CRISPR-Cas Controls Cryptic Prophages

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ABSTRACT

The bacterial archetypal adaptive immune system, CRISPR-Cas, is thought to be non-functional in the best-studied bacterium, *Escherichia coli* K-12. Instead, we demonstrate here that the *E. coli* CRISPR-Cas system is active and inhibits its nine defective (i.e., cryptic) prophages. Specifically, deactivation of CRISPR-Cas via deletion of *cas2*, which encodes one of the two conserved CRISPR-Cas proteins, reduces growth by 40%, increases cell death by 700%, and prevents persister cell resuscitation; hence, CRISPR-Cas serves to inhibit the remaining deleterious effects of these cryptic prophages. Consistently, seven of the 13 *E. coli* spacers contain matches to the cryptic prophages, and, after excision, CRISPR-Cas cleaves cryptic prophage CP4-57 and DLP-12 DNA. Moreover, we determine that the key genes in these cryptic prophages that CRISPR-Cas represses by cleaving the excised DNA include lysis protein YdfD of Qin and lysis protein RzoD of DLP-12. Therefore, we report the novel results that (i) CRISPR-Cas is active in *E. coli* and (ii) CRISPR-Cas is used to tame cryptic prophages; i.e., unlike with active lysogens, CRISPR-Cas and cryptic prophages may stably exist.

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INTRODUCTION

Along with restriction/modification¹ and toxin/antitoxin (TA) systems², prokaryotes utilize clustered, regularly-interspaced, short palindromic repeats (CRISPR) and CRISPR-associated (Cas)³ proteins to combat phages. These systems are interrelated in that we found some Cas proteins are derived from TA systems; for example, *Sulfolobus solfataricus* Cas2 is structurally similar to antitoxin GhoS of the *Escherichia coli* GhoT/GhoS TA system⁴. In addition, in a manner similar to our discovery² that toxins of TA systems inhibit phage by degrading host mRNA after the toxin is activated by the attacking phage when the phage shuts down transcription (e.g., Hok/Sok inhibits T4 phage), some Cas proteins induce host dormancy rather than degrading phage DNA to inhibit phage propagation⁵. Also, TA systems have been found to stabilize CRISPR-Cas systems by making them addictive to the host⁶.

Although CRISPR-Cas systems exclude both external lytic and temperate (lysogenic) phages⁷, CRISPR-Cas systems of lysogens that target their own integrated prophages decrease long-term fitness and either the cell dies or the phage is lost^{7,8}. Also, the class I-E³ CRISPR-Cas system of *E. coli* is not related to immunity for external phages⁹ and is thought to be inactive in the wild-type strain¹⁰, due to repression by H-NS, although it is functional when induced¹¹. To date, the relationship of CRISPR-Cas to cryptic prophages; i.e., those phage remnants that are unable to form lytic particles, has not been investigated.

Up to 20% of bacterial genomes may contain stable phage DNA¹², and for *E. coli*, we discovered that its nine cryptic prophages are not genomic junk but instead encode genes for proteins that increase resistance to sub-lethal concentrations of quinolone and β -lactam antibiotics as well as protect the cell from osmotic, oxidative, and acid stresses¹³. Although these cryptic prophages do not help a subpopulation of cells weather extreme stress by entering into the dormant state known as persistence¹⁴, these phage remnants facilitate the resuscitation of these persister cell via nutrient sensing¹⁵. Therefore, the bacterial cell can capture the genome of its former parasite to both combat stress¹³ as well as to revive from dormancy¹⁵.

Here we explore the role of the native *E. coli* CRISPR-Cas systems in the regulation of cryptic prophages. We find that the CRISPR-Cas system is required for inhibiting these phage fossils since, if CRISPR-Cas is inactivated, cells die due to activation of the lysis proteins YdfD of Qin and RzoD from

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DLP-12. Hence, we discovered CRISPR-Cas is active in *E. coli* and serves to regulate its phage fossils.

