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2	Competition between mobile genetic elements drives optimization of a phage-encoded			
3	CRISPR-Cas system: Insights from a natural arms-race			
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17 Abstract

18 CRISPR-Cas systems function as adaptive immune systems by acquiring nucleotide sequences called spacers that mediate sequence-specific defense against competitors. Uniquely, the 19 20 phage ICP1 encodes a Type I-F CRISPR-Cas system that is deployed to target and overcome 21 PLE, a mobile genetic element with anti-phage activity in Vibrio cholerae. Here, we exploit the 22 arms race between ICP1 and PLE to examine spacer acquisition and interference under 23 laboratory conditions to reconcile findings from wild populations. Natural ICP1 isolates encode 24 multiple spacers directed against PLE, but we find that single spacers do not equally interfere 25 with PLE mobilization. High-throughput sequencing to assay spacer acquisition reveals that 26 ICP1 can also acquire spacers that target the V. cholerae chromosome. We find that targeting 27 the V. cholerae chromosome proximal to PLE is sufficient to block PLE and propose a model in which indirect chromosomal spacers are able to circumvent PLE by Cas2-3-mediated 28 29 processive degradation of the V. cholerae chromosome before PLE mobilization. Generally, 30 laboratory acquired spacers are much more diverse than the subset of spacers maintained by 31 ICP1 in nature, showing how evolutionary pressures can constrain CRISPR-Cas targeting in 32 ways that are often not appreciated through in vitro analyses. 33 Introduction 34 Phages often vastly outnumber their bacterial hosts in a variety of environments (1). As 35

such, bacteria have evolved numerous mechanisms for phage defense, including adaptive
immunity via clustered regularly interspaced short palindromic repeats (CRISPR) and CRISPRassociated (Cas) proteins (2,3). CRISPR-Cas systems are composed of a CRISPR array—a
series of "spacers" of foreign sequence alternating with repeats that are transcribed into
CRISPR RNAs (crRNAs)—and CRISPR-associated (Cas) genes. Together with crRNAs, Cas
proteins defend against foreign nucleic acids, such as the genome of an infecting phage,
through a three-step process: adaptation, crRNA expression, and interference. During

adaptation, a foreign DNA fragment is incorporated into the CRISPR array to provide a 43 44 molecular memory of the challenges that the host cell has faced. This CRISPR array is expressed and processed into individual crRNAs, which complex with Cas proteins and survey 45 the cell for complementary invading nucleotides. Upon finding a complementary sequence, 46 47 termed protospacer, a Cas nuclease is recruited to the site to mediate interference by cleaving the substrate, ultimately leading to the destruction of the invader (4.3). Across CRISPR-Cas 48 containing bacteria and archaea, Class 1 Type I CRISPR-Cas systems employing a Cas3 49 50 enzyme for DNA unwinding and degradation (5), are the most prevalent (6). 51 CRISPR-Cas systems do not discriminate between horizontally acquired traits based on 52 fitness gain or loss. Hence, CRISPR-Cas systems are equally capable of halting harmful 53 invading phage DNA as they are halting beneficial mobile genetic elements, including those encoding antibiotic resistance and pathogenicity genes (7–9). As such, some pathogens only 54 55 have alternative anti-phage defense systems (10). For example, the currently circulating biotype 56 of epidemic Vibrio cholerae, the causative agent of the diarrheal disease cholera, does not rely 57 on CRISPR-Cas for phage defense (11). Instead, V. cholerae evolved to use phage inducible chromosomal island-like elements (PLEs) to defend against the prevalent lytic phage, ICP1 58 59 (12). PLEs are mobile genetic elements that reside integrated in the small chromosome (chromosome II) of V. cholerae (12). During ICP1 infection of PLE(+) V. cholerae, PLE excises 60 from the host chromosome, replicates to high copy and is horizontally transduced to naïve 61 62 neighboring cells, all the while inhibiting phage replication through unknown mechanisms (Fig. 63 1).

In order to overcome the anti-phage activity encoded by *V. cholerae* PLE, some ICP1
isolates use a Type I-F CRISPR-Cas system that directly targets PLE (Fig. 1), making the
CRISPR-Cas system essential for the phage to form plaques on PLE(+) *V. cholerae* (13). Type
I-F systems are composed of three Csy proteins that make up the Csy complex along with
Cas6f, a protein involved in crRNA processing (14). This complex interacts with the processed

69 crRNA to search DNA for a complementary protospacer with an appropriate self versus non-self 70 discrimination sequence, known as the protospacer adjacent motif (PAM) (15). Upon finding a match with an appropriate PAM, the trans-acting Cas2-3 fusion protein is recruited to degrade 71 the target DNA. In addition to exonuclease activity, Cas2-3 is a processive helicase in vitro, 72 73 allowing for continued degradation of the target DNA (16,17). Recently, sequence analysis identified phages that are predicted to encode CRISPR arrays and/or Cas genes (18,19); 74 75 however, ICP1 is the only phage shown to encode a fully functional CRISPR-Cas system 76 (12, 13).

77 As is true when CRISPR-Cas is harnessed by a prokaryotic host for genome defense, 78 the ICP1-encoded CRISPR-Cas system is tasked with targeting and degrading a hostile mobile 79 genetic element. However, there are additional challenges associated with a phage encoding 80 and relying on CRISPR-Cas for its own survival. The ICP1 infection cycle occurs over a 20 81 minute period, and current data suggest that ICP1 synthesizes its CRISPR-Cas machinery de 82 novo upon infection of V. cholerae (13). PLE is induced to excise within minutes of infection through interactions with an early phage-encoded gene product (20). Thus, in order to 83 overcome PLE, CRISPR synthesis and interference must outpace a rapidly replicating target. 84 85 ICP1 and V. cholerae are consistently co-isolated from patient stool samples in regions where cholera is endemic such as Bangladesh (12,21,22). Five genetically distinct PLE variants 86 in V. cholerae have appeared in temporally discrete waves across cholera epidemics (12). 87 88 Previous analysis revealed that ICP1-encoded CRISPR-Cas can adapt and acquire new 89 spacers against PLE under laboratory conditions (13), however the rules governing spacer 90 acquisition and targeting efficacy for this system are not known. Further, recent comparative genomics of 18 ICP1 isolates collected from Bangladesh between 2001-2012 found that 50% 91 carry CRISPR-Cas (23), however the contemporary state of circulating ICP1 and V. cholerae 92 93 PLE in the region are not known.

