323 observed for our original, N-terminal His6-tagged variant (compare AcrIIA22 concentrations in

Figures 5B and 5C). This difference in nicking activity is also reflected in plasmid protection phenotypes observed *in vivo*; only C-terminally tagged AcrIIA22, but not N-terminally tagged AcrIIA22, protected a plasmid from SpyCas9 attack (Supplemental Figure 10F). Thus, our studies find a strong correlation between AcrIIA22's nicking and plasmid protection activities.

328 If AcrIIA22 encoded a true nickase, we hypothesized that we might be able to abrogate this 329 activity via point mutations in putative catalytic residues. Therefore, we searched for individual 330 point mutations that impaired nicking activity in vitro. If such mutants existed, they would allow us to test our hypothesis that AcrIIA22 is a nickase. Reasoning that acidic amino acids were likely to 331 be important catalytic residues<sup>33</sup>, we individually changed each aspartic acid and glutamic acid in 332 AcrIIA22 to an alanine. Hypothesizing that AcrIIA22's in vitro biochemical activity would correlate 333 with its anti-Cas9 function in vivo, we tested whether these alanine variants still inhibited SpyCas9 334 335 in our plasmid protection assay. Of the 11 mutants tested, D14A stood out. This mutant showed clear SpyCas9-dependent plasmid loss, with a >250-fold reduction in plasmid retention compared 336 337 to a wild-type AcrIIA22 control (Figure 6A).

338 Purification of the D14A mutant (via a C-terminal twin-strep tag) revealed that it displayed similar expression level, purification yield, oligometric size distribution and solution behavior as 339 wild-type AcrIIA22 (Figure 6B, Supplemental Figure 10E), indicating that the mutant protein is still 340 341 properly folded. The D14A mutant was substantially impaired for nicking activity compared to the 342 wild-type AcrIIA22 protein (Figures 6C, Supplemental Figure 10G), consistent with its diminished anti-Cas9 activity in vivo (Figure 6A). Unlike previous observations with the 2-aa deletion mutant 343 344 (Figure 4B), the reduction in nicking for the D14A mutant is unlikely to be the result of oligomeric differences between it and wild-type AcrIIA22 (Figure 6B). Instead, we speculate that D14 may 345 346 contribute to AcrIIA22's nicking activity, as two D14 residues from different AcrIIA22 monomers 347 sit very near to one another in our proposed tetramer, such that they may be stabilized via the 348 presence of a divalent cation under physiological conditions (Supplemental Figure 8F).

349 Our surveys of divergent AcrIIA22 homologs also revealed a naturally occurring AcrIIA22 350 homolog with diminished function in vivo (AcrIIA22a, Fig. 3B). Despite encoding for a protein that differs by only two amino acids from the original sequence (V3I and R30K), acrIIA22a was >450-351 fold less effective at protecting a plasmid from SpyCas9 than acrIIA22 (Fig. 3B). We examined 352 353 whether this loss of SpyCas9 protection correlated with loss of nicking activity, just like the D14A mutant. Upon purification, a twin-strep-tagged AcrII22a protein eluted with an SEC profile identical 354 355 to that of AcrIIA22, suggesting a similar oligomeric state (Figure 6B). Yet, AcrIIA22a exhibited substantially less nicking activity that wild-type AcrIIA22 in vitro (Figure 6C). In our proposed 356 357 AcrIIA22 tetramer, R30 likely forms a hydrogen bond with the C-terminus of a diagonal monomer, 358 raising the possibility that the R30K variant alters the protein's conformational plasticity or mediates other allosteric effects (Supplemental Figure 8G). As with D14A, the partial loss of 359 nicking activity seen for AcrIIA22a (Figure 6C) correlated with a partial loss of plasmid protection 360 against SpyCas9 (Figure 3B). Thus, we describe two closely related AcrIIA22 variants, one 361 engineered and one naturally occurring, whose nicking activity in vitro corresponds directly to 362 plasmid protection in vivo. From these data, along with our other in vitro and in vivo findings, we 363 364 conclude that acrIIA22 encodes a nickase protein that relieves the torsional stress of supercoiled plasmids. 365

### 366 AcrIIA22's nicking activity indirectly impairs SpyCas9

367 Having established that AcrIIA22 is a DNA nickase, we next investigated how this biochemical activity may enable its inhibition of SpyCas9 without directly binding the Cas protein. We therefore 368 tested the consequences of expressing AcrIIA22 on a target plasmid in the presence of SpyCas9. 369 370 As before, we began by comparing overnight plasmid purifications of a target plasmid expressing 371 AcrIIA22 and a null mutant with an early stop codon as a negative control. For both genotypes, we subjected the acrIIA22-encoding plasmid to SpyCas9 targeting during bacterial growth. We 372 were unable to recover the negative control target plasmid after overnight growth, implying that 373 374 this target plasmid was eliminated by SpyCas9 (Figure 7A). In contrast, SpyCas9 did not eliminate 375 a target plasmid that expressed full-length AcrIIA22 (Figure 7A), consistent with AcrIIA22's 376 capacity to protect against SpyCas9 (Figure 1C).

To be effective, a CRISPR-Cas system must eliminate its target at a faster rate than the target 377 378 can replicate<sup>34</sup>. Our findings raised the possibility that AcrIIA22 modifies a target plasmid into a SpyCas9-resistant conformation to win this 'kinetic race' against SpyCas9, potentially shifting the 379 380 equilibrium to favor plasmid persistence instead of elimination. To test this kinetic race model, we asked whether a plasmid that had been pre-treated with AcrIIA22 could resist digestion by 381 SpyCas9 in vitro. Therefore, we purified the open-circle plasmid that resulted from AcrIIA22 pre-382 treatment and determined how efficiently it was cleaved by SpyCas9 compared to an unmodified, 383 384 supercoiled plasmid (Figure 7B). SpyCas9 showed a clear preference for cleaving the supercoiled substrate versus the AcrIIA22-treated open-circle plasmid (Figures 7C, 7D, 7E), consistent with 385 previous reports<sup>13-16</sup>. An open-circle plasmid pre-treated with the nickase Nb.BssSI was similarly 386 recalcitrant to SpyCas9 digestion (Figures 7C, 7D). Taken together, our findings suggest that 387 388 relieving DNA torsion provides the mechanistic explanation for AcrIIA22's ability to inhibit SpyCas9 in vivo. 389

### 390 Discussion

In this study, we identify and characterize *acrIIA22*, a previously undescribed gene that can 391 antagonize SpyCas9. We show that AcrIIA22 homologs are common in genomes and prophages 392 393 of CAG-217 bacteria, which have a high prevalence of Cas9 homologs. Using a combination of structural and biochemical studies, we show that AcrIIA22 acts by nicking supercoiled DNA to 394 relieve torsional stress on a target plasmid, and that this activity correlates with protection against 395 396 SpyCas9 in vivo and in vitro. This torsion-based model for SpyCas9 inhibition helps explain why AcrIIA22 protects plasmids more effectively than phage Mu against SpyCas9. Because plasmids 397 398 are maintained as circular, extrachromosomal elements, they are more likely to undergo torsional change when nicked than the dsDNA genome of phage Mu, which is injected as linear DNA and 399 spends significant time integrated into *E. coli*'s genome<sup>35</sup>. Additionally, linear DNA experiences 400 much lower torsional stress and therefore is less susceptible than supercoiled plasmids to 401 cleavage by SpyCas9<sup>15</sup>. This difference also likely explains why AcrIIA22 failed to protect a linear 402 403 dsDNA substrate from SpyCas9 during our earlier in vitro experiments (Supplemental Figure 7).

Previous *in vitro* experiments indicate that Cas9 requires a higher degree of negative supercoiling than type I CRISPR-Cas systems to provide the free energy needed for R-loop formation<sup>13</sup>. Similarly, *in vivo* observations have shown that DNA supercoiling promotes the recruitment of SpyCas9 to its target site in bacteria<sup>14</sup>. Based on these published findings, we speculate that Cas9 may be particularly susceptible to changes in DNA torsion among CRISPR-Cas systems. Thus, factors that modify DNA torsion, like AcrIIA22, could provide a general means to protect against Cas9 or other enzymes with a strong preference for negative supercoils. 411 Taken together, our data implicate DNA topology as a new battleground in the evolutionary arms race between CRISPR-Cas systems and MGEs. Because DNA topology is dynamically 412 regulated in phages, plasmids and other MGEs, many topology-modifying factors already exist in 413 414 these genomes. Our findings suggest that at least some of these factors could have secondary 415 effects on CRISPR-Cas activity and thus prove useful in the context of a molecular arms race<sup>31,36</sup>. For instance, though not studied in the context of bacterial defense systems, the fitness of phage 416 T4 is improved via the expression of an accessory protein that modifies DNA supercoiling and the 417 propensity of R-loops to form<sup>37</sup>. Other phages, such as the intrinsically Cas9-resistant phage T5<sup>38</sup>, 418 419 incorporate regular nicks into their genome, the function of which has eluded description for over 40 years<sup>39</sup>. Additionally, conjugative plasmids were recently shown to evade CRISPR-Cas in 420 *Vibrio cholerae* by the action of homologs of the recombination proteins Red $\beta$  and  $\lambda$ Exo<sup>40</sup>. Based 421 422 on our findings, we hypothesize that phages and MGEs targeted by Cas9 exploit factors that 423 modify DNA topology as a general tactic to evade host immunity.

424 Functional selections like ours are biased towards identifying genes that work well in heterologous contexts. For example, even though AcrIIA22 is encoded on the genome of a 425 genetically intractable bacterium, we could identify it using a functional metagenomic selection for 426 427 SpyCas9 antagonism in E. coli. Although we have characterized its activities in E. coli and in vitro, 428 we cannot be certain that AcrIIA22 functions similarly in its native context. Little is known about the life cycle of native CAG-217 phages, though many dsDNA phage genomes undergo circular, 429 topologically-constrained stages during their replicative cycles<sup>41</sup>, during which AcrIIA22 might act 430 431 to specifically overcome Cas9 immunity. Alternatively, AcrIIA22 may enable Cas9 evasion as a 432 secondary function related to some other activity. Comparative genomics (Figure 2) and structural homology to a proposed recombination-stimulating protein of phage T5 suggest a potential role 433 434 for AcrIIA22 in recombination, a process which has recently been shown to promote CRISPR-Cas evasion<sup>40</sup>. 435

Nevertheless, the heterologous behavior of AcrIIA22 in E. coli is clearly sufficient for SpyCas9 436 437 antagonism in vivo and its nicking activity can protect plasmids from SpyCas9 in vitro. Furthermore, AcrIIA22 mutants that are defective for nicking in vitro (Figure 6C, Supplemental 438 Figure 9B) are orders of magnitude less effective at protecting a plasmid from SpyCas9 in vivo 439 (Figures 3B, 6A, Supplemental Figure 9A). This indicates that modest changes in nicking activity 440 can have major consequences for plasmid survival, which is consistent with our kinetic race model 441 442 (Figure 7B) and previous observations that non-linear equilibrium dynamics determine whether 443 an MGE withstands CRISPR-Cas immunity<sup>34</sup>.

444 Our results suggest that other proteins that affect DNA torsion may also enable Cas9 445 antagonism. For example, in addition to AcrIIA22, the Nb.BssSI nickase was capable of protecting a plasmid from SpyCas9 in vitro. Yet, despite the regular occurrence of nickases in nature, 446 functional selections for anti-Cas9 activity have not previously recovered such enzymes<sup>12,42</sup>. We 447 448 speculate that AcrIIA22 was identified from a metagenomic library because it treads a balance between activity and toxicity in E. coli; its nicking activity is high enough to antagonize SpyCas9 449 in a kinetic race, but not so high that it would be toxic to the host cell (Supplemental Figure 1). 450 451 Such a balance could result from the inherent activity of the enzyme or via some form of regulation, either direct or indirect. AcrIIA22's activity is probably also regulated in its native 452 context to avoid secondary impacts on other essential processes. Potential forms of regulation 453 454 include sequence preference, oligomerization, or transient interactions with Cas9 or other host 455 factors (Figures 4B, 4C). Studies of other phage- and bacterial-encoded nickases may provide 456 further insight into whether AcrIIA22 proteins have additional properties that render them 457 especially well-suited to antagonize Cas9.

Is AcrIIA22 a true anti-CRISPR? AcrIIA22 lacks features that are typical of conventional Acrs, 458 459 such as the ability to bind Cas proteins or to inhibit CRISPR-Cas activity as a purified protein. However, other Acr proteins also lack these features. For example, the well-characterized 460 461 SpyCas9 antagonist AcrIIA1 does not inhibit purified SpyCas9, but instead stimulates Cas9 462 degradation *in vivo*<sup>24</sup>. Similarly, AcrIIA7 does not appear to bind SpyCas9 but can nevertheless inhibit it via an unknown mechanism in vitro<sup>42</sup>. Indeed, anti-CRISPR proteins are defined by a 463 common strategy and outcome rather than by a common biochemical mechanism. Our finding 464 that AcrIIA22 is encoded by prophages as a single gene that strongly protects plasmids and 465 partially protects phages from SpyCas9 (Figure 3B, Supplemental Figure 3) makes it much more 466 similar to other Acrs<sup>23</sup> and distinct from non-canonical CRISPR-Cas evasion strategies like DNA 467 glucosylation<sup>6</sup>. 468

Although it can protect phage Mu from SpyCas9, AcrIIA22 does not appear to provide the 469 470 same potency of Cas9 inhibition as some other characterized Acrs. However, potent inhibition is 471 not a pre-requisite for effective anti-CRISPR activity. In nature, multiple phages can cooperate to overcome Cas9 immunity by each contributing some anti-CRISPR protein to overcome a common 472 473 foe<sup>43,44</sup>. These dynamics can favor weak anti-CRISPRs over strong ones, as the latter permits a 474 higher incidence of cheater phages (those without anti-CRISPRs) to persist in mixed phage populations<sup>45</sup>. Thus, even in cases where AcrIIA22 only weakly inhibits Cas9 (Supplemental 475 476 Figure 3), it may nonetheless confer substantial benefit. Additionally, slowing down Cas9 cleavage could increase the time and probability for escape mutants to arise (e.g. Cas9 target-477 site variants<sup>1</sup>, deletion mutants<sup>40</sup>), allow for additional Acr expression<sup>43,44</sup>, or permit further 478 479 genome replication to overwhelm CRISPR-Cas immunity<sup>34</sup>. This phenomenon – weak inhibition giving rise to long-term resistance - is reproducibly observed in cases of strong selective 480 481 pressure. For instance, in the context of antibiotic resistance, the expression of QNR pentapeptide 482 proteins by many human pathogens can provide low-level drug tolerance, extend survival, and 483 allow time for additional mutations to develop that completely resist quinolone antibiotics<sup>46</sup>.

484 As the use of functional metagenomics to study phage-bacterial conflicts grows more 485 common, many novel genes and mechanisms for CRISPR-Cas inhibition are likely to be described<sup>12,42</sup>. Like AcrIIA22, which has no homology to any previously described anti-CRISPR 486 and lacks other genetic signatures used for acr discovery (e.g., linkage with helix-turn-helix 487 transcription factors)<sup>47,48</sup>, these new genes may not exhibit canonical Acr behaviors. It is inevitable 488 that these discoveries will lead to a more nuanced understanding of the arms race between 489 490 CRISPR-Cas systems and MGEs. These findings will also reveal undiscovered strategies for molecular antagonism and new battlegrounds in the age-old conflict between bacteria and their 491 492 phages.

#### 493 Methods

#### 494 Plasmid protection assay

All plasmid protection assays were done in Escherichia coli (strain: NEB Turbo). As 495 described previously<sup>12</sup>, SpyCas9 was expressed via the arabinose-inducible promoter pBAD on 496 497 a CloDF13-based plasmid marked with a spectinomycin resistance cassette. The SpyCas9 498 construct, called pSpyCas9\_crA, was designed to eliminate a target vector with a kanamycin 499 resistance cassette. This target vector also expressed a gene-of-interest (e.g., an acr) via the 500 doxycycline-inducible pLtetO-1 promoter (Supplemental Table 4). We induced expression from 501 the target vector via depression of the TetR transcription factor with doxycycline (we generically named this vector pZE21 tetR; Supplemental Table 4). IPTG was used in samples with the target 502 vector to ensure high levels of TetR expression (which was driven by the lac promoter) and thus 503 inducible control of our gene of interest. Unless noted in Supplemental Table 5, all genes, 504 505 including each alanine mutant depicted in Figure 6A, were synthesized by Synbio technologies and cloned directly into pZE21 tetR for functional testing. 506

507 Cultures of each sample were grown overnight at 37°C with shaking at 220 rpm in lysogeny broth (LB; 10 g/L casein peptone, 10 g/L NaCl, 5 g/L ultra-filtered yeast powder) containing 508 spectinomycin 50 µg/ml, kanamycin 50 µg/ml, and 0.5mM IPTG. These growth conditions kept 509 510 both SpyCas9 and the gene of interest in uninduced states. The next morning, overnight cultures were diluted 1:50 into LB broth containing spectinomycin (at 50 µg/ml), kanamycin (at 50 µg/ml), 511 512 0.5mM IPTG, and doxycycline 100 ng/ml to induce the gene of interest. Cultures were grown at 37°C on a roller drum to mid-log phase (for approximately 1.5 hours to OD600 of 0.3-0.6). Once 513 514 cells reached mid-log phase, they were diluted to OD600 value of 0.01 into two media types: (a) 515 LB containing spectinomycin 50 µg/ml, 0.5mM IPTG, and doxycycline 100 ng/ml, and (b) LB containing spectinomycin 50 µg/ml, 0.5mM IPTG, doxycycline 100 ng/ml, and 0.2% (L) arabinose. 516 517 These media induced either the gene of interest alone, or both the gene of interest and SpyCas9, 518 respectively. Each sample was grown in triplicate in a 96 well plate in a BioTek Cytation 3 plate reader. After 6 hours of growth at 37°C with shaking at 220 rpm, each sample was diluted ten-519 520 fold and plated on two types of media: (a) LB spectinomycin 50 µg/ml + 0.5mM IPTG or (b) LB spectinomycin 50 µg/ml, kanamycin 50 µg/ml, 0.5mM IPTG. Plates were incubated at 37°C 521 522 overnight. Then, colonies were counted to determine the fraction of colony forming units (cfus) that maintained kanamycin resistance (and thus the target vector). All figures depicting these data 523 show the log-transformed proportion of Kan<sup>R</sup>/total cfu, with or without SpyCas9 induction. The 524 525 growth curves in Supplemental Figure 1 match the experiment depicted in Figure 1C for the uninduced SpyCas9 samples. For the uninduced orf 1 control samples, doxycycline was omitted 526 from media throughout the experiment. Growth rates referenced in the text and in Supplemental 527 528 Figure 1 were calculated using the slope of the OD600 growth curves during log phase, following a natural log transformation. 529

To test AcrIIA22 function against a panel of Cas9 and Cas12 orthologs in Figure 3C, we 530 used a slightly modified, three-plasmid setup. As before, spyCas9, nmCas9, fnCas12 and 531 *lbCas12* were encoded in a CloDF13-based plasmid with a spectinomycin resistance cassette. 532 533 Expression of the Cas effector was controlled by promoter J23100 and a theophylline riboswitch. 534 The accompanying gRNAs were encoded in a separate set of plasmids called pDual4 under an arabinose expression system, in a p15A-based plasmid and a chloramphenicol resistance 535 cassette (Supplemental Table 4). The gRNAs in the different pDual4 constructs were 536 537 programmed to target the kanamycin-marked target plasmid in the same manner as pSpyCas9\_crA. All assays were done in *Escherichia coli* (strain: NEB Turbo) following the same
plasmid protection assay described previously. However, in this case, we induced expression of
the different Cas effectors and gRNAs, by adding 2 mM theophylline and 0.2% (L) arabinose,
respectively, to the media.

### 542 Impact of AcrIIA22 on GFP expression

We swapped spyCas9 for eqfp in our CloDF13-based plasmid and co-expressed AcrIIA22 543 544 to determine if AcrIIA22 impacted expression from this construct. If AcrIIA22 influenced CloDF13's copy number or the transcription of spyCas9, we anticipated that it would also impact GFP levels 545 546 in this construct (pCloDF13 GFP; Supplemental Table 4). To perform this experiment, we co-547 transformed pCloDF13 GFP and pZE21 tetR encoding acrIIA22 into E. coli Turbo. Single colonies were picked into 4 ml of LB containing spectinomycin at 50 µg/ml ('spec50') and 548 kanamycin at 50 µg/ml ('kan50') and 0.5mM IPTG and grown overnight at 37°C shaking at 549 220rpm. The next morning the overnight culture was diluted 1:50 into both LB spec50 Kan50 + 550 0.5mM IPTG with or without doxycycline (to induce acrIIA22) and grown at 37°C for about 1.5 551 552 hours to mid-log phase (OD600 0.2-0.6). The OD600 was measured, and all samples were diluted 553 to OD600 of 0.01 in two media types: (a) LB spec50 + kan50 + 0.5mM IPTG + 0.2% arabinose (inducing gfp only) or (b) LB spec50 + kan50 + 0.5mM IPTG + 0.2% arabinose + 100ng/ml 554 555 doxycycline (inducing *qfp* and *acrIIA22*). A volume of 200 µl of each sample was then transferred to a 96-well plate in triplicate and GFP fluorescence was measured every 15 minutes for 24 hours 556 (GFP was excited using 485 nm light and emission detected via absorbance at 528 nm). In 557 558 parallel, we included control samples that lacked the kanamycin-marked plasmid and varied whether doxycycline was added or not (at 100 ng/ml). In these control samples, we noticed that 559 560 doxycycline slightly diminished GFP expression (it is possible that sub-lethal levels of the 561 antibiotic may still depress translation). Thus, we normalized GFP fluorescence measurements in 562 our experiment with AcrIIA22 to account for this effect in all samples containing doxycycline. These normalized fluorescence measurements are shown in Supplemental Figure 2B. 563

### 564 Western blots to determine AcrIIA22's impact on SpyCas9 expression

We grew overnight cultures of E. coli Turbo that expressed pSpyCa9 crNT and 565 pZE21 tetR encoding a gene of interest (Supplemental Tables 4, 5) in LB spec50 + kan50 + 566 567 0.5mM IPTG. The next morning, we diluted these cultures 1:100 in 4ml of either (a) LB spec50 + kan50 + 0.5mM IPTG or (b) LB spec50 + kan50 + 0.5mM IPTG + 100 ng/ml doxycycline (to induce 568 the gene of interest). We included samples that expressed either acrIIA22 or gfp as a gene of 569 570 interest. In all SpyCas9 constructs, we used a crRNA that did not target our plasmid backbone (pSpyCa9\_crNT) to ensure that acrIIA22 expression remained high and its potential impact on 571 SpyCas9 expression levels would be most evident. All samples were grown for two hours at 37°C 572 573 to reach mid-log phase (OD600 0.3 to 0.5) and transferred into media that contained 0.2% arabinose to induce SpyCas9. At transfer, volumes were normalized by OD600 value to ensure 574 that an equal number of cells were used (diluted to a final OD600 of 0.05 in the arabinose-575 576 containing medium). This second medium either contained or lacked 100 ng/ml doxycycline to control expression of acrIIA22 or gfp, as with the initial media. Throughout this experiment, we 577 included a control strain that lacked pZE21 tetR and only expressed SpyCas9. Kanamycin and 578 579 doxycycline were omitted from its growth media. For this control strain, we also toggled the 580 addition of arabinose in the second growth medium to ensure that positive and negative controls 581 for SpyCas9 expression were included in our experiment. After three hours and six hours of 582 SpyCas9 induction, OD600 readings were again taken and these values used to harvest an equal

number of cells per sample (at three hours, OD600 values were between 0.76 and 0.93 and
0.75ml to 0.9ml volumes harvested; at six hours 0.4ml was uniformly harvested as all absorbance
readings were approximately 1.6).

586 All samples were centrifuged at 4100g to pellet cells, resuspended in 100 µl of denaturing lysis buffer (12.5 mM Tris-HCl, pH 6.8; 4% SDS), and passed through a 25 gauge needle several 587 times to disrupt the lysate. Samples were then boiled at 100°C for 10 minutes, spun at 13,000 588 589 rpm at 4°C for 15 minutes and the supernatants removed and frozen at -20°C. The next day, 12 µl of lysate was mixed with 4 µl of 4x sample buffer (200 mM Tris-HCl, 8% SDS, 40% glycerol, 590 591 200 mM DTT, and 0.05% bromophenol blue) and boiled at 100°C for 10 minutes. Then, 10 µl sample was loaded onto a BioRad Mini-Protean "any KD Stain Free TGX" gel (cat. #4569035) 592 593 and run at 150V for 62 minutes. To verify that equivalent amounts of each sample were run, gels 594 were visualized on a BioRad chemidoc for total protein content. Protein was then transferred to a 0.2 µM nitrocellulose membrane using the Bio-Rad Trans-Blot Turbo system (25 V, 1.3 A for 10 595 596 min). We then washed membranes in PBS/0.1% Triton-X before incubating them with a mixture 597 of the following two primary antibodies, diluted in in LI-COR Odyssey Blocking Solution (cat. #927-40000): (i) monoclonal anti-SpyCas9, Diagenode cat. #C15200229-50, diluted 1:5,000; (ii) 598 polyclonal anti-GAPDH, GeneTex cat. # GTX100118, diluted 1:5,000. The GAPDH antibody 599 600 served as a loading control and a second check to ensure equal protein levels were run. Membranes were left shaking overnight at 4°C, protected from light. Then, membranes were 601 washed four times in PBS/0.1% Triton-X (ten-minute washes) before they were incubated for 30 602 603 minutes at room temperature with a mixture of secondary antibodies conjugated to infrared dyes. 604 Both antibodies were diluted 1:15,000 in LI-COR Odyssey Blocking Solution. To detect SpyCas9, the following secondary antibody was used: IR800 donkey, anti-mouse IgG, LI-COR cat# 926-605 606 32212. To detect GAPDH, IR680 goat, anti-rabbit IgG, LI-COR cat# 926-68071 was used. Blots were imaged on a LI-COR Odyssey CLx after three additional washes. 607

### 608 Phage plaquing assay

609 We grew overnight cultures of *E. coli* Turbo expressing pSpyCa9 crMu and pZE21 tetR encoding a gene of interest (Supplemental Tables 4, 5) at 37°C in LB spec50 + kan50 + 0.5 mM 610 IPTG. Genes of interest were either acrIIA4, *qfp*, or acrIIA22. The pSpyCas9 construct targeted 611 612 phage Mu and was previously demonstrated to confer strong anti-phage immunity in this system<sup>12</sup>. A control strain expressing pZE21-tetR-gfp and SpyCas9 crNT (which encoded a 613 614 CRISPR RNA that does not target phage Mu) was grown similarly. The next morning, all cultures were diluted 50-fold into LB spec50 + kan50 + 0.5 mM IPTG + 5 mM MgCl2 and grown at 37°C 615 for three hours. Then, doxycycline was added to a final concentration of 100 ng/ml to induce the 616 gene of interest. Two hours later, SpyCas9 was induced by adding a final concentration of 0.2% 617 w/v arabinose. Two hours after that, cultures were used in soft-agar overlays on one of two media 618 types, discordant for arabinose, to either maintain SpyCas9 expression or let it fade as arabinose 619 620 was diluted in top agar and consumed by the host bacteria (per Supplemental Figure S2). Top and bottom agar media were made with LB spec50 + kan50 + 0.5 mM IPTG + 5 mM MgCl2. In 621 cases where SpyCas9 expression was maintained, arabinose was also added at a final 622 623 concentration of 0.02% to both agar types. Top agar was made using 0.5% Difco agar and bottom agar used a 1% agar concentration. For the plaquing assay, 100 µl of bacterial culture was mixed 624 with 3 ml of top agar, allowed to solidify, and ten-fold serial dilutions of phage Mu spotted on top 625 626 using 2.5 µl droplets. After the droplets dried, plates were overturned and incubated at 37°C 627 overnight before plaques were imaged the following day.