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RESULTS

CRISPR-Cas increases growth. To inactivate CRISPR-Cas in *E. coli*, the *cas2* deletion was chosen since Cas2 is conserved in almost all CRSIPR-Cas systems¹⁶ as part of the adaptor complex and serves as the structural scaffold of the Cas1/Cas2 complex which cleaves the source, protospacer DNA, and the CRISPR array¹⁷. We found the *E. coli* K-12 *cas2* mutant grows 40% slower in rich medium compared to the wild-type strain (specific growth rate of $0.79 \pm 0.21/h$ vs. $1.3 \pm 0.11/h$, respectively). Similarly, in minimal glucose (0.4 wt%) medium, deletion of *cas2* also reduces growth by 33% (0.42 \pm 0.02/h vs. $0.62 \pm$ 0.02/h, respectively).

CRISPR-Cas increases single-cell resuscitation. Moreover, since CRSIPR-Cas increases growth, we tested for its effect on persister cell resuscitation using single-cell microscopy. Persister cell resuscitation is germane in that the dormant cells are highly stressed and have limited resources for their revival via activation of hibernating ribosomes^{18,19}; for example, we have shown inhibiting ATP synthesis leads to a 5,000-fold increase in persister cell formation²⁰. Since we discovered facile means for converting the whole population of cells into persister cells¹⁸⁻²⁰ that has been used by at least 17 independent labs to date with various bacterial species²¹, these stressed cells are an excellent model for testing the effects of CRISPR-Cas on *E. coli* physiology.

Here, we found inactivation of CRISPR-Cas via *cas2* nearly completely prevents persister cell resuscitation, while the wild-type cell has 52% resuscitation (**Fig. 1A, Table S1**). Hence, CRSIPR-Cas is active in *E. coli* and plays key roles in its growth and recovery from extreme stress.

CRISPR-Cas prevents cell death. To explore how CRISPR-Cas increases growth (including persister cell resuscitation), we checked for death in resuscitating persister cells of the *cas2* deletion strain using the Live/Dead stain. We found that inactivating CRISPR-Cas leads to a 7-fold increase in death in resuscitating cells (**Fig. 1B, Table S2**). In addition, there were 34-fold more cells termed "ghosts" that lack cytosolic material (**Fig. S1**) and are likely dead but have intact membranes so are not stained by the propidium iodide

dye we use. Corroborating these results, there was 11-fold and 5-fold more death for stationary- and exponential-phase cells, respectively, when CRISPR-Cas was inactivated. These results indicate inactivating CRISPR-Cas leads to cell death.

CRISPR-Cas increases growth by repressing cryptic prophage genes. We hypothesized that since CRISPR-Cas systems inhibit some phage lysogens^{7,8}, the *E. coli* system may be preventing cell death by repressing some of the cryptic prophage genes. To test this hypothesis, we first examined the *E. coli* CRISPR-Cas system for spacers related to the nine cryptic prophages. *E. coli* K-12 CRISPR-Cas contains 13 spacers^{11,23}, each containing 32 or 33 nt²⁴, between *iap* and *cas2*. Between the 14 29-nt repeat sequences (5'-GTGTTCCCCGCATCAGCGGGGATAAACCG), we found that six of the 13 spacers contain 12 to 16 nt of perfect matches to seven of the nine cryptic prophages (DLP-12, CP4-57, CPS-53, CP4-6, Rac, Qin, and e14) (**Fig. 2A**). In general, spacer lengths vary from 21 to 72 nt²⁵ with perfect complementarity of 6 to 12 nt¹⁰. Furthermore, the two predicted protospacer adjacent motif (PAM) sites for cryptic phage DLP-12, 5'-GAA (consensus I-E site) and 5'CAT (**Fig. S2**), support rapid and reduced interference in *E. coli*, respectively²⁶. Moreover, we found these spacers cannot be deleted, which shows they are essential for *E. coli* K-12. Together, these results suggest CRISPR-Cas regulates the *E. coli* cryptic prophages.

The presence of the cryptic prophage-related spacers and the cell death seen upon inactivating CRISPR-Cas suggested that *E. coli* CRISPR-Cas suggested may be repressing some of the cryptic prophage lysis genes; hence, we checked for derepression of the cryptic prophage lysis genes in these seven cryptic prophages, specifically *ydfD* (Qin), *hokD* (Qin), *ypjF* (CP4-57), *essD* (DLP-12), and *rzoD* (DLP-12) via quantitative reverse transcription polymerase chain reaction (qRT-PCR). We found that inactivation of CRISPR-Cas via the *cas2* deletion results in activation of *ydfD* by 35-fold and activation of *rzoD* by 183-fold in resuscitating persister cells (**Fig. 1C**). Moreover, *rzoD* was activated 1024-fold in exponentially-growing cells (**Fig. 1C**). RzoD is a putative DLP-12 lysis lipoprotein that we previously showed was toxic through its interaction with toxin Hha²⁷ of the Hha/TomB TA system²⁸. YdfD of Qin cryptic prophage has been shown to lyse cells when induced²⁹. Hence, CRSIPR-Cas represses at least two *E. coli* proteins RzoD and YdfD that can reduce cell growth.