Here, we provide an up-to-date understanding of the genomic variants of ICP1 and PLE 94 95 circulating in Bangladesh. We find that natural ICP1 isolates encode multiple anti-PLE spacers and experimentally validate that increased PLE targeting by ICP1 is required to fully abolish 96 PLE mobilization. Significantly, using a high-throughput spacer acquisition assay and 97 98 experimental validation, we show that noncanonical PAMs and indirect protospacers in the V. 99 cholerae small chromosome can unexpectedly provide protection against PLE. Our results 100 support a model in which ICP1-encoded CRISPR-Cas that is directed against the V. cholerae 101 small chromosome is in a race to reach PLE before it excises from the chromosome to exert its 102 anti-phage activity. Taken together, our study highlights the differences between interference 103 competent spacers under laboratory conditions and those that are selected for in nature to provide mechanistic insight into the evolutionary pressures governing the interactions between 104 105 epidemic V. cholerae and its longstanding battle with the predatory phage ICP1.

106

107 Methods.

108 Strains, growth conditions and genomic analysis.

109 Phage, bacterial strains and plasmids used in this study are listed in Supplementary Tables 1-3. 110 Bacteria were routinely grown at 37 °C on lysogeny broth (LB) agar or in LB broth with aeration. Media was supplemented with ampicillin (50 µg/ml), kanamycin (75 µg/ml), spectinomycin (100 111 µg/ml), and/or streptomycin (100 µg/ml) when appropriate. Phage susceptibility was determined 112 113 by standard soft agar overlays as described (11) and phage plague spot plates were performed 114 as described previously (20). Cholera stool samples collected and stored at the ICDDR,B 115 between 2015-2017 were probed for the presence of phage by standard soft agar overlays, and 116 V. cholerae isolates were recovered by plating on Thiosulfate Citrate Bile Salts Sucrose selective media (Difco). ICP1 specific primers (13,22) and PLE specific primers (Supplementary 117 118 Table 4) were used for preliminary screening of isolates from stool samples. The presence of CRISPR-Cas in ICP1 and PLE in V. cholerae was validated by whole genome sequencing. 119

120 Genomic libraries were generated using NEBNext Ultra II DNA Library preparation kit for 121 Illumina (New England Biolabs), according to the manufacturer's recommended protocols. Paired-end sequencing (2 x 150 bp) was performed on an Illumina HiSeq4000 (University of 122 123 California, Berkeley QB3 Core Facility). Sequencing assembly/mapping and detection of 124 CRISPR was performed as described (23). The V. cholerae clinical isolate KDS1 genome was 125 sequenced on Illumina HiSeq4000, PacBio Sequel and Oxford Nanopore MinION sequencers (University of California, Berkeley QB3 Core Facility). Assembly of KDS1 sequences was 126 127 performed using the canu assembler v1.6 (24) to combine the PacBio and Oxford Nanopore 128 reads into genomic scaffolds for the large and small chromosomes using default settings and an 129 expected genome size of 4033460bp. This generated two scaffolds of the expected sizes for each chromosome which were then polished with the Illumina paired-end sequences using Pilon 130 131 v1.22 (25) with the "fix all" command to generate a high-quality genomic assembly in a fasta 132 format of both chromosomes (Supplementary File 1).

133 V. cholerae mutants were constructed by natural transformation as described (26). Mutations in ICP1 were generated using CRISPR-Cas mediated genome engineering with the 134 V. cholerae classical biotype Type I-E system as described (11) (Supplementary Table 3). 135 136 Engineered phage +/- Cas1 D244A with a spacer 9 were validated by plaquing on a permissive PLE 1 host and determining the frequency of phage with a newly acquired spacer by calculating 137 the efficiency of plaquing on the permissive PLE 1 host to a PLE 1 host with the protospacer 138 139 deleted. Examination of PLE replication and transduction during phage infection was described 140 as reported previously (12).

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142 High throughput spacer acquisition, data processing and analyses

143 Three independent experiments were performed as follows: A 50 mL culture of PLE 1 V.

144 *cholerae* was grown to $OD_{600} = 0.3$ and infected with ICP1_2011_A Δ S9 (13) at an MOI of 1.

145 Infected cells were incubated for 90 minutes at 37 °C with aeration, at which point lysis was

146 observed. The lysate was treated with chloroform and centrifuged to remove bacterial debris. 147 Phage were precipitated with 10% (w/v) polyethylene glycol (PEG) 8000 at 4°C overnight. Phage pellets were collected by centrifugation at 4°C and the passaging was repeated as 148 above. After three passages, the resulting pools were plated on the PLE 1^{S8*} host to select for 149 150 phage with expanded arrays that allow plague formation. Phage DNA libraries were generated 151 by homopolymer tail-mediated PCR (HTM-PCR) as previously described (27). As ICP1 2011 A possesses only a single functional CRISPR array (Fig. 1b), the expanded phage CRISPR 1 152 153 array was amplified from genomic DNA libraries by PCR using custom barcoded primers 154 (Supplementary Table 4) to sequence the leader proximal spacer. 50bp single-end sequencing 155 was performed on an Illumina HiSeq 2500 (Tufts University Core Facility) using a custom 156 sequencing primer. The resulting reads as fast files (Supplementary File 2) were mapped to 157 the large and small chromosome of V. cholerae strain KDS1 using Bowtie v1.2.2 (28) with a 158 seed length of 31 and allowing for 0 max total mismatches which ensured that spacer to 159 protospacer matches were 100% identical. These mappings were performed in two parallel ways: first, to obtain all possible spacer mapping locations regardless of the number of identical 160 protospacer targets (i.e. translucent spacers in Fig 3b) and second, restricting max alignments 161 162 to 1 which only mapped spacers with exactly one unique mapping location across both chromosomes. With a custom Python script (https://git.io/fNVgZ) we extracted the PAM 163 sequences and GG PAM slippage locations from the restricted unique mappings. We also used 164 165 this script to generate spacer mapping location graphs for both set of mappings using 166 Biopython's GenomeDiagram module (29).