### 628 Identification of AcrIIA22 homologs and hypervariable genomic islands

629 We searched for AcrIIA22 homologs in three databases: NCBI nr, IMG/VR, and a set of assembled contigs from 9,428 diverse human microbiome samples<sup>18</sup>. Accession numbers for the 630 631 NCBI homologs are indicated on the phylogenetic tree in Figure 3A. We retrieved AcrIIA22 homologs via five rounds of an iterative PSI-BLAST search against NCBI nr performed on October 632 2<sup>nd</sup>, 2017. In each round of searching, at least 90% of the guery protein (the original AcrIIA22 hit) 633 634 was covered, 88% of the subject protein was covered, and the minimum amino acid identity of an alignment was 23% (minimum 47% positive residues; e-value ≤ 0.001). Only one unique AcrIIA22 635 homolog was identified in IMG/VR (from several different phage genomes) via a blastp search 636 against the July, 2018 IMG/VR proteins database (using default parameters). This homolog was 637 also found in other databases and its amino acid sequence is identical to that of AcrIIA22b (Figure 638 639 3A).

Most unique AcrIIA22 homologs were identified in the assembly data of over 9,400 human 640 microbiomes performed by Pasolli and colleagues<sup>18</sup>. These data are grouped into multiple 641 642 datasets: (i) the raw assembly data, and (ii) a set of unique species genome bins (SGBs), which were generated by first assigning species-level phylogenetic labels to each assembly and then 643 selecting one representative genome assembly per species. We identified AcrIIA22 homologs 644 645 using several queries against both databases. First, we performed a tblastn search against the SGB database using the AcrIIA22 sequence as a query, retrieving 141 hits from 137 contigs. A 646 manual inspection of the genome neighborhoods for these hits revealed that most homologs 647 648 originated from a short, hypervariable genomic island; some homologs were encoded by 649 prophages. No phage-finding software was used to identify prophages; they were apparent from 650 a manual inspection of the gene annotations that neighbored acrIIA22 homologs (see the section 651 entitled "Annotation and phylogenetic assignment of metagenomic assemblies" for details).

652 To find additional examples of AcrIIA22 homologs and of these genomic islands, we then 653 queried the full raw assembly dataset. To do so without biasing for *acrIIA22*-encoding sequences, 654 we used the *purF* gene that flanked *acrIIA22*-encoding genomic islands as our initial guery sequence. Specifically, we used the *purF* gene from contig number 1 in Supplemental Table 3; 655 its sequence is also in Supplemental Table 5. To consider only the recent evolutionary history of 656 657 this locus, we required all hits have ≥98% nucleotide identity and required all hits to be larger than 15 kilobases in length to ensure sufficient syntenic information. From these contigs, we further 658 659 filtered for those that had  $\geq$ 98% nucleotide identity to *radC*, the gene which flanked the other end of acrIIA22-encoding genomic islands. Again, we used the variant from contig number 1 in 660 Supplemental Table 3; its sequence is also in Supplemental Table 5. In total, this search yielded 661 258 contig sequences; nucleotide sequences and annotations for these contigs are provided in 662 Supplemental Dataset 5. We then searched for acrIIA22 homologs in these sequences using 663 tblastn, again observing them in genomic islands and prophage genomes (which were assembled 664 as part of the 258 contigs). In total, this search revealed 320 acrIIA22 homologs from 258 contigs. 665 The 258 genomic islands from these sequences were retrieved manually by extracting all 666 nucleotides between the purF and radC genes. These extracted sequences were then clustered 667 668 at 100% nucleotide identity with the sequence analysis suite Geneious Prime 2020 v1.1 to identify 669 128 unique genomic islands.

Altogether, our two searches yielded 461 AcrIIA22 sequences from these metagenomic databases that spanned 410 contig sequences. The 461 AcrIIA22 homologs broke down into two groups: 410 clustered with genomic island-like sequences whereas 51 clustered with prophage673 like homologs. In nature, the relative prevalence of AcrIIA22 in genomic islands or prophages may 674 not be accurately reflected by these numbers because we never directly searched for prophageencoded homologs. We then combined these 461 AcrIIA22 sequences with those from NCBI and 675 676 IMG/VR and clustered the group on 100% amino acid identity to reveal 30 unique proteins. To achieve this, we used the software cd-hit<sup>49</sup> with the following parameters: -d 0 -g 1 -aS 1.0 -c 1.0. 677 These 30 sequences were numbered to match one of their parent contigs (as indicated in 678 679 Supplemental Table 3) and used to create the phylogenetic tree depicted in Figure 3A. For 680 AcrIIA22 homologs found outside NCBI, the nucleotide sequences and annotations of their parent 681 contigs can be found in Supplemental Datasets 1 and 2. For NCBI sequences, accession numbers are shown in Figure 3A. The gene sequences used in functional assays (Figure 3B) 682 683 have been reprinted in Supplemental Table 5, for convenience.

### 684 Annotation and phylogenetic assignment of metagenomic assemblies

Contig sequences from IMG/VR, the Pasolli metagenomic assemblies, and some NCBI 685 entries lacked annotations, making it difficult to make inferences about acrIIA22's genomic 686 687 neighborhood. To facilitate these insights, we annotated all contigs as follows. We used the genefinder MetaGeneMark<sup>50</sup> to predict open reading frames (ORFs) using default parameters. We 688 then used their amino acid sequences in a profile HMM search with HMMER3<sup>51</sup> against 689 690 TIGRFAM<sup>52</sup> and Pfam<sup>53</sup> profile HMM databases. The highest scoring profile was used to annotate each ORF. We annotated these contigs to facilitate genomic neighborhood analyses for acrIIA22; 691 these are not intended to provide highly accurate functional predictions of their genes. Thus, we 692 693 erred on the side of promiscuously assigning gene function; our annotations should therefore be treated with appropriate caution. A visual inspection of these annotated contigs made apparent 694 695 several examples of acrIIA22-encoding prophages (we noticed 35-40 kilobase insertions in some 696 contigs that were otherwise nearly identical to those without prophages). We were confident that these insertions were prophages because they contained mostly co-linear genes with key phage 697 functions annotated. As a simple means to sample this phage diversity, we manually extracted 698 699 nine examples of these prophage sequences (their raw sequences and annotated genomes can 700 be found in Supplemental Datasets 3 and 4). Annotations were imported into the sequence analysis suite Geneious Prime 2020 v1.1 for manual inspection of genome neighborhoods. 701

We used the genome taxonomy database (GTDB) convention for all sequences discussed 702 in this manuscript<sup>54</sup>. In part, this was because all *acrIIA22* genomes are found in clostridial 703 genomes, which are notoriously polyphyletic in NCBI taxonomies (for instance, species in the 704 NCBI genus *Clostridium* appear in 121 GTDB genera and 29 GTDB families)<sup>55</sup>. All SGBs that we 705 706 retrieved from the Pasolli assemblies were assigned taxonomy as part of that work and were 707 called Clostridium sp. CAG-217. Similarly, NCBI assemblies that encoded the most closely acrIIA22 homologs to our original hit were assigned to the GTDB genus CAG-217<sup>54,55</sup>. The raw 708 709 assembly data from the Pasolli database was not assigned a taxonomic label but was nearly identical in nucleotide composition to the CAG-217 contigs (Figure 2, Supplemental Figure 4, 710 Supplemental Datasets 1 and 2). Therefore, we also refer to these sequences as originating in 711 CAG-217 genomes but take care to indicate which assemblies have been assigned a rigorous 712 taxonomy and which ones for which taxonomy has been inferred in this fashion (Supplemental 713 Table 3). 714

### 715 **Comparing genes in genomic islands to phage genomes**

716 We first examined the annotated genes within each of the 128 unique genomic islands. 717 Manual inspection revealed 54 unique gene arrangements that differed in gene content and 718 orientation. We then selected one representative from each arrangement and extracted amino 719 acid sequences from each encoded gene (n=506). Next, we collapsed these 506 proteins into orthologous groups by clustering at 65% amino acid using cd-hit with the following parameters: -720 d 0 -g 1 -aS 0.95 -c 0.65. These cluster counts were used to generate the histogram depicted in 721 722 Supplemental Figure 4C. To determine which protein families may also be phage-encoded, we 723 gueried the longest representative from each cluster with at least two sequences against the 724 database of nine CAG-217 phages described in the section entitled "Annotation and phylogenetic assignment of metagenomic assemblies". We used tblastn with default parameters to perform this 725 726 search, which revealed that some proteins in the CAG-217 genomic islands have homologs in 727 prophage genomes that are out-of-frame with respect to the MetaGeneMark annotations depicted 728 in Supplemental Figure 4A.

### 729 Phylogenetic tree of AcrIIA22 homologs

The 30 unique AcrIIA22 homologs we retrieved were used to create the phylogeny depicted in Figure 3A. These sequences were aligned using the sequence alignment tool in the sequence analysis suite Geneious Prime 2020 v1.1. This alignment is provided as Supplemental Dataset 6. From this alignment, the phylogenetic tree in Figure 3A was generated using PhyML with the LG substitution model<sup>56</sup> and 100 bootstraps. Coloration and tip annotations were then added in Adobe Illustrator.

# 736 Identification of CRISPR-Cas systems and Acrs in CAG-217 assemblies

737 To determine the type and distribution of CRISPR-Cas systems and Acrs in CAG-217 genomes, we downloaded all assembly data for the 779 SGBs assigned to CAG-217 in Pasolli 738 et.  $al^{18}$  (bin 4303). We then predicted CRISPR-Cas systems for all 779 assemblies in bulk using 739 740 the command line version of the CRISPR-Cas prediction suite, cctyper<sup>57</sup>. Specifically, we used 741 version 1.2.1 of cctyper with the following options: --prodigal meta --keep\_tmp. To identify type II-A Acrs, we first downloaded representative sequences for each of the 21 experimentally 742 743 confirmed type II-A Acrs from the unified resource for tracking anti-CRISPRs<sup>58</sup>. We then used tblastn to guery these proteins against the 779 CAG-217 genome bins and considered any hit 744 745 with e-value better than 0.001 (which included all hits with >30% identity across 50% of the query). To check if these Acrs were present in acrIIA22-encoding phages, we performed an identical 746 tblastn search, but this time using the set of nine acrIIA22-encoding prophages as a database. 747

# 748 Recombinant protein overexpression and purification

The AcrIIA22 protein and its mutants were codon optimized for *E. coli* (Genscript or SynBio 749 750 Technologies) and the gene constructs were cloned into the pET15HE or pET15b plasmid<sup>12</sup> to contain an N-terminal, thrombin-cleavable 6XHistidine (His6) tag. These plasmids differ by only a 751 752 few bases just upstream of the N-terminal thrombin tag. For purified, twin-strep tagged proteins, 753 constructs were cloned into a modified pET15b that lacks the N-terminal tag but instead has a C-754 terminal twin-strep tag (Supplemental Table 4). Constructs were transformed and overexpressed 755 in BL21 (DE3) RIL or BL21 (DE3) pLysS E. coli cells. A 10 mL overnight culture (grown in LB + 100 µg/mL ampicillin) was diluted 100-fold into the same media and grown at 37°C with shaking 756 757 to an OD600 of 0.8 for His6-tagged constructs and 0.3 for twin-strep-tagged constructs. 758 Expression was then induced with 0.5 mM IPTG. For His6-tagged constructs, the culture was

shaken for an additional 3 hours at 37°C; twin-strep-tagged constructs were induced at 16C for
22 hours. Cells were harvested by centrifugation and the pellet stored at -20°C.

Cell pellets for His6-tagged constructs were resuspended in 25 mM Tris, pH 7.5, 300 mM
 NaCl, 20 mM imidazole; twin-strep tagged constructs were resuspended in Tris 100nM 8.0 pH,
 150mM NaCl, 1mM EDTA. Cells were lysed by sonication on ice. The lysate was centrifuged in
 an SS34 rotor at 18,000 rpm for 25 minutes, followed by filtering through a 5 µm syringe filter for
 the His6-tagged constructs and a 0.45 µM syringe filter for the twin-strep-tagged constructs.

766 To purify His6-tagged constructs, the clarified lysate was bound using the batch method to Ni-767 NTA agarose resin (Qiagen) at 4°C for 1 hour. The resin was transferred to a gravity column 768 (Biorad), washed with >50 column volumes of Lysis Buffer, and eluted with 25 mM Tris, pH 7.5, 769 300 mM NaCl, 200 mM imidazole. The protein was diluted with 2 column volumes of 25 mM Tris, 770 pH 7.5 and purified on a HiTrapQ column (GE Healthcare) using a 20 mL gradient from 150 mM 771 to 1 M NaCl in 25 mM Tris, pH 7.5. Peak fractions were pooled, concentrated, and buffer exchanged into 200 mM NaCl, 25 mM Tris, pH 7.5 using an Amicon Ultra centrifugal filter with a 772 773 3,000 molecular weight cutoff (Millipore, UFC900324), then cleaved in an overnight 4°C incubation with biotinylated thrombin (EMD Millipore). Streptavidin agarose slurry (Novagen) was 774 775 incubated with cleaved protein at 4°C for 30 minutes to remove thrombin. The sample was then 776 passed through a 0.22 µm centrifugal filter and loaded onto a HiLoad 16/60 Superdex 200 prep 777 grade size exclusion column (Millipore Sigma) equilibrated in 25 mM Tris, pH 7.5, 200 mM NaCl. The peak fractions were pooled, concentrated, and confirmed for purity by SDS-PAGE before use 778 779 in most assays. Figure 4B depicts size exclusion chromatography data generated for thrombincleaved AcrIIA22 variants generated using a Superdex75 16/60 (GE HealthCare) column with 25 780 781 mM Tris, pH 7.5, 200mM NaCl. To correlate nicking activity with protein content across fractions 782 (Supplemental Figure 10B), we collected 13 fractions that span the entire elution peak as well as fractions without AcrIIA22 protein. The protein gel shown in Supplemental Figures 10A and 10B 783 was loaded with 5ul of each concentrated fraction. 784

785 For two additional proteins, we also performed similar Ni-NTA-based purifications of His6tagged constructs, with small deviations from the protocol described in the preceding paragraph. 786 787 Recombinant AcrIIA4 was purified similarly to other His6-tagged Acr proteins but with the 788 following deviations, as previously described<sup>12</sup>. IPTG was used at 0.2 mM and cells were harvested after 18 hours of induction at 18°C. Thrombin cleavage also occurred at 18°C. This 789 790 untagged version was used to help generate Supplemental Figure 6. Peak fractions for all proteins were pooled, concentrated, flash frozen as single-use aliquots in liquid nitrogen, and stored at 791 -80°C. SpyCas9 was expressed in E. coli from plasmid pMJ806 (addgene #39312) to contain a 792 793 TEV-cleavable N-terminal 6XHis-MBP tag and was purified as described previously<sup>12</sup> with 794 sequential steps of purification consisting of Ni-NTA affinity chromatography, TEV cleavage, 795 Heparin HiTrap chromatography, and SEC. The protein was stored in a buffer consisting of 200 796 mM NaCl, 25 mM Tris (pH 7.5), 5% glycerol, and 2 mM DTT. Again, peak fractions were pooled, 797 concentrated, and flash frozen as single-use aliquots.

We also purified AcrIIA22 and AcrIIA4 constructs with a C-terminal twin-strep tag. The protein was expressed and lysed as described above and purified according to the manufacturer's guidelines (IBA Life Sciences, Inc.). Clarified lysates were passed over Strep-Tactin-Sepharose resin using a gravity filtration column. The flow through was passed over the resin a second time. The column was washed with a minimum of 20 column volumes of buffer W, followed by elution in buffer E (150 mM NaCl, 100 mM Tris, pH 8.0 mM, 1 EDTA, 2.5 mM desthiobiotin). The eluted 804 protein was purified over a HiTrapQ column (GE Healthcare) using a 40 mL gradient from 150 805 mM to 0.5 M NaCl in 25 mM Tris, pH 7.5. Peak fractions were pooled and then purified again via 806 size exclusion chromatography with a Biorad Enrich SEC650 10x300mm column in 150mM NaCl. 807 25 mM Tris, pH 7.5. These elution data are shown for AcrIIA22 and its variants in Figure 6B. Fractions were collected across the elution peak and confirmed for purity via silver stain 808 (Supplemental Figure 10E), per manufacturer's recommendations (Thermo Fisher Cat. No. 809 24612). For these proteins, we chose fraction number four to carry forward, as it eluted at 810 approximately four times the monomer's molecular weight, consistent with our proposed tetramer, 811 812 which is depicted in Figure 4C. Protein was then concentrated and flash frozen as single-use aliquots for later use. 813

### 814 X-ray crystallography and structural analyses

An AcrIIA22 crystal was grown using 14mg/mL protein via the hanging drop method using 815 200mM ammonium nitrate, 40% (+/-)-2-methyl-2,4-pentanediol (MPD, Hampton Research), 816 817 10mM MgCl2 as a mother liquor. Diffraction data was collected at the Argonne National 818 Laboratory Structural Biology Center synchrotron facility (Beamline 19BM). Data was processed with HKL2000 in space group P4332, then built and refined using COOT<sup>59</sup> and PHENIX<sup>60</sup>. The 819 completed 2.80Å structure was submitted to the Protein Data Bank with PDB Code 7JTA. The 820 821 detailed PDB validation report is provided (Supplemental Dataset 7). We submitted this finished 822 coordinate file to the PDBe PISA server (Protein Data Bank Europe, Protein Interfaces, Surfaces and Assemblies: http://pdbe.org/pisa/) which uses free energy and interface contacts to calculate 823 likely multimeric assemblies<sup>27</sup>. The server calculated tetrameric, dimeric and monomeric 824 825 structures to be thermodynamically stable in solution. The tetrameric assembly matches the 826 molecular weight expected from the size exclusion column elution peak and is the most likely guaternary structure as calculated by the PISA server. The tetramer gains -41.8 kcal/mol free 827 828 energy by solvation when formed and requires an external driving force of 3.1 kcal/mol to disassemble it according to PISA  $\Delta G$  calculations. 829

### 830 sgRNA generation

The single-guide RNA (sgRNA) for use in *in vitro* experiments was generated as described previously<sup>12</sup>. We made the dsDNA template via one round of thermal cycling (98°C for 90 s, 55°C for 15 s, 72°C for 60 s) in 50 µl reactions. We used the Phusion PCR polymerase mix (NEB) containing 25 pmol each of the following two oligo sequences; the sequence that binds the protospacer on our pIDTsmart target vector is underlined:

- 836 (i) GAAATTAATACGACTCACTATAGG<u>TAATGAAATAAGATCACTAC</u>GTTTTAGAGCT
   837 AGAAATAGCAAGTTAAAATAAGGCTAGTCCG
- 838 (ii) AAAAAAGCACCGACTCGGTGCCACTTTTTCAAGTTGATAACGGACTAGCCTTAT 839 TTTAACTTGC.

The dsDNA templates were then purified using an Oligo Clean and Concentrator Kit (ZymoResearch) before quantification via the Nanodrop. Single-guide RNA (sgRNA) was transcribed from this double-stranded DNA (dsDNA) template by T7 RNA polymerase using Megashortscript Kit (Thermo Fisher #AM1354). Reactions were then treated with DNAse, extracted via phenol-chloroform addition and then chloroform addition, ethanol precipitated, resuspended in RNase free water, quantified by Nanodrop, analyzed for quality on 15% acrylamide/TBE/UREA gels, and frozen at  $-20^{\circ}$ C.

### 847 Pulldown assay using twin-strep-tagged AcrIIA22 and AcrIIA4

The same buffer, consisting of 200 mM NaCl, 25 mM Tris (pH 7.5), was used for pulldowns 848 and to dilute proteins. As a precursor to these assays, 130 pmol SpyCas9 and sgRNA were 849 incubated together at room temperature for 15 minutes where indicated. SpyCas9, with or without 850 851 pre-complexed sgRNA, was then incubated with 230 pmol AcrIIA4 or 320 pmol AcrIIA22 for 25 852 minutes at room temperature. Subsequently, 50 µl of a 10% slurry of Strep-Tactin Resin (IBA 853 Lifesciences #2-1201-002), which was pre-equilibrated in binding buffer, was added to the binding 854 reactions, and incubated at 4°C on a nutator for 45 minutes. Thereafter, all incubations and 855 washes were carried out at 4°C or on ice. Four total washes of this resin were performed, which 856 included one tube transfer. Washes proceeded via centrifugation at 2000 rpm for one minute, 857 aspiration of the supernatant with a 25-gauge needle, and resuspension of the beads in 100 µl 858 binding buffer. Strep-tagged proteins were eluted via suspension in 40 µl of 1x BXT buffer (100 859 mM Tris-CI, 150 mM NaCI, 1 mM EDTA, 50 mM Biotin, pH 8.0) and incubated for 15 min at room 860 temperature. After centrifugation, 30 µl of supernatant was removed and mixed with 4X reducing sample buffer (Thermo Fisher). Proteins were then separated by SDS PAGE on BOLT 4-12% 861 862 gels in MES buffer (Invitrogen) and visualized by Coomassie staining.

### 863 SpyCas9 linear DNA cleavage assay

All SpyCas9 cleavage reactions using linear DNA were performed in cleavage buffer<sup>61</sup> 864 865 (20mM Tris HCI (pH7.5), 5% glycerol, 100mM KCI, 5mM MgCl2, 1mM DTT). In preparation for 866 these reactions, all proteins were diluted in 30 mM NaCl / 25 mM Tris, pH 7.4 / 2.7mM KCl, 867 whereas all DNA and sgRNA reagents were diluted in nuclease-free water. Where indicated, SpyCas9 (0.36 µM) was incubated with sgRNA (0.36 µM) for 10 minutes at room temperature. 868 Before use, sgRNA was melted at 95°C for five minutes and then slowly cooled at 0.1 °C/s to 869 870 promote proper folding. SpyCas9 (either pre-complexed with sgRNA or not, as indicated in 871 Supplemental Figure 7) was then incubated for 10 minutes at room temperature with AcrIIA4 (2.9 872 µM) or AcrIIA22 at each of the following concentrations: [23.2, 11.6, 5.8, and 2.9 µM]. As 873 substrate, the plasmid pIDTsmart was linearized by restriction digest and used at a final 874 concentration of 3.6 nM. The reaction was initiated by the addition of this DNA substrate either in 875 isolation or in combination with sgRNA (0.36 µM) as indicated in Supplemental Figure 7. Reactions were immediately moved to a 37°C incubator and the reaction stopped after fifteen 876 877 minutes via the addition of 0.2% SDS/100 mM EDTA and incubation at 75°C for five minutes. 878 Samples were then run on a 1.5% TAE agarose gel at 120V for 40 minutes. Densitometry was 879 used to calculate the proportion of DNA cleaved by SpyCas9; band intensities were quantified using the BioRad ImageLab software v5.0. 880

### 881 In vivo assay to assess impact of AcrIIA22 on plasmid topology

In all experiments, cultures were first grown overnight at  $37^{\circ}$ C with shaking at 220 rpm in LB with 0.5mM IPTG and, if included, spectinomycin at 50 µg/mL, and kanamycin at 50 µg/mL. For each sample with a SpyCas9-expressing plasmid (*e.g.* Figure 7A), overnight cultures were grown with spectinomycin and kanamycin and diluted 1:50 into LB with 0.5mM IPTG,

spectinomycin (at 50 µg/mL), and, where indicated, doxycycline (at 100 ng/mL, to induce *acr*s). 886 887 Cultures were grown at 37°C with shaking at 220 rpm. If required, 0.2% (L)-arabinose was added after two hours of growth to induce spyCas9 expression. The next morning, cultures were 888 centrifuged at 4100g and plasmids purified using a miniprep kit (Qiagen). We measured the 889 890 concentration of dsDNA in each miniprep using the Qubit-4 fluorometer and the associated 891 dsDNA high sensitivity assay kit (Invitrogen). For each sample with a SpyCas9-expressing 892 plasmid, 150ng of DNA was digested with the restriction enzyme HinclI (NEB) per manufacturer's 893 recommendations, except that digests were incubated overnight before being stopped by heating 894 at 65°C for 20 minutes. This restriction enzyme will cut once, only in the SpyCas9 plasmid, to linearize it. This allowed us to visualize the SpyCas9 plasmid as a single band, which allowed us 895 to identify bands from *acrIIA22*-encoding undigested plasmids more easily. It also served as an 896 internal control for plasmid DNA that is unaffected by SpyCas9 targeting or AcrIIA22 expression 897 (Supplemental Figure 2). Following restriction digest, 30ng of sample was analyzed via gel 898 electrophoresis using a 0.7% TAE-agarose gel run at 120V for 30 minutes. 899

In samples that lacked a SpyCas9-expressing plasmid (*e.g.* Figure 5A), overnight cultures were grown with kanamycin and diluted into LB. Where required, 0.5mM IPTG and doxycycline at 100 ng/mL were added to induce the gene of interest. The next morning, cultures were centrifuged at 4100*g* and plasmids purified using a miniprep kit (Qiagen). The concentration of dsDNA in each miniprep was measured using the Qubit-4 fluorometer and the associated dsDNA high sensitivity assay kit (Invitrogen). Then, 30ng of purified plasmid was directly analyzed by gel electrophoresis using a 0.7% TAE-agarose gel run at 120V for 30 minutes.

### 907 In vitro AcrIIA22 plasmid nicking assay

908 Except for the divalent cation experiment, all reactions were performed using NEB buffer 3.1 (100 mM NaCl, 50 mM Tris-HCl, pH 7.9, 10 mM MqCl2, 100 µg/mL BSA). To determine cation 909 preference, the same reaction buffer was re-created, but MgCl2 was omitted. All proteins were 910 diluted in 130 mM NaCl, 25 mM Tris, pH 7.4, 2.7 mM KCl. DNA was diluted in nuclease-free water. 911 In the cation preference experiment, 60 µM His6-AcrIIA22 and 6 nM of purified pIDTsmart plasmid 912 913 DNA were used. All other reactions were set up with AcrIIA22 constructs and concentrations 914 indicated in figure panels and captions. In the cation preference experiment (Supplemental Figure 11A), reactions were started by adding 10 mM of the indicated cation. All other reactions were 915 initiated via the addition of 2 nM pIDTsmart plasmid DNA. In these cases, reactions were 916 immediately transferred to a 37°C incubator. At 0.5, 1, 2, 4, 6, or 20-hour timepoints, a subset of 917 the reaction was removed and run on a 1.5% TAE agarose gel at 120V for 30 minutes. For the 918 919 fractionation experiment depicted in Supplemental Figure 10B, 5ul of each concentrated fraction was used in a 15ul reaction volume and the reaction was incubated for 24 hours at 37°C. For the 920 921 cation preference experiment, only the 2-hour timepoint was considered and the reaction was stopped via the addition of NEB loading buffer and 100 mM EDTA. In this case, DNA was 922 923 visualized on a 1% TBE gel run for 60 minutes at 110V. Densitometry was used to calculate the 924 proportion of DNA in each topological form via band intensities quantified using the BioRad 925 ImageLab software v5.0.

### 926 SpyCas9 cleavage kinetics assay

All cleavage reactions were performed in the cleavage buffer<sup>61</sup> containing 20mM Tris HCl (pH7.5), 5% glycerol, 100mM KCl, 5mM MgCl2, 1mM DTT. In preparation for these reactions, all proteins were diluted in 30 mM NaCl / 25 mM Tris, pH 7.4 / 2.7mM KCl, whereas all DNA and sgRNA reagents were diluted in nuclease-free water.