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Excision of DLP-12 is not regulated by CRISPR-Cas. Since inactivation of CRSIPR-Cas leads to extraordinary derepression of the DLP-12 *rzoD* lysis gene, we checked for increased excision with the *cas2* deletion strain using qPCR. We found the *cas2* deletion has little impact on DLP-12 excision. We also tested the effect of CRISPR-Cas on two other cryptic prophages with significant excision (CP4-57 and e14)¹³, and also found no effect of deleting *cas2* (Table S3). Hence, CRISPR-Cas does not affect cryptic prophage excision.

CRISPR-Cas cleaves DLP-12 and CP4-57 DNA after excision. Since DLP-12 has significant excision in *E. coli*¹³ but inactivation of CRISPR-Cas does not alter its excision, we hypothesized that CRISPR-Cas repressed lysis gene *rzoD* by cleaving DLP-12 DNA after excision. Using qRT-PCR with resuscitating persister cells, we found that the *cas2* deletion leads to 40-fold increase in DLP-12 DNA that includes spacer 3 as well a 184-fold increase in CP4-57 DNA that includes spacer 6 (**Fig. 2B**). Therefore, CRISPR-Cas actively cleaves excised cryptic prophage DNA to regulate the lysis genes of the captured phage foe.

DISCUSSION

Our results reveal a new role for CRISPR-Cas systems: regulation of phage fossils. The evidence for this includes that inactivating CRISPR-Cas by deleting *cas2* (i) reduces growth by 40%, (ii) nearly eliminates resuscitation from the persister state (**Fig. 2**), (iii) causes ghost cell formation and cell death (**Fig. 1**), (iv) derepresses the cryptic prophage lysis genes *ydfD* and *rzoD*, and (v) increases CP4-57 and DLP-12 cryptic prophage DNA after their excision. Since there is no change in excision of DLP-12 upon inactivating CRISPR-Cas, our results show the mechanism for regulating the lysis genes *ydfD* and *rzoD* of the cryptic prophages is via CRISPR-Cas cleavage of the excised prophages. Supporting this, the *E. coli* K-12 class 1-E CRISPR-Cas system cleaves dsDNA³⁰. What remains to be determined is the mechanism by which CRISPR-Cas cleaves only the excised cryptic prophage rather than genomic DNA; perhaps this is related to the relatively inefficient PAM and seeding regions for some of the spacers (**Fig. S2**).

This new function for CRISPR-Cas is likely general and may explain why many species appear to have inactive CRISPR-Cas systems as was previously thought for *E. coli*¹⁰; i.e., instead of protecting cells from

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external phages, CRISPR-Cas systems may also control resident cryptic prophages which are prevalent. Critically, our results provide the first example where it is beneficial for the host to have an active CRISPR-Cas system that targets *inactive* integrated phages (i.e., cryptic prophages) since previous reports show targeting *active* temperate phages is deleterious; i.e., either the cell dies or the phage is lost^{7,8,31}. Since *E. coli* cryptic prophages like rac have been present in its genome for 4.5 million years³², the active K-12 CRISPR-Cas system is stable with the cryptic prophages; in fact, there has been little change in the *E. coli* spacers for at least 42,000 years³³.

Our results also indicate that, although the cryptic prophages are stable and the cell makes use of the genetic tools encoded by its former foe to combat myriad stresses¹³ and to sense nutrients prior to exiting the persister state¹⁵, the source of these tools must be elegantly regulated by CRISPR-CAS since they often harbor deleterious membrane lysis proteins like YdfD and RzoD. Similarly, host Rho has been found recently to silence cryptic prophage toxin/antitoxin systems through transcription termination³⁴, and H-NS silences cryptic prophages through 65 binding sites³⁵. Therefore, phages may be captured by the host, but they must be tamed, and this now includes repression by CRISPR-Cas.