167

168 Results

169 **ICP1-encoded CRISPR-Cas is fixed in the natural phage population**

We set out to compare ICP1 and PLE from contemporary cholera patient stool samples
to previously identified isolates from the International Centre for Diarrhoeal Disease Research,

172 Bangladesh (ICDDR,B) in Dhaka, Bangladesh (13,21). We isolated eight new ICP1 isolates 173 from cholera patient stool samples collected between 2015-2017 and found that all isolates 174 harbor CRISPR-Cas. Thus it appears that ICP1 isolates lacking CRISPR have not been identified in Bangladesh since 2006 (23). Analysis of the CRISPR arrays indicates a strong 175 176 selection for spacers specifically targeting PLE (Fig. 1b, Supplementary Table 5), supporting the 177 function of the ICP1-encoded CRISPR-Cas system as a counter-attack against the anti-phage island PLE (13). To evaluate if the fixation of CRISPR in ICP1 is necessitated by co-circulating 178 179 PLE in epidemic V. cholerae, we determined the prevalence of PLE over the same near two-180 decade long period in Dhaka. Combined with previous analyses (12,21), we observed an 181 increase in the prevalence of PLE(+) V. cholerae in epidemic sampling over time (Fig. 1c). Of 182 note is the high prevalence of PLE 1 V. cholerae over the past 6 years, indicating that this 183 variant of the anti-phage island is currently dominating the epidemic landscape in Dhaka. 184 Despite the relatively long period over which PLE 1 has been dominant in Dhaka, and 185 consistent with previous results (12,21), whole genome sequencing of eight PLE 1 V. cholerae isolates showed that PLE 1 is 100% identical at the nucleotide level in all strains. 186 187

188 Multiple spacers increase ICP1 CRISPR-Cas mediated PLE interference

All of the natural phage we isolated encode multiple CRISPR spacers against PLE (Fig. 189 1b); however, previous work revealed that only one functional spacer is required for ICP1 to 190 191 overcome PLE-mediated anti-phage activity as evaluated by plague formation (13). Conversely, 192 a single spacer against the PLE did not prevent transduction of PLE (12). To investigate the 193 consequences of varying spacer number and identity on PLE transduction and replication, we 194 used co-isolated ICP1 and PLE 1 V. cholerae obtained from a cholera patient sample in 2011 (13). This ICP1 isolate harbors two spacers (spacers 8 and 9) at the leading edge of the 195 196 CRISPR 1 array that target PLE 1. We also used an isogenic phage with a spontaneous loss of spacer 9 (13), as well as one that acquired an additional 10th spacer targeting PLE *in vitro* (Fig. 197

198 2a). Despite the ability to overcome PLE and form plaques, spacer 8 targeting was not sufficient 199 to decrease PLE transduction during ICP1 infection relative to an untargeted control (Fig. 2b). In comparison, two anti-PLE spacers decreased PLE transduction during ICP1 infection and three 200 201 spacers completely abolished PLE transduction, showing that increased CRISPR targeting by 202 ICP1 has a stronger anti-PLE effect. To evaluate potential differences between spacer 8 and spacer 9 on PLE targeting, we used PLE 1 with a protospacer mutation (PLE 1^{PS8*}) that inhibits 203 spacer 8-mediated PLE targeting (13). Strikingly, just spacer 9 targeting PLE alone was able to 204 205 decrease PLE transduction to the same level as when two spacers were targeting PLE (Fig. 2b). 206 We next analyzed the copy number of PLE during infection with the ICP1 encoding one 207 or two targeting spacers to identify if the differences in reducing PLE transduction were due to 208 differences in PLE copy number (Fig 2c). In the absence of ICP1 CRISPR targeting, PLE 209 replicates to high copy number, which facilitates horizontal transmission. Targeting with only 210 one spacer was sufficient to significantly decrease PLE replication, and in agreement with the 211 transduction data, spacer 9 had a stronger inhibitory effect on PLE replication than spacer 8. 212 Altogether, these results demonstrate that not all spacers selected in nature equally interfere with PLE mobilization and that increasing the number of spacers provides enhanced capacity of 213 214 ICP1 to interfere with PLE.

215

Interference-driven spacer acquisition in ICP1 reveals indirect targets and non-canonical
 PAMs

Since spacer composition variability in nature was lower than we expected (Fig 1b), we next set out to experimentally sample the repertoire of spacers that ICP1 can acquire to overcome PLE. Low-throughput experiments previously demonstrated that ICP1 can acquire new spacers targeting the PLE under laboratory conditions without the need to overexpress *cas* genes (13). To further analyze the natural process of interference-driven spacer acquisition in this system, we performed high-throughput sequencing of expanded CRISPR arrays of phage 224 selected on PLE 1 V. cholerae. We infected PLE 1 V. cholerae with ICP1 containing spacer 8 225 (Fig 2a), and the recovered lysate was probed for ICP1 progeny with newly acquired spacers that allowed for plaque formation on a PLE 1^{PS8*} host. Illumina sequencing of the leader-226 227 proximal spacer in CRISPR 1 allowed us to sample over 10⁶ acquired spacers in each replicate 228 experiment (Supplementary Table 6). In order to accurately map the spacers to the PLE 1 V. 229 cholerae host, we performed complete whole-genome sequencing and assembly of the bacterial genome (Supplementary File 1). As was previously reported (12), we found that PLE 1 was 230 231 integrated in a V. cholerae repeat (VCR), of which over 100 repeats intersperse the V. cholerae 232 small chromosome in a gene-capture region, the superintegron (30). In total, 96% of the 233 acquired spacers mapped to PLE, while, interestingly, the other 4% mapped to V. cholerae 234 chromosomes (Supplementary Table 6).

235 Mapping of the spacers to the small chromosome showed a pattern of strand bias that 236 reflected previous observations in primed acquisition experiments performed in other Type I-F 237 systems (31), with a distribution of acquired spacers 5' of the protospacer on the non-targeted 238 strand and 3' of the protospacer on the targeted strand (Fig. 3a, Supplementary Fig. 2). The distribution of spacers acquired 5' of the protospacer on the nontargeted were split between the 239 240 small chromosomal region proximal to the PLE 1 integration site (Fig. 3b), as well as the 3' end of PLE. Acquired spacers mapping to the V. cholerae chromosome were not evenly distributed 241 between the large and small chromosome, but instead ~90% of the chromosomal spacers 242 243 mapped to the small chromosome (Supplementary Table 6, Fig. 3b). Spacers that mapped to 244 the large chromosome were restricted to a mu-like region (Fig. 3c), which was duplicated in this 245 strain and was also in the small chromosome proximal to PLE (Fig. 3b). Acquired spacers 246 mapped uniformly throughout the superintegron, however, this is likely an artifact as the superintegron is highly repetitive. When considering spacers that map to a single site in the 247 248 small chromosome, we observed an obvious bias for acquired spacers mapping closer to the 249 PLE integration site (Fig. 3b).