Purified pIDTsmart plasmid was pre-treated with either AcrIIA22, the nickase Nb.BssSI 931 (NEB), or no enzyme. For the AcrIIA22 pre-treatment, 3.1 µg of plasmid was incubated with 230 932 µM AcrIIA22 and the plasmid nicked as described previously. Plasmid nicking with Nb.BssSI was 933 carried out via manufacturer's recommendations (NEB). Both reactions were incubated at 37 °C 934 935 for 2 hours. To isolate the nicked plasmid, samples were then run on a 1.5% agarose gel for 2 936 hours and the open-circle form of the plasmid was excised and purified using the Zymo Research Gel DNA Recovery Kit. Untreated plasmid was also purified via gel extraction. Plasmid vield was 937 938 quantified using a Nanodrop.

939 To determine SpyCas9's substrate preference, we incubated each pre-treated plasmid 940 substrate with SpyCas9 and assayed for the appearance of a linearized plasmid as indication of 941 SpyCas9 digestion. In all cases, SpyCas9 was used at a final concentration of 0.32 µM. All 942 reaction components except dsDNA were added on ice, following which SpyCas9 was complexed with equimolar levels of its sgRNA for ten minutes at room temperature. Before addition to the 943 944 reaction, sgRNA was melted at 95°C for five minutes and then slowly cooled at 0.1 °C/s to promote 945 proper folding. To begin the reaction, DNA substrate was added to the reaction mix at a final 946 concentration of 2 nM and the samples moved immediately to 37 °C. At each timepoint, a subset 947 of the reaction was removed, and digestion stopped with 0.2% SDS/100 mM EDTA and by incubating at 75°C for 5 minutes. Samples were run on a 1.5% TAE gel at 120V for 40 minutes 948 and densitometry was used to calculate the proportion of DNA in each topological form via band 949 950 intensities quantified with the BioRad ImageLab software v5.0.

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### 963 **Competing Interests**

All authors declare no significant competing financial, professional, or personal interests that might have influenced the performance or presentation of the work described in this manuscript.

#### 966 Main Figure Captions

Figure 1. Functional selection reveals a metagenomic contig encoding a novel SpyCas9 967 inhibitor. (A) A plasmid protection assay was used to reveal SpyCas9 inhibition. In this assay, 968 969 plasmids without SpyCas9 inhibitors are cleaved by Cas9 and do not give rise to Kan<sup>R</sup> colonies, whereas those encoding inhibitors withstand SpyCas9 attack and yield Kan<sup>R</sup> colonies. (**B**) The 970 contig F01A 4 protects a plasmid from SpyCas9 attack but an early stop codon in orf 1 ( $\Delta$ 1) 971 972 eliminates this phenotype. Stop codons in orf 2 or orf 3 ( $\Delta 2$  and  $\Delta 3$ ) have no effect. Thus, we 973 conclude that orf 1 is necessary for inhibition of SpyCas9. Asterisks depict statistically significant 974 differences in plasmid retention between the indicated genotype and an empty vector control in 975 SpyCas9-inducing conditions (Student's t-test, p<0.002, n=3); ns indicates no significance. All p-976 values were corrected for multiple hypotheses using Bonferroni's method. (C) Expression of orf 1 (which we name acrIIA22) is sufficient for SpyCas9 antagonism, protecting a plasmid as 977 978 effectively as acrIIA4. Asterisks are as in panel B but relate to the GFP negative control rather than to an empty vector. The individual numerical values that underlie the summary data in this 979 980 figure may be found as supporting information file SI\_Data.

Figure 2. AcrIIA22 homologs are found in hypervariable regions of prophage and bacterial 981 982 genomes in the unnamed clostridial genus, CAG-217. (A) We show a schematic representation of an *acrIIA22* homolog embedded in a prophage genome, which is integrated into 983 a bacterial genome (contig #57). We can delineate precise boundaries of the inserted prophage 984 985 based on comparison to a near-identical bacterial contig (contig #55). Prophage genes are colored by functional category, according to the legend at the top. Bacterial genes are colored 986 light gray. (B) Homologs of acrIIA22 are depicted in diverse genomic islands, including Contig #1, 987 whose sequence includes a portion identical to F01A 4, the original metagenomic contig we 988 989 recovered. All acrIIA22 homologs in these loci are closely related but their adjacent genes are 990 different, unrelated gene families (depicted by different colors). Genomic regions flanking these 991 hypervariable islands, including genes immediately adjacent to these islands (purF and radC, in 992 bold outlines), are nearly identical to one another (≥98% nucleotide identity). Contigs are 993 numbered to indicate their descriptions in Supplemental Table 3, which contains their metadata, taxonomy, and sequence retrieval information. All sequences and annotations may also be found 994 995 in Supplemental Datasets 1 and 2. (C) We propose an evolutionary model for the origin of the acrIIA22-encoding hypervariable genomic islands depicted in panel B. We propose that acrIIA22 996 997 moved via prophage integration into a bacterial genomic locus but remained following an 998 incomplete prophage excision event. Its neighboring genes subsequently diversified via horizontal 999 exchange with additional phage genomes without these phage genomes inserting into the locus. 1000 Supplemental Figure 4 depicts a more detailed version of the genomic data underlying this model.

1001 Figure 3. Several AcrIIA22 homologs in the CAG-217 clostridial genus can inhibit SpyCas9. (A) A phylogeny of all unique AcrIIA22 homologs identified from metagenomic and NCBI 1002 1003 databases. Phylogenetic classifications were assigned corresponding to the GTDB naming convention (Methods). Prophage sequences are shaded brown and homologs from hypervariable 1004 1005 bacterial genomic islands are shaded yellow. Sequences obtained from NCBI are labeled with protein accession numbers. In other cases, AcrIIA22 homologs are numbered to match their 1006 1007 contig-of-origin (Supplemental Table 3). In some cases, more than one AcrIIA22 homolog is found on the same contig ('gi' or 'p' indicates its presence in a hypervariable genomic island or prophage 1008 1009 genome, respectively). Circles at nodes indicate bootstrap support  $\geq 0.75$ . Dashed boxes 1010 separate sequences identified from CAG-217 versus Eubacterium R bacterial genera. Filled 1011 areen circles indicate homoloas that were tested for their ability to inhibit SpvCas9 in the plasmid 1012 protection assay in panel B. These homologs have been named with 'a', 'b', or 'c' suffixes to distinguish them from the original AcrIIA22 metagenomic hit; their amino acid identity to the 1013 1014 original hit is shown in parentheses. (B) Several homologs of AcrIIA22 in CAG-217 genomes inhibit SpyCas9. Asterisks depict statistically significant differences in plasmid retention under 1015 SpyCas9-inducing conditions between the indicated sample and a null mutant with an early stop 1016 1017 codon in *acrIIA22*, as indicated in the legend at right (ns indicates no significance; p > 0.05). All 1018 p-values were corrected for multiple hypotheses using Bonferroni's method (Student's t-test, n=3). 1019 (C) AcrIIA22 inhibits divergent Cas9 proteins from Streptococcus pyogenes (SpyCas9) or Neisseria meningitidis (NmCas9) but not Cas12 proteins from Lachnospiraceae bacterium 1020 1021 (LbCas12) or Francisella novicida (FnCas12). As in panel B, green bars indicate samples with 1022 expression of the indicated Cas nuclease while unexpressed controls are depicted with gray lines. For Cas-expressing samples, significance was determined via a Student's t-test (n=3) and 1023 denoted as follows: '\*', p≤0.05 ; '\*\*' p≤0.001 ; 'ns' no significance. Due to slight differences in the 1024 1025 plasmid protection assay in panel C compared to panel B, A22 was re-tested against SpyCas9 to 1026 confirm activity (Methods). The individual numerical values that underlie the summary data in this 1027 figure may be found as supporting information file SI\_Data.

1028 Figure 4. AcrIIA22 is an oligomeric PC4-like protein. (A) AcrIIA22's crystal structure reveals a 1029 homodimer of two four-stranded  $\beta$ -sheets. (B) AcrIIA22 elutes as an oligomer that is approximately four times the predicted molecular mass of its monomer, which is 7 kDa. The gray, 1030 1031 dashed trace depicts protein standards of the indicated molecular weight, in kDa. The orange 1032 trace depicts the elution profile of a two-amino acid C-terminal AcrIIA22 truncation mutant that is predicted to disrupt oligomerization. (C) Ribbon diagram of a proposed AcrIIA22 tetramer which 1033 1034 requires binding between anti-parallel β-strands at the C-termini of AcrIIA22 monomers to form extended, concave β-sheets. The putative oligomerization interface is indicated by the regions 1035 1036 highlighted in yellow and the dashed box, and is detailed further in panel F. Each monomer in the 1037 proposed tetramer is labeled with lower-case Roman numerals (i-iv). (D)  $\beta$ -sheet topology and orientation in AcrIIA22 (blue) resemble that of PC4-like family proteins (in gray, PDB:4BG7 from 1038 1039 phage T5). (E) A monomer of AcrIIA22 (in blue, PDB:7JTA) is structurally similar to a PC4-like 1040 single-stranded DNA binding protein, which is proposed to promote recombination in phage T5 1041 (in gray, PDB:4BG7, Z-score=6.2, matched residues 15%), except for a missing C-terminal alpha 1042 helix. (F) A putative oligomerization interface between the C-termini of two AcrIIA22 monomers 1043 from panel (C) is shown in more detail. Dashed lines indicate potential hydrogen bonds between 1044 the polypeptide backbones. This interface occurs twice in the putative tetramer, between redhued and blue-hued monomers in panel C. 1045

Figure 5. AcrIIA22 nicks supercoiled plasmids in vivo and in vitro. (A) Gel electrophoresis of 1046 1047 plasmids purified from overnight E. coli cultures expressing either acrIIA22, or a null mutant with 1048 an early stop codon, or neither. Compared to the null mutant, more plasmid runs in a slowly migrating, open-circle conformation (OC) rather than supercoiled plasmid (SC) with the wild-type 1049 acrIIA22 allele, suggesting that acrIIA22 may impact plasmid topology. %SC indicates the 1050 percentage of DNA in the supercoiled form for each sample. (B) N-terminally His6-tagged 1051 AcrIIA22 nicks supercoiled plasmids in vitro. (C) C-terminally twin-strep-tagged AcrIIA22 nicks 1052 supercoiled plasmids in vitro with higher specific activity than shown in panel B (compare protein 1053 1054 concentrations). Original, uncropped versions of images depicted in figure may be found in the 1055 supporting information file, SI raw images.

1056 Figure 6. Impaired nicking activity of AcrIIA22 variants in vitro correlates with lower 1057 SpyCas9 inhibition in vivo. (A) Alanine mutagenesis of acidic amino acid residues (glutamic acid or aspartic acid) in AcrIIA22 reveals that D14 is important for plasmid protection against 1058 1059 SpyCas9. Asterisks depict statistically significant differences in plasmid retention under SpyCas9-1060 inducing and non-inducing conditions, per the legend at right. The D14A mutant is significantly impaired, the E4A mutant is slightly impaired, whereas all other mutants are not impaired for 1061 plasmid protection against SpyCas9 compared to an uninduced control. All p-values were 1062 corrected for multiple hypotheses using Bonferroni's method (Student's t-test, n=3). (B) AcrIIA22 1063 1064 (black), AcrIIA22a (dark gray), and a D14A mutant (light gray) all elute with similar oligomer profiles via SEC. The dashed trace depicts protein standards of the indicated molecular weight, 1065 1066 in kDa. (C) AcrIIA22a and the D14A mutant are impaired for nicking relative to AcrIIA22. All 1067 experiments were performed in triplicate, with standard deviations indicated by dashed lines (in most cases, the data points obscure these error bars). Asterisks denote cases where AcrIIA22 is 1068 1069 significantly different than both AcrIIA22a and the D14A mutant after correcting for multiple 1070 hypotheses (Student's t-test, n=3, Bonferroni correction). A single asterisk (\*) means that adjusted p-values for both comparisons are below 0.05. A double asterisk (\*\*) means that adjusted p-1071 1072 values are both below 0.005. Supplemental Figures 10G and 10H show representative gels for 1073 these nicking experiments. The individual numerical values that underlie the summary data in this 1074 figure may be found as supporting information file SI\_Data.

1075 Figure 7. Nicking by AcrIIA22 protects plasmids from SpyCas9 in vivo and in vitro. (A) Gel 1076 electrophoresis of plasmids purified from overnight E. coli cultures expressing either wildtype 1077 acrIIA22 or a mutant with an early stop codon ('null'). In these cultures, SpyCas9 was expressed from a second plasmid, which was linearized via a unique restriction site before electrophoresis. 1078 1079 The acrIIA22-encoding plasmids are indicated with the 'pTarget' label. OC, open-circle; SC, 1080 supercoiled. The '%pTarget' figure indicates the fraction of total DNA attributable to pTarget. 1081 quantified by densitometry analysis. In cases with complete pTarget elimination, all DNA comes 1082 from the SpyCas9 expression plasmid, and thus these bands are more pronounced. However, in the presence of wildtype acrIIA22, pTarget is protected from SpyCas9-mediated cleavage and 1083 1084 makes up 43% of total plasmid DNA. (B) We present a schematic of the experimental design for the data depicted in panel C. The experiment tests whether SpyCas9 preferentially cleaves a 1085 1086 supercoiled or open-circle plasmid target in vitro. Though both plasmid substrates will be 1087 linearized following SpyCas9 cleavage, linear DNA will accumulate more readily with a preferred substrate. (C) Plasmid purifications from overnight cultures were either left unmodified or pre-1088 1089 treated with one of two nickase enzymes, AcrIIA22 or Nb.BssSI, following which each substrate was digested with SpyCas9 in vitro. The percentage of DNA in the linear form is guantified below 1090 1091 the gel, which indicates complete SpyCas9 cleavage. Linear, open-circle (OC), and supercoiled 1092 (SC) plasmid forms are indicated along with the left of the gel, and reaction components below 1093 the gel. SpyCas9 cuts DNA strands sequentially; incomplete digestions with supercoiled substrates produce open-circle plasmids if only one strand has been cleaved (e.g. lane 5). Pre-1094 1095 nicked plasmids, by either AcrIIA22 or Nb.BssSI, are less susceptible to linearization via SpyCas9 1096 cleavage. (D) Endpoint measurements indicate that SpyCas9 more efficiently linearizes supercoiled plasmids than substrates nicked with either AcrIIA22 or Nb.BssSI (Student's t-test, 1097 1098 n=3). (E) A time course experiment demonstrates that SpyCas9 more efficiently linearizes 1099 supercoiled plasmids than AcrIIA22-treated substrates. An asterisk (\*) denotes significant 1100 differences between AcrIIA22-treated and untreated substrates (Student's t-test, p < 0.05, n=3).

1101 The individual numerical values and original images for the data presented in this figure may be 1102 found in the supporting information files SI\_Data and SI\_raw\_images, respectively.

# 1103 Supplemental Figure Captions

Supplemental Figure 1. Orf 1 (acrIIA22) confers mild toxicity in E. coli. Growth rates with 1104 1105 orf 1 induction (green) are 7% lower than those without orf 1 induction (orange). The cfu data shown in Figure 1C were generated from the same experiment depicted here (samples were 1106 1107 removed after six hours of growth to determine these cfu counts). Thus, these data demonstrate 1108 that anti-SpyCas9 activity occurs under conditions with mild orf\_1 toxicity. Growth curves are 1109 shown for samples without SpyCas9 induction to ensure that orf 1 toxicity is not mitigated due to 1110 elimination of its plasmid. Points indicate averages from three replicates. Standard deviations at each timepoint are so small that the error bars do not exceed the bounds of the data point. The 1111 1112 individual numerical values that underlie the summary data in this figure may be found as supporting information file SI\_Data. 1113

Supplemental Figure 2. Orf\_1 (acrIIA22) does not impact SpyCas9 expression. (A) A 1114 1115 schematic description of the experimental design shown in panel (B) is presented. If ORF 1 1116 prevented transcription from pCas9 or altered its copy number, we would expect expression of the orf 1 gene to deplete the level of green fluorescence observed from a construct that replaces 1117 1118 the spycas9 gene with gfp. (B) Fluorescence measurements for the experiment depicted in panel A show that ORF\_1 does not impact GFP expression throughout an E. coli growth curve. Points 1119 1120 indicate averages from three replicates, error bars indicate standard deviation. A western blot shows no depletion of SpyCas9 expression as a function of ORF 1 or GFP expression in growing 1121 1122 *E. coli* cultures at three hours (**C**) or six hours (**D**). As an internal control, GAPDH expression was also detected. The individual numerical values and original images for the data presented in this 1123 1124 figure may be found in the supporting information files SI\_Data and SI\_raw\_images, respectively.

1125 Supplemental Figure 3. AcrilA22 only modestly protects Mu phages against SpyCas9. Mu phage fitness was measured by plaguing on E. coli in the presence of gfp, acrIIA22, or acrIIA4 1126 via serial ten-fold dilutions. Bacterial clearing (black) occurs when phage Mu overcomes SpyCas9 1127 1128 immunity and lyses E. coli. In (A) and in (B), SpyCas9 with a Mu-targeting crRNA confers 1129 substantial protection against phage Mu relative to a non-targeting (n.t.) control, in both conditions 1130 tested. These conditions are depicted at left, with the only difference being whether SpyCas9 was only expressed in liquid growth prior to phage infection (panel A) or expressed both in liquid media 1131 1132 and in solid media throughout infection (panel B). When expressed from a second plasmid, the 1133 positive control acrIIA4 significantly enhances Mu fitness by inhibiting SpyCas9 in all conditions in trans. Though acrIIA22 confers protection against SpyCas9 compared to gfp (negative control), 1134 this effect is milder than with *acrIIA4* and dependent on SpyCas9 expression. 1135

1136 Supplemental Figure 4. AcrIIA22 homologs are found in hypervariable regions of prophage 1137 and bacterial genomes in the CAG-217clostridial genus. (A) Homologs of acrIIA22 are depicted in three related prophage genomes, integrated at three different genomic loci, revealed 1138 1139 by a comparison of prophage-bearing contigs (#57, #56, #37) relative to unintegrated contigs (#55, #58, #17 respectively), which are otherwise nearly identical. Prophage genes are colored 1140 1141 by functional category, according to the legend at the left of panel A. Genes immediately adjacent 1142 to acrIIA22 (solid boxes) vary across phages, despite strong relatedness across much of the 1143 prophage genomes. Bacterial genes are colored gray, except for contig #17, which is also 1144 depicted in panel B, below. (B) Homologs of acrIIA22 are depicted in diverse genomic islands,

1145 including Contig #1, whose sequence includes a portion that is identical to the original 1146 metagenomic contig we recovered (F01A 4). All acrIIA22 homologs in these loci are closely 1147 related but differ in their adjacent genes, which often have homologs in the prophages depicted 1148 in panel A (dashed boxes). Bacterial genomic regions flanking these hypervariable islands are 1149 nearly identical to one another and to prophage integration locus #3, as shown by homology to contig #17 from panel A. Contigs are numbered to indicate their descriptions in Supplemental 1150 Table 3, which contains their metadata, taxonomy, and sequence retrieval information. All 1151 1152 sequences and annotations can also be found in Supplemental Datasets 1 and 2. (C) We tabulate 1153 the prevalence of various protein families (clustered at 65% amino acid identity) in a set of 54 unique genomic islands. Each of these islands is flanked by the conserved genes purF and radC 1154 1155 but contains a different arrangement of encoded genes. Domain-level annotations are indicated 1156 below each protein family (unk; unknown function). Gene symbols above each protein family are 1157 colored and lettered to indicate their counterparts or homologs in panels A and B. The phage capsid icon indicates sequences with homologs in prophage genomes. (D) An evolutionary model 1158 1159 for the origin of the acrIIA22-encoding hypervariable genomic islands depicted in panel B is shown. This panel is reprinted from Figure 2C, for continuity. We propose that acrIIA22 moved 1160 via a phage insertion into a bacterial genomic locus, remained following an incomplete prophage 1161 1162 excision event, and its neighboring genes subsequently diversified via horizontal exchange with additional phage genomes. The individual numerical values that underlie the summary data in 1163 1164 this figure may be found as supporting information file SI Data.

1165 Supplemental Figure 5. Genomic proximity of acrIIA22 homologs to other acr genes. An 1166 acrIIA22-encoding prophage like the one depicted in Figure 2A and those in Supplemental Figure 4A is shown. This prophage encodes for a homolog of the previously described SpyCas9 inhibitor 1167 acrIIA17 within one kilobase of an acrIIA22 homolog. Sequence relatedness between the depicted 1168 acrIIA17 gene and the originally discovered acrIIA17 is shown<sup>22</sup>. Because phages often encode 1169 1170 multiple acrs in the same locus, the co-localization of acrIIA17 with acrIIA22 is consistent with the 1171 latter gene functioning natively to inhibit CRISPR-Cas activity. Prophage genes are colored by 1172 functional category, per the legend and as in Supplemental Figure 4A. Contigs are numbered to 1173 indicate their descriptions in Supplemental Table 3, which contains their metadata, taxonomy, and sequence retrieval information. All sequences and annotations can also be found in 1174 1175 Supplemental Datasets 1 and 2.

1176 Supplemental Figure 6. AcrIIA22 does not strongly bind SpyCas9. SpyCas9 and sgRNA were pre-incubated before mixing with a twin-strep (TS) tagged AcrIIA22 or AcrIIA4. SpyCas9 without 1177 sgRNA was also used. Strep-Tactin pulldowns on AcrIIA4 also pulled down SpyCas9 pre-1178 incubated with sgRNA, as previously reported<sup>12</sup>. Similar pulldowns with AcrIIA22 indicate little to 1179 no interaction with SpyCas9, regardless of whether sgRNA was used. These images depict total 1180 protein content visualized by Coomassie stain. Reaction components are indicated below the gel 1181 1182 image. Asterisks (\*) and dagger (†) symbols indicate AcrIIA4 and AcrIIA22 protein bands that run at slightly different positions than expected due to gel distortion. Original, uncropped versions of 1183 1184 images depicted in figure may be found in the supporting information file, SI\_raw\_images.

Supplemental Figure 7. AcrIIA22 does not protect linear DNA from SpyCas9 cleavage. (A) A schematic cartoon depicts the experiment in panel (B). SpyCas9 was pre-incubated with sgRNA targeting linear DNA. Then, Acr candidates were added. Subsequently, cleavage reactions were performed, and the DNA products visualized by gel electrophoresis. (B) We show the products of the reactions described in panel A for the inhibitors AcrIIA22 and AcrIIA4. SpyCas9 activity is

greatly inhibited by AcrIIA4 but unaffected by AcrIIA22, as indicated by the proportion of cleaved 1190 1191 DNA product. Reaction components are depicted atop the gel image, with molar equivalents 1192 relative to SpyCas9 indicated. The percent of DNA substrate cleaved by SpyCas9 is guantified 1193 below each lane. (C) We perform a similar experiment as in panel A, except candidate Acrs were 1194 incubated with SpyCas9 before sqRNA addition. Reactions were begun via the simultaneous addition of sgRNA and linear dsDNA instead of just dsDNA. (D) The products of the reactions 1195 described in panel C for AcrIIA22 and AcrIIA4 inhibitors are shown. SpyCas9 activity is inhibited 1196 1197 by AcrIIA4 but unaffected by AcrIIA22, as indicated by the proportion of cleaved DNA product. 1198 The data depicted in this figure are not directly comparable to those in figure 7, due to methodological differences and because the preparations of SpyCas9 used in each experiment 1199 1200 exhibited different activities. Original, uncropped versions of images depicted in figure may be 1201 found in the supporting information file, SI raw images.

Supplemental Figure 8. AcrIIA22 resembles a PC4-like protein. (A) We present a ribbon 1202 1203 diagram of a proposed AcrIIA22 tetramer, which requires binding between anti-parallel β-strands 1204 at the C-termini of AcrIIA22 monomers to form extended, concave  $\beta$ -sheets. This putative oligomerization interface is indicated by the regions highlighted in yellow. Each monomer in the 1205 1206 proposed tetramer is labeled with lower-case Roman numerals (i-iv). (B) Space filling model of 1207 the tetrameric AcrIIA22 structure from panel A, with relative charge depicted, highlighting a groove (dashed line with arrowhead) that may accommodate nucleic acids (based on analogy to other 1208 PC4-like proteins). (C) AcrIIA22 monomers (i) and (ii) from the tetramer in panel A likely interact 1209 1210 via a series of hydrophobic interactions, as indicated by the predominantly non-polar sidechains 1211 colored in yellow. The boxed region highlights residue D14, which is important for nicking activity and plasmid protection against SpyCas9, and is enlarged in panel F. (D) In conventional PC4-like 1212 1213 family proteins, such as the putative single-stranded DNA binding protein from phage T5 depicted 1214 in gray (PDB:4BG7), the same topology of outward facing, concave  $\beta$ -sheets are instead 1215 stabilized via interactions between opposing  $\alpha$ -helices (depicted in opaque light blue). (E) An 1216 overlay of  $\beta$ -sheets from AcrIIA22 (blue, PDB:7JTA) and the phage T5 PC4-like protein (gray, PDB:4BG7) illustrates their similar topologies. (F) Two D14 residues in loop regions of AcrIIA22 1217 1218 monomers (i) and (ii) are in close proximity. These residues are important for nicking activity and may bind divalent cations in cells under physiological pH. (G) A close view of a putative salt bridge 1219 between R30 of monomers (i) / (ii) and the peptide backbone of the C-terminus of monomers (iv) 1220 1221 / (iii), respectively. AcrIIA22 monomers are colored as described in panel A.

1222 Supplemental Figure 9. A 2-aa truncation mutant of AcrIIA22 is impaired for SpyCas9 1223 inhibition and nicking activity. (A) An in vivo plasmid protection assay. Asterisks depict 1224 statistically significant differences in plasmid retention under SpyCas9-inducing conditions with either wild-type AcrIIA22, a null mutant with an early stop codon, a 2-aa truncation, or a negative 1225 control gfp gene (adj. p < 0.005, Student's t-test, n=3). The truncation mutant retains mild but 1226 1227 severely impaired activity, as it protects a plasmid from SpyCas9 more effectively than a null mutant (p = 0.012) or GFP control (p = 0.015). All p-values were corrected for multiple hypotheses 1228 1229 using Bonferroni's method. (B) The 2-aa truncation mutant is impaired for nicking in vitro, relative 1230 to wild-type AcrIIA22. In both cases, 25µM of protein was used following NiNTA-based purification of an N-terminal, His6-tagged construct. An asterisk (\*) denotes significant differences between 1231 AcrIIA22-treated and untreated substrates (Student's t-test, p < 0.05, n=3). Standard deviations 1232 1233 are indicated by dashed lines (in most cases, the data points obscure these error bars). The 1234 individual numerical values that underlie the summary data in this figure may be found as 1235 supporting information file SI\_Data.

