

1 **Cryptic prophage-encoded small protein DicB protects *Escherichia coli* from phage**
2 **infection by inhibiting inner membrane receptor proteins**

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9 Running title: DicB inhibits mannose phosphotransferase system proteins

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14 **Abstract**

15 Bacterial genomes harbor cryptic prophages that have lost genes required for induction,
16 excision from host chromosomes, or production of phage progeny. *Escherichia coli* K12 strains
17 contain a cryptic prophage Qin that encodes a small RNA, DicF, and small protein, DicB, that
18 have been implicated in control of bacterial metabolism and cell division. Since DicB and DicF
19 are encoded in the Qin immunity region, we tested whether these gene products could protect
20 the *E. coli* host from bacteriophage infection. Transient expression of the *dicBF* operon yielded
21 cells that were ~100-fold more resistant to infection by λ phage than control cells, and the
22 phenotype was DicB-dependent. DicB specifically inhibited infection by λ and other phages that
23 use ManYZ membrane proteins for cytoplasmic entry of phage DNA. In addition to blocking
24 ManYZ-dependent phage infection, DicB also inhibited the canonical sugar transport activity of
25 ManYZ. Previous studies demonstrated that DicB interacts with MinC, an FtsZ polymerization
26 inhibitor, causing MinC localization to mid-cell and preventing Z ring formation and cell division.
27 In strains producing mutant MinC proteins that do not interact with DicB, both DicB-dependent
28 phenotypes involving ManYZ were lost. These results suggest that DicB is a pleiotropic
29 regulator of bacterial physiology and cell division, and that these effects are mediated by a key
30 molecular interaction with the cell division protein MinC.

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34 **Importance**

35 Temperate bacteriophages can integrate their genomes into the bacterial host chromosome and
36 exist as prophages whose gene products play key roles in bacterial fitness and interactions with
37 eukaryotic host organisms. Most bacterial chromosomes contain “cryptic” prophages that have
38 lost genes required for production of phage progeny but retain genes of unknown function that
39 may be important for regulating bacterial host physiology. This study provides such an example
40 – where a cryptic prophage-encoded product can perform multiple roles in the bacterial host and
41 influence processes including metabolism, cell division, and susceptibility to phage infection.
42 Further functional characterization of cryptic prophage-encoded functions will shed new light on
43 host-phage interactions and their cellular physiological implications.

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45 Introduction

46 Bacteriophages are abundant in the environment with an estimated 10^{31} bacteriophage
47 (phage) particles, and outnumber their bacterial hosts by a factor of 10 to 1 (1, 2). They are
48 found in all ecosystems that harbor bacteria and play a vital role in driving bacterial evolution
49 (3). Based on their lifecycles, phages can be broadly classified as virulent or temperate. Virulent
50 phages use a lytic lifecycle wherein they infect bacterial hosts, use the host cell's resources to
51 make more phage particles and ultimately lyse the cell to release progeny virions into the
52 environment. Temperate phages can grow using a lytic lifecycle, or alternatively can undergo
53 lysogeny – integrating their genomes at a specific attachment site in the host chromosome and
54 remaining stably associated with the host. The bacterium with an integrated phage genome
55 (prophage) is called a lysogen. Changes in host metabolic conditions or external environmental
56 triggers can induce the prophage, which then excises out of the host chromosome and resumes
57 a lytic lifecycle (4, 5).

58 Nearly half of all sequenced bacterial genomes have been found to contain at least one
59 prophage, with many genomes containing multiple prophages (6). Lysogeny comes at a cost to
60 the bacterial host due to the extra burden of replication of prophage DNA and the threat of
61 lysogen induction which is lethal to the host cell. On the other hand, there are many well-
62 documented examples of lysogenic conversion, where prophage-encoded products confer new
63 and advantageous characteristics on the host (7, 8). Many prophages carry virulence genes that
64 contribute to the pathogenicity of a bacterial host, *e.g.*, phage-encoded Shiga toxin in *E. coli*
65 O157 strains (9), phage-encoded Diphtheria toxin in *Corynebacterium diphtheriae* (10), and
66 neurotoxin in *Clostridium botulinum* (11). Prophage-encoded toxins, host cell invasion factors
67 and serum resistance proteins promote various aspects of the infection processes carried out by
68 bacterial pathogens (7). Another well-documented benefit of prophages is superinfection
69 immunity. In a mixed population of lysogens and other bacteria, if a prophage becomes induced
70 and lyses a host cell, the active phage particles released will only infect and lyse the non-

71 lysogens, while the lysogens are protected by the prophage-encoded immunity functions (5).
72 Less well-characterized at a mechanistic level are examples of prophage genes that increase
73 the host's ability to grow under different environmental or stress conditions (12–14).