250 Consistent with CRISPR⁺ ICP1 isolates from nature (Supplementary Figure 1), the majority (~70%) of the spacers acquired experimentally targeted protospacers in PLE 1 that 251 252 were flanked by a 3' GA PAM (Fig. 3d). However, ~30% of protospacers in PLE had non-253 canonical PAMs, and of those, the majority were GG or GT. Previous CRISPR acquisition 254 studies in Type I-F systems indicate that alternative PAMs can be explained by a "slippage" 255 event (31,32). To identify putative slippage events, we analyzed the sequences adjacent to GG PAMs and found that 45% of GG PAMs have a canonical GA within 3 nucleotides of the PAM 256 257 position, suggesting that the ICP1 acquisition machinery has a propensity to slip (Fig. 4a). 258 We next wanted to determine if these non-canonical PAMs are functional for PLE 259 interference. To do so, we engineered ICP1 to encode a single spacer reflective of an 260 experimentally acquired spacer with the most common non-canonical PAMs: either a GG or GT 261 (Fig. 3d) and evaluated plaque formation on PLE 1 V. cholerae. Despite relying on a non-262 canonical PAM, we found that ICP1 is able to target those protospacers and overcome PLE, 263 albeit at a lower efficiency than when targeting a protospacer with a canonical GA PAM (Fig. 264 4b). Even when no canonical PAM was within +/- 3 nt, ICP1 was still able to overcome PLE targeting a protospacer with a GT PAM. As PAM mutations are frequently a source for primed 265 266 acquisition (33), we tested if the observed residual CRISPR activity was due to further spacer 267 acquisition and interference. We constructed a Cas1 D244A mutation, which disrupts a conserved metal coordinating residue to inhibit spacer acquisition (32) (Supplementary Fig. 3) 268 269 and tested if plague formation was altered (Fig 4b). We observed no difference in the efficiency 270 of plaque formation between the Cas1 mutants and the parental phage, suggesting that the 271 ICP1 CRISPR-Cas system is more tolerant of divergent PAMs during infection than previously 272 characterized (13).

273

274 **Protospacers in the small chromosome facilitate ICP1 CRISPR-Cas-mediated PLE**

275 interference

276 In our spacer acquisition experiment we identified a subset of spacers that target a mu-277 like region in the V. cholerae large chromosome (Fig. 3c), suggesting that CRISPR targeting of the mu-like region was advantageous in overcoming PLE. To test the role of the mu-like region 278 279 protospacer in PLE interference, we isolated ICP1 that had acquired a spacer that targets the mu-like region and was able to form plaques on PLE 1^{PS8^*} (Fig 5a). Since assembly of the V. 280 281 cholerae genome revealed that the mu-like region was present in each chromosome (Fig 5a) we wanted to evaluate if targeting the mu-like region per se was allowing for plaque formation, or if 282 283 the chromosomal context was important in allowing for CRISPR-meditated interference with 284 PLE. To test this difference, we generated a single knockout of the mu-like region in the large 285 chromosome and a double knockout in both chromosomes. ICP1 CRISPR-mediated 286 interference with PLE was abolished in the double knockout, however, knocking out the mu-like 287 region in the large chromosome had no effect on ICP1 plaque formation (Fig 5b). These results 288 show that CRISPR targeting of the V. cholerae large chromosome is dispensable for phage 289 overcoming PLE, while targeting the small chromosome is sufficient to overcome PLE activity. 290

291 When CRISPR goes off target: going the distance to maintain interference

292 As processivity of Cas2-3 has been demonstrated in vitro (17), we speculated that ICP1 targeting of the small chromosome proximal to PLE interferes with PLE anti-phage activity by 293 the processive degradation of PLE along with the chromosome; however, PLE excises from the 294 295 chromosome early during ICP1 infection (20). This timing suggests that CRISPR targeting and 296 Cas2-3 processive degradation of the small chromosome would have to happen prior to PLE 297 excision and would therefore likely be distance dependent. In support of this hypothesis, 298 experimentally acquired spacers mapping to the small chromosomal clustered proximal to PLE (Fig. 3b). To test the impact of targeting at increasing distances from PLE, we engineered ICP1 299 300 to possess CRISPR arrays containing only one spacer drawn from the experimental acquisition pool that targets the small chromosome at varying distances away from PLE. We then assayed 301

302 the ability of these engineered phage to overcome PLE and form plagues (Fig. 6a). As a 303 positive control, ICP1 engineered with a spacer that targets internal to PLE formed robust and equal plagues on PLE(-) and PLE 1 hosts. In comparison, phage with a spacer that targets far 304 (>400 kb) from PLE were unable to form plagues on PLE 1. Conversely, ICP1 that target a 305 306 protospacer only 0.5, 1.5 or 2.5 kb from PLE were able to efficiently overcome PLE and form 307 plaques. Phage targeting protospacers at intermediate distances away from PLE (>20 kb) demonstrated weak plaque formation on PLE 1. Surprisingly, we observed that ICP1 with some 308 309 spacers targeting relatively far from PLE (53 and 46kb away) were still able to form robust 310 plaques on PLE 1 (Fig 6a). While all of the spacers selected for this assay had one perfect 311 protospacer match in the chromosome (and have a GA PAM), we identified >100 putative 312 promiscuous target sites for these spacers which would bring the chromosomal target much 313 closer to PLE 1, which may explain these phage's ability to overcome PLE. To test if spacer 314 acquisition had a role in plaque formation, we engineered Cas1 deficient ICP1 in each CRISPR 315 proficient chromosomal targeting phage and assayed for plague formation on the PLE 1 host. 316 Despite being unable to acquire spacers (Supplementary Fig. 3), the phage retained the same plaquing phenotype. We quantified the weaker plaque formation observed when ICP1 targets 317 318 >20 kb away from PLE 1 by measuring plaque size compared to PLE (-) V. cholerae (Fig 6b). As compared to phage with PLE internal and PLE proximal spacers, phage with chromosomal 319 320 spacers targeting >20 kb away from PLE had significantly limited plague size. These results 321 indicate that some PLE-mediated anti-phage activity is retained when CRISPR-Cas is directed 322 at increasing distances from PLE in the small chromosome.