1236 Supplemental Figure 10. AcrIIA22 nicks supercoiled plasmids. (A) A Coomassie stain of an 1237 N-terminally His6-tagged AcrIIA22 construct shows no co-purifying proteins. (B) The nicking activity for this protein preparation (bottom) correlates with the intensity of the Coomassie-stained 1238 1239 protein band across purification fractions (top). In each lane, supercoiled (SC) plasmid DNA 1240 represents the un-nicked fraction whereas open circle (OC) and linear DNA have been nicked at least once. (C) This panel is a quantification of the experiment depicted in panel B across all 13 1241 fractions collected. (D) His6-AcrIIA22 nicks supercoiled plasmids in a time and concentration 1242 dependent manner. A decrease in the proportion of supercoiled plasmid DNA indicates nicking 1243 1244 activity, as depicted in Figure 5B. (E) A silver stain of a C-terminally twin-strep-tagged AcrIIA22 construct shows no co-purifying proteins. Equal volumes of each protein fraction were loaded in 1245 1246 each lane, for all samples. Fraction 4 was concentrated and used for all in vitro experiments. (F) 1247 A C-terminal, but not N-terminal twin-strep tag is compatible with AcrIIA22's ability to protect a target plasmid from SpyCas9 elimination in vivo. Statistically significant differences in plasmid 1248 retention between SpyCas9-inducing and non-inducing conditions were determined via a 1249 Student's t-test (n=3); '\*\*' indicates p≤0.001. All p-values were adjusted for multiple hypotheses 1250 using the Bonferroni correction. (G) The D14A mutation in AcrIIA22 impairs nicking activity. Over 1251 1252 time, the wild-type AcrIIA22-twin-strep construct consistently converts a higher fraction of plasmid DNA from its supercoiled (SC) form to an open-circle (OC) conformation than a D14A mutant. 1253 Control plasmids include a miniprepped sample and sample pre-treated with the commercial 1254 1255 nickase, Nb.BssSI. Reaction times are indicated to the right of each image. (H) AcrIIA22a (Figure 1256 3B) is impaired for nicking activity relative to AcrIIA22. As in panel G, both constructs were purified via C-terminal twin-strep tags. The individual numerical values and original images for the data 1257 1258 presented in this figure may be found in the supporting information files SI\_Data and 1259 SI raw images, respectively.

1260 Supplemental Figure 11. Divalent cations influence AcrIIA22's nicking activity. (A) We

1261 present the impact of different divalent cations on AcrIIA22's nicking activity, which is highest

with Mg<sup>2+</sup>, Mn<sup>2+</sup>, and Co<sup>2+</sup>. OC, open-circle plasmid form. SC, supercoiled plasmid. (**B**) The

1263 open-circle plasmid product persists through phenol-chloroform extraction following AcrIIA22

1264 treatment, indicating that it directly results from AcrIIA22's nicking activity.

### References

1265	1	Barrangou, R. et al. CRISPR provides acquired resistance against viruses in
1266		prokaryotes. Science <b>315</b> , 1709-1712, doi:10.1126/science.1138140 (2007).
1267	2	Marraffini, L. A. & Sontheimer, E. J. CRISPR interference limits horizontal gene transfer
1268		in staphylococci by targeting DNA. Science 322, 1843-1845,
1269		doi:10.1126/science.1165771 (2008).
1270	3	Deveau, H. et al. Phage response to CRISPR-encoded resistance in Streptococcus
1271		thermophilus. <i>J Bacteriol</i> <b>190</b> , 1390-1400, doi:10.1128/JB.01412-07 (2008).
1272	4	Mendoza, S. D. et al. A bacteriophage nucleus-like compartment shields DNA from
1273		CRISPR nucleases. Nature 577, 244-248, doi:10.1038/s41586-019-1786-y (2020).
1274	5	Malone, L. M. et al. A jumbo phage that forms a nucleus-like structure evades CRISPR-
1275		Cas DNA targeting but is vulnerable to type III RNA-based immunity. Nat Microbiol 5, 48-
1276		55, doi:10.1038/s41564-019-0612-5 (2020).
1277	6	Bryson, A. L. et al. Covalent Modification of Bacteriophage T4 DNA Inhibits CRISPR-
1278		Cas9. <i>mBio</i> 6, e00648, doi:10.1128/mBio.00648-15 (2015).
1279	7	Stanley, S. Y. & Maxwell, K. L. Phage-Encoded Anti-CRISPR Defenses. Annu Rev
1280		Genet 52, 445-464, doi:10.1146/annurev-genet-120417-031321 (2018).
1281	8	Trasanidou, D. et al. Keeping crispr in check: diverse mechanisms of phage-encoded
1282		anti-crisprs. FEMS Microbiol Lett, doi:10.1093/femsle/fnz098 (2019).
1283	9	Davidson, A. R. et al. Anti-CRISPRs: Protein Inhibitors of CRISPR-Cas Systems. Annu
1284		<i>Rev Biochem</i> <b>89</b> , 309-332, doi:10.1146/annurev-biochem-011420-111224 (2020).
1285	10	Wiegand, T., Karambelkar, S., Bondy-Denomy, J. & Wiedenheft, B. Structures and
1286		Strategies of Anti-CRISPR-Mediated Immune Suppression. Annu Rev Microbiol 74, 21-
1287		37, doi:10.1146/annurev-micro-020518-120107 (2020).
1288	11	Hatfull, G. F. Dark Matter of the Biosphere: the Amazing World of Bacteriophage
1289		Diversity. J Virol 89, 8107-8110, doi:10.1128/JVI.01340-15 (2015).
1290	12	Forsberg, K. J. et al. Functional metagenomics-guided discovery of potent Cas9
1291		inhibitors in the human microbiome. <i>eLife</i> <b>8</b> , e46540, doi:10.7554/eLife.46540 (2019).
1292	13	Szczelkun, M. D. et al. Direct observation of R-loop formation by single RNA-guided
1293		Cas9 and Cascade effector complexes. Proc Natl Acad Sci U S A 111, 9798-9803,
1294		doi:10.1073/pnas.1402597111 (2014).
1295	14	Farasat, I. & Salis, H. M. A Biophysical Model of CRISPR/Cas9 Activity for Rational
1296		Design of Genome Editing and Gene Regulation. <i>PLoS Comput Biol</i> <b>12</b> , e1004724,
1297		doi:10.1371/journal.pcbi.1004724 (2016).
1298	15	Ivanov, I. E. et al. Cas9 interrogates DNA in discrete steps modulated by mismatches
1299		and supercoiling. Proc Natl Acad Sci U S A 117, 5853-5860,
1300		doi:10.1073/pnas.1913445117 (2020).
1301	16	Tsui, T. K. M., Hand, T. H., Duboy, E. C. & Li, H. The Impact of DNA Topology and
1302		Guide Length on Target Selection by a Cytosine-Specific Cas9. ACS Synth Biol 6, 1103-
1303		1113, doi:10.1021/acssynbio.7b00050 (2017).
1304	17	Paez-Espino, D. et al. IMG/VR v.2.0: an integrated data management and analysis
1305		system for cultivated and environmental viral genomes. Nucleic acids research 47,
1306		D678-D686, doi:10.1093/nar/gky1127 (2019).
1307	18	Pasolli, E. et al. Extensive Unexplored Human Microbiome Diversity Revealed by Over
1308		150,000 Genomes from Metagenomes Spanning Age, Geography, and Lifestyle. Cell
1309		<b>176</b> , 649-662 e620, doi:10.1016/j.cell.2019.01.001 (2019).
1310	19	Dobrindt, U., Hochhut, B., Hentschel, U. & Hacker, J. Genomic islands in pathogenic and
1311		environmental microorganisms. Nat Rev Microbiol 2, 414-424, doi:10.1038/nrmicro884
1312		(2004).

1313	20	Juhas, M. et al. Genomic islands: tools of bacterial horizontal gene transfer and
1314		evolution. FEMS Microbiol Rev 33, 376-393, doi:10.1111/j.1574-6976.2008.00136.x
1315		(2009).
1316	21	Makarova, K. S. et al. Evolutionary classification of CRISPR-Cas systems: a burst of
1317		class 2 and derived variants. Nat Rev Microbiol 18, 67-83, doi:10.1038/s41579-019-
1318		0299-x (2020).
1319	22	Mahendra, C. et al. Broad-spectrum anti-CRISPR proteins facilitate horizontal gene
1320		transfer. Nat Microbiol 5, 620-629, doi:10.1038/s41564-020-0692-2 (2020).
1321	23	Bondy-Denomy, J., Pawluk, A., Maxwell, K. L. & Davidson, A. R. Bacteriophage genes
1322		that inactivate the CRISPR/Cas bacterial immune system. Nature 493, 429-432,
1323		doi:10.1038/nature11723 (2013).
1324	24	Osuna, B. A. et al. Listeria Phages Induce Cas9 Degradation to Protect Lysogenic
1325		Genomes. Cell Host Microbe 28, 31-40 e39, doi:10.1016/j.chom.2020.04.001 (2020).
1326	25	Song, Y. et al. High-resolution comparative modeling with RosettaCM. Structure 21,
1327		1735-1742, doi:10.1016/j.str.2013.08.005 (2013).
1328	26	Janowski, R. & Niessing, D. The large family of PC4-like domains - similar folds and
1329	20	functions throughout all kingdoms of life. RNA Biol <b>17</b> , 1228-1238,
1330		doi:10.1080/15476286.2020.1761639 (2020).
1331	27	Krissinel, E. & Henrick, K. Inference of macromolecular assemblies from crystalline
1332		state. <i>J Mol Biol</i> <b>372</b> , 774-797, doi:10.1016/j.jmb.2007.05.022 (2007).
1333	28	Steigemann, B., Schulz, A. & Werten, S. Bacteriophage T5 encodes a homolog of the
1334	20	eukaryotic transcription coactivator PC4 implicated in recombination-dependent DNA
1335		replication. <i>J Mol Biol</i> <b>425</b> , 4125-4133, doi:10.1016/j.jmb.2013.09.001 (2013).
1336	29	Werten, S. Identification of the ssDNA-binding protein of bacteriophage T5: Implications
1337	20	for T5 replication. <i>Bacteriophage</i> <b>3</b> , e27304, doi:10.4161/bact.27304 (2013).
1338	30	Werten, S. <i>et al.</i> High-affinity DNA binding by the C-terminal domain of the
1339	50	transcriptional coactivator PC4 requires simultaneous interaction with two opposing
1340		unpaired strands and results in helix destabilization. <i>J Mol Biol</i> <b>276</b> , 367-377,
1340		doi:10.1006/jmbi.1997.1534 (1998).
1341	31	Dorman, C. J. & Ni Bhriain, N. CRISPR-Cas, DNA Supercoiling, and Nucleoid-
1342	51	Associated Proteins. <i>Trends Microbiol</i> <b>28</b> , 19-27, doi:10.1016/j.tim.2019.08.004 (2020).
1343	32	Xiong, X. <i>et al.</i> SspABCD–SspE is a phosphorothioation-sensing bacterial defence
1344	52	system with broad anti-phage activities. <i>Nature Microbiology</i> <b>5</b> , 917-928,
1345		doi:10.1038/s41564-020-0700-6 (2020).
	22	Yang, W. Nucleases: diversity of structure, function and mechanism. Q Rev Biophys 44,
1347	33	1-93, doi:10.1017/S0033583510000181 (2011).
1348 1349	34	Vink, J. N. A. <i>et al.</i> Direct Visualization of Native CRISPR Target Search in Live Bacteria
	34	Reveals Cascade DNA Surveillance Mechanism. <i>Mol Cell</i> <b>77</b> , 39-50 e10,
1350		doi:10.1016/j.molcel.2019.10.021 (2020).
1351	25	Harshey, R. M. Transposable Phage Mu. <i>Microbiol Spectr</i> <b>2</b> ,
1352	35	10.1128/microbiolspec.MDNA1123-0007-2014, doi:10.1128/microbiolspec.MDNA3-
1353		•
1354	26	0007-2014 (2014).
1355	36	Westra, E. R. <i>et al.</i> CRISPR immunity relies on the consecutive binding and degradation
1356		of negatively supercoiled invader DNA by Cascade and Cas3. <i>Mol Cell</i> <b>46</b> , 595-605, doi:10.1016/j.moloci.2012.02.018 (2012)
1357	27	doi:10.1016/j.molcel.2012.03.018 (2012).
1358	37	Mattenberger, Y., Silva, F. & Belin, D. 55.2, a phage T4 ORFan gene, encodes an
1359		inhibitor of Escherichia coli topoisomerase I and increases phage fitness. <i>PLoS One</i> <b>10</b> , 0124200, doi:10.1271/journal.page.0124200 (2015)
1360	20	e0124309, doi:10.1371/journal.pone.0124309 (2015).
1361	38	Ramirez-Chamorro, L., Boulanger, P. & Rossier, O. Strategies for Bacteriophage T5
1362		Mutagenesis: Expanding the Toolbox for Phage Genome Engineering. <i>Frontiers in</i>
1363		<i>Microbiology</i> <b>12</b> , 816 (2021).

1364	39	Johnston, J. V., Nichols, B. P. & Donelson, J. E. Distribution of "minor" nicks in
1365		bacteriophage T5 DNA. <i>J Virol</i> <b>22</b> , 510-519 (1977).
1366	40	Roy, D., Huguet, K. T., Grenier, F. & Burrus, V. IncC conjugative plasmids and
1367		SXT/R391 elements repair double-strand breaks caused by CRISPR–Cas during
1368		conjugation. Nucleic Acids Research, doi:10.1093/nar/gkaa518 (2020).
1369	41	Weigel, C. & Seitz, H. Bacteriophage replication modules. FEMS Microbiol Rev 30, 321-
1370		381, doi:10.1111/j.1574-6976.2006.00015.x (2006).
1371	42	Uribe, R. V. et al. Discovery and Characterization of Cas9 Inhibitors Disseminated
1372		across Seven Bacterial Phyla. Cell Host Microbe 25, 233-241 e235,
1373		doi:10.1016/j.chom.2019.01.003 (2019).
1374	43	Borges, A. L. et al. Bacteriophage Cooperation Suppresses CRISPR-Cas3 and Cas9
1375		Immunity. Cell 174, 917-925 e910, doi:10.1016/j.cell.2018.06.013 (2018).
1376	44	Landsberger, M. et al. Anti-CRISPR Phages Cooperate to Overcome CRISPR-Cas
1377		Immunity. Cell 174, 908-916 e912, doi:10.1016/j.cell.2018.05.058 (2018).
1378	45	Chevallereau, A. et al. Exploitation of the Cooperative Behaviors of Anti-CRISPR
1379		Phages. Cell Host Microbe 27, 189-198 e186, doi:10.1016/j.chom.2019.12.004 (2020).
1380	46	Hooper, D. C. & Jacoby, G. A. Mechanisms of drug resistance: quinolone resistance.
1381		Ann N Y Acad Sci 1354, 12-31, doi:10.1111/nyas.12830 (2015).
1382	47	Pawluk, A., Davidson, A. R. & Maxwell, K. L. Anti-CRISPR: discovery, mechanism and
1383		function. Nat Rev Microbiol, doi:10.1038/nrmicro.2017.120 (2017).
1384	48	Borges, A. L., Davidson, A. R. & Bondy-Denomy, J. The Discovery, Mechanisms, and
1385		Evolutionary Impact of Anti-CRISPRs. Annu Rev Virol 4, 37-59, doi:10.1146/annurev-
1386		virology-101416-041616 (2017).
1387	49	Li, W. & Godzik, A. Cd-hit: a fast program for clustering and comparing large sets of
1388		protein or nucleotide sequences. Bioinformatics 22, 1658-1659,
1389		doi:10.1093/bioinformatics/btl158 (2006).
1390	50	Zhu, W., Lomsadze, A. & Borodovsky, M. Ab initio gene identification in metagenomic
1391		sequences. Nucleic Acids Res 38, e132, doi:10.1093/nar/gkq275 (2010).
1392	51	Finn, R. D., Clements, J. & Eddy, S. R. HMMER web server: interactive sequence
1393		similarity searching. Nucleic Acids Res <b>39</b> , W29-37, doi:10.1093/nar/gkr367 (2011).
1394	52	Haft, D. H. et al. TIGRFAMs: a protein family resource for the functional identification of
1395	-	proteins. Nucleic Acids Res 29, 41-43 (2001).
1396	53	Bateman, A. et al. The Pfam protein families database. Nucleic Acids Res 28, 263-266
1397		(2000).
1398	54	Parks, D. H. <i>et al.</i> A complete domain-to-species taxonomy for Bacteria and Archaea.
1399		Nat Biotechnol, doi:10.1038/s41587-020-0501-8 (2020).
1400	55	Parks, D. H. <i>et al.</i> A standardized bacterial taxonomy based on genome phylogeny
1401		substantially revises the tree of life. Nat Biotechnol 36, 996-1004, doi:10.1038/nbt.4229
1402		(2018).
1403	56	Le, S. Q. & Gascuel, O. An Improved General Amino Acid Replacement Matrix.
1404		Molecular Biology and Evolution <b>25</b> , 1307-1320, doi:10.1093/molbev/msn067 (2008).
1405	57	Russel, J., Pinilla-Redondo, R., Mayo-Muñoz, D., Shah, S. A. & Sørensen, S. J.
1406	0.	CRISPRCasTyper: An automated tool for the identification, annotation and classification
1407		of CRISPR-Cas loci. <i>bioRxiv</i> , 2020.2005.2015.097824, doi:10.1101/2020.05.15.097824
1408		(2020).
1400	58	Bondy-Denomy, J. <i>et al.</i> A Unified Resource for Tracking Anti-CRISPR Names. <i>The</i>
1400		<i>CRISPR Journal</i> <b>1</b> , 304-305, doi:10.1089/crispr.2018.0043 (2018).
1411	59	Emsley, P., Lohkamp, B., Scott, W. G. & Cowtan, K. Features and development of Coot.
1412	00	Acta Crystallogr D Biol Crystallogr <b>66</b> , 486-501, doi:10.1107/S0907444910007493
1412		(2010).
1413		

- 1414 60 Adams, P. D. et al. PHENIX: a comprehensive Python-based system for macromolecular
- 1415 structure solution. Acta Crystallogr D Biol Crystallogr 66, 213-221,
- 1416 doi:10.1107/S0907444909052925 (2010).
- 1417 61 Harrington, L. B. et al. A Broad-Spectrum Inhibitor of CRISPR-Cas9. Cell,
- 1418 doi:10.1016/j.cell.2017.07.037 (2017).

Data collection		
Space Group	P4332	
Cell Dimensions		
a, b, c (Å)	128.56, 128.56, 128.56	
α, β, γ (°)	90.0, 90.0, 90.0	
Resolution (Å)	50.00 - 2.80	
R <sub>merge</sub>	0.106 (0.906)	
I/σ <sub>1</sub>	17.4 (2.6)	
Completeness (%)	98.7 (100.0)	
Redundancy	10.4 (10.7)	
CC 1/2	0.837	
Refinement		
No. Reflections	9334	
R <sub>work</sub> (R <sub>free</sub> ) (%)	22.2 (24.6)	
No. Complex in ASU	2	
No. atoms		
Protein	810	
Heteroatoms	50	
Water	3	
B-factor	82.82	
R.m.s deviations		
Bond lengths (Å)	0.003	
Bond angles ( <sup>0</sup> )	0.610	
Ramachandran		
Preferred (%)	98.15	
Allowed (%)	1.85	
Outliers (%)	0	

 Table 1. Structural features of AcrIIA22.

### Other supplemental materials.

**Supplemental Table 1.** Whether known anti-CRISPRs can bind Cas proteins or inhibit their cleavage activity as purified proteins.

Supplemental Table 2. PC4-like proteins with structural homology to AcrIIA22.

**Supplemental Table 3.** Descriptions of all sequences used in this study. All sequences and annotations are also available as supplemental data.

Supplemental Table 4. Plasmids used in this study.

Supplemental Table 5. Gene sequences used in this study.

SI\_Data. All raw data for main and supplemental figures depicted in this study (as a spreadsheet).

**SI\_raw\_images**. Full gel images for all cropped gels depicted in this study, compiled into a .pdf document.

**Supplemental Dataset 1**. 68 contigs sequences referenced in the manuscript with Pfam, TIGRFAM, and AcrIIA22 homolog annotations (in genbank format).

Supplemental Dataset 2. 68 contigs sequences referenced in the manuscript (in fasta format).

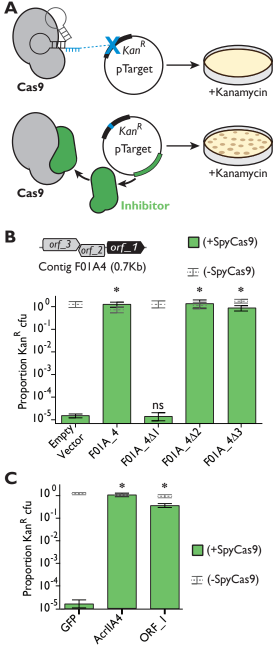
**Supplemental Dataset 3**. Nine AcrIIA22-encoding prophage sequences referenced in the manuscript with Pfam, TIGRFAM, and AcrIIA22 homolog annotations (in genbank format).

**Supplemental Dataset 4**. Nine AcrIIA22-encoding prophage sequences referenced in the manuscript (in fasta format).

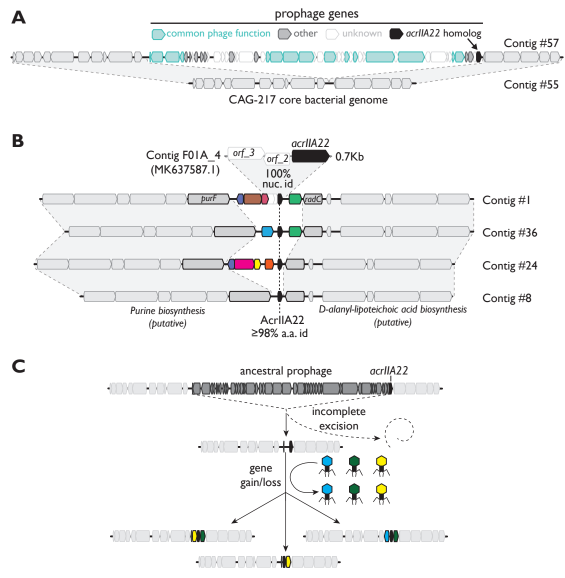
**Supplemental Dataset 5**. All metagenomic contigs with  $\geq$ 98% nucleotide identity to *acrIIA22*-associated genes, *purF* and *radC*. Pfam, TIGRFAM, and AcrIIA22 homolog annotations are also provided (file in genbank format).

**Supplemental Dataset 6**. Amino acid sequence alignment of 30 AcrIIA22 homologs (in fasta format).

Supplemental Dataset 7. The detailed PDB validation report for AcrIIA22's crystal structure.



**Figure 1.** Functional selection reveals a metagenomic contig encoding a novel SpyCas9 inhibitor. (A) A plasmid protection assay was used to reveal SpyCas9 inhibition. In this assay, plasmids without SpyCas9 inhibitors are cleaved by Cas9 and do not give rise to Kan<sup>R</sup> colonies, whereas those encoding inhibitors withstand SpyCas9 attack and yield Kan<sup>R</sup> colonies. (B) The contig F01A\_4 protects a plasmid from SpyCas9 attack but an early stop codon in *orf\_1 (\Delta1*) eliminates this phenotype. Stop codons in *orf\_2* or *orf\_3 (\Delta2* and  $\Delta$ 3) have no effect. Thus, we conclude that *orf\_1* is necessary for inhibition of SpyCas9. Asterisks depict statistically significant differences in plasmid retention between the indicated genotype and an empty vector control in SpyCas9-inducing conditions (Student's t-test, p<0.002, n=3); ns indicates no significance. All p-values were corrected for multiple hypotheses using Bonferroni's method. (C) Expression of *orf\_1* (which we name *acrIIA22*) is sufficient for SpyCas9 antagonism, protecting a plasmid as effectively as *acrIIA4*. Asterisks are as in panel B but relate to the GFP negative control rather than to an empty vector. The individual numerical values that underlie the summary data in this figure may be found as supporting information file SI Data.



hypervariable genomic islands

Figure 2. AcrIIA22 homologs are found in hypervariable regions of prophage and bacterial genomes in the unnamed clostridial genus, CAG-217. (A) We show a schematic representation of an acrIIA22 homolog embedded in a prophage genome, which is integrated into a bacterial genome (contig #57). We can delineate precise boundaries of the inserted prophage based on comparison to a nearidentical bacterial contig (contig #55). Prophage genes are colored by functional category, according to the legend at the top. Bacterial genes are colored light gray. (B) Homologs of acrIIA22 are depicted in diverse genomic islands, including Contig #1, whose sequence includes a portion identical to F01A 4, the original metagenomic contig we recovered. All acrIIA22 homologs in these loci are closely related but their adjacent genes are different, unrelated gene families (depicted by different colors). Genomic regions flanking these hypervariable islands, including genes immediately adjacent to these islands (purF and radC, in bold outlines), are nearly identical to one another (≥98% nucleotide identity). Contigs are numbered to indicate their descriptions in Supplemental Table 3, which contains their metadata, taxonomy, and sequence retrieval information. All sequences and annotations may also be found in Supplemental Datasets 1 and 2. (C) We propose an evolutionary model for the origin of the acrIIA22encoding hypervariable genomic islands depicted in panel B. We propose that acrIIA22 moved via prophage integration into a bacterial genomic locus but remained following an incomplete prophage excision event. Its neighboring genes subsequently diversified via horizontal exchange with additional phage genomes without these phage genomes inserting into the locus. Supplemental Figure 4 depicts a more detailed version of the genomic data underlying this model.

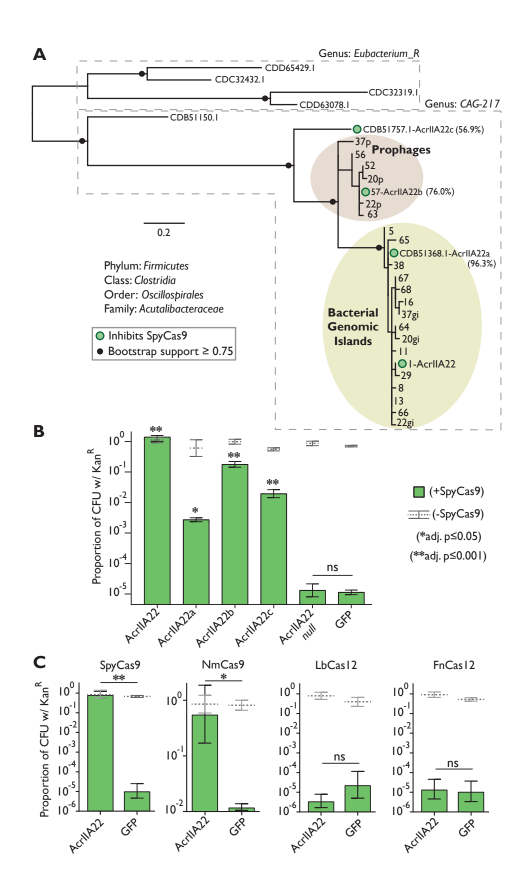


Figure 3. Several AcrIIA22 homologs in the CAG-217 clostridial genus can inhibit SpyCas9. (A) A phylogeny of all unique AcrIIA22 homologs identified from metagenomic and NCBI databases. Phylogenetic classifications were assigned corresponding to the GTDB naming convention (Methods). Prophage sequences are shaded brown and homologs from hypervariable bacterial genomic islands are shaded yellow. Sequences obtained from NCBI are labeled with protein accession numbers. In other cases, AcrIIA22 homologs are numbered to match their contig-of-origin (Supplemental Table 3). In some cases, more than one AcrIIA22 homolog is found on the same contig ('gi' or 'p' indicates its presence in a hypervariable genomic island or prophage genome, respectively). Circles at nodes indicate bootstrap support  $\ge$  0.75. Dashed boxes separate sequences identified from CAG-217 versus *Eubacterium* R bacterial genera. Filled green circles indicate homologs that were tested for their ability to inhibit SpyCas9 in the plasmid protection assay in panel B. These homologs have been named with 'a', 'b', or 'c' suffixes to distinguish them from the original AcrIIA22 metagenomic hit; their amino acid identity to the original hit is shown in parentheses. (B) Several homologs of AcrIIA22 in CAG-217 genomes inhibit SpyCas9. Asterisks depict statistically significant differences in plasmid retention under SpyCas9-inducing conditions between the indicated sample and a null mutant with an early stop codon in acrIIA22, as indicated in the legend at right (ns indicates no significance; p > 0.05). All p-values were corrected for multiple hypotheses using Bonferroni's method (Student's t-test, n=3). (C) AcrIIA22 inhibits divergent Cas9 proteins from Streptococcus pyogenes (SpyCas9) or Neisseria meningitidis (NmCas9) but not Cas12 proteins from Lachnospiraceae bacterium (LbCas12) or Francisella novicida (FnCas12). As in panel B, green bars indicate samples with expression of the indicated Cas nuclease while unexpressed controls are depicted with gray lines. For Cas-expressing samples, significance was determined via a Student's t-test (n=3) and denoted as follows: '\*', p≤0.05 ; '\*\*' p≤0.001 ; 'ns' no significance. Due to slight differences in the plasmid protection assay in panel C compared to panel B, A22 was re-tested against SpyCas9 to confirm activity (Methods). The individual numerical values that underlie the summary data in this figure may be found as supporting information file SI Data.