MATERIALS AND METHODS

Bacteria and growth conditions. Bacteria (**Table S4**) were cultured in lysogeny broth³⁶ at 37°C. pCA24N-based plasmids³⁷ were retained in overnight cultures via chloramphenicol (30 μ g/mL), and kanamycin (50 μ g/mL) was used for deletion mutants, where applicable.

Spacer knockout. To attempt to delete the CRISPR-Cas spacer region, the one-step inactivation method for single gene deletions³⁸ was utilized in which primers that included the sequences flanking the spacer region (**Table S5**) were used with plasmid pKD4 to allow for insertion of the kanamycin resistance cassette (1.5 kb) and FRT sites.

Persister cells. Exponentially-growing cells (turbidity of 0.8 at 600 nm) were converted nearly completely to persister cells^{18,20} by adding rifampicin (100 μg/mL) for 30 min to stop transcription, centrifuging, and adding LB with ampicillin (100 μg/mL) for 3 h to lyse non-persister cells. To remove ampicillin, cells were

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washed twice with 0.85% NaCl then re-suspended in 0.85% NaCl. Persister concentrations were enumerated via a drop assay³⁹.

Single-cell persister resuscitation. Persister cells (5 μL) were added to 1.5% agarose gel pads containing M9 glucose (0.4 wt%) medium⁴⁰, and single-cell resuscitation was visualized at 37°C via a light microscope (Zeiss Axio Scope.A1, bl_ph channel at 1000 ms exposure). For each condition, at least two independent cultures were used with 150 to 300 individual cells used per culture.

Membrane integrity assay. To determine membrane integrity, the persister cells were analyzed with the LIVE/DEAD BacLight Bacterial Viability Kit (Molecular Probes, Inc., Eugene, OR, catalog number L7012). The fluorescence signal was analyzed via a Zeiss Axioscope.A1 using excitation at 485 nm and emission at 530 nm for green fluorescence and using excitation at 485 nm and emission at 630 nm for red fluorescence.

qRT-PCR. To quantify transcription from the cryptic prophage lytic genes, RNA was isolated from persister cells that were resuscitated by adding M9 glucose (0.4%) medium for 10 min then washed with 0.85% NaCl and from exponential cells grown to a turbidity of 0.8; samples were cooled rapidly using ethanol/dry ice in the presence of RNA Later. RNA was isolated using the High Pure RNA Isolation Kit (Roche). The following qRT-PCR thermocycling protocol was used with the iTaqTM universal SYBR® Green One-Step kit (Bio-Rad): 95 °C for 5 min; 40 cycles of 95 °C for 15 s, 60 °C for 1 min for two replicate reactions for each sample/primer pair. The annealing temperature was 60°C for all primers (**Table S5**).

qPCR. To quantify prophage excision and the levels of DNA flanking the CRISPR-Cas cleavage sites, total DNA (100 ng) was isolated from exponentially-growing and persister resuscitating cells using an UltraClean Microbial DNA Isolation Kit (Mo Bio Laboratories). Excised cryptic prophage was quantified using primers for each prophage excisionase (**Table S5**) that only yield a PCR product upon prophage excision, and the relative amount of each target gene was determined using reference gene *purM*. The level of cryptic prophage flanking the CRISPR-Cas cleave site was quantified using primers that flank each site (**Table S5**). The qPCR reaction performed using CFX96 Real Time System. The reaction and analysis was conducted using the StepOne Real-Time PCR System (Bio-Rad).