To control for differences in spacer sequences, we also varied the location of the PLE and tested the ability of ICP1 with a single chromosomal spacer targeting the small chromosome to interfere with PLE 1. Following ICP1-mediated transduction, PLE 1 integrates into the *V. cholerae* repeat (VCR) of the new host (12). We collected a pool of PLE 1 transductants where PLE was integrated at varying distances from the chromosomal protospacer and challenged these strains with ICP1. As a control, we determined that all of the
tested PLE 1 *V. cholerae* hosts were susceptible to ICP1 CRISPR-Cas interference when ICP1
possessed a PLE internal spacer (Supplementary Fig. 4); consistent with our earlier finding,
PLE integrated at an increasing distance away from the protospacer was less susceptible to
ICP1-encoded CRISPR interference (Fig. 6c).

333

334 Discussion

Our results reveal that the latest front in the ongoing arms race between contemporary 335 336 isolates of epidemic V. cholerae and its predator ICP1 necessitate the persistence of the ICP1-337 encoded Type I-F CRISPR-Cas system to counter PLE-mediated anti-phage activity (Fig 1). By using a high-throughput spacer acquisition assay, we gained insight into the full range of 338 339 spacers that can combat PLE. Interestingly, our experimental findings on acquisition and 340 interference do not reflect the rather limited diversity of spacers that ICP1 maintains against 341 PLE in nature. These results highlight that not all spacers are equally proficient for interference, and that coupled analysis of these competing mobile genetic elements from nature reveals the 342 evolutionary benefits of a particular complement of spacers more so than laboratory-based 343 344 studies. Despite a lack of clear evidence indicating where the ICP1-encoded CRISPR-Cas 345 system originated, it serves as a tractable model through which we can examine the biology of an endogenous Type I-F CRISPR-Cas system against its cognate foe. 346

Co-culture studies competing phage against CRISPR-Cas proficient bacterial hosts demonstrated that mutational escape by phage is limited by bacterial populations that have heterogenous CRISPR arrays (34). Here, we see that PLE 1 is highly conserved over time, even when co-circulating with CRISPR proficient ICP1. In light of previous suggestions, the diversity of CRISPR arrays in ICP1 populations may limit the success of PLE escape mutants. Surprisingly, however, we see very little diversity in the spacer composition of ICP1 CRISPR arrays with the same minimal spacers being conserved in phage circulating for over eight years (Fig. 1b). Likewise, CRISPR-proficient ICP1 isolated from nature always encoded more than
one spacer against PLE, which would be expected to limit CRISPR escape mutations. It may be
that there is limited room for genetic drift in the PLE genome, permitting ICP1 to streamline its
CRISPR array, keeping only the most efficient spacers while also maintaining an advantageous
genome size.

359 Akin to studies of bacterial Type I-F CRISPR-Cas mediated interference with plasmid transformation and conjugation (35), we similarly see that the spacer sequence and quantity of 360 361 spacers in the array have a role in ICP1's ability to abolish PLE spread (Fig. 2). This may be 362 due to differences in crRNA abundance or stability, or sequence dependent subtleties that dictate interference potential, as has been proposed previously (36). Despite spacer 9's 363 364 improved interference with PLE mobilization compared to spacer 8, we still observed a slight 365 defect in plaque size when comparing engineered phage with only spacer 9 relative to a PLE(-) 366 host (Fig. 6b), suggesting that even this improved spacer alone is not sufficient to fully 367 overcome PLE-mediated anti-phage activity. By encoding a seemingly redundant set of spacers targeting PLE, ICP1 increases its ability to overcome PLE and limit PLE spread in the 368 environment. 369

370 As expected, the majority of spacers acquired in our high-throughput acquisition assay directly target PLE (Fig. 3a). Analysis of natural ICP1 isolates recovered from cholera patient 371 stool samples shows that the phage-encoded CRISPR-Cas system recognizes a GA PAM, 372 373 (Supplementary Fig. 1) which, although atypical for Type I-F systems (37), has been confirmed 374 through single mutations to a C in both positions (13). Notably, we found that ICP1 was able to 375 incorporate spacers that targeted non-canonical PAMs (Fig. 3d) and that these spacers can 376 suffice for PLE interference (Fig. 4b). In comparison to another high throughput spacer acquisition assay in a Type I-F system, which found >90% of all protospacers flanked by the 377 378 canonical PAM (31), it appears that the phage-encoded system is less discriminating with only 70% of protospacers flanked by the expected PAM. However, targeting a protospacer with a 379

380 non-canonical PAM reduced the efficiency of plaquing compared to the canonical PAM (Fig 4b). 381 As such, in nature ICP1 targeting a protospacer with a non-canonical PAM would not be able to completely interfere with PLE and thus would be selected against. This hypothesis is 382 additionally supported by the observation that very few non-canonical PAM protospacers were 383 384 associated with indirect targets in the small chromosome. As these chromosomal spacers are themselves less proficient for interference (Fig. 6a and 6b), the added disadvantage of targeting 385 386 a protospacer with a non-canonical PAM likely tips the balance in favor of PLE, likely explaining 387 the lower abundance of these spacers in our selection experiments.

388 Despite the presence of spacers that target the V. cholerae large chromosome in the 389 high-throughput spacer acquisition assay (Fig. 3c), we show that targeting this chromosome is 390 dispensable for CRISPR interference of PLE (Fig. 5b). Interestingly, two of the natural ICP1 isolates contain a spacer that targets a gene on the V. cholerae large chromosome (Fig. 1b). 391 392 We speculate that this spacer was acquired from a V. cholerae strain possessing a duplication 393 or rearrangement that is not represented in currently sequenced isolates, in which the protospacer was in the small chromosome proximal to PLE, allowing the phage to overcome 394 395 PLE activity. However, this spacer does not seem to be maintained in the phage population, 396 likely due to diminished PLE interference relative to PLE-direct spacers as we experimentally 397 observed.