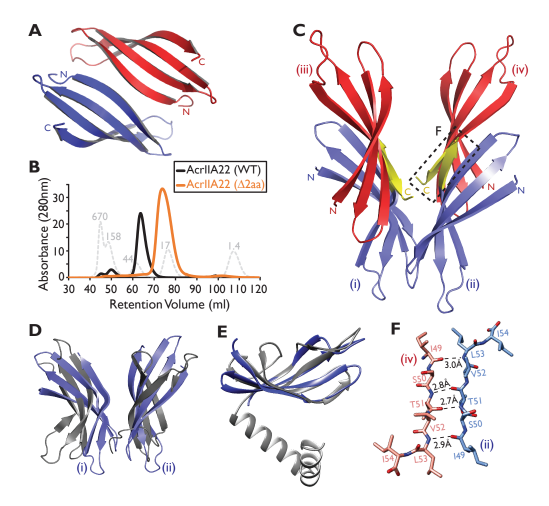
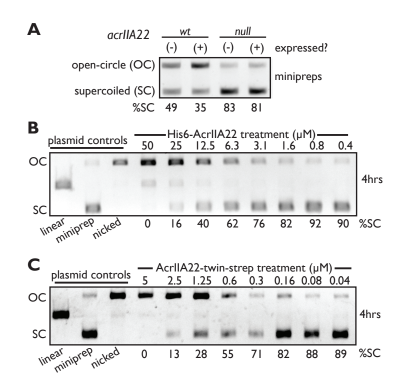


Figure 4. AcrIIA22 is an oligomeric PC4-like protein. (A) AcrIIA22's crystal structure reveals a homodimer of two four-stranded  $\beta$ -sheets. (B) AcrIIA22 elutes as an oligomer that is approximately four times the predicted molecular mass of its monomer, which is 7 kDa. The gray, dashed trace depicts protein standards of the indicated molecular weight, in kDa. The orange trace depicts the elution profile of a two-amino acid C-terminal AcrIIA22 truncation mutant that is predicted to disrupt oligomerization. (C) Ribbon diagram of a proposed AcrIIA22 tetramer which requires binding between anti-parallel β-strands at the C-termini of AcrIIA22 monomers to form extended, concave  $\beta$ -sheets. The putative oligomerization interface is indicated by the regions highlighted in yellow and the dashed box, and is detailed further in panel F. Each monomer in the proposed tetramer is labeled with lower-case Roman numerals (i-iv). (D)  $\beta$ -sheet topology and orientation in AcrIIA22 (blue) resemble that of PC4-like family proteins (in gray, PDB:4BG7 from phage T5). (E) A monomer of AcrIIA22 (in blue, PDB:7JTA) is structurally similar to a PC4-like single-stranded DNA binding protein, which is proposed to promote recombination in phage T5 (in gray, PDB:4BG7, Z-score=6.2, matched residues 15%), except for a missing C-terminal alpha helix. (F) A putative oligomerization interface between the C-termini of two AcrIIA22 monomers from panel (C) is shown in more detail. Dashed lines indicate potential hydrogen bonds between the polypeptide backbones. This interface occurs twice in the putative tetramer, between red-hued and blue-hued monomers in panel C.



**Figure 5.** AcrIIA22 nicks supercoiled plasmids *in vivo* and *in vitro*. (A) Gel electrophoresis of plasmids purified from overnight *E. coli* cultures expressing either acrIIA22, or a null mutant with an early stop codon, or neither. Compared to the null mutant, more plasmid runs in a slowly migrating, open-circle conformation (OC) rather than supercoiled plasmid (SC) with the wild-type *acrIIA22* allele, suggesting that *acrIIA22* may impact plasmid topology. %SC indicates the percentage of DNA in the supercoiled form for each sample. (B) N-terminally His6-tagged AcrIIA22 nicks supercoiled plasmids *in vitro*. (C) C-terminally twin-strep-tagged AcrIIA22 nicks supercoiled plasmids *in vitro* with higher specific activity than shown in panel B (compare protein concentrations). Original, uncropped versions of images depicted in figure may be found in the supporting information file, SI\_raw\_images.

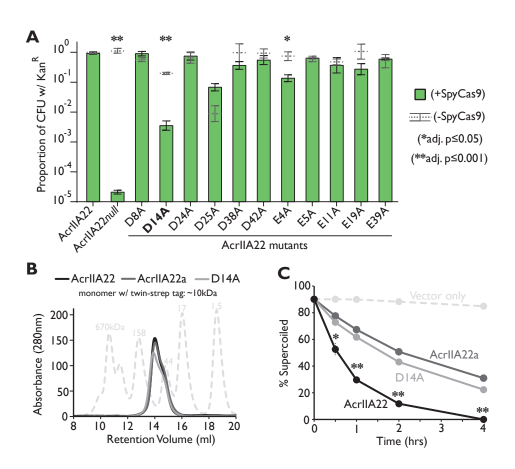


Figure 6. Impaired nicking activity of AcrIIA22 variants in vitro correlates with lower SpyCas9 inhibition in vivo. (A) Alanine mutagenesis of acidic amino acid residues (glutamic acid or aspartic acid) in AcrIIA22 reveals that D14 is important for plasmid protection against SpyCas9. Asterisks depict statistically significant differences in plasmid retention under SpyCas9-inducing and non-inducing conditions, per the legend at right. The D14A mutant is significantly impaired, the E4A mutant is slightly impaired, whereas all other mutants are not impaired for plasmid protection against SpyCas9 compared to an uninduced control. All p-values were corrected for multiple hypotheses using Bonferroni's method (Student's t-test, n=3). (B) AcrIIA22 (black), AcrIIA22a (dark gray), and a D14A mutant (light gray) all elute with similar oligomer profiles via SEC. The dashed trace depicts protein standards of the indicated molecular weight, in kDa. (C) AcrIIA22a and the D14A mutant are impaired for nicking relative to AcrIIA22. All experiments were performed in triplicate, with standard deviations indicated by dashed lines (in most cases, the data points obscure these error bars). Asterisks denote cases where AcrIIA22 is significantly different than both AcrIIA22a and the D14A mutant after correcting for multiple hypotheses (Student's ttest, n=3, Bonferroni correction). A single asterisk (\*) means that adjusted p-values for both comparisons are below 0.05. A double asterisk (\*\*) means that adjusted p-values are both below 0.005. Supplemental Figures 10G and 10H show representative gels for these nicking experiments. The individual numerical values that underlie the summary data in this figure may be found as supporting information file SI Data.

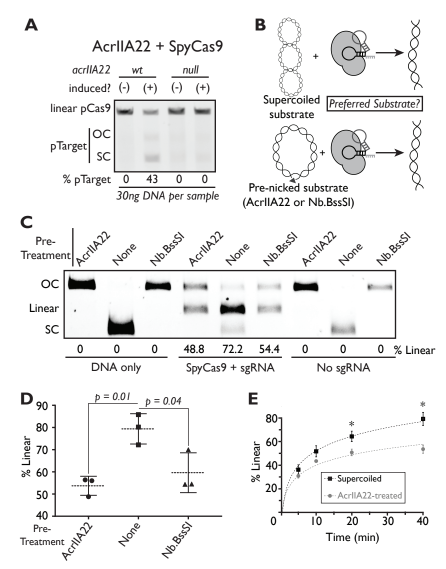
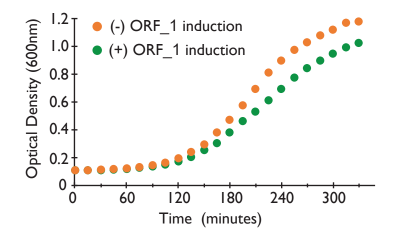
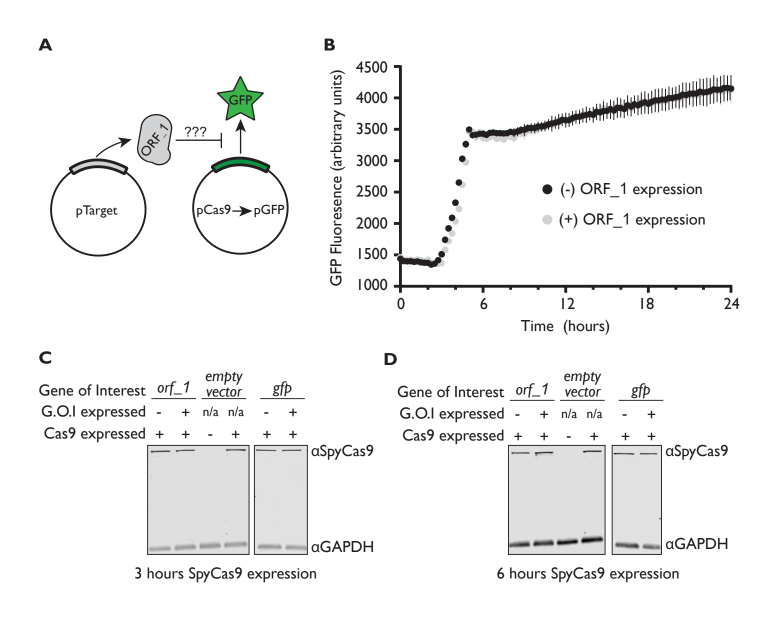


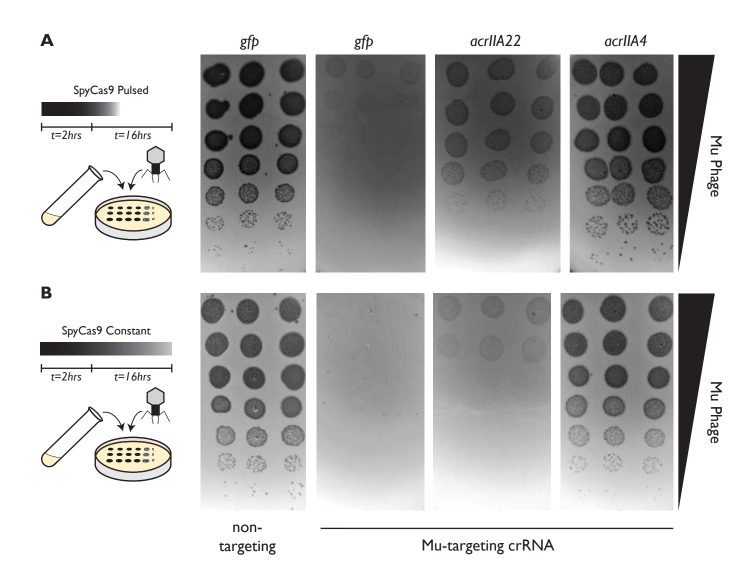
Figure 7. Nicking by AcrIIA22 protects plasmids from SpyCas9 in vivo and in vitro. (A) Gel electrophoresis of plasmids purified from overnight E. coli cultures expressing either wildtype acrIIA22 or a mutant with an early stop codon ('null'). In these cultures, SpyCas9 was expressed from a second plasmid, which was linearized via a unique restriction site before electrophoresis. The acrIIA22-encoding plasmids are indicated with the 'pTarget' label. OC, open-circle; SC, supercoiled. The '%pTarget' figure indicates the fraction of total DNA attributable to pTarget, quantified by densitometry analysis. In cases with complete pTarget elimination, all DNA comes from the SpyCas9 expression plasmid, and thus these bands are more pronounced. However, in the presence of wildtype acrIIA22, pTarget is protected from SpyCas9-mediated cleavage and makes up 43% of total plasmid DNA. (B) We present a schematic of the experimental design for the data depicted in panel C. The experiment tests whether SpyCas9 preferentially cleaves a supercoiled or open-circle plasmid target in vitro. Though both plasmid substrates will be linearized following SpyCas9 cleavage, linear DNA will accumulate more readily with a preferred substrate. (C) Plasmid purifications from overnight cultures were either left unmodified or pre-treated with one of two nickase enzymes, AcrIIA22 or Nb.BssSI, following which each substrate was digested with SpyCas9 in vitro. The percentage of DNA in the linear form is quantified below the gel, which indicates complete SpyCas9 cleavage. Linear, opencircle (OC), and supercoiled (SC) plasmid forms are indicated along with the left of the gel, and reaction components below the gel. SpyCas9 cuts DNA strands sequentially; incomplete digestions with supercoiled substrates produce open-circle plasmids if only one strand has been cleaved (e.g. lane 5). Pre-nicked plasmids, by either AcrIIA22 or Nb.BssSI, are less susceptible to linearization via SpyCas9 cleavage. (D) Endpoint measurements indicate that SpyCas9 more efficiently linearizes supercoiled plasmids than substrates nicked with either AcrIIA22 or Nb.BssSI (Student's t-test, n=3). (E) A time course experiment demonstrates that SpyCas9 more efficiently linearizes supercoiled plasmids than AcrIIA22-treated substrates. An asterisk (\*) denotes significant differences between AcrIIA22-treated and untreated substrates (Student's t-test, p < 0.05, n=3). The individual numerical values and original images for the data presented in this figure may be found in the supporting information files SI Data and SI\_raw\_images, respectively.



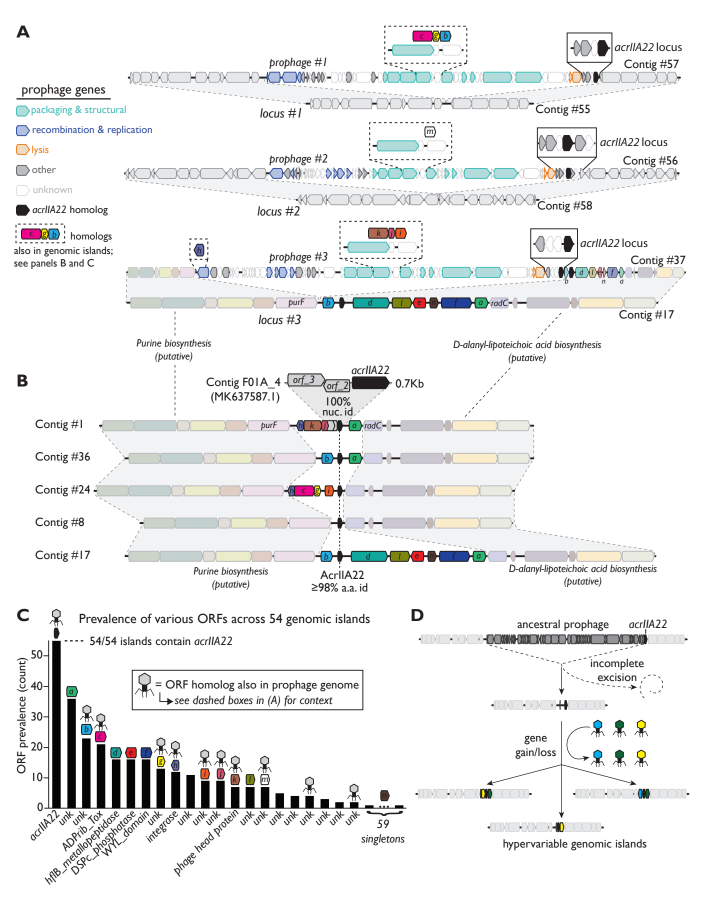
**Supplemental Figure 1**. *Orf\_1 (acrIIA22) confers mild toxicity in E. coli*. Growth rates with *orf\_1* induction (green) are 7% lower than those without *orf\_1* induction (orange). The cfu data shown in Figure 1C were generated from the same experiment depicted here (samples were removed after six hours of growth to determine these cfu counts). Thus, these data demonstrate that anti-SpyCas9 activity occurs under conditions with mild *orf\_1* toxicity. Growth curves are shown for samples without SpyCas9 induction to ensure that *orf\_1* toxicity is not mitigated due to elimination of its plasmid. Points indicate averages from three replicates. Standard deviations at each timepoint are so small that the error bars do not exceed the bounds of the data point.



**Supplemental Figure 2**. *Orf\_1 (acrIIA22)* does not impact SpyCas9 expression. (A) A schematic description of the experimental design shown in panel (B) is presented. If ORF\_1 prevented transcription from pCas9 or altered its copy number, we would expect expression of the *orf\_1* gene to deplete the level of green fluorescence observed from a construct that replaces the *spycas9* gene with *gfp*. (B) Fluorescence measurements for the experiment depicted in panel A show that ORF\_1 does not impact GFP expression throughout an *E. coli* growth curve. Points indicate averages from three replicates, error bars indicate standard deviation. A western blot shows no depletion of SpyCas9 expression as a function of ORF\_1 or GFP expression in growing *E. coli* cultures at three hours (C) or six hours (D). As an internal control, GAPDH expression was also detected.

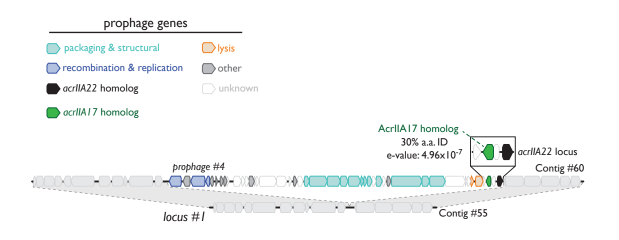


**Supplemental Figure 3**. AcrIIA22 only modestly protects Mu phages against SpyCas9. Mu phage fitness was measured by plaquing on *E. coli* in the presence of *gfp*, *acrIIA22*, or *acrIIA4* via serial ten-fold dilutions. Bacterial clearing (black) occurs when phage Mu overcomes SpyCas9 immunity and lyses *E. coli*. In (A) and in (B), SpyCas9 with a Mu-targeting crRNA confers substantial protection against phage Mu relative to a non-targeting (n.t.) control, in both conditions tested. These conditions are depicted at left, with the only difference being whether SpyCas9 was only expressed in liquid growth prior to phage infection (panel A) or expressed both in liquid media and in solid media throughout infection (panel B). When expressed from a second plasmid, the positive control *acrIIA4* significantly enhances Mu fitness by inhibiting SpyCas9 in all conditions *in trans*. Though *acrIIA22* confers protection against SpyCas9 compared to *gfp* (negative control), this effect is milder than with *acrIIA4* and dependent on SpyCas9 expression.

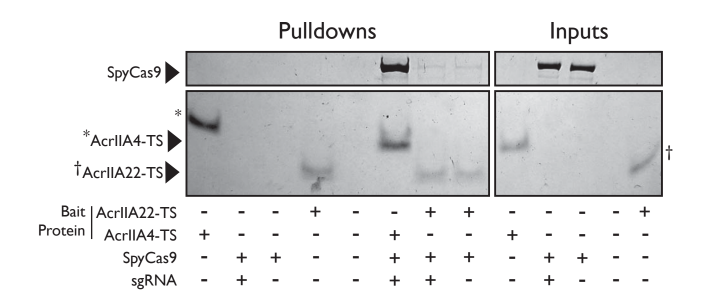


Supplemental Figure 4.

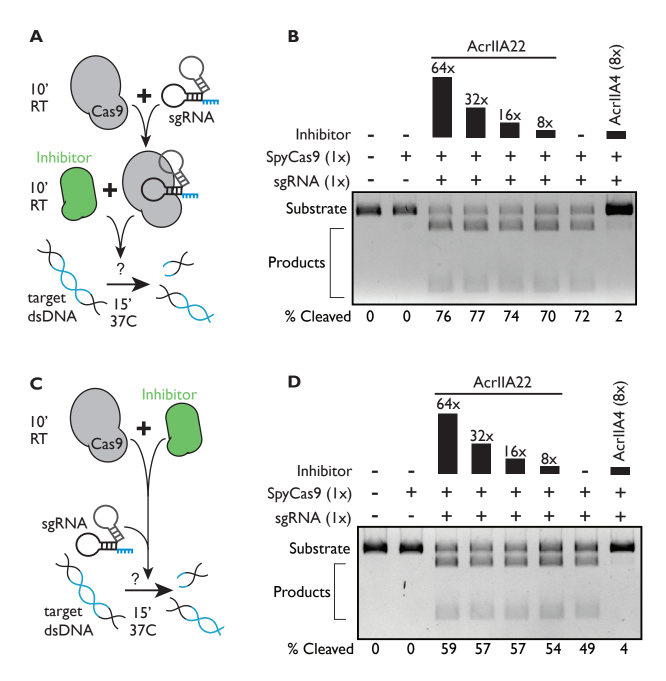
Supplemental Figure 4. AcrIIA22 homologs are found in hypervariable regions of prophage and bacterial genomes in the CAG-217 clostridial genus. (A) Homologs of acrIIA22 are depicted in three related prophage genomes, integrated at three different genomic loci, revealed by a comparison of prophage-bearing contigs (#57, #56, #37) relative to unintegrated contigs (#55, #58, #17 respectively), which are otherwise nearly identical. Prophage genes are colored by functional category, according to the legend at the left of panel A. Genes immediately adjacent to acrIIA22 (solid boxes) vary across phages, despite strong relatedness across much of the prophage genomes. Bacterial genes are colored gray, except for contig #17, which is also depicted in panel B, below. (B) Homologs of acrIIA22 are depicted in diverse genomic islands, including Contig #1, whose sequence includes a portion that is identical to the original metagenomic contig we recovered (F01A 4). All acrIIA22 homologs in these loci are closely related but differ in their adjacent genes, which often have homologs in the prophages depicted in panel A (dashed boxes). Bacterial genomic regions flanking these hypervariable islands are nearly identical to one another and to prophage integration locus #3, as shown by homology to contig #17 from panel A. Contigs are numbered to indicate their descriptions in Supplemental Table 3, which contains their metadata, taxonomy, and sequence retrieval information. All sequences and annotations can also be found in Supplemental Datasets 1 and 2. (C) We tabulate the prevalence of various protein families (clustered at 65% amino acid identity) in a set of 54 unique genomic islands. Each of these islands is flanked by the conserved genes purF and radC but contains a different arrangement of encoded genes. Domain-level annotations are indicated below each protein family (unk; unknown function). Gene symbols above each protein family are colored and lettered to indicate their counterparts or homologs in panels A and B. The phage capsid icon indicates sequences with homologs in prophage genomes. (D) An evolutionary model for the origin of the *acrIIA22*-encoding hypervariable genomic islands depicted in panel B is shown. This panel is reprinted from Figure 2C, for continuity. We propose that acrIIA22 moved via a phage insertion into a bacterial genomic locus, remained following an incomplete prophage excision event, and its neighboring genes subsequently diversified via horizontal exchange with additional phage genomes.



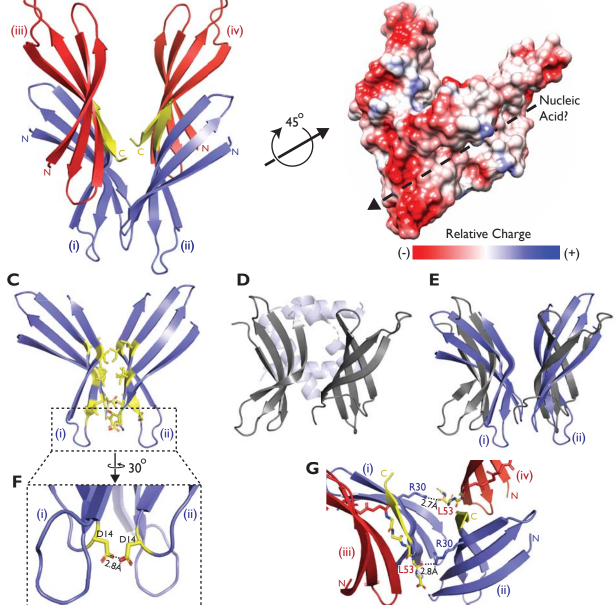
**Supplemental Figure 5**. **Genomic proximity of** *acrllA22* **homologs to other** *acr* **genes.** An *acrllA22*-encoding prophage like the one depicted in Figure 2A and those in Supplemental Figure 4A is shown. This prophage encodes for a homolog of the previously described SpyCas9 inhibitor *acrllA17* within one kilobase of an *acrllA22* homolog. Sequence relatedness between the depicted *acrllA17* gene and the originally discovered acrllA17 is shown<sup>22</sup>. Because phages often encode multiple *acrs* in the same locus, the co-localization of *acrllA17* with *acrllA22* is consistent with the latter gene functioning natively to inhibit CRISPR-Cas activity. Prophage genes are colored by functional category, per the legend and as in Supplemental Figure 4A. Contigs are numbered to indicate their descriptions in Supplemental Table 3, which contains their metadata, taxonomy, and sequence retrieval information. All sequences and annotations can also be found in Supplemental Datasets 1 and 2.



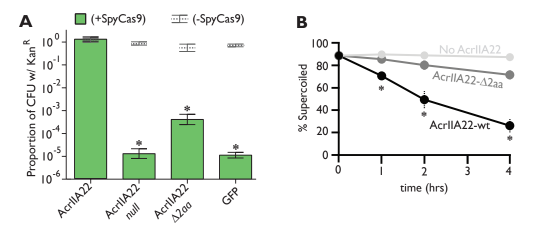
**Supplemental Figure 6**. AcrIIA22 does not strongly bind SpyCas9. SpyCas9 and sgRNA were pre-incubated before mixing with a twin-strep (TS) tagged AcrIIA22 or AcrIIA4. SpyCas9 without sgRNA was also used. Strep-Tactin pulldowns on AcrIIA4 also pulled down SpyCas9 pre-incubated with sgRNA, as previously reported<sup>12</sup>. Similar pulldowns with AcrIIA22 indicate little to no interaction with SpyCas9, regardless of whether sgRNA was used. These images depict total protein content visualized by Coomassie stain. Reaction components are indicated below the gel image. Asterisks (\*) and dagger (†) symbols indicate AcrIIA4 and AcrIIA22 protein bands that run at slightly different positions than expected due to gel distortion.