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FIGURE LEGENDS

- **Fig. 1. Inactivating CRISPR-Cas eliminates persister cell resuscitation by activating cryptic prophage lytic proteins, causing cell death.** (**A**) Single cell persister resuscitation for wild-type BW25113 and the *cas2* mutant after 6 hours on 0.4 wt% glucose. Black arrows indicate cells that resuscitate, and the scale bar indicates 10 μm. Cells were observed using light microscopy (Zeiss Axio Scope.A1). Representative results from two independent cultures are shown, and tabulated cell numbers are shown in **Table S1**. (**B**) LIVE/DEAD staining of resuscitating persister cells shows the *cas2* mutation causes cell death. DF is dark field, SYTO9 is a membrane permeable stain for nucleic acids (green), and PI is propidium iodide, which is a membrane impermeable stain for the nucleic acids of dead cells (red). Tabulated cell numbers are shown in **Table S2**. (**C**) The *cas2* mutation derepresses cryptic prophage lysis genes *ydfD* (in resuscitating persister cells) and *rzoD* (in both resuscitating persister cells and exponentially-growing cells). Lytic genes from five cryptic prophages were checked by qRT-PCR: *ydfD* (Qin), *hokD* (Qin), *ypjF* (CP4-57), *essD* (DLP-12), and *rzoD* (DLP-12).
- **Fig. 2.** Inactivating CRISPR-Cas increases excised cryptic prophage DNA. (A) The 14 repeat (R, hexagon) and 13 spacer (squares) sequences of the CRISPR-Cas system (from the *iap* to *cas2* part of the *E. coli* genome) showing the cryptic prophage spacer matches (red text) and prophage DNA protospacer sequences (blue text) which includes matches to seven of the nine cryptic prophages (DLP-12, CP4-57, CPS-53, CP4-6, rac, Qin, and e14). The potential PAM sequences are shown in **Fig. S2**. (B) The *cas2* mutation increases cryptic prophage excised DNA that flanks the cleavage sites, as determined by qPCR.

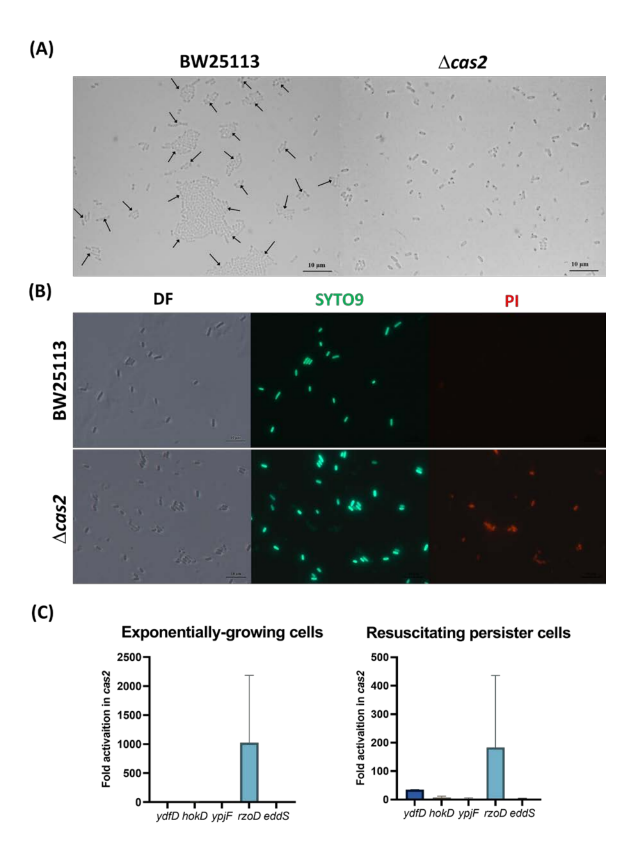
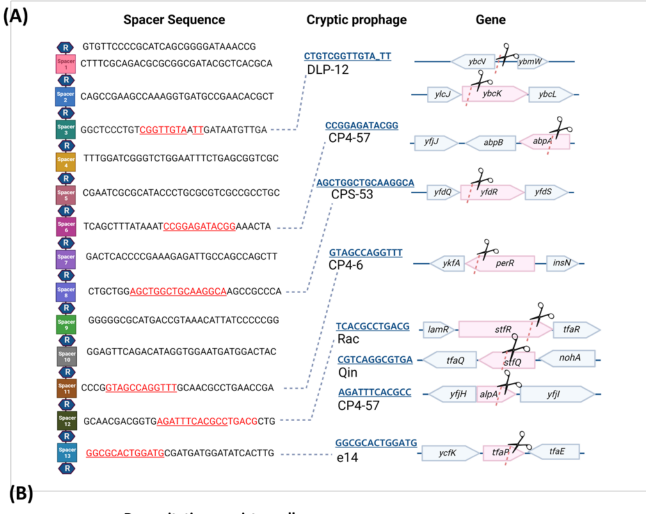