74 Growing evidence suggests that in many genomes, most of the resident prophages are
75 cryptic (defective), having suffered mutations that leave them unable to excise from the host
76 chromosome, lyse host cells, or produce infectious phage particles (15–18). A recent study
77 identified and characterized orthologous prophages that were integrated in an ancestral host
78 genome and subsequently passed down vertically with the host chromosome in *E. coli* and
79 *Salmonella* (16). Most of these prophages showed evidence of loss of large portions of the
80 original prophage genome, but the remaining genes were under purifying selection (16). These
81 results suggest that certain prophage genes are selected for during host evolution because they
82 encode products that are advantageous to the host under some condition. The cryptic
83 prophages of *E. coli* K12 have been associated with several host phenotypes, including biofilm
84 formation, stress sensitivity, and antibiotic resistance (19). To understand the molecular basis of
85 cryptic prophage-associated phenotypes, functional characterization of prophage-encoded
86 genes is essential.

87 In *E. coli* K12, the cryptic prophage Qin carries an operon encoding a small protein,
88 DicB, and small RNA (sRNA), DicF, that both function as cell division inhibitors (20–25). The
89 sRNA DicF represses *ftsZ* translation by directly base pairing with the *ftsZ* mRNA near the
90 Shine Dalgarno sequence (24, 25). DicF also regulates other mRNAs that encode a variety of
91 regulatory and metabolic functions (25). The 62-amino acid protein DicB inhibits cell division by
92 directly interacting with MinC and recruiting it to the septum via interactions with the septal
93 protein ZipA, where MinC stimulates depolymerization of the Z ring, resulting in cell
94 filamentation (23, 26–28). The region immediately upstream of the *dicBF* operon encodes *dicA*
95 and *dicC* and is similar in sequence and structural arrangement to the lambdoid phage immunity
96 locus. DicA is analogous to the P22 phage C2 repressor and DicC to the P22 Cro repressor

97 (29). DicA represses the *dicBF* operon promoter (which is similar to the λ phage P_L promoter)
98 and the natural condition(s) leading to induction of the operon are unknown (29). DicB and DicF
99 are conserved in many strains of *E. coli*, and interestingly, many strains of pathogenic *E. coli*
100 possess multiple cryptic prophages encoding *dicBF* operons (25, 30, 31).

101 In this study, we have identified a role for the *E. coli dicBF* operon in resistance to
102 bacteriophage infection. Short-term expression of the *dicBF* operon promotes *E. coli* resistance
103 to λ phage infection. The resistance phenotype is primarily attributable to DicB. DicB does not
104 affect λ phage adsorption to host cells. Instead, our results suggest that DicB inhibits injection of
105 λ DNA into the cytoplasm through the inner membrane proteins ManYZ, which are components
106 of the mannose phosphotransferase system. Consistent with an effect of DicB on ManYZ
107 activity, we found that growth of *dicB*-expressing cells on minimal media with mannose as the
108 sole carbon source was strongly inhibited. Our results suggest that products encoded by the
109 *dicBF* operon, found in cryptic prophages in many *E. coli* and *Shigella* strains, can impact
110 bacterial physiology, including by altering cells' susceptibility to bacteriophage infection. We
111 postulate that this may be a common reason why certain cryptic prophage genes are retained in
112 host chromosomes.

113 **Materials and methods**

114 **Strain construction and media.** All strains and phages used in this study are summarized in
115 Table S1 and oligonucleotides (from Integrated DNA Technologies) are listed in Table S2. The
116 strains used in this study are derivatives of *E. coli* K12 strains MG1655 and BW25113.
117 Chromosomal mutations were constructed using λ red recombination method as described in
118 (32–34) or moved into the required strain background using P1 transduction (35).