CRISPR targeting of the V. cholerae small chromosome can overcome PLE, but our 398 399 results suggest a model in which there is a limit to the distance over which processive Cas2-3 400 degradation can occur to reach the PLE prior to excision (Fig. 6d), an action which occurs within 401 five minutes of ICP1 infection that is directed by an early-expressed ICP1 protein (20). The limit 402 of processivity appears to be around a distance of 23 kb (Fig. 6a and 6c), at which point either Cas2-3 is unable to continue to process along the V. cholerae chromosome or PLE excises 403 404 before interference occurs. In vitro studies of Cas3 from Type I-E systems have demonstrated 405 Cas3 translocation velocities of 89 to 300 bases per second and average processivities

406 between 12 to 19 kb (38,39), however, the functional role and limitations of processivity in vivo 407 are not known. Our results are the first indications of Cas2-3 processivity in vivo, with over 22 kb from a distal chromosomal protospacer over which CRISPR-Cas can maintain activity to 408 overcome PLE. As this event must occur within five minutes of ICP1 initiating infection, the 409 410 estimated processivity of ICP1 Cas2-3 is within the range of what has been reported for Type I-E Cas3, which is especially remarkable given the complexity of the crowded intracellular 411 412 environment compared to simplified in vitro systems. In comparison to other Cas nucleases like Cas9, which introduces a single double-413 414 stranded break (40,41), Cas2-3 degrades DNA as it translocates away from the protospacer (17), making it more likely to destroy and thus interfere with its target. This predicted advantage 415 may account for the increased prevalence of Type I systems for phage defense (42). In the 416 context of the battle between ICP1 and PLE, this processivity permits interference even with an 417 indirect CRISPR target and has important implications for harnessing CRISPR-Cas in 418 419 biotechnology and medicine. Since the characterization of the ICP1-encoded CRISPR-Cas system, phage engineered with CRISPR-Cas systems to target virulent, antibiotic resistant 420 bacteria have been assayed for therapeutic applications (43,44), showing the value of 421 422 innovating from natural systems to overcome disparate biological problems. 423

424 Additional Information

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430 Ethics

- 431 The collection of cholera patient stools were approved by the ICDDR,B institutional review
- 432 board. All samples were deidentified and written informed consent was obtained from adult
- 433 participants and from the guardians of children.
- 434 Data Accessibility
- The datasets supporting this article have been uploaded as part of the Supplementary Material.
- 436 Authors' Contributions
- 437 ACM, KNL and KDS carried out the molecular lab work. AA developed and implemented tools
- 438 for sequence data analysis. MA coordinated the collection of clinical specimens. ACM, MA and
- 439 KDS conceived of the study. All authors participated in data analysis. ACM and KDS wrote the
- 440 manuscript with input from all authors and all authors gave final approval for publication.

441 **Competing Interests**

- 442 We declare we have no competing interests.
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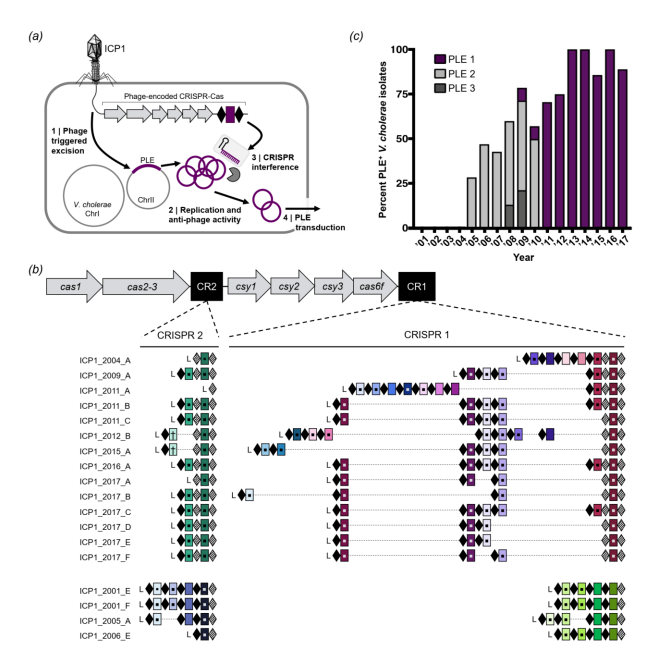
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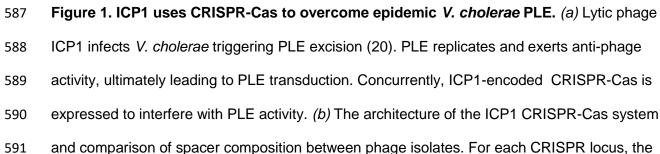
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- 582
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- 584

585 Figures





- repeat (28 bp) and spacer (32 bp) content is detailed as black diamonds and colored rectangles,
- respectively. Repeats that match the repeat consensus (13) are shown in solid diamonds, and
- 594 degenerate repeats are indicated in hatched black diamonds. An AT-rich leader sequence (L)
- 595 precedes each CRISPR locus. Identical spacers shared between isolates are shown as
- 596 rectangles with identical colors. Spacers containing a square target PLE, and spacers
- 597 containing a cross target the V. cholerae large chromosome. (c) Percentage of V. cholerae
- isolates harboring PLE recovered from epidemic sampling at the ICDDR,B over time (n=230
- 599 strains analyzed).