Figure 1



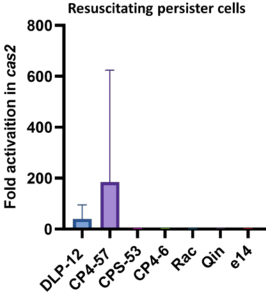


Figure 2

Supporting Information

CRISPR-Cas Controls Cryptic Prophages

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Table S1. Inactivating CRISPR-Cas eliminates persister cell resuscitation on glucose agarose gel pads. Single persister cells were observed using light microscopy (Zeiss Axio Scope.A1). The total number and waking number of persister cells are shown after 6 hours on 0.4 wt% glucose. Fold-change in waking is relative to BW25113. These results are the combined observations from two independent experiments (independent culture results separated by "/"), and standard deviations are shown. The microscope images are shown in **Fig. 1.**

	Total cells	Waking cells	% waking	Fold- change
BW25113	504/382	199/248	52 ± 18	1
$\Delta cas 2$	516/677	6/15	1.7 ± 0.7	-30.9

Table S2. Inactivating CRISPR-Cas causes cell death and ghost cell formation. Persister cells were washed with PBS twice, resuscitated by M9 0.4% glucose for 10 min and stained with LIVE/DEAD reagents. Exponential cells were grown to a turbidity of 0.8 (at 600 nm) and stationary cells were grown to turbidity of 2.0. The ghost cells in the persister population were visualized using a Zeiss Axioscope.A1microscope. The results are the combined observations from two independent experiments (independent culture results separated by "/"). The microscope images are shown in **Fig. 1B**, **Fig. S1**, and **Fig. S3**.

	Strains	Total cells	Dead cells	Ghost cells	% Ghost cells	fold- change ghost cells	% dead	fold- change dead cells
Resuscitated	BW25113	52/115	2/7	0/1	0.6	1	5.4	1
	$\Delta cas 2$	142/106	54/38	38/13	20.6	34	37.1	6.9
Exponential	BW25113	153/147	0/2	-	-	-	0.7	1
	$\Delta cas 2$	59/40	1/2	-	-	-	3.4	4.9
Stationary	BW25113	492/123	1/0				0.1	1
	∆cas2	1136/1736	14/18	-	-	-	1.13	11.2

Table S3. CRISPR-Cas does not affect cryptic prophage excision in stationary cells.

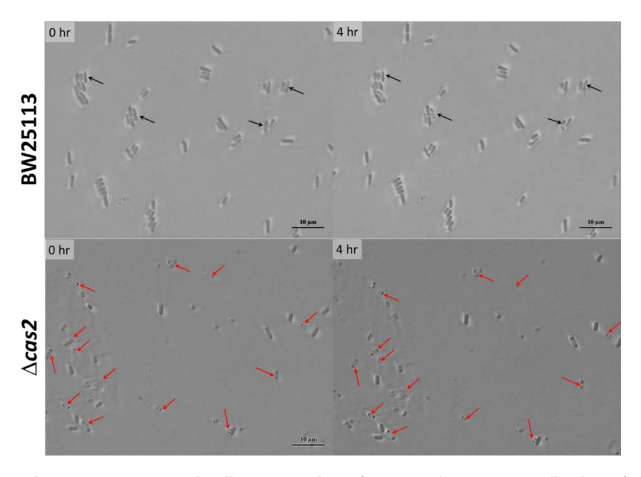
Gene	(house]	<i>rM</i> keeping ne)	CP ²	1 -57	e14		DLP-12	
Strain	WT	Cas2	WT	cas2	WT	cas2	WT	cas2
СТ	9.30	9.70	30.45	30.37	16.30	17.45	30.58	30.05
<u></u>	± 0.13	± 0.18	± 1.73	± 0.68	± 0.06	± 0.51	± 0.49	± 0.5
ΔCΤ			21.16	20.67	7.00	8.16	21.29	20.35
ΔС1			± 1.73	± 0.70	± 0.15	± 0.54	± 0.51	± 0.53
ΔΛСΤ				-0.48		1.16		-0.94
				± 0.7		± 0.54		±0.53
fold				1.40		-2.23		1.92

Table S4. E. coli bacterial strains and plasmids utilized.