119 Construction of strain DB240, which has a P_{lac} promoter inserted upstream of *ydfA*
120 replacing the native *dicBF* operon promoter, is described in (25). Oligonucleotides O-PR185 and
121 O-PR186 were used to amplify the kanamycin resistance gene from pKD13 and the PCR
122 product was recombined into the chromosome of DB240 using λ red functions produced by

123 pSIM6 (34). The resulting $\Delta dicB::kan$ strain was called PR163. The kanamycin cassette was
124 removed using pCP20 to create $\Delta dicB::scar$ strain PR165.

125 A $\Delta manXYZ::kan$ deletion was moved into DJ624 and DB240 by P1 transduction from
126 YS208 (36) to create PR187 and PR191 respectively. MinC mutants with single amino acid
127 changes E156A and R172A (37) were constructed by first inserting a $kan-araC-P_{BAD}-ccdB$ PCR
128 product in the *minC* gene in strain DJ624. Oligonucleotides O-PR209/O-PR210 (for E156A) and
129 O-PR205/O-PR206 (for R172A) were used to amplify the $kan-ccdB$ region of strain YS243 and
130 the PCR product was recombined into DJ624 (pSIM6) to generate strains PR178 and PR179,
131 respectively. Oligonucleotides O-PR211 and O-PR212 (containing the E156A mutation) were
132 used to amplify a segment of DNA from the control strain DJ480 to generate PCR product with
133 the desired mutations for *minC* E156A and recombined into PR178 pSIM6 to generate strain
134 PR180. Oligonucleotides O-PR213 and O-PR214 (containing the R172A mutation) were used to
135 generate a PCR product similarly with mutations for *minC* R172A and recombined into PR179
136 pSIM6 to create strain PR181. A P_{lac} promoter replacing the promoter of the *dicBF* operon, was
137 introduced in PR180 and PR181 to generate strains PR182 and PR183 respectively.

138 *E. coli* K12 strains were grown in LB medium at 37°C on a rotary shaker. All phage
139 dilutions were made in TM buffer containing 10mM Tris-HCl and 10mM MgSO₄, and phage
140 infections were carried out using the same buffer. For phage infections, the top agar was made
141 with equal parts of LB agar and TM buffer, unless specified otherwise (38). Top agar was added
142 to the infection mixture and plated on to LB agar plates.

143 **Phage propagation.** New stocks of each phage were prepared as described in Rotman *et al.*
144 (38). Plating cultures were prepared by growing DJ480 in TB medium with 5 mM MgSO₄ (and
145 0.2% maltose exclusively for λ stock preparation) until late log phase, after which an equal
146 amount of TM buffer was added and the mixture was vortexed vigorously. Old phage stocks
147 were titered on the prepared plating culture and plated using top agar, made of equal parts TB

148 agar and TM buffer, onto TB agar plates and incubated overnight at 37°C. The next day, a
149 single individual plaque was punched out and incubated in TM buffer at room temperature for 1-
150 2 hours with occasional vortexing. Between 10 and 30 µl of the single plaque eluate was mixed
151 with 300 µl of DJ480 plating culture and incubated at 37°C for 15 minutes. 3 ml TB/TM top agar
152 was added and plated onto TB plates for incubation at 37°C. After 3 to 7 hours when the lysis
153 was confluent, the plate was overlaid with 5ml TM buffer overnight at room temperature. The
154 TM buffer containing phage was collected in the morning and 4ml fresh TM buffer was added to
155 the plate and kept at room temperature. After 8 hours, the remaining TM containing phage was
156 collected and the combined eluate was centrifuged to pellet the agar and cells down. The
157 supernatant was transferred into a fresh tube, 50ul chloroform was added and the fresh phage
158 lysate was stored at 4°C.

159 **Efficiency of Plaquing (EOP) assay.** The strains used in this experiment were precultured
160 overnight in LB and subcultured in LB medium to ensure all the strains were in the same state of
161 growth when phage infection was carried out. After one hour of sub-culturing (when OD₆₀₀ was
162 ~0.1-0.2), IPTG was added to a final concentration of 0.5 mM to induce the P_{lac} promoter. After
163 one hour with IPTG induction, the cells were washed and resuspended in LB medium. Final
164 OD₆₀₀ was measured, and 1 ml of the culture was centrifuged and resuspended in 1 ml TM
165 buffer. 100 µl of phage dilution was added to 100 µl of bacteria from the previous step and
166 incubated for 10 minutes at 37°C. After 10 minutes, prewarmed 3ml LB top agar was added to
167 the mixture and plated onto LB agar plates. The plates were incubated overnight at 37°C and
168 the plaques were counted. EOP was calculated as: (phage titer on test strain in pfu/ml)/(phage
169 titer on control strain in pfu/ml) (39, 40).