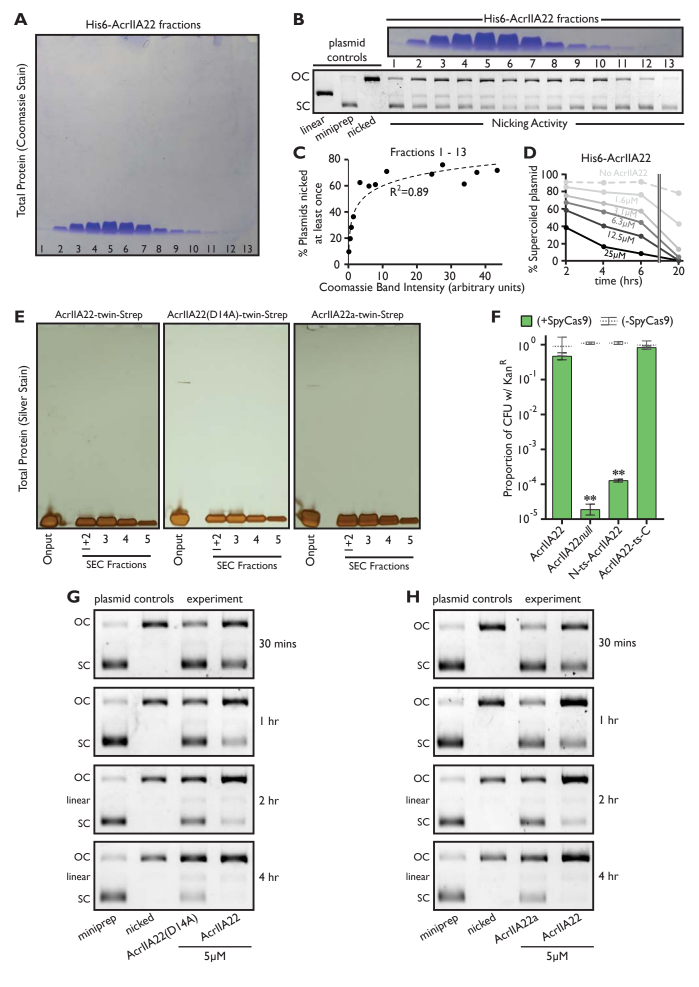
Supplemental Figure 7. AcrIIA22 does not protect linear DNA from SpyCas9 cleavage. (A) A schematic cartoon depicts the experiment in panel (B). SpyCas9 was pre-incubated with sgRNA targeting linear DNA. Then, Acr candidates were added. Subsequently, cleavage reactions were performed, and the DNA products visualized by gel electrophoresis. (B) We show the products of the reactions described in panel A for the inhibitors AcrIIA22 and AcrIIA4. SpyCas9 activity is greatly inhibited by AcrIIA4 but unaffected by AcrIIA22, as indicated by the proportion of cleaved DNA product. Reaction components are depicted atop the gel image, with molar equivalents relative to SpyCas9 indicated. The percent of DNA substrate cleaved by SpyCas9 is quantified below each lane. (C) We perform a similar experiment as in panel A, except candidate Acrs were incubated with SpyCas9 before sqRNA addition. Reactions were begun via the simultaneous addition of sgRNA and linear dsDNA instead of just dsDNA. (D) The products of the reactions described in panel C for AcrIIA22 and AcrIIA4 inhibitors are shown. SpyCas9 activity is inhibited by AcrIIA4 but unaffected by AcrIIA22, as indicated by the proportion of cleaved DNA product. The data depicted in this figure are not directly comparable to those in figure 7, due to methodological differences and because the preparations of SpyCas9 used in each experiment exhibited different activities.



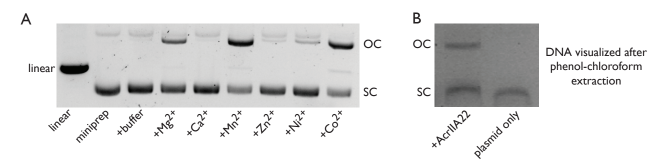
Supplemental Figure 8. AcrIIA22 resembles a PC4-like protein. (A) We present a ribbon diagram of a proposed AcrIIA22 tetramer, which requires binding between anti-parallel β-strands at the C-termini of AcrIIA22 monomers to form extended, concave  $\beta$ -sheets. This putative oligomerization interface is indicated by the regions highlighted in yellow. Each monomer in the proposed tetramer is labeled with lower-case Roman numerals (i-iv). (B) Space filling model of the tetrameric AcrIIA22 structure from panel A, with relative charge depicted, highlighting a groove (dashed line with arrowhead) that may accommodate nucleic acids (based on analogy to other PC4-like proteins). (C) AcrIIA22 monomers (i) and (ii) from the tetramer in panel A likely interact via a series of hydrophobic interactions, as indicated by the predominantly non-polar sidechains colored in yellow. The boxed region highlights residue D14, which is important for nicking activity and plasmid protection against SpyCas9, and is enlarged in panel F. (D) In conventional PC4-like family proteins, such as the putative single-stranded DNA binding protein from phage T5 depicted in gray (PDB:4BG7), the same topology of outward facing, concave  $\beta$ -sheets are instead stabilized via interactions between opposing  $\alpha$ -helices (depicted in opaque light blue). (E) An overlay of  $\beta$ -sheets from AcrIIA22 (blue, PDB:7JTA) and the phage T5 PC4-like protein (gray, PDB:4BG7) illustrates their similar topologies. (F) Two D14 residues in loop regions of AcrIIA22 monomers (i) and (ii) are in close proximity. These residues are important for nicking activity and may bind divalent cations in cells under physiological pH. (G) A close view of a putative salt bridge between R30 of monomers (i) / (ii) and the peptide backbone of the C-terminus of monomers (iv) / (iii), respectively. AcrIIA22 monomers are colored as described in panel A.



Supplemental Figure 9. A 2-aa truncation mutant of AcrIIA22 is impaired for SpyCas9 inhibition and nicking activity. (A) An *in vivo* plasmid protection assay. Asterisks depict statistically significant differences in plasmid retention under SpyCas9-inducing conditions with either wild-type AcrIIA22, a null mutant with an early stop codon, a 2-aa truncation, or a negative control *gfp* gene (adj. p < 0.005, Student's t-test, n=3). The truncation mutant retains mild but severely impaired activity, as it protects a plasmid from SpyCas9 more effectively than a null mutant (p = 0.012) or GFP control (p = 0.015). All p-values were corrected for multiple hypotheses using Bonferroni's method. (B) The 2-aa truncation mutant is impaired for nicking *in vitro*, relative to wild-type AcrIIA22. In both cases,  $25\mu$ M of protein was used following NiNTA-based purification of an N-terminal, His6-tagged construct. An asterisk (\*) denotes significant differences between AcrIIA22-treated and untreated substrates (Student's t-test, p < 0.05, n=3). Standard deviations are indicated by dashed lines (in most cases, the data points obscure these error bars).



Supplemental Figure 10. AcrIIA22 nicks supercoiled plasmids. (A) A Coomassie stain of an N-terminally His6-tagged AcrIIA22 construct shows no co-purifying proteins. (B) The nicking activity for this protein preparation (bottom) correlates with the intensity of the Coomassie-stained protein band across purification fractions (top). In each lane, supercoiled (SC) plasmid DNA represents the un-nicked fraction whereas open circle (OC) and linear DNA have been nicked at least once. (C) This panel is a quantification of the experiment depicted in panel B across all 13 fractions collected. (D) His6-AcrIIA22 nicks supercoiled plasmids in a time and concentration dependent manner. A decrease in the proportion of supercoiled plasmid DNA indicates nicking activity, as depicted in Figure 5B. (E) A silver stain of a C-terminally twin-strep-tagged AcrIIA22 construct shows no co-purifying proteins. Equal volumes of each protein fraction were loaded in each lane, for all samples. Fraction 4 was concentrated and used for all *in vitro* experiments. (F) A C-terminal, but not N-terminal twin-strep tag is compatible with AcrIIA22's ability to protect a target plasmid from SpyCas9 elimination in vivo. Statistically significant differences in plasmid retention between SpyCas9-inducing and non-inducing conditions were determined via a Student's t-test (n=3); '\*\*' indicates p≤0.001. All p-values were adjusted for multiple hypotheses using the Bonferroni correction. (G) The D14A mutation in AcrIIA22 impairs nicking activity. Over time, the wild-type AcrIIA22-twin-strep construct consistently converts a higher fraction of plasmid DNA from its supercoiled (SC) form to an open-circle (OC) conformation than a D14A mutant. Control plasmids include a miniprepped sample and sample pre-treated with the commercial nickase, Nb.BssSI. Reaction times are indicated to the right of each image. (H) AcrIIA22a (Figure 3B) is impaired for nicking activity relative to AcrIIA22. As in panel G, both constructs were purified via C-terminal twin-strep tags.



**Supplemental Figure 11. Divalent cations influence AcrIIA22's nicking activity.** (**A**) We present the impact of different divalent cations on AcrIIA22's nicking activity, which is highest with Mg<sup>2+</sup>, Mn<sup>2+</sup>, and Co<sup>2+</sup>. OC, open-circle plasmid form. SC, supercoiled plasmid. (**B**) The open-circle plasmid product persists through phenol-chloroform extraction following AcrIIA22 treatment, indicating that it directly results from AcrIIA22's nicking activity.

Acr	Binds cognate Cas protein?	Inhibit as pure proteins?	References			
AcrIIA1	Yes	No	(Osuna et al., 2020)			
AcrIIA2	Yes	Yes	(Jiang et al., 2019; Liu et al., 2019)			
AcrIIA3	unknown	unknown	(Rauch et al., 2017)			
AcrIIA4	Yes	Yes	(Dong et al., 2017; Shin et al., 2017; Yang and Patel, 2017)			
AcrIIA5	Yes	Yes	(An et al., 2020; Garcia et al., 2019; Song et al., 2019)			
AcrIIA6	Yes	Yes	(Fuchsbauer et al., 2019)			
AcrIIA7	No	Yes	(Uribe et al., 2019)			
AcrIIA8	Yes	Yes	(Uribe et al., 2019)			
AcrIIA9	Yes	Yes	(Uribe et al., 2019)			
AcrIIA10	Yes	Yes	(Uribe et al., 2019)			
AcrIIA11	Yes	Yes	(Forsberg et al., 2019)			
AcrIIA12	probable	Yes	(Eitzinger et al., 2020; Osuna et al., 2020)			
AcrIIA13	unknown	Yes	(Watters et al., 2020)			
AcrIIA14	unknown	Yes	(Watters et al., 2020)			
AcrIIA15	unknown	Yes	(Watters et al., 2020)			
AcrIIA16	Yes	Yes	(Mahendra et al., 2020)			
AcrIIA17	Yes	No	(Mahendra et al., 2020)			
AcrIIA18	Yes	No	(Mahendra et al., 2020)			
AcrIIA19	Yes	No	(Mahendra et al., 2020)			
AcrIIA20	unknown	Yes	(Eitzinger et al., 2020)			
AcrIIA21	unknown	Yes	(Eitzinger et al., 2020)			
AcrIIA22	No	No	This study			
AcrIIA23	unknown	unknown	(Varble et al., 2020)			
AcrIIC1	Yes	Yes	(Pawluk et al., 2016)			
AcrIIC2	Yes	Yes	(Pawluk et al., 2016)			
AcrIIC3	Yes	Yes	(Pawluk et al., 2016)			
AcrIIC4	Yes	Yes	(Lee et al., 2018)			
AcrIIC5	Yes	Yes	(Lee et al., 2018)			
AcrVA1	Yes	Yes	(Knott et al., 2019b; Watters et al., 2018; Zhang et al., 2019)			
AcrVA2	unknown	unknown	(Marino et al., 2018)			
AcrVA3	unknown	unknown	(Marino et al., 2018)			

Supplemental Table 1. Whether all known anti-CRISPRs can bind Cas proteins or inhibit their cleavage activity as purified proteins.

AcrVA4	Yes	Yes	(Knott et al., 2019a; Knott et al., 2019b; Watters et al., 2018; Zhang et al., 2019)
AcrVA5	transiently	Yes	(Knott et al., 2019b; Watters et al., 2018; Zhang et al., 2019)
AcrVIA1(Lse)	Yes	Yes	(Meeske et al., 2020)
AcrVIA1(Lwa)	Yes	unknown	(Lin et al., 2020)
AcrVIA2	Yes	unknown	(Lin et al., 2020)
AcrVIA3	Yes	unknown	(Lin et al., 2020)
AcrVIA4	Yes	unknown	(Lin et al., 2020)
AcrVIA5	Yes	unknown	(Lin et al., 2020)
AcrVIA6	Yes	unknown	(Lin et al., 2020)
AcrVIA7	unknown	unknown	(Lin et al., 2020)
AcrIB1	unknown	unknown	(Lin et al., 2020)
AcrIC1	unknown	unknown	(Leon et al., 2020)
AcrIC2	probable	unknown	(Leon et al., 2020)
AcrIC3	unknown	unknown	(Leon et al., 2020)
AcrIC4	probable	unknown	(Leon et al., 2020)
AcrIC5	probable	unknown	(Leon et al., 2020)
AcrIC6	unknown	unknown	(Leon et al., 2020)
AcrIC7	probable	unknown	(Leon et al., 2020)
AcrIC8	probable	unknown	(Leon et al., 2020)
AcrID1	Yes	unknown	(He et al., 2018)
AcrIE1	Yes	unknown	(Pawluk et al., 2017)
AcrIE2	unknown	unknown	(Pawluk et al., 2014)
AcrIE3	probable	unknown	(Stanley, 2018)
AcrIE4	unknown	unknown	(Pawluk et al., 2014)
AcrIE5	unknown	unknown	(Pawluk et al., 2014)
AcrIE6	unknown	unknown	(Pawluk et al., 2014)
AcrIE7	unknown	unknown	(Pawluk et al., 2014)
AcrIE4-IF7	unknown	unknown	(Marino et al., 2018)
AcrIE8	unknown	unknown	(Pinilla-Redondo et al., 2020)
AcrIF1	Yes	unknown	(Bondy-Denomy et al., 2015; Chowdhury et al., 2017; Guo et al., 2017)
AcrIF2	Yes	unknown	(Bondy-Denomy et al., 2015; Chowdhury et al., 2017; Guo et al., 2017)
AcrIF3	Yes	unknown	(Bondy-Denomy et al., 2015; Wang et al., 2016a; Wang et al., 2016b)
AcrIF4	Yes	unknown	(Bondy-Denomy et al., 2015)
AcrIF5	unknown	unknown	(Bondy-Denomy et al., 2013)

AcrIF6	Yes	Yes	(Zhang et al., 2020)
AcrIF7	Yes	unknown	(Hirschi et al., 2020)
AcrIF8	Yes	Yes	(Zhang et al., 2020)
AcrIF9	Yes	Yes	(Hirschi et al., 2020; Zhang et al., 2020)
AcrIF10	Yes	unknown	(Guo et al., 2017)
AcrIF11	unknown	unknown	(Marino et al., 2018)
AcrIF12	unknown	unknown	(Marino et al., 2018)
AcrIF13	unknown	unknown	(Marino et al., 2018)
AcrIF14	unknown	unknown	(Marino et al., 2018)
AcrIF15	probable	unknown	(Pinilla-Redondo et al., 2020)
AcrIF16	unknown	unknown	(Pinilla-Redondo et al., 2020)
AcrIF17	unknown	unknown	(Pinilla-Redondo et al., 2020)
AcrIF18	probable	unknown	(Pinilla-Redondo et al., 2020)
AcrIF19	unknown	unknown	(Pinilla-Redondo et al., 2020)
AcrIF20	unknown	unknown	(Pinilla-Redondo et al., 2020)
AcrIF21	unknown	unknown	(Pinilla-Redondo et al., 2020)
AcrIF22	unknown	unknown	(Pinilla-Redondo et al., 2020)
AcrIF23	unknown	unknown	(Pinilla-Redondo et al., 2020)
AcrIF24	unknown	unknown	(Pinilla-Redondo et al., 2020)
AcrIII-1	No (degrades	No	(Athukoralage et al., 2020)
	CA4 second		
AcrIIIB1	messenger) Yes	unknown	(Bhoobalan-Chitty et al., 2019)
ACHIDT	162	UTIKITUWIT	

	Structural Homolog	Function		Similarity	to AcrIIA	.22
PDBID	Name	DNA/RNA Binding*	Zscore	r.m.s.d.	n-align	% A.A. ID
4bg7	PC4 putative transcriptional coactivator p15	DNA	6.2	2.5	54	15
3k44	<i>D. melanogaster</i> Pur-α	DNA/RNA	5.9	2.6	47	9
5fgp	Pur-α repeat I and II from <i>D. melanogaster</i>	DNA/RNA	5.6	2.1	48	8
3n8b	Pur-α from <i>B. burgdorferi</i>	DNA/RNA	5	2.8	48	6
2gje	Mitochondrial RNA Binding Protein (T. brucei)	RNA	4.9	2.5	52	8
5zkl	Protein of unknown function SP_0782, S. pneumoniae	DNA	4.7	3.6	52	12
5fgo	<i>D. melanogaster</i> Pur-α repeat III	No info	4.5	2.7	44	14
1pcf	Replication & transcription cofactor PC4 CTD	DNA	4.5	2.5	45	7
2ltt	Putative Uncharacterized Protein YDBC	DNA	4.5	2.8	50	12
4bhm	MoSub1-DNA PC4 transcription cofactor	DNA	3.9	2.8	45	4
3cm1	SSGA-like sporulation specific cell division protein	No info	2.8	3.7	47	13
1I3a	Transcription factor PBF-2 (P24, WHY1)	DNA	2.8	5	48	8
4ntq	Anti-toxin Cdil, <i>E. cloacae</i>	No info	2.7	3	49	12
3n1k	WHY2 transcription factor, S. tuberosum	DNA	2.6	2.8	52	4

Supplemental Table 2. PC4-like proteins with structural homology to AcrIIA22

\*RNA/DNA binding data from (Janowski and Niessing, 2020).

**Supplemental Table 3.** All sequences used in this study. Sequence names and databases are indicated. All sequences and annotations are also available as supplemental data. Sequences retrieved from Pasolli *et al.* refer to the following study: (Pasolli et al., 2019).

Contig No.	Sequence Name	How Used?	NCBI Nuc. ID	Pasolli et al SGB?	Pasolli et al Raw Assembly?	Pasolli Reconstructed Genome Name	SGB id	GTDB Taxonomy	External Data Available at:
1	4303_LiJ_2014V1.UC63- 0bin.67_NODE_112_leng th_95405_cov_4.60675	Figure 2, find gene functions from 54 unique genomic loci; Figure 3 Acr Seq	n/a	Yes	No	LiJ_2014V1. UC63- 0bin.67	4303	d_Bacteria; p_Firmicutes_A; c_Clostridia; o_Oscillospirales; f_Acutalibacteraceae; g_CAG-217	http://segatalab.cibio .unitn.it/data/Pasolli_ et_al.html
2	Bengtsson- PalmeJ_2015TRAVELRE S9_NODE_4_length_53858 0_cov_9.43148	find gene functions from 54 unique genomic loci	n/a	No	Yes	n/a	n/a	d_Bacteria; p_Firmicutes_A; c_Clostridia; o_Oscillospirales; f_Acutalibacteraceae; g_CAG-217 (inferred)	http://segatalab.cibio .unitn.it/data/Pasolli_ et_al.html
3	ChengpingW_2017AS9ra w_NODE_922_length_2766 4_cov_3.49089	find gene functions from 54 unique genomic loci	n/a	No	Yes	n/a	n/a	d_Bacteria; p_Firmicutes_A; c_Clostridia; o_Oscillospirales; f_Acutalibacteraceae; g_CAG-217 (inferred)	http://segatalab.cibio .unitn.it/data/Pasolli_ et_al.html
4	CosteaPI_2017SID713B0 25-11-0- 0_NODE_4_length_351620 _cov_7.46108	find gene functions from 54 unique genomic loci	n/a	No	Yes	n/a	n/a	d_Bacteria; p_Firmicutes_A; c_Clostridia; o_Oscillospirales; f_Acutalibacteraceae; g_CAG-217 (inferred)	http://segatalab.cibio .unitn.it/data/Pasolli_ et_al.html
5	BritoIL_2016M1.64.ST_N ODE_47_length_140472_co v_9.49805	find gene functions from 54 unique genomic loci; Figure 3 Acr Seq	n/a	No	Yes	n/a	n/a	d_Bacteria; p_Firmicutes_A; c_Clostridia; o_Oscillospirales; f_Acutalibacteraceae; g_CAG-217 (inferred)	http://segatalab.cibio .unitn.it/data/Pasolli_ et_al.html
6	BritoIL_2016M2.57.ST_N ODE_3_length_405636_cov _14.0428	find gene functions from 54 unique genomic loci	n/a	No	Yes	n/a	n/a	d_Bacteria; p_Firmicutes_A; c_Clostridia; o_Oscillospirales; f_Acutalibacteraceae; g_CAG-217 (inferred)	http://segatalab.cibio .unitn.it/data/Pasolli_ et_al.html
7	BritoIL_2016WL.14.ST_N ODE_13_length_259523_co v_10.8408	find gene functions from 54 unique genomic loci	n/a	No	Yes	n/a	n/a	d_Bacteria; p_Firmicutes_A; c_Clostridia; o_Oscillospirales; f_Acutalibacteraceae; g_CAG-217 (inferred)	http://segatalab.cibio .unitn.it/data/Pasolli_ et_al.html
8	ChengpingW_2017AS67r aw_NODE_2_length_43917 7_cov_9.00174	Figure 2, find gene functions from 54 unique genomic loci; Figure 3 Acr Seq	n/a	No	Yes	n/a	n/a	d_Bacteria; p_Firmicutes_A; c_Clostridia; o_Oscillospirales; f_Acutalibacteraceae; g_CAG-217 (inferred)	http://segatalab.cibio .unitn.it/data/Pasolli_ et_al.html
9	CM_madagascarA90_04 _1FE_NODE_125_length_8 1453_cov_9.00904	find gene functions from 54 unique genomic loci	n/a	No	Yes	n/a	n/a	d_Bacteria; p_Firmicutes_A; c_Clostridia; o_Oscillospirales; f_Acutalibacteraceae; g_CAG-217 (inferred)	http://segatalab.cibio .unitn.it/data/Pasolli_ et_al.html
10	CM_madagascarV12_01 _2FE_NODE_5_length_202 628_cov_9.50435	find gene functions from 54 unique genomic loci	n/a	No	Yes	n/a	n/a	d_Bacteria; p_Firmicutes_A; c_Clostridia; o_Oscillospirales; f_Acutalibacteraceae; g_CAG-217 (inferred)	http://segatalab.cibio .unitn.it/data/Pasolli_ et_al.html
11	CosteaPI_2017SID713A0 46-11-0- 0_NODE_322_length_6900 0_cov_4.32987	find gene functions from 54 unique genomic loci; Figure 3 Acr Seq	n/a	No	Yes	n/a	n/a	d_Bacteria; p_Firmicutes_A; c_Clostridia; o_Oscillospirales; f_Acutalibacteraceae; g_CAG-217 (inferred)	http://segatalab.cibio .unitn.it/data/Pasolli_ et_al.html
12	CosteaPI_2017SID713A0 45-11-0- 0_NODE_78_length_64886 _cov_3.97493	find gene functions from 54 unique genomic loci	n/a	No	Yes	n/a	n/a	d_Bacteria; p_Firmicutes_A; c_Clostridia; o_Oscillospirales; f_Acutalibacteraceae; g_CAG-217 (inferred)	http://segatalab.cibio .unitn.it/data/Pasolli_ et_al.html
13	CosteaPI_2017SID713A0 04-11-0- 0_NODE_1_length_647860 _cov_14.1013	find gene functions from 54 unique genomic loci; Figure 3 Acr Seq	n/a	No	Yes	n/a	n/a	d_Bacteria; p_Firmicutes_A; c_Clostridia; o_Oscillospirales; f_Acutalibacteraceae; g_CAG-217 (inferred)	http://segatalab.cibio .unitn.it/data/Pasolli_ et_al.html
14	CosteaPI_2017peacemak er-11-60-	find gene functions from 54 unique genomic loci	n/a	No	Yes	n/a	n/a	d_Bacteria; p_Firmicutes_A; c_Clostridia; o_Oscillospirales; f_Acutalibacteraceae; g_CAG-217 (inferred)	http://segatalab.cibio .unitn.it/data/Pasolli_ et_al.html

	0_NODE_48_length_49378 _cov_15.5445								
15	CosteaPI_2017SID713A0 63-11-90- 0_NODE_2082_length_169 60_cov_2.98527	find gene functions from 54 unique genomic loci	n/a	No	Yes	n/a	n/a	d_Bacteria; p_Firmicutes_A; c_Clostridia; o_Oscillospirales; f_Acutalibacteraceae; g_CAG-217 (inferred)	http://segatalab.cibio .unitn.it/data/Pasolli_ et_al.html
16	CosteaPI_2017SID713A0 88-11-0- 0_NODE_89_length_11329 7_cov_5.77445	find gene functions from 54 unique genomic loci; Figure 3 Acr Seq	n/a	No	Yes	n/a	n/a	d_Bacteria; p_Firmicutes_A; c_Clostridia; o_Oscillospirales; f_Acutalibacteraceae; g_CAG-217 (inferred)	http://segatalab.cibio .unitn.it/data/Pasolli_ et_al.html
17	CosteaPI_2017SID713A0 62-11-0- 0_NODE_38_length_19219 6_cov_4.03099	Find gene functions from 54 unique genomic loci	n/a	No	Yes	n/a	n/a	d_Bacteria; p_Firmicutes_A; c_Clostridia; o_Oscillospirales; f_Acutalibacteraceae; g_CAG-217 (inferred)	http://segatalab.cibio .unitn.it/data/Pasolli_ et_al.html
18	CosteaPI_2017SID713B0 51-11-0- 0_NODE_14_length_29861 9_cov_7.20988	find gene functions from 54 unique genomic loci	n/a	No	Yes	n/a	n/a	d_Bacteria; p_Firmicutes_A; c_Clostridia; o_Oscillospirales; f_Acutalibacteraceae; g_CAG-217 (inferred)	http://segatalab.cibio .unitn.it/data/Pasolli_ et_al.html
19	FengQ_2015SID31872_N ODE_2_length_392843_cov _5.93617	find gene functions from 54 unique genomic loci	n/a	No	Yes	n/a	n/a	d_Bacteria; p_Firmicutes_A; c_Clostridia; o_Oscillospirales; f_Acutalibacteraceae; g_CAG-217 (inferred)	http://segatalab.cibio .unitn.it/data/Pasolli_ et_al.html
20	FengQ_2015SID530258_ NODE_5_length_350476_c ov_17.595	find gene functions from 54 unique genomic loci; Figure 3 Acr Seq	n/a	No	Yes	n/a	n/a	d_Bacteria; p_Firmicutes_A; c_Clostridia; o_Oscillospirales; f_Acutalibacteraceae; g_CAG-217 (inferred)	http://segatalab.cibio .unitn.it/data/Pasolli_ et_al.html
21	FengQ_2015SID530373_ NODE_21_length_272157_ cov_9.73468	find gene functions from 54 unique genomic loci	n/a	No	Yes	n/a	n/a	d_Bacteria; p_Firmicutes_A; c_Clostridia; o_Oscillospirales; f_Acutalibacteraceae; g_CAG-217 (inferred)	http://segatalab.cibio .unitn.it/data/Pasolli_ et_al.html
22	HeQ_2017SZAXPI02956 1- 52_NODE_1_length_50275 2_cov_8.09488	find gene functions from 54 unique genomic loci; Figure 3 Acr Seq	n/a	No	Yes	n/a	n/a	d_Bacteria; p_Firmicutes_A; c_Clostridia; o_Oscillospirales; f_Acutalibacteraceae; g_CAG-217 (inferred)	http://segatalab.cibio .unitn.it/data/Pasolli_ et_al.html
23	HeQ_2017SZAXPI02957 5- 90_NODE_229_length_949 18_cov_3.79903	find gene functions from 54 unique genomic loci	n/a	No	Yes	n/a	n/a	d_Bacteria; p_Firmicutes_A; c_Clostridia; o_Oscillospirales; f_Acutalibacteraceae; g_CAG-217 (inferred)	http://segatalab.cibio .unitn.it/data/Pasolli_ et_al.html
24	KarlssonFH_2013S463_ NODE_1_length_570037_c ov_16.3973	Figure 2, find gene functions from 54 unique genomic loci	n/a	No	Yes	n/a	n/a	d_Bacteria; p_Firmicutes_A; c_Clostridia; o_Oscillospirales; f_Acutalibacteraceae; g_CAG-217 (inferred)	http://segatalab.cibio .unitn.it/data/Pasolli_ et_al.html
25	LiJ_201402.UC12- 1_NODE_323_length_4999 5_cov_5.04395	find gene functions from 54 unique genomic loci	n/a	No	Yes	n/a	n/a	d_Bacteria; p_Firmicutes_A; c_Clostridia; o_Oscillospirales; f_Acutalibacteraceae; g_CAG-217 (inferred)	http://segatalab.cibio .unitn.it/data/Pasolli_ et_al.html
26	LiJ_2014V1.FI02_NODE_ 274_length_84286_cov_3.4 9253	find gene functions from 54 unique genomic loci	n/a	No	Yes	n/a	n/a	d_Bacteria; p_Firmicutes_A; c_Clostridia; o_Oscillospirales; f_Acutalibacteraceae; g_CAG-217 (inferred)	http://segatalab.cibio .unitn.it/data/Pasolli_ et_al.html
27	LiJ_2017H1M413815_NO DE_71_length_81514_cov_ 18.301	find gene functions from 54 unique genomic loci	n/a	No	Yes	n/a	n/a	d_Bacteria; p_Firmicutes_A; c_Clostridia; o_Oscillospirales; f_Acutalibacteraceae; g_CAG-217 (inferred)	http://segatalab.cibio .unitn.it/data/Pasolli_ et_al.html
28	LiJ_2017H2M514909_NO DE_68_length_69076_cov_ 10.283	find gene functions from 54 unique genomic loci	n/a	No	Yes	n/a	n/a	d_Bacteria; p_Firmicutes_A; c_Clostridia; o_Oscillospirales; f_Acutalibacteraceae; g_CAG-217 (inferred)	http://segatalab.cibio .unitn.it/data/Pasolli_ et_al.html
29	LiuW_2016SRR3992969 _NODE_1149_length_1899 _9_cov_8.45033	find gene functions from 54 unique genomic loci; Figure 3 Acr Seq	n/a	No	Yes	n/a	n/a	d_Bacteria; p_Firmicutes_A; c_Clostridia; o_Oscillospirales; f_Acutalibacteraceae; g_CAG-217 (inferred)	http://segatalab.cibio .unitn.it/data/Pasolli_ et_al.html