Strains and Plasmids	Features	Source
Strains		
BW25113	rrnB3 Δ lacZ4787 hsdR514 Δ (araBAD)567 Δ (rhaBAD)568 rph-1	41
BW25113 Δcas2	$\Delta cas2$, Km ^R	41
Plasmids		
pCA24N	Cm^R ; $lacI^q$	42
pCA24N_cas2	Cm^R ; $lacI^q$, P_{T5-lac} :: $cas2^+$	42
pKD4	FRT::Kan ^R ::FRT, Amp ^R	38

Table S5. Primers used in this study for qPCR and qRT-PCR. * indicates excision primers.

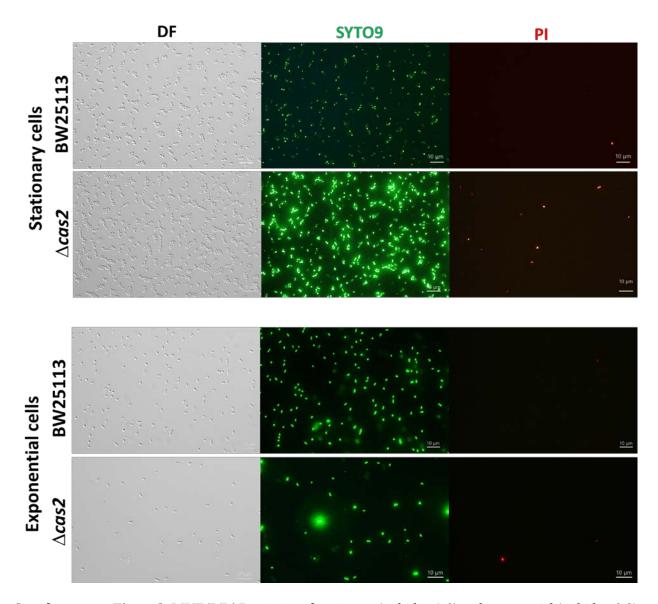
Gene	Sequence (5'-'3)
Spacer knockou	
DS	F:TTATGCGGATAATGCTACCTCTGGTGAAGGAGTTGGCGAAGGCGTCTTGA
	GTGTAGGCTGGAGCTGCTTC
	R:CCCCGGTAGATTTGGATGGTTTAAGGTTGGTGTCTTTTTTACCTGTTTGA
	CATATGAATATCCTCCTTAG
Spacer	F: GAAGGAGTTGGCGAAGGCGTCTTGA
	R: GTTGGTGTCTTTTTACCTGTTTGA
qRT-PCR	
purM	F: CGGTGTTGATATTGACGCGGG
	R: CAGCACGGGTTCACGATATTTTTG
ydfD (Qin)	F: TCAGCATTTGTGCTTGTTCTG
	R: CTGCCGGGATTTCGATATTA
hokD (Qin)	F: GCCCTGATCGTCATCTGTTT
	R: AGCTGTGAAGACAGCGACCT
<i>ypjF</i> (CP4-57)	F: TATGGCCTGACACTGAACGA
	R: GTTGTTGCCACGAAAATCCT
rzoD (DLP-12)	F: TGCTCTGCGTGATGATGTTG
	R: TCTCTCTGAGGGTGAAATAATCC
essD (DLP-12)	F: CTCACAGTGGGCAGCAATAG
	R: TATTCACCTCTCGCAGCCTT
qPCR	
*CP4-57	F: AAGCATGTAGTACCGAGGATGTAGG
	R: TATGTCTCCTCACCGTCTGGTCGG
*e14	F: GTGCAAACATCGGTGACGAA
	R: TTCAGCAGCTTAGCGCCTTC
*DLP-12	F: CAAAAGCCATTGACTCAGCAAGG
	R: CGGATAAGACGGCATAAATGA
purM 400	F: CTGATTGCACTCGGTTCCAG
1	R: CGTTTTCACCGTTGGCATTG
Spacer 3-1	F: CATGCAATTACAACATCAGGGTA
~ [R: GCCATTGGTAAAACCTTCCA
Spacer 3-2	F: AAATTCCGAAAAAGCTCCTGA
~	R: TGCTTTCAAGATTATGGCGTA
Spacer 6	F: CACGAAAGCCAGCCTATTCC
Spacer o	R: CCGCTGTTTCTCCCAGG
Spacer 8	F: GCCAGCATAATGAGATCGGC
Бриссі б	R: TTTTACCCAAACTCAGCGCG
Spacer 11	F: TGTGCAGTTGTACCAGTGGA
Spacer 11	R: CAACCCAGCAAAGTTTCGGA
Spacer 12-1	F: TGTATGTGCCCCGGTGTTAT
Spacer 12-1	R: GCAGATGAAGGCGCATTACA
Spacer 12-2	F: AGCTTTACACCTCGGCTCAT
Spacer 12-2	R: CCGGAACTCTTGTGTTGGTG
Spacer 12 2	
Spacer 12-3	F: TCTACAGGGAAAGGACGACC
Cmagar 12	R: CTCTGCAACCAAGTGAACCA
Spacer 13	
Spacer 13	F: ACAACCACTATCGCCCCTTT R:GGTAAGGCTGCATTGGGAAG