170 **Efficiency of center of infection (ECOI) assay.** The strains were prepared for infection as
171 described for EOP assay. The only difference was in the last step of sample preparation, where
172 the cells were resuspended in TM buffer with 0.5 mM IPTG to induce P_{lac-dicBF} during phage
173 infection. The procedure followed for the ECOI assay was based on that described in Moineau,

174 *et al.* (41). *λvir* lysates were added to 500 μ l of prepared strains at a multiplicity of infection
175 (MOI) of 0.1 or less and incubated at 37°C for 10 minutes. The infection mixture was washed
176 with TM buffer containing 0.5 mM IPTG to remove unadsorbed phages and resuspended in 500
177 μ l of fresh buffer. The infected cells were diluted in TM/IPTG buffer and 100 μ l of each dilution
178 of strains was added to 100 μ l of DJ480 cells in TM buffer, LB/TM top agar was added to this
179 mixture and plated onto LB agar plates. The plates were incubated overnight at 37°C, and the
180 plaques arising from each individual infection were observed and counted. ECOI is calculated
181 as (number centers of infection/ml from test strain) \times 100/(number centers of infection/ml from
182 control strain) (41).

183 **One step growth curve.** The samples were prepared for infection as described for ECOI
184 assays. The one-step growth curve experiment was designed based on (41). After
185 resuspending the cells, *λvir* was added at an MOI of 0.1 or less to 500 μ l of cells and incubated
186 for 10 minutes at 37°C. The infection mixture was washed to remove unadsorbed phages and
187 resuspended in 500 μ l of TM buffer with 0.5 mM IPTG. The strains were diluted 1:10,000 for
188 DJ480 and 1:1000 for DB240 (*P_{lac}-dicBF*) to a final volume of 20 ml in LB with 10 mM MgSO₄
189 and 0.5 mM IPTG in flasks and incubated in a 37°C water bath. Immediately, 100 μ l was
190 withdrawn from the flask and added to 100 μ l phage-sensitive DJ480 cells in TM buffer (for lawn
191 formation), added to prewarmed top agar, and plated onto LB agar plates. The first time point
192 was 30 minutes after the start of infection. The same procedure was repeated for each time
193 point. The burst size was calculated as (phage titer at 100 minutes – initial titer at 30 minutes)/
194 initial titer at 30 minutes. The latent period was calculated as the mid-point of the exponential
195 phase of the growth curve (41).

196 **Adsorption assay.** The procedure described above for ECOI assays was followed and after
197 strains were infected with *λvir* at an MOI of 0.1 and allowed to adsorb for 10 minutes at 37°C,
198 the strains were centrifuged for 5 minutes at 13,000 rpm to pellet cells and adsorbed phage. 100
199 μ l of the supernatant was removed and dilutions were made in TM buffer. 10 μ l of each dilution

200 was added to 100 μ l DJ480 (phage-sensitive) cells in TM buffer and incubated for 10 minutes at
201 37°C. This mixture was plated onto LB plates using top agar and incubated overnight at 37°C.
202 Control titer was calculated using the same procedure as above using the same amount of
203 phage required for MOI of 0.1 added to 500 μ l TM buffer (no bacteria). Percentage of adsorption
204 was calculated as (control titer – residual titer)*100/ control titer (39, 42).

205 **Growth on minimal medium plates with different sugars.** For growth assays, M63 minimal
206 medium plates with sugars (glucose, fructose, mannose, N-acetyl glucosamine, and
207 glucosamine) at a final concentration of 0.2% were prepared without or with 0.025 mM IPTG.
208 The strains were streaked on plates and incubated for 44 hours at 37°C. By visual inspection,
209 strains were scored for growth with +++ denoting normal growth, ++ and + denoting decreasing
210 growth and – as no growth.