	LiuW_2016SRR3992984	find gone functions						dBacteria; pFirmicutes_A; cClostridia;	http://segatalab.cibio
30		find gene functions	n/a	No	Yes	2/2	2/2		
30	_NODE_127_length_61384	from 54 unique	n/a	INO	res	n/a	n/a	o_Oscillospirales; f_Acutalibacteraceae;	.unitn.it/data/Pasolli_
	_cov_18.0593	genomic loci						g_CAG-217 (inferred)	et_al.html
	LiuW_2016SRR3993014	find gene functions						d_Bacteria; p_Firmicutes_A; c_Clostridia;	http://segatalab.cibio
31	_NODE_8_length_143441_	from 54 unique	n/a	No	Yes	n/a	n/a	<ul> <li>o_Oscillospirales; f_Acutalibacteraceae;</li> </ul>	.unitn.it/data/Pasolli_
	cov_89.3981	genomic loci						g_CAG-217 (inferred)	et_al.html
	QinJ_2012NOM001_NO	find gene functions						dBacteria; pFirmicutes_A; cClostridia;	http://segatalab.cibio
32	DE_179_length_28679_cov	from 54 unique	n/a	No	Yes	n/a	n/a	o_Oscillospirales; f_Acutalibacteraceae;	.unitn.it/data/Pasolli_
	2.87521	genomic loci						g_CAG-217 (inferred)	et_al.html
	QinJ_2012T2D-	find gene functions						dBacteria; pFirmicutes_A; cClostridia;	http://segatalab.cibio
33	050_NODE_25_length_192	from 54 unique	n/a	No	Yes	n/a	n/a	o_Oscillospirales; f_Acutalibacteraceae;	.unitn.it/data/Pasolli_
55	521_cov_10.1129	genomic loci	n/a		103	n/a	Π/a	g_CAG-217 (inferred)	et_al.html
	VatanenT_2016G78791_	0							http://segatalab.cibio
		find gene functions		NI-	Mar	- 1-	- 1-	d_Bacteria; p_Firmicutes_A; c_Clostridia;	
34	NODE_43_length_22491_c	from 54 unique	n/a	No	Yes	n/a	n/a	oOscillospirales; fAcutalibacteraceae;	.unitn.it/data/Pasolli_
	ov_6.98654	genomic loci						g_CAG-217 (inferred)	et_al.html
	XieH_2016YSZC12003_3	find gene functions						d_Bacteria; p_Firmicutes_A; c_Clostridia;	http://segatalab.cibio
35	5392_NODE_87_length_19	from 54 unique	n/a	No	Yes	n/a	n/a	<ul> <li>Oscillospirales; f_Acutalibacteraceae;</li> </ul>	.unitn.it/data/Pasolli_
	6476_cov_13.3023	genomic loci						gCAG-217 (inferred)	et_al.html
	XieH_2016YSZC12003_3	Figure 2, find gene						dBacteria; pFirmicutes_A; cClostridia;	http://segatalab.cibio
36	5563_NODE_11_length_35	functions from 54	n/a	No	Yes	n/a	n/a	o_Oscillospirales; f_Acutalibacteraceae;	.unitn.it/data/Pasolli_
	3850_cov_18.1068	unique genomic loci						g_CAG-217 (inferred)	et_al.html
		Figure 2, find gene							
		functions from 54							
	XieH_2016YSZC12003_3	unique genomic loci,						d Bacteria; p Firmicutes A; c Clostridia;	http://segatalab.cibio
37	6005_NODE_238_length_9	source of orf1-	n/a	No	Yes	n/a	n/a	o_Oscillospirales; f_Acutalibacteraceae;	.unitn.it/data/Pasolli_
57	9923 cov 5.91259		n/a	INU	165	n/a	11/a	g CAG-217 (inferred)	
	9923_000_5.91259	encoding phage						g_CAG-217 (interted)	et_al.html
		genome; Figure 3							
		Acr Seq							
		find gene functions							
	XieH_2016YSZC12003_3	from 54 unique						d Bacteria; p Firmicutes A; c Clostridia;	http://segatalab.cibio
38	6794_NODE_1_length_781	genomic loci, source	n/a	No	Yes	n/a	n/a	o_Oscillospirales; f_Acutalibacteraceae;	.unitn.it/data/Pasolli_
	521_cov_10.2961	of orf1-encoding	n/a	110	100	11/4	n/a	g_CAG-217 (inferred)	et_al.html
	321_000_10.2001	phage genome;						g_0A0 217 (intened)	Ct_al.nam
		Figure 3 Acr Seq							
	XieH_2016YSZC12003_3	find gene functions						d_Bacteria; p_Firmicutes_A; c_Clostridia;	http://segatalab.cibio
39	7133_NODE_3_length_676	from 54 unique	n/a	No	Yes	n/a	n/a	o_Oscillospirales; f_Acutalibacteraceae;	.unitn.it/data/Pasolli_
	817_cov_24.9073	genomic loci						g CAG-217 (inferred)	et_al.html
	XieH_2016YSZC12003_3	find gene functions						d_Bacteria; p_Firmicutes_A; c_Clostridia;	http://segatalab.cibio
40	7322_NODE_5_length_601	from 54 unique	n/a	No	Yes	n/a	n/a	o_Oscillospirales; f_Acutalibacteraceae;	.unitn.it/data/Pasolli_
Ť	737_cov_115.712	genomic loci	1.0			1,70	1.70	g_CAG-217 (inferred)	et_al.html
	XieH_2016YSZC12003_3	find gene functions						d_Bacteria; p_Firmicutes_A; c_Clostridia;	http://segatalab.cibio
41	7399_NODE_3_length_598	from 54 unique	n/a	No	Yes	n/a	n/a	o_Oscillospirales; f_Acutalibacteraceae;	.unitn.it/data/Pasolli
41		genomic loci	n/d	INU	162	ıı/a	n/a		et_al.html
	430_cov_49.9887	0						g_CAG-217 (inferred)	
40	XieH_2016YSZC12003_3	find gene functions		NI -	V		- 1-	d_Bacteria; p_Firmicutes_A; c_Clostridia;	http://segatalab.cibio
42	7878_NODE_8_length_402	from 54 unique	n/a	No	Yes	n/a	n/a	o_Oscillospirales; f_Acutalibacteraceae;	.unitn.it/data/Pasolli_
	183_cov_76.149	genomic loci						g_CAG-217 (inferred)	et_al.html
	YuJ_2015_SZAXPI003435	find gene functions						dBacteria; pFirmicutes_A; cClostridia;	http://segatalab.cibio
43	-	from 54 unique	n/a	No	Yes	n/a	n/a	o_Oscillospirales; f_Acutalibacteraceae;	.unitn.it/data/Pasolli_
10	11_NODE_1_length_77221	genomic loci	1u			1,70	1.// 4	g_CAG-217 (inferred)	et_al.html
	8_cov_11.5924	genomic loci							et_ai.iiuiii
	YuJ_2015SZAXPI015230	find gong functions						d Postaria: p. Eirmicutas Aug. Clastridia:	http://pogetalah.aikia
	-	find gene functions		NI -	V		- 1-	d_Bacteria; p_Firmicutes_A; c_Clostridia;	http://segatalab.cibio
44	16_NODE_32_length_1743	from 54 unique	n/a	No	Yes	n/a	n/a	o_Oscillospirales; f_Acutalibacteraceae;	.unitn.it/data/Pasolli_
	49_cov_17.3543	genomic loci						gCAG-217 (inferred)	et_al.html
		find gene functions						dBacteria; pFirmicutes_A; cClostridia;	http://segatalab.cibio
45	ZeeviD_2015PNP_DietInt	from 54 unique	n/a	No	Yes	n/a	n/a	o_Oscillospirales; f_Acutalibacteraceae;	.unitn.it/data/Pasolli_
10	ervention_11_NODE_16_le	genomic loci	1.0			1,70	1.70	g_CAG-217 (inferred)	et_al.html
		genomic loci			1		1		et_ati

	ngth_97163_cov_10.0000_I D_22997								
46	ZeeviD_2015PNP_Main_ 234_NODE_10_length_202 229_cov_13.9987_ID_1807 29	find gene functions from 54 unique genomic loci	n/a	No	Yes	n/a	n/a	d_Bacteria; p_Firmicutes_A; c_Clostridia; o_Oscillospirales; f_Acutalibacteraceae; g_CAG-217 (inferred)	http://segatalab.cibio .unitn.it/data/Pasolli_ et_al.html
47	ZeeviD_2015PNP_Main_ 294_NODE_20_length_208 110_cov_20.9981_ID_1060 95	find gene functions from 54 unique genomic loci	n/a	No	Yes	n/a	n/a	d_Bacteria; p_Firmicutes_A; c_Clostridia; o_Oscillospirales; f_Acutalibacteraceae; g_CAG-217 (inferred)	http://segatalab.cibio .unitn.it/data/Pasolli_ et_al.html
48	ZeeviD_2015PNP_Main_ 390_NODE_33_length_137 723_cov_10.9985_ID_4647 5	find gene functions from 54 unique genomic loci	n/a	No	Yes	n/a	n/a	d_Bacteria; p_Firmicutes_A; c_Clostridia; o_Oscillospirales; f_Acutalibacteraceae; g_CAG-217 (inferred)	http://segatalab.cibio .unitn.it/data/Pasolli_ et_al.html
49	ZeeviD_2015PNP_Main_ 578_NODE_20_length_138 741_cov_8.9988_ID_13256 3	find gene functions from 54 unique genomic loci	n/a	No	Yes	n/a	n/a	d_Bacteria; p_Firmicutes_A; c_Clostridia; o_Oscillospirales; f_Acutalibacteraceae; g_CAG-217 (inferred)	http://segatalab.cibio .unitn.it/data/Pasolli_ et_al.html
50	ZeeviD_2015PNP_Main_ 741_NODE_13_length_214 417_cov_12.0572_ID_9167 9	find gene functions from 54 unique genomic loci	n/a	No	Yes	n/a	n/a	d_Bacteria; p_Firmicutes_A; c_Clostridia; o_Oscillospirales; f_Acutalibacteraceae; g_CAG-217 (inferred)	http://segatalab.cibio .unitn.it/data/Pasolli_ et_al.html
51	ZellerG_2014CCIS03857 607ST-4- 0_NODE_542_length_3529 1_cov_2.7674	find gene functions from 54 unique genomic loci	n/a	No	Yes	n/a	n/a	d_Bacteria; p_Firmicutes_A; c_Clostridia; o_Oscillospirales; f_Acutalibacteraceae; g_CAG-217 (inferred)	http://segatalab.cibio .unitn.it/data/Pasolli_ et_al.html
52	ZellerG_2014CCIS22958 137ST-20- 0_NODE_40_length_18149 3_cov_7.91373	find gene functions from 54 unique genomic loci, source of orf1-encoding phage genome; Figure 3 Acr Seq	n/a	No	Yes	n/a	n/a	d_Bacteria; p_Firmicutes_A; c_Clostridia; o_Oscillospirales; f_Acutalibacteraceae; g_CAG-217 (inferred)	http://segatalab.cibio .unitn.it/data/Pasolli_ et_al.html
53	XieH_2016YSZC12003_3 5635_NODE_109_length_1 56568_cov_5.12141	find gene functions from 54 unique genomic loci	n/a	No	Yes	n/a	n/a	d_Bacteria; p_Firmicutes_A; c_Clostridia; o_Oscillospirales; f_Acutalibacteraceae; g_CAG-217 (inferred)	http://segatalab.cibio .unitn.it/data/Pasolli_ et_al.html
54	ZeeviD_2015PNP_Main_ 85_NODE_182_length_529 97_cov_7.0000_ID_133080	find gene functions from 54 unique genomic loci	n/a	No	Yes	n/a	n/a	d_Bacteria; p_Firmicutes_A; c_Clostridia; o_Oscillospirales; f_Acutalibacteraceae; g_CAG-217 (inferred)	http://segatalab.cibio .unitn.it/data/Pasolli_ et_al.html
55	4303_HeQ_2017SZAXPI 029570- 85bin.1_NODE_2_length _608092_cov_26.3259	Figure 2	n/a	Yes	No	HeQ_2017S ZAXPI029570- 85bin.1	4303	d_Bacteria; p_Firmicutes_A; c_Clostridia; o_Oscillospirales; f_Acutalibacteraceae; g_CAG-217	http://segatalab.cibio .unitn.it/data/Pasolli_ et_al.html
56	4303_CosteaPI_2017SID 713B074-11-90- 0bin.57_NODE_18_lengt h_238289_cov_5.37382	Source of orf1- encoding phage genome; Figure 3 Acr Seq	n/a	Yes	No	CosteaPI_2017 SID713B074 -11-90- 0bin.57	4303	d_Bacteria; p_Firmicutes_A; c_Clostridia; o_Oscillospirales; f_Acutalibacteraceae; g_CAG-217	http://segatalab.cibio .unitn.it/data/Pasolli_ et_al.html
57	Clostridiales_bacterium_isol ate_CIM:MAG_317_1 contig_8085	Figure 2, source of orf1-encoding phage genome	QAL M010 0000 2.1	No	No	n/a	n/a	d_Bacteria; p_Firmicutes_A; c_Clostridia; o_Oscillospirales; f_Acutalibacteraceae; g_CAG-217 (inferred)	NCBI Genbank
58	TPA_asm:_Ruminococcace ae_bacterium_isolate_UBA8 277_contig_226	Figure 3 Acr Seq	DPD R010 0001 0.1	No	No	n/a	n/a	d_Bacteria; p_Firmicutes_A; c_Clostridia; o_Oscillospirales; f_Acutalibacteraceae; g_CAG-217 (inferred)	NCBI Genbank
59	KarlssonFH_2013S424_ NODE_2_length_526279_c ov_9.22761	Source of orf1- encoding phage genome	n/a	No	Yes	n/a	n/a	d_Bacteria; p_Firmicutes_A; c_Clostridia; o_Oscillospirales; f_Acutalibacteraceae; g_CAG-217 (inferred)	http://segatalab.cibio .unitn.it/data/Pasolli_ et_al.html

60	XieH_2016YSZC12003_3 6696_NODE_1_length_776 477_cov_39.8546	Source of orf1- encoding phage genome, AcrIIA17 encoding phage (figure S5)	n/a	No	Yes	n/a	n/a	d_Bacteria; p_Firmicutes_A; c_Clostridia; o_Oscillospirales; f_Acutalibacteraceae; g_CAG-217 (inferred)	http://segatalab.cibio .unitn.it/data/Pasolli_ et_al.html
61	XieH_2016YSZC12003_3 7308R1_NODE_3_length_7 17276_cov_26.9646	source of orf1- encoding phage genome	n/a	No	Yes	n/a	n/a	d_Bacteria; p_Firmicutes_A; c_Clostridia; o_Oscillospirales; f_Acutalibacteraceae; g_CAG-217 (inferred)	http://segatalab.cibio .unitn.it/data/Pasolli_ et_al.html
62	ZellerG_2014CCIS88007 743ST-4- 0_NODE_31_length_21091 0_cov_8.07406	source of orf1- encoding phage genome	n/a	No	Yes	n/a	n/a	d_Bacteria; p_Firmicutes_A; c_Clostridia; o_Oscillospirales; f_Acutalibacteraceae; g_CAG-217 (inferred)	http://segatalab.cibio .unitn.it/data/Pasolli_ et_al.html
63	4303_QinN_2014LD- 22bin.75_NODE_22_leng th_329763_cov_10.7401	Figure 3 Acr Seq	n/a	Yes	No	QinN_2014L D-22bin.75	4303	d_Bacteria; p_Firmicutes_A; c_Clostridia; o_Oscillospirales; f_Acutalibacteraceae; g_CAG-217	http://segatalab.cibio .unitn.it/data/Pasolli_ et_al.html
64	ZellerG_2014CCMD2596 3797ST-21- 0_NODE_9_length_356111 _cov_10.1715	Figure 3 Acr Seq	n/a	No	Yes	n/a	n/a	d_Bacteria; p_Firmicutes_A; c_Clostridia; o_Oscillospirales; f_Acutalibacteraceae; g_CAG-217 (inferred)	http://segatalab.cibio .unitn.it/data/Pasolli_ et_al.html
65	ZellerG_2014CCIS41222 843ST-4- 0_NODE_17_length_26713 3_cov_14.7383	Figure 3 Acr Seq	n/a	No	Yes	n/a	n/a	d_Bacteria; p_Firmicutes_A; c_Clostridia; o_Oscillospirales; f_Acutalibacteraceae; g_CAG-217 (inferred)	http://segatalab.cibio .unitn.it/data/Pasolli_ et_al.html
66	FengQ_2015SID530168_ NODE_20_length_224404_ cov_6.02914	Figure 3 Acr Seq	n/a	No	Yes	n/a	n/a	d_Bacteria; p_Firmicutes_A; c_Clostridia; o_Oscillospirales; f_Acutalibacteraceae; g_CAG-217 (inferred)	http://segatalab.cibio .unitn.it/data/Pasolli_ et_al.html
67	FengQ_2015_SID530041_ NODE_7_length_421742_c ov_9.32571	Figure 3 Acr Seq	n/a	No	Yes	n/a	n/a	d_Bacteria; p_Firmicutes_A; c_Clostridia; o_Oscillospirales; f_Acutalibacteraceae; g_CAG-217 (inferred)	http://segatalab.cibio .unitn.it/data/Pasolli_ et_al.html
68	FengQ_2015SID31223_N ODE_13_length_228767_co v_7.50553	Figure 3 Acr Seq	n/a	No	Yes	n/a	n/a	d_Bacteria; p_Firmicutes_A; c_Clostridia; o_Oscillospirales; f_Acutalibacteraceae; g_CAG-217 (inferred)	http://segatalab.cibio .unitn.it/data/Pasolli_ et_al.html

**Supplemental Table 4**. Plasmids used in this study. Supplemental Table S5 indicates genes expressed from pZE21\_tetR.

Plasmid	crRNA promoter, sequence (5'-3')	Notes	Refs	Purpose
pZE21_tetR	n/a	Contains tetR behind pLac promoter for inducible expression of candidate Acrs. Targeted by crRNA_A; PAM = AGG.	(Forsberg et al., 2019)	Expressing genes to test <i>in vivo</i> anti- CRISPR activity
pSpyCas9 _crA	pJ23100, GTTCATTCAGGGCAC CGGAC	Arabinose-inducible SpyCas9 with pZE21 targeting pZE21_tetR	(Forsberg et al., 2019)	Target pZE21_tetR for elimination with SpyCas9
pSpyCas9 _crMu	pJ23100, GTAATACTTGTCCCGC AAAG	Mu-targeting spacer for phage Mu immunity testing. Otherwise identical to pSpyCas9_crA	(Forsberg et al., 2019)	Phage Mu immunity testing
pSpyCa9_crNT	pJ23100, GAACGAAAAGCTGCG CCGGG	non-targeting spacer used as control. Otherwise identical to pSpyCas9_crA	(Forsberg et al., 2019)	Phage Mu immunity testing, Western blots
pCloDF13_GFP	pJ23100, GAACGAAAAGCTGCG CCGGG	<i>eGFP</i> gene replaces <i>spyCas9</i> in pSpyCas9_crA		Measure generic protein expression from pSpyCas9 expression vector
pIDTsmart	n/a	Plasmid used for <i>in-vitro</i> SpyCas9 digestion. Sequence available at: <u>https://www.idtdna.com/pages/products/genes-</u> and-gene-fragments/custom-gene-synthesis		Plasmid template for <i>in-vitro</i> nuclease reactions
pET15b/HE	n/a	Novagen Cat. No. 69661-3; pET15 variants 'b' and 'HE' differ only by a few bases upstream of the N-terminal thrombin cut site		Protein purification
pSpyCas9_Fig3C	n/a	J23100 promoter expressing a theophylline inducible SpyCas9, used in Figure 3C	(Uribe et al., 2019)	Testing AcrIIA22 activity against a panel of Cas9 and Cas12 effector nucleases
pNmCas9_Fig3C	n/a	J23100 promoter expressing a theophylline inducible NmCas9, used in Figure 3C	(Uribe et al., 2019)	Testing AcrIIA22 activity against a panel of Cas9 and Cas12 effector nucleases
pLbCas12_Fig3C	n/a	J23100 promoter expressing a theophylline inducible LbCas12, used in Figure 3C	(Uribe et al., 2019)	Testing AcrIIA22 activity against a panel of Cas9 and Cas12 effector nucleases
pFnCas12_Fig3C	n/a	J23100 promoter expressing a theophylline inducible FnCas12, used in Figure 3C	(Uribe et al., 2019)	Testing AcrIIA22 activity against a panel of Cas9 and Cas12 effector nucleases
pDual4_Spy	P <sub>BAD</sub> , GTTCATTCAGGGCAC CGGAC	Arabinose inducible gRNA for SpyCas9 targeting pZE21	(Uribe et al., 2019)	Testing AcrIIA22 activity against a panel of Cas9 and Cas12 effector nucleases
pDual4_Nm	P <sub>BAD</sub> , GAACACGGCGGCATC AGAGC	Arabinose inducible gRNA for NmCas9 targeting pZE21	(Uribe et al., 2019)	Testing AcrIIA22 activity against a panel of Cas9 and Cas12 effector nucleases
pDual4_Lb	P <sub>BAD</sub> , TCAAGACCGACCTGT CCGGTGCCCTGAATG	Arabinose inducible gRNA for LbCas12 targeting pZE21	(Uribe et al., 2019)	Testing AcrIIA22 activity against a panel of Cas9 and Cas12 effector nucleases
pDual4_Fn	P <sub>BAD</sub> , TCAAGACCGACCTGT CCGGTGCCCTGAATG	Arabinose inducible gRNA for FnCas12 targeting pZE21	(Uribe et al., 2019)	Testing AcrIIA22 activity against a panel of Cas9 and Cas12 effector nucleases

## **Supplemental Table 5**. Gene sequences used in this study.

Gene Name	Sequence	Notes
acrIIA22wt	atggtagtagaagagacgcgggatttagccgaaactgcggattgtgtagtgatcgaagccattttagtggatgacggattgcgttacagacag	The italicized six base pairs were deleted in the ∆2aa truncation mutant via Q5 site-directed mutagenesis (NEB).
acrIIA22- null	atggtagtagaagagacgcgggatttagccgaaactgcggattgtgtagtgatcgaagccattt <u>A</u> agtggatgacggattgcgttacagacagctttctgtcggcatcaaa gacgaaaacggcgacattattcgtatcgtccctatttcaaccgttctgatctag	Mutation to introduce early stop codon via Q5 site-directed mutagenesis (NEB). Indicated in bold, capitalized, underline
acrIIA22a	atggtcatagaagagacgcgggatttagctgaaactgcggattgtgtagtgatcgaagccattttagtggatgacggattgcgttacaaacagctttccgtcggcatcaaaga cgaaaacggtgacattattcgtatcgtccctatttcaaccgttctgatctag	Same amino acid sequence as NCBI protein CDB51368.1; synthesized by GenScript and cloned into pZE21_tetR
acrllA22b	atgattgtggaagataccaaagatttggttgaaactgcggactatgtgatcatcgaagctgttttagtggatgatggattgcgttacaaacaa	Synthesized by GenScript and cloned into pZE21_tetR
acrIIA22c	atgaaaatgattgtggaagatacgaaagatctggtagaaacggacgattatgtaatcattgaagcgactttgtcagagggcgatttgttgttgtgcaaattgccgtgggcattc gcaacgaagtgggcgacattgttcgtattattcccatttccaccaacccaatctaa	Same amino acid sequence as NCBI protein CDB51757.1; synthesized by GenScript and cloned into pZE21_tetR
ts-acrIIA22	atg <b>tggagtcatccacaatttgagaag</b> ggaggaggcagtggaggaggcagt <b>ggaggaagtgcctggagccacccgcagttcgaaaaa</b> ggcagtggtggtggt agtggtggaggaatggtagtagaagagacgcgggatttagccgaaactgcggattgtgtagtgatcgaagccattttagtggatgacggattgcgttacagacag	N-terminal twin-strep (ts) tagged AcrIIA22. The tag is indicated in bold italics, linker regions are only italicized; synthesized by GenScript and cloned into pZE21_tetR for functional testing.
acrIIA22-ts	atggtagtagaagagacgcgggatttagccgaaactgcggattgtgtagtgatcgaagccattttagtggatgacggattgcgttacagacag	C-terminal twin-strep (ts) tagged AcrIIA22. The tag is indicated in bold italics, linker regions are only italicized; synthesized by GenScript and cloned into pZE21_tetR or pET15 for functional testing or protein purification, respectively.
purF	atgttcgatagtttgcacgaggaatgcggtgttttcggcgtatttgaaaatcagaccactacggtggcccagacggcgtatctggctctgtttgccttgcagcacagagggcag gagagttgcggcattgccgtgaatgacgacggcgtgtttcgccaccatcggggcgacggactggtgccggatgtgtttagcaaggagcagctggctg	Flanks <i>acrIIA22</i> -encoding bacterial genomic islands. Used as bait to retrieve additional examples of this locus for genomic and evolutionary analyses.
radC	atgcgtgccgcttatctgcaaggcggcggcggcgacgctatgccggaccaccagttgctggaattgctgctgtccatcagcattccccgcagagatgtaaagcccattgcctatg cgctcattaaccgcttcggctcgctggagcaggtgtttgccgccgggcgcagcagatctgcaacaagtgccgggcgtcggcgaacagaccgccgtacagattct gctggta cgggatctgaaccggcggatccatcaaaatcaaaacaaac	Flanks <i>acrIIA22</i> -encoding bacterial genomic islands. Used as bait to retrieve additional examples of this locus for genomic and evolutionary analyses.
acrIIA4	atgaatattaacgatttgatccgtgagattaagaataaggattatactgtcaaattgtccgggacagattccaattctattacacaattaatcatccgtgtgaataacgatggtaa tgagtatgtcatctctgaatcagaaaacgagagcatcgtagaaaagttcatcagtgccttcaagaacgggtggaaccaagagtatgaagatgaggaggaattttacaatg atatgcagacaattacgcttaaatcagaattgaatt	Discovered by (Rauch et al., 2017); synthesized by GenScript and cloned into pZE21_tetR
spyCas9	atggataagaaatactcaataggcttagatatcggcacaaatagcgtcggatgggcggtgatcactgatgaatataaggttccgtctaaaaagttcaaggttctgggaaata cagaccgccacagtatcaaaaaaaatcttataggggctcttttatttgacagtggagagacagcggaagcggctgttccaaacggacagctgtagaaggtatacacgt cggaagaatcgtatttgttatctacaggagatttttcaaatgagatggcgaaagtagatgatagttctttcatcgacttgaagagtctttttggtggaagaagacaagaagcat gaacgtcatcctatttttggaaatatagtagatgaagtgctatatcatgagaaatatccaactatctat	The sequence was amplified from Addgene plasmid #48645 (Esvelt <i>et al.</i> , 2013) for use in this study as described in (Forsberg <i>et al.</i> , 2019).