Supplementary Figure 1. Single cell persister waking of BW25113 $\Delta cas2$. Persister cell waking of BW25113 and BW25113 $\Delta cas2$ on M9 0.4% glucose agar plates incubated at 37°C for 4 hours. Black arrows indicate waking cells, and red arrows indicate ghost cells. Scale bars indicate 10 μ m. Representative results from two independent cultures are shown.

```
Spacer 3:
          5'-GGCTCCCTGTCGGTTGTAATTGATAATGTTGA-'3
       3'-ATGTATGTAGACAGCCAACATAAAGGGAGGTCTTA-'5:DLP-12
Spacer 3:
          5'-GGCTCCCTGTCGGTTGTAATTGATAATGTTGA-'3
        3'-TTCAGAAGAGACAGCCAAATTACACAGCGGTCTGT-'5: DLP-12
Spacer 6: 5'-TCAGCTTTATAAATCCGGAGATACGGAAACTA-'3
       3'-TGTAACGGCAGTCTAACGGCCTCTATGCCCTAGTC-'5:CP4-57
          5'-CTGCTGGAGCTGGCTGCAAGGCAAGCCGCCCA-'3
               3'-GACGCCGTGGTCGACCGACGTTTCGTGGCGTACAA-'5:CPS-53
Spacer 11: 5'-CCCGGTAGCCAGGTTTGCAACGCCTGAACCGA-'3
                3'-AGTAAGTCATCGGTCCAAAACTTGTGTAGTTCGCG-'5:CP4-6
Spacer 12: 5'-GCAACGACGGTGAGATTTCACGCCTGACGCTG-'3
                 3'-GAGTTAGGTAGGCTGTCTAAAGTGCGGCTCGTAAG-'5:CP4-57
Spacer 12: 5'-GCAACGACGGTGAGATTTCACGCCTGACGCTG-'3
                         3'-GGTCGCTGCGTATAGTCCCTAGTGCGGACTGCCTA-'5:Rac
Spacer 12: 5'-GCAACGACGGTGAGATTTCACGCCTGACGCTG-'3
       3'-GGTCGCTGCGTATAGTCCCTAGTGCGGACTGCCTA-'5:Qin
Spacer 13: 5'-GGCGCACTGGATGCGATGATGGATATCACTTG-'3
       3'-AGACCGCGTGACCTACATAGTTGGTCAAGGTCACG-'5 :e14
```

Supplementary Figure 2. Predicted base-pairing interactions between each of the CRISPR spacers (i.e., spacers 3, 6, 8, 11, 12, and 13) and a protospacer within the cryptic phages (DLP-12, CP4-57, CPS-53, CP4-6, CP4-57, Rac, Qin, and e14). Note, some spacers match more than one region within a particular cryptic prophage. The PAMs are indicated by bold/green highlighted text, and yellow highlight indicates active PAMs²⁶ (3'TTC; i.e., 5'GAA, is the consensus PAM for *E coli*). The complementary nt between the spacers and cryptic prophage protospacers is indicated by blue text and vertical lines.



Supplementary Figure 3. LIVE/DEAD staining of stationary (turbidity 2.0) and exponential (turbidity 0.8) cells shows the *cas2* mutation causes cell death. DF is dark field, SYTO9 is a membrane permeable stain for nucleic acids (green), and PI is propidium iodide, which is a membrane impermeable stain for the nucleic acids of dead cells (red). Tabulated cell numbers are shown in **Table S2**.