211 **Results**

212 **Transient induction of the *dicBF* operon protects against λ phage infection.** The region of
213 Qin prophage containing the *dicBF* operon (Fig. 1A) resembles the immunity regions of P22 and
214 other lambdoid phages (29). While functions have not been identified for most of the products of
215 the *dicBF* operon, DicB (a small protein) and DicF (a small RNA) have been shown to inhibit cell
216 division (20, 22–25). We showed previously that DicF post-transcriptionally regulates a variety
217 of genes involved in cell division, growth and metabolism (25). Given their position in the
218 immunity region of the prophage genome, and the fact that the characterized gene products
219 impact cell physiology, we hypothesized that products of the *dicBF* operon could cause changes
220 in the host cell that promote resistance to phage infection. We tested this by comparing phage
221 infections of control and *dicBF*-expressing cells. Since the conditions under which the *dicBF*
222 operon is normally expressed are not fully understood, we used an inducible expression system
223 described previously (25) where we replaced the *dicBF* operon promoter with a P_{lac} promoter at
224 the native locus. In addition to the P_{lac} -*dicBF* strain, we used strains with deletions of different
225 genes in the operon (Fig. 1B).

226 We measured phage infection of these strains by Efficiency of Plaquing (EOP) assays,
227 initially using phage λ . In this assay, the phage is titered on all bacterial strains and the titer in
228 plaque forming units (pfu)/ml is calculated for each strain. The EOP is defined as: phage titer on
229 test strain/phage titer on control strain. The control strain lacked the P_{lac} promoter. Strains with
230 the P_{lac} promoter driving *dicBF* expression were exposed to IPTG for 60 minutes. Then, strains
231 were infected with λ_{vir} (a λ mutant that can only complete the lytic cycle during infection of host
232 cells) and plated for titer as described in Materials and Methods. The EOP for the P_{lac} -*dicBF*
233 strain was 0.04 (Fig. 1B), meaning that rate of infection of the *dicBF*-expressing strain was only
234 4% relative to the control strain. This result suggested that transient expression of the *dicBF*
235 operon conferred resistance to infection by λ_{vir} . To further characterize the basis for this
236 phenotype, we deleted *dicF* and *dicB*, singly and in combination because previous studies
237 identified growth or cell division phenotypes associated with these genes (23, 25). Phenotypes
238 of deletion mutants demonstrated that *dicB* played the most prominent role in the resistance
239 phenotype (Fig. 1B). Deletion of *dicB* alone or *dicB* in combination with *dicF* restored the EOP of
240 λ_{vir} to nearly that of control. In contrast, deletions of *dicF* alone had a minimal effect on the
241 resistance phenotype (Fig. 1B). We also carried out infections with wild-type λ phage, and saw
242 similar results for EOP on control, *dicBF*-expressing and deletion mutant strains (Fig. S1).

243 Because previous studies showed that ectopic expression of the *dicBF* operon impairs
244 growth of the host strain, we reasoned that poor growth of test strains could influence the results
245 of EOP assays. To more accurately assess the outcome of a phage infection on cells
246 expressing the *dicBF* operon, we conducted center of infection (COI) assays. For this assay,
247 strains (Fig. 1B) were induced with 0.5 mM IPTG and λ_{vir} infection was carried out at a
248 multiplicity of infection (MOI) of 0.1. After adsorption of phage to test strains, the unadsorbed
249 phages were removed by washing and the infected test cells were diluted and mixed with the
250 phage-sensitive control strain. Productive infections of the test strain are detected as plaques
251 (centers of infection) on the phage-sensitive control strain. The efficiency of λ_{vir} forming centers