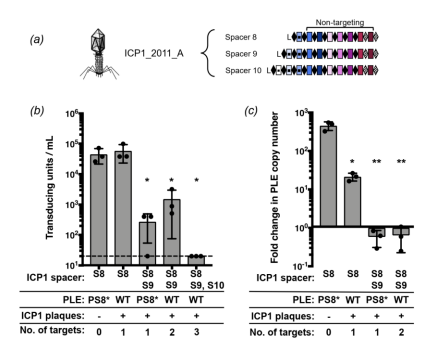
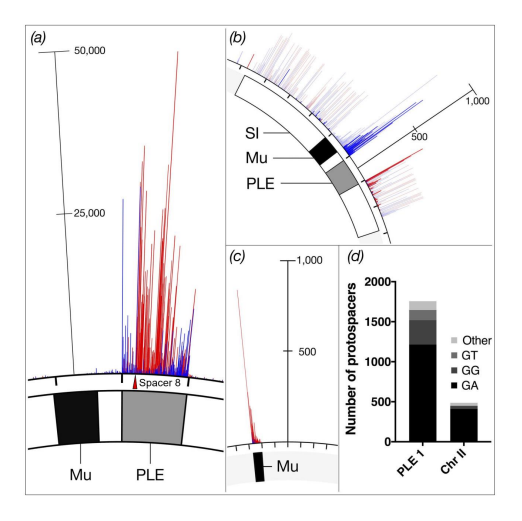


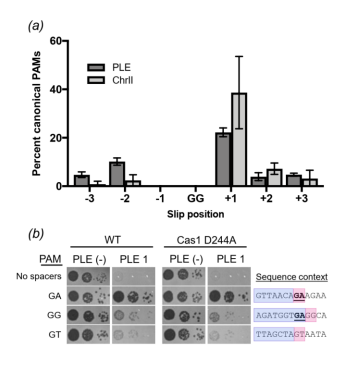


Figure 2. CRISPR can limit horizontal transmission of PLE. (*a*) ICP1_2011_A with anti-PLE spacers S8, S9 and S10 tested in panels b and c. (*b*) PLE transduction after infection with ICP1 with 0,1,2 or 3 spacers. The dashed line indicates the limit of detection for this assay. A single spacer is necessary and sufficient to permit lytic growth of ICP1 on PLE 1 *V. cholerae* as seen by equal plaque formation. (*c*) PLE replication 20 minutes after infection with ICP1 with 0,1 or 2 spacers as determined by qPCR. For panels b and c, error bars indicate standard deviations of biological triplicates. Significance was determined by T Test, * p<0.05, ** p<0.005.



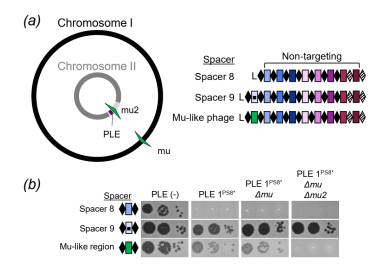
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Figure 3. High-throughput interference driven spacer acquisition mapping. (a) The 611 locations of the ICP1 CRISPR leader-proximal spacer on V. cholerae small chromosome. The 612 613 location of the interference-efficient spacer (S8) is indicated with the red triangle. (b) Spacer locations on the V. cholerae small chromosome (PLE mappings not shown for clarity). Uniquely 614 615 mapped spacers are shown in solid blue or red, while translucent bars show mapping of 616 spacers to all possible locations. (c) Spacer locations on the large chromosome. For panels a, b 617 and c, spacers on the plus and minus strand are indicated in red and blue, respectively. The 618 scale bar measures the number of mapped spacers, and the tick marks around the chromosome are in 18kb intervals. The white box represents the superintegron (SI), the black 619 box is the mu-like region and the grey box is PLE 1. (d) Proportion of unique protospacers with 620 621 a GA or other dinucleotide PAM sequence in PLE or in the small chromosome.



623

Figure 4. Characterizing non-canonical PAMs. (*a*) Frequency of canonical GA PAM +/- 3nt from non-canonical GG PAM across all data sets. (*b*) Tenfold dilutions of ICP1 engineered to contain a spacer that targets PLE 1 with a non-canonical PAM spotted on *V. cholerae* PLE(-) or PLE 1 lawns showing the ability of different phage strains to form plaques (dark spots, zones of killing) (left). Sequence context (right) of the region adjacent to the PAM. The protospacer is boxed in purple and PAM is boxed in pink. The consensus canonical PAM GA is bolded and underlined.



632

633 Figure 5. ICP1 CRISPR-targeting of the small chromosome facilitates PLE interference.

634 (a) Cartoon (left) of the V. cholerae large and small chromosomes. The superintegron is shown

in light grey, the PLE is shown in purple. The two mu-like regions in the large and small

636 chromosome are shown in green arrows. ICP1_2011_A CRISPR variants (right) used to test the

role of targeting sites. (b) Tenfold dilutions of ICP1 with the spacers indicated spotted on V.

638 *cholerae* lawns showing the ability of different phage strains to form plaques.

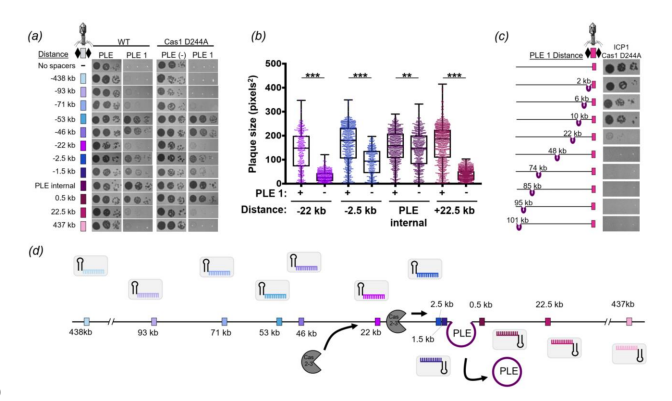
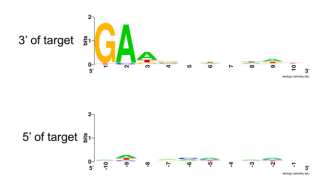




Figure 6. Interference potential of spacers directed to the small chromosome is

642 dependent on proximity to PLE. (a) Tenfold dilutions ICP1 engineered with a chromosomal protospacer +/- Cas1 spotted on lawns of V. cholerae. (b) Plague size of ICP1 variants plated 643 644 with V. cholerae. The distance of the chromosomal protospacer from PLE 1 integration site is 645 indicated. Significance was determined by Mann-Whitney U Test, **p<0.005, ***p<0.0001. (c) 646 Tenfold dilutions of ICP1 engineered with a chromosomal protospacer spotted on lawns of V. cholerae harboring PLE in different locations in the chromosome. (d) Model of race between 647 ICP1 Cas2-3 processive degradation of the V. cholerae chromosome and ICP1-mediated PLE 648 excision. Csy complexes (grey boxes) with crRNAs (colored) search for a complementary 649 650 protospacer (colored rectangles, experimentally assessed in panel a). Cas2-3 (dark grey) is recruited to the protospacer and processively degrades the DNA towards PLE (purple). ICP1 is 651 able to form plaques when Cas2-3 degrades PLE before PLE excises from the chromosome, 652 which occurs within 5 minutes of ICP1 infection. 653