	aatcaattatttgaagaaaaccctattaacgcaagtggagtagatgctaaagcgattctttct	
	ggtgagaagaaaaatggcttatttgggaatctcattgctttgtcattgggtttgacccctaatttaaatcaaattttgatttggcagaagatgctaaattacagctttcaaaagata	
	cttacgatgatgattagataatttattggcgcaaattggagatcaatatgctgatttgtttttggcagctaagaatttatcagatgctattttactttcagatatcctaagagtaaatac	
	tgaaataactaaggctcccctatcagcttcaatgattaaacgctacgatgaacatcatcaagacttgactcttttaaaagctttagttcgacaacaacttccagaaaagtataa	
	agaaatcttttttgatcaatcaaaaaacggatatgcaggttatattgatgggggagctagccaagaagaattttataaatttatcaaaccaattttagaaaaaatggatgg	
	gaggaattattggtgaaactaaatcgtgaagatttgctgcgcaagcaa	
	gacaagaagacttttatccattttaaaagacaatcgtgagaagattgaaaaaatcttgacttttcgaattccttattatgttggtccattggcgcgtggcaatagtcgttttgcatgg	
	atgactcggaagtctgaagaaacaattaccccatggaattttgaagaagttgtcgataaaggtgcttcagctcaatcatttattgaacgcatgacaaactttgataaaaatttc	
	aggtgaacagaagaaagccattgttgatttactcttcaaaacaaatcgaaaagtaaccgttaagcaattaaaagaagattatttcaaaaaaatagaatgttttgatagtgttg	
	aaatttcaggagttgaagatagatttaatgcttcattaggtacctacc	
	atattgtttaacattgaccttatttgaagatagggagatgattgaggaaagacttaaacatatgctcacctctttgatgataaggtgatgaaacagcttaaacgtcgccgttat	
	actggttggggacgtttgtctcgaaaattgattaatggtattagggataagcaatctggcaaaacaatattagattttttgaaatcagatggttttgccaatcgcaattttatgcagc	
	tgatccatgatgatagtttgacatttaaagaagacattcaaaaagcacaagtgtctggacaaggcgatagtttacatgaacatattgcaaatttagctggtagccctgctattaa	
	aaaaggtattttacagactgtaaaagttgttgatgaattggtcaaagtaatggggcggcataagccagaaaatatcgttattgaaatggcacgtgaaaatcagacaactcaa	
	aagggccagaaaaattcgcgagagcgtatgaaacgaatcgaagaaggtatcaaagaattaggaagtcagattcttaaagagcatcctgttgaaaatactcaattgcaaa	
	atgaaaagctctatctctattatctccaaaatggaagagacatgtatgt	
	taaagacgattcaatagacaataaggtcttaacgcgttctgataaaaatcgtggtaaatcgggataacgttccaagtgaagaagtagtcaaaaagatgaaaaactattggag	
	a caacttc taa acgcca agttaat cactca acgt a agtttg at a atttaacga a agctg a acgt gg aggtttg a gt ga actt ga ta a a gct gg ttt a taa acgcc a att gg tt a cache a gct gg a gt gg ttt a taa acgcc a att gg tt a cache a gct gg a gt gg a gt gg a gt gg a gt gg a g	
	gaaactcgccaaatcactaagcatgtggcacaaattttggatagtcgcatgaatactaaatacgatgaaaatgataaacttattcgagaggttaaagtgattaccttaaaatcc	
	aaattagttictgacticcgaaaaagatticcaattctataaagtacgtgagattaacaattaccatcatgcccatgatgcgtatctaaatgccgtcgttggaactgctttgattaag	
	aaatatccaaaacttgaatcggagttgtctatggtgattataaagtttatgatgttcgtaaaatgattgctaagtctgagcaagaaataggcaaagcaaccgcaaaatatttctt	
	ttactctaatatcatgaacttcttcaaaacagaaattacacttgcaaatggagagattcgcaaacgccctctaatcgaaactaatggggaaactggagaaattgtctgggata	
	aagggcgagattttgccacagtgcgcaaagtattgtccatgccccaagtcaatattgtcaagaaaacagaagtacagacag	
	aaagaaattcggacaagcttattgctcgtaaaaaagactgggatccaaaaaaatatggtggttttgatagtccaacggtagcttattcagtcctagtggttgctaaggtggaa	
	aaagggaaatcgaagaagttaaaatccgttaaagagttactagggatcacaattatggaaagaagttcctttgaaaaaaatccgattgactttttagaagctaaaggatata	
	aggaagttaaaaaagacttaatcattaaactacctaaatatagtctttttgagttagaaaacggtcgtaaacggatgctggctagtgccggagaattacaaaaaggaaatga	
	gctggctctgccaagcaaatatgtgaattttttatatttagctagtcattatgaaaagttgaagggtagtccagaagataacgaacaaaaacaattgtttgt	
	g cattatttagatgagattattgagcaaatcagtgaattttctaagcgtgttattttagcagatgccaatttagataaagttcttagtgcatataacaaaca	
	acgtgaacaagcagaaaatattattcatttatttacgttgacgaatcttggagctcccgctgcttttaaatattttgatacaacaattgatcgtaaacgatatacgtctacaaaaga	
	agttttagatgccactcttatccatcaatccatcactggtctttatgaaacacgcattgatttgagtcagctaggaggtgactga	
	atggccgccttcaagcccaaccccatcaactacatcctgggcctggacatcggcatcgccagcgtgggctgggccatggtggagatcgacgaggacgagaaccccatc	
	tgcctgatcgacctgggtgtgcgcgtgttcgagcgcgctgaggtgcccaagactggtgacagtctggctatggctcgccggcttgctcgttcgt	
	cgcgctcaccgccttctgcgcgctcgccgcctgctgaagcgcgagggtgtgctgcaggctgccgacttcgacgagaacggcctgatcaagagcctgcccaacactccttg	
	gcagctgcgcgctgccgctctggaccgcaagctgactcctctggagtggagcgccgtgctgctgccacctgatcaagcaccgcggctacctgagccagcgcaagaacga	
	gggcgagaccgccgacaaggagctgggtgctctgctgaagggcgtggccgacaacgcccacgccctgcagactggtgacttccgcactcctgctgagctggccctgaa	
	caagttcgagaaggagagcggccacatccgcaaccagcgcggcgactacagccacaccttcagccgcaaggacctgcaggccgagctgatcctgctgttcgagaag	
	cagaaggagttcggcaacccccacgtgagcggcggcctgaaggagggcatcgagaccctgctgatgacccagcgccccgccctgagcggcgacgccgtgcagaag	
	atgctggggccactgcaccttcgagccagccgaggcccaaggccgccaagaacacctacaccgccgagcgcttcatctggctgaccaagctgaacaacctgcgcatcctg	
	gagcagggcagcgagcgccccctgaccgacaccgagcgcgccaccctgatggacgagccctaccgcaagagcaagctgacctacgcccaggcccgcaagctgct	
	gggtctggaggacaccgccttcttcaagggcctgcgctacggcaaggacaacgccgaggccagcaccctgatggagatgaaggcctaccacgccatcagccgcgccc	
	tggagaaggagggcctgaaggacaagaagactctctgaacctgagccccgagctgcaggacgagatcggcaccgccttcagcctgttcaagaccgacgaggacat	
		The sequence was amplified from Addgene
nmCarl	cgtgcccctgatggagcagggcaagcgctacgacgaggcctgcgccgagatctacggcgaccactacggcaagaagaacaccgaggagaagatctacctgcctcct	plasmid #48646 (Esvelt <i>et al.</i> , 2013). The
nmCas9		sequence of NmCas9 was cloned in place of
	catcgagaccgcccgcgaggtgggcaagagcttcaaggaccgcaaggagatcgagaagcgccaggaggagaaccgcaaggacggccggc	the cas9 gene in pSpyCas9_Fig3C.
	gttccgcgagtacttccccaacttcgtgggcgagcccaagagcaaggacatcctgaagctgcgcctgtacgagcagcagcagcaggcaagtgcctgtacagcggcaagga	
	gatcaacctgggccgcctgaacgagaagggctacgtggagatcgaccacgccctgcccttcagccgcacctgggacgacagcttcaacaacaaggtgctggtgctggg	
	cagcgagaaccagaaccagggcaaccagaccccctacgagtacttcaacggcaaggacaacagccgcgagtggcaggagttcaaggcccgcgtggagaccagcc	
	gcttcccccgcagcaagaagcagcgcatcctgctgcagaagttcgacgaggacggcttcaaggagcgcaacctgaacgacacccgctacgtgaaccgcttcctgtgcc	
	agttcgtggccgaccgcctgcgcctgaccggcaagggcaaggagcgtgttcgccagcaacggccagatcaccaacctgctgcgcggcttctgggggcctgcgcaag	
	gtgcgcgccgagaacgaccgccaccacgccctggacgccgtggtggtggcggccgcagcaccgtggccatgcagcagaagatcacccgcttcgtgcgctacaaggagat	
	gaacgccttcgacggtaaaaccatcgacaaggagaccggcgaggtgctgcaccagaagacccacttcccccagccctgggagttcttcgcccaggaggtgatgatgctcg	
	cgtgttcggcaagcccgacggcaagcccgagttcgaggaggccgacacccccgagaagctgcgcaccctgctggccgagaagctgagcagccgtcgggccgtg	
	cacgagtacgtgactcctctgttcgtgagccgcgcccccaacgcaagatgagcggtcagggtcacatggagagccgtgaagagcgccaagcgcctggacgagggcgt	
	gagcgtgctgcgcgtgcccctgacccagctgaaggtgaaggacctggagaagatggtgaaccgcgagcgcgagcccaagctgtacgaggccctgaaggccccgctg	
	gaggcccacaaggacgaccccgccaaggccttcgccgagcccttctacaagtacgacaaggccggcaaccgccaccagcaggtgaaggccgtgcgcgtggagcag	
	gtgcagaagaccggcgtgtgggtgcgcaaccacaacggcatcgccgacaacgccaccatggtgcgcgtggacgtgttcgagaagggcgacaagtactacctggtgcc	

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Inccast2 <ul> <li></li></ul>			
#dpcase_cope_cope_cope_cope_cope_cope_cope_cop			
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IbCas12         Iggaatictitugagegingtagegaatticgaaggingtagegaatticticaaggingtagegaatticacticaaggingtagegaatticacticaaggingtagegaatticacticaaggingtagegaatticacticaaggingtagegaatticacticaaggingtagegaatticacticaaggingtagegaatticacticaaggingtagegaatticacticaaggingtagegaatticacticaaggingtagegaatticacticaaggingtagegaatticacticaaggingtagegaatticacticaaggingtagegaatticacticaaggingtagegaatticacticaaggingtagegaatticacticaaggingtagegaatticactigaatgingtagegaatticactigaatgingtagegaatticactigaatgingtagegaatticaatticaaggingtagegaatticactigaatgingtagegaatticactigaatgingtagegaatticactigaatgingtagegaatticactigaatgingtagegaatticaatticaaggingtaggingtagegaatticactigaatgingtagegaatticaatticaaggingtaggingtagegaatticactigaatgingtagegaatticactigaatgingtagegaatticactigaatgingtagegaatticaaggingtaggingtagegaatticaaggingtaggingtagegaatticaaggingtaggingtagegaatticaaggingtaggingtagegaatticaagginggingtaggingingingtagegaatticaagginggingingtaggingtagginggingtaggi		gcaacgaaggatacaaaagtctgttcaagaaggatataattgagacaattttgccagagttcctcgatgacaaggacgagattgcgctggtcaattcgttcaacggattcac	
iahttrogsteate:agaaggitasegaagtitasegaagtitasegattitasegattitaggaaggitasegattitasegattigggaaggitasegattitasegattigggaaggitasegattitasegattigggaaggitasegattitasegattigggaaggitasegattiggaaggitasegattitasegattigggaaggitasegattiggaaggitageaggitaggit		aacagcattcacaggcttctttgataatcgggaaaatatgttctctgaggaggcaaagtccacttctattgcgttcaggtgtatcaatgagaatctcactaggtacatttccaaca	
iahttrogtcatcasagagitaticagedigtatagiggtiggtiggtiggtiggtiggtiggtiggti		tggatatctttgagaaggttgacgcaatttttgacaagcacgaagttcaggagattaaggagaagatcctcaattccgattatgacgttgaggacttcttcgaaggtgagtttttt	
IbCas12         ceasagecpasecajasgicasageticasageticatageticasageticasageticasageticasageticasageticaseticasticaseticageticageticaseticaseticagecogeticase			
InCas12          ctggagidpit/cgggaitac/tcicataagigattic/cgggaitac/tcicatagagaagaagigaga/tgicagigattic/cgggaitac/tgigagagattic/cggagagaagigaga/tgicagagagattic/cggagagagagagagagagagagagagagagagagaga			
assicg/ccg/cctut/ccaagataaagataagata/ccaagataagatac/ccaagatagatagatagatagatagatagatagatagata			
IbCas12       citicitiacigagaatiacicaaagagaaagittaaagittaagaatiaagittagittaagittagittaagittagittaagittagittaagitta			
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## **References for Supplemental Tables.**

An, S.Y., Ka, D., Kim, I., Kim, E.-H., Kim, N.-K., Bae, E., and Suh, J.-Y. (2020). Intrinsic disorder is essential for Cas9 inhibition of anti-CRISPR AcrIIA5. Nucleic Acids Research *48*, 7584-7594.

Athukoralage, J.S., McMahon, S.A., Zhang, C., Gruschow, S., Graham, S., Krupovic, M., Whitaker, R.J., Gloster, T.M., and White, M.F. (2020). An anti-CRISPR viral ring nuclease subverts type III CRISPR immunity. Nature *577*, 572-575.

Bhoobalan-Chitty, Y., Johansen, T.B., Di Cianni, N., and Peng, X. (2019). Inhibition of Type III CRISPR-Cas Immunity by an Archaeal Virus-Encoded Anti-CRISPR Protein. Cell *179*, 448-458 e411.

Bondy-Denomy, J., Garcia, B., Strum, S., Du, M., Rollins, M.F., Hidalgo-Reyes, Y., Wiedenheft, B., Maxwell, K.L., and Davidson, A.R. (2015). Multiple mechanisms for CRISPR-Cas inhibition by anti-CRISPR proteins. Nature *526*, 136-139.

Bondy-Denomy, J., Pawluk, A., Maxwell, K.L., and Davidson, A.R. (2013). Bacteriophage genes that inactivate the CRISPR/Cas bacterial immune system. Nature 493, 429-432.

Chowdhury, S., Carter, J., Rollins, M.F., Golden, S.M., Jackson, R.N., Hoffmann, C., Nosaka, L., Bondy-Denomy, J., Maxwell, K.L., Davidson, A.R., *et al.* (2017). Structure Reveals Mechanisms of Viral Suppressors that Intercept a CRISPR RNA-Guided Surveillance Complex. Cell *169*, 47-57 e11.

Dong, Guo, M., Wang, S., Zhu, Y., Wang, S., Xiong, Z., Yang, J., Xu, Z., and Huang, Z. (2017). Structural basis of CRISPR-SpyCas9 inhibition by an anti-CRISPR protein. Nature *546*, 436-439.

Eitzinger, S., Asif, A., Watters, K.E., Iavarone, A.T., Knott, G.J., Doudna, J.A., and Minhas, F. (2020). Machine learning predicts new anti-CRISPR proteins. Nucleic Acids Res *48*, 4698-4708.

Esvelt, K.M., Mali, P., Braff, J.L., Moosburner, M., Yaung, S.J., and Church, G.M. (2013). Orthogonal Cas9 proteins for RNA-guided gene regulation and editing. Nat. Methods *10*, 1116–1123.

Forsberg, K.J., Bhatt, I.V., Schmidtke, D.T., Javanmardi, K., Dillard, K.E., Stoddard, B.L., Finkelstein, I.J., Kaiser, B.K., and Malik, H.S. (2019). Functional metagenomics-guided discovery of potent Cas9 inhibitors in the human microbiome. Elife 8.

Fuchsbauer, O., Swuec, P., Zimberger, C., Amigues, B., Levesque, S., Agudelo, D., Duringer, A., Chaves-Sanjuan, A., Spinelli, S., Rousseau, G.M., *et al.* (2019). Cas9 Allosteric Inhibition by the Anti-CRISPR Protein AcrIIA6. Mol Cell *76*, 922-937 e927.

Garcia, B., Lee, J., Edraki, A., Hidalgo-Reyes, Y., Erwood, S., Mir, A., Trost, C.N., Seroussi, U., Stanley, S.Y., Cohn, R.D., *et al.* (2019). Anti-CRISPR AcrIIA5 Potently Inhibits All Cas9 Homologs Used for Genome Editing. Cell Rep *29*, 1739-1746 e1735.

Guo, T.W., Bartesaghi, A., Yang, H., Falconieri, V., Rao, P., Merk, A., Eng, E.T., Raczkowski, A.M., Fox, T., Earl, L.A., *et al.* (2017). Cryo-EM Structures Reveal Mechanism and Inhibition of DNA Targeting by a CRISPR-Cas Surveillance Complex. Cell *171*, 414-426 e412.

He, F., Bhoobalan-Chitty, Y., Van, L.B., Kjeldsen, A.L., Dedola, M., Makarova, K.S., Koonin, E.V., Brodersen, D.E., and Peng, X. (2018). Anti-CRISPR proteins encoded by archaeal lytic viruses inhibit subtype I-D immunity. Nat Microbiol *3*, 461-469.

Hirschi, M., Lu, W.T., Santiago-Frangos, A., Wilkinson, R., Golden, S.M., Davidson, A.R., Lander, G.C., and Wiedenheft, B. (2020). AcrIF9 tethers non-sequence specific dsDNA to the CRISPR RNA-guided surveillance complex. Nat Commun *11*, 2730.

Janowski, R., and Niessing, D. (2020). The large family of PC4-like domains - similar folds and functions throughout all kingdoms of life. RNA Biol *17*, 1228-1238.

Jiang, F., Liu, J.J., Osuna, B.A., Xu, M., Berry, J.D., Rauch, B.J., Nogales, E., Bondy-Denomy, J., and Doudna, J.A. (2019). Temperature-Responsive Competitive Inhibition of CRISPR-Cas9. Mol Cell *73*, 601-610 e605.

Knott, G.J., Cress, B.F., Liu, J.J., Thornton, B.W., Lew, R.J., Al-Shayeb, B., Rosenberg, D.J., Hammel, M., Adler, B.A., Lobba, M.J., et al. (2019a). Structural basis for AcrVA4 inhibition of specific CRISPR-Cas12a. Elife 8.

Knott, G.J., Thornton, B.W., Lobba, M.J., Liu, J.J., Al-Shayeb, B., Watters, K.E., and Doudna, J.A. (2019b). Broad-spectrum enzymatic inhibition of CRISPR-Cas12a. Nat Struct Mol Biol 26, 315-321.

Lee, J., Mir, A., Edraki, A., Garcia, B., Amrani, N., Lou, H.E., Gainetdinov, I., Pawluk, A., Ibraheim, R., Gao, X.D., *et al.* (2018). Potent Cas9 Inhibition in Bacterial and Human Cells by AcrIIC4 and AcrIIC5 Anti-CRISPR Proteins. mBio *9*.

Leon, L.M., Park, A.E., Borges, A.L., Zhang, J.Y., and Bondy-Denomy, J. (2020). Mobile element warfare via CRISPR and anti-CRISPR in <em&gt;Pseudomonas aeruginosa&lt;/em&gt. bioRxiv, 2020.2006.2015.151498.

Lin, P., Qin, S., Pu, Q., Wang, Z., Wu, Q., Gao, P., Schettler, J., Guo, K., Li, R., Li, G., et al. (2020). CRISPR-Cas13 Inhibitors Block RNA Editing in Bacteria and Mammalian Cells. Mol Cell 78, 850-861 e855.

Liu, L., Yin, M., Wang, M., and Wang, Y. (2019). Phage AcrIIA2 DNA Mimicry: Structural Basis of the CRISPR and Anti-CRISPR Arms Race. Mol Cell 73, 611-620 e613.

Mahendra, C., Christie, K.A., Osuna, B.A., Pinilla-Redondo, R., Kleinstiver, B.P., and Bondy-Denomy, J. (2020). Broad-spectrum anti-CRISPR proteins facilitate horizontal gene transfer. Nat Microbiol *5*, 620-629.

Marino, N.D., Zhang, J.Y., Borges, A.L., Sousa, A.A., Leon, L.M., Rauch, B.J., Walton, R.T., Berry, J.D., Joung, J.K., Kleinstiver, B.P., *et al.* (2018). Discovery of widespread Type I and Type V CRISPR-Cas inhibitors. Science.

Meeske, A.J., Jia, N., Cassel, A.K., Kozlova, A., Liao, J., Wiedmann, M., Patel, D.J., and Marraffini, L.A. (2020). A phage-encoded anti-CRISPR enables complete evasion of type VI-A CRISPR-Cas immunity. Science *369*, 54-59.

Osuna, B.A., Karambelkar, S., Mahendra, C., Christie, K.A., Garcia, B., Davidson, A.R., Kleinstiver, B.P., Kilcher, S., and Bondy-Denomy, J. (2020). Listeria Phages Induce Cas9 Degradation to Protect Lysogenic Genomes. Cell Host Microbe 28, 31-40 e39.

Pasolli, E., Asnicar, F., Manara, S., Zolfo, M., Karcher, N., Armanini, F., Beghini, F., Manghi, P., Tett, A., Ghensi, P., *et al.* (2019). Extensive Unexplored Human Microbiome Diversity Revealed by Over 150,000 Genomes from Metagenomes Spanning Age, Geography, and Lifestyle. Cell *176*, 649-662 e620.

Pawluk, A., Amrani, N., Zhang, Y., Garcia, B., Hidalgo-Reyes, Y., Lee, J., Edraki, A., Shah, M., Sontheimer, E.J., Maxwell, K.L., *et al.* (2016). Naturally Occurring Off-Switches for CRISPR-Cas9. Cell *167*, 1829-1838 e1829.

Pawluk, A., Bondy-Denomy, J., Cheung, V.H., Maxwell, K.L., and Davidson, A.R. (2014). A new group of phage anti-CRISPR genes inhibits the type I-E CRISPR-Cas system of Pseudomonas aeruginosa. MBio *5*, e00896.

Pawluk, A., Shah, M., Mejdani, M., Calmettes, C., Moraes, T.F., Davidson, A.R., and Maxwell, K.L. (2017). Disabling a Type I-E CRISPR-Cas Nuclease with a Bacteriophage-Encoded Anti-CRISPR Protein. MBio 8.

Pinilla-Redondo, R., Shehreen, S., Marino, N.D., Fagerlund, R.D., Brown, C.M., Sørensen, S.J., Fineran, P.C., and Bondy-Denomy, J. (2020). Discovery of multiple anti-CRISPRs uncovers anti-defense gene clustering in mobile genetic elements. bioRxiv, 2020.2005.2022.110304.

Rauch, B.J., Silvis, M.R., Hultquist, J.F., Waters, C.S., McGregor, M.J., Krogan, N.J., and Bondy-Denomy, J. (2017). Inhibition of CRISPR-Cas9 with Bacteriophage Proteins. Cell *168*, 150-158 e110.

Shin, J., Jiang, F., Liu, J.-J., Bray, N.L., Rauch, B.J., Baik, S.H., Nogales, E., Bondy-Denomy, J., Corn, J.E., and Doudna, J.A. (2017). Disabling Cas9 by an anti-CRISPR DNA mimic. Science Advances *3*.

Song, G., Zhang, F., Zhang, X., Gao, X., Zhu, X., Fan, D., and Tian, Y. (2019). AcrIIA5 Inhibits a Broad Range of Cas9 Orthologs by Preventing DNA Target Cleavage. Cell Rep *29*, 2579-2589 e2574.

Stanley, S.Y. (2018). An Investigation of Bacteriophage Anti-CRISPR and Anti-CRISPR Associated Proteins. In Department of Molecular Genetics (<u>http://hdl.handle.net/1807/97883</u>: University of Toronto), pp. 120.

Uribe, R.V., van der Helm, E., Misiakou, M.A., Lee, S.W., Kol, S., and Sommer, M.O.A. (2019). Discovery and Characterization of Cas9 Inhibitors Disseminated across Seven Bacterial Phyla. Cell Host Microbe *25*, 233-241 e235.

Varble, A., Campisi, E., Euler, C.W., Fyodorova, J., Rostøl, J.T., Fischetti, V.A., and Marraffini, L.A. (2020). Integration of prophages into CRISPR loci remodels viral immunity in <em&gt;Streptococcus pyogenes&lt;/em&gt. bioRxiv, 2020.2010.2009.333658.

Wang, J., Ma, J., Cheng, Z., Meng, X., You, L., Wang, M., Zhang, X., and Wang, Y. (2016a). A CRISPR evolutionary arms race: structural insights into viral anti-CRISPR/Cas responses. Cell Res 26, 1165-1168.

Wang, X., Yao, D., Xu, J.G., Li, A.R., Xu, J., Fu, P., Zhou, Y., and Zhu, Y. (2016b). Structural basis of Cas3 inhibition by the bacteriophage protein AcrF3. Nat Struct Mol Biol 23, 868-870.

Watters, K.E., Fellmann, C., Bai, H.B., Ren, S.M., and Doudna, J.A. (2018). Systematic discovery of natural CRISPR-Cas12a inhibitors. Science.

Watters, K.E., Shivram, H., Fellmann, C., Lew, R.J., McMahon, B., and Doudna, J.A. (2020). Potent CRISPR-Cas9 inhibitors from Staphylococcus genomes. Proc Natl Acad Sci U S A *117*, 6531-6539.

Yang, H., and Patel, D.J. (2017). Inhibition Mechanism of an Anti-CRISPR Suppressor AcrIIA4 Targeting SpyCas9. Mol Cell 67, 117-127 e115.

Zetsche, B., Gootenberg, J.S., Abudayyeh, O.O., Slaymaker, I.M., Makarova, K.S., Essletzbichler, P., Volz, S.E., Joung, J., van der Oost, J., Regev, A., et al. (2015). Cpf1 Is a Single RNA-Guided Endonuclease of a Class 2 CRISPR-Cas System. Cell.

Zhang, H., Li, Z., Daczkowski, C.M., Gabel, C., Mesecar, A.D., and Chang, L. (2019). Structural Basis for the Inhibition of CRISPR-Cas12a by Anti-CRISPR Proteins. Cell Host Microbe 25, 815-826 e814.

Zhang, K., Wang, S., Li, S., Zhu, Y., Pintilie, G.D., Mou, T.C., Schmid, M.F., Huang, Z., and Chiu, W. (2020). Inhibition mechanisms of AcrF9, AcrF8, and AcrF6 against type I-F CRISPR-Cas complex revealed by cryo-EM. Proc Natl Acad Sci U S A *117*, 7176-7182.