252 of infection (ECOI) is calculated as (number centers of infection/ml from test strain) x
253 100/(number centers of infection/ml from control strain). The ECOI for λ_{vir} on $P_{lac-dicBF}$ cells
254 was 3% (Fig. 1C). This result is similar to the results of EOP assays (Fig. 1B), suggesting that
255 the growth characteristics of the $P_{lac-dicBF}$ test strain did not impact the experimental outcome.
256 Deletion of *dicB*, alone or in combination with *dicF*, restored the ECOI to ~80%. The $\Delta dicF$
257 strain gave an ECOI of 5% (Fig. 1C). These results are again consistent with our EOP
258 experiments (Fig.1B) implicating DicB as the major player in the phage resistance phenotype.
259 **The *dicBF* operon promotes resistance against λ , but not other phages.** To determine if
260 transient expression of *dicBF* conferred resistance to other phages, we conducted infection
261 experiments using control and $P_{lac-dicBF}$ strains with nine different lytic and temperate phages
262 (Fig. 2). In this experiment, the EOP of λ_{vir} on the $P_{lac-dicBF}$ strain was 0.016 or 1.6%
263 compared to the control strain, which was the lowest of the nine phages tested (Fig. 2). Partial
264 resistance was observed for T3 phage, which had an EOP of 0.14 on $P_{lac-dicBF}$ cells. However,
265 the EOP for the remaining seven phages, including phages $\phi 80$ and HK97, which are closely
266 related to λ , was similar to control cells (Fig. 2). These results suggest DicB does not provide a
267 broad-spectrum of resistance against bacteriophages.
268 **Effect of *dicBF* expression on λ phage growth.** The classical experiment to study the growth
269 cycle of phage in bacteria is the one-step growth curve, as described by Ellis and Delbrück (43).
270 They observed a latent period where numbers of phage recovered from infected cells remained
271 low as new phage particles were being synthesized inside the host cell. After the latent period is
272 the “burst” where numbers of infectious phage particles increase rapidly as the phage life cycle
273 is completed and cells are lysed to release mature progeny. We conducted one-step growth
274 curves for λ_{vir} on control and *dicBF*-expressing strains, essentially as described above for ECOI
275 experiments over a time course following infection. λ_{vir} was added at an MOI of 0.1 to control
276 and $P_{lac-dicBF}$ cells resuspended in TM buffer. After phage adsorption, the cells were washed to
277 remove unadsorbed phage and the phage-host complexes were diluted into fresh medium with

278 IPTG (see Materials and Methods). At each time-point, the number of infectious phage particles
279 in each culture was calculated by removing samples and plating for PFU on a phage-sensitive
280 control strain.

281 As expected based on previous results (Figs. 1B, C), the ECOI for λvir on $P_{lac-dicBF}$
282 cells was reduced by almost 2 logs compared with the control strain at the early time points, and
283 the reduced numbers of phage produced by $P_{lac-dicBF}$ cells persisted across the phage growth
284 curve (Fig. 3). The latent period for $P_{lac-dicBF}$ cells (75 min.) was ~10 min. longer than for
285 control cells (65 min.) (Fig. 3). The calculated burst sizes were 343 phage/ $P_{lac-dicBF}$ cell
286 compared to 169 phage/control cell. This increase in burst size in $dicBF$ -expressing cells is
287 likely due to filamentation of cells caused by DicB and DicF. It has been shown before that
288 filamenting cells produce more phage than normal size cells (44, 45). Importantly, we observed
289 that the ~3% of phages that escaped DicB-mediated resistance followed a growth curve similar
290 to that of phages growing on control cells. Collectively, these data led us to hypothesize that
291 DicB affects an early step of the phage life cycle like adsorption or DNA injection, since phages
292 that escape this DicB effect complete a relatively normal life-cycle.

293 **The $dicBF$ operon does not affect phage adsorption to host cells.** To test if expression of
294 the $dicBF$ operon affects the first step of phage infection, we tested the ability of λvir to adsorb
295 to host cells expressing this operon. During the ECOI experiment, once phage infection was
296 carried out with cells in TM buffer at an MOI of 0.1 and incubated at 37°C for 10 mins, the
297 phage-cell mixture was centrifuged and the supernatant containing the unadsorbed phages was
298 removed. This supernatant was titered on phage-sensitive control cells by standard plaque
299 assay (residual titer). Control titer was calculated using the same procedure as above, with
300 phage added to TM buffer instead of bacterial cells in the first step of the experiment. The
301 percentage of adsorption was calculated as $(\text{control titer} - \text{residual titer}) * 100 / \text{control titer}$.

302 The adsorption of λvir to strains expressing the $dicBF$ operon was the same as
303 adsorption to the control strain (Table 1). Notably, while λvir and HK97 both adsorb to the same

304 outer membrane receptor, LamB (46, 47), the effect of *dicBF* expression on the EOP of these
305 two phages is significantly different – with reduced EOP only for *λvir* (Fig. 2). These
306 observations strongly suggest that DicB does not affect the phage life cycle at the step of
307 adsorption to host cells.

308 **Recombinant λ phages with the host range region of $\phi 80$ are not affected by DicB.** The
309 genomes of λ phage