- 654
- 655



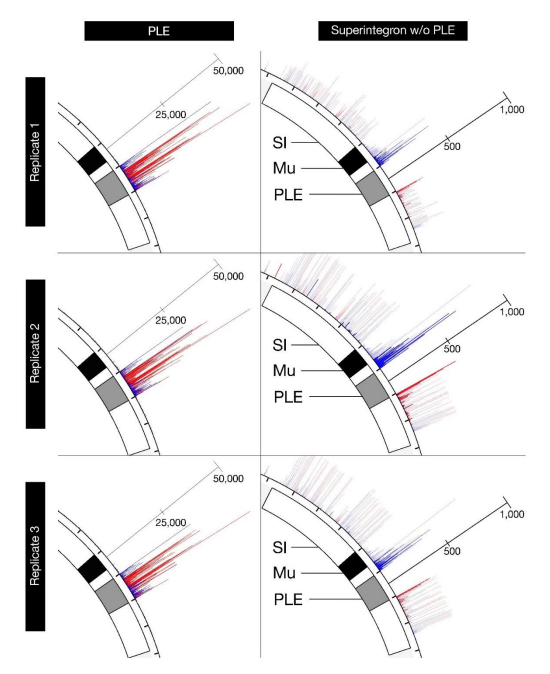
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657 Supplementary Figure 1. Sequence logo of PAMs of natural ICP1 isolates. The PAM

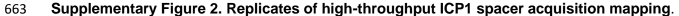
658 sequence of the ICP1-encoded CRISPR-Cas system. Alignment of flanking sequence of all

known targets of spacers found in natural ICP1 isolates. Sequence logos were generated using

- 660 WebLogo (45).
- 661



662



664 Spacer locations of the most leader-proximal spacers on the plus and minus strand are 665 indicated in red and blue, respectively. Uniquely mapped spacers are shown in solid blue or red,

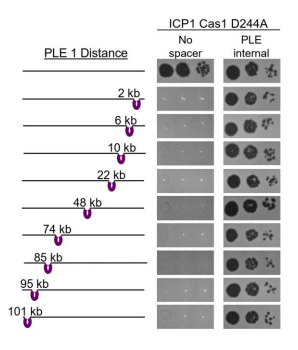
- 666 while translucent bars show mapping of spacers to all possible locations. The scale bars
- 667 measure the number of mapped spacers, and the tick marks around the chromosome are in
- 18kb intervals. Replicate 2 is the same data as in Figure 3a and b.
- 669

a)		(b)	
Unconserved	1 2 3 4 5 6 7 8 9 10 Conserved		
ICP1 P. atrosepticu P. aeruginosa Y. pestis Consistency	10 20 30 40 50 MQ KQIJTEQKRN MYILSRCKVL VKNGQVCHLH ED ED FOUTT MDNPFSPSDL KTILHSKRAN VYYLQHCRIL VNGGRVEYVT EEGNQSLYWN MDD-ISPSEL KTILHSKRAN LYYLQHCRVL VNGGRVEYVT EGNQSLYWN MENATHSSDL KTILHSKRAN LYYLQHCRVL VNGGRVEYVT DEGRHSHYWN MENATHSSDL KTILHSKRSN IYYLEYCRVL VNGGRVEYVT DEGRQSLYWN 630322535 6*57857 6*67889* 77777585 7852273677	olaquing	10 ⁻² 10 ⁻³ 10 ⁻⁴
ICP1 P. atrosepticu P. aeruginosa Y. pestis Consistency	60 70 80 90 100 VPYANTVFIG LAEGTSITNE AMSMLAANGV IVFWTKGGGY DMFAAD 1 IPIANTSVVM LGTGTSVTQA AMREFARACV MVGFCGGGGT PLFAANEAEV 1 IPIANTSIL LGTGTSITQA AMREFARACV LVGFCGGGGT PLFAANEAEV 1 IPIANTTVIM LGTGTSVTQA AMREFARACV LVGFCGGGGT PLFAANDVEV 9 6***5384 *76***9*76 **666*666**7*5766***6 68*8*73335	Efficiency of plaquing	
ICP1 P. atrosepticu P. aeruginosa Y. pestis Consistency	110		10 ⁻⁸ WT Cas1 D244A
ICP1 P. atrosepticu P. aeruginosa Y. pestis Consistency	160		
ICP1 P. atrosepticus P. aeruginosa Y. pestis Consistency	210		
ICP1	260. Z270. 280. 290. 300		
	IAVLHCKTRR GGLVFDVADI IKISILDEDK FIAMBCURSK TEFRAV LAVLHCKTRR GGLVFDVADI IKDSLIDPQA FIAMBCDEE RDFRQACIDN LSVLHCKTRR GGLVFDVADI IKDSLIDPQA FIAMQCDEE QEFRQRCISG 758***7*** ****7****8 **6789**6* *48*56*467 58*875*742		
ICP1 P. atrosepticu P. aeruginosa Y. pestis Consistency	FDKNDI AYL INNIKRICME NSDVHQE 7 FQQSEALDVM IGSLQDVAST LSQVGR- LSRAQALDFM IDTLKDVAQR STVSA FQRTEALDVM IDGIKETAAL CSQVPR- 756466*648 *548755643 2746230		

670

671 Supplementary Figure 3. Conserved residue in Cas1 is necessary for spacer acquisition.

- (a) Praline alignment (46) of Cas1 from ICP1 and other Type I-F systems in the organism
- 673 indicated. Black arrow indicates conserved residue mutated in ICP1. (b) Spacer acquisition was
- 674 measured by the number of plaques on a PLE(+) host without a protospacer relative to the
- number of plaques on a PLE(+) host with a functional protospacer. Dashed line indicates level
- of detection. Error bars indicate standard deviation of three independent replicates.





- 679 Supplementary Figure 4. PLE transduced in unique sites in *V. cholerae* chromosome can
- 680 **be overcome by ICP1-CRISPR mediated interference.** Tenfold dilutions ICP1 without any
- spacer or with a PLE targeting spacer spotted on lawns of *V. cholerae*. Transductants used are
- the same as in Figure 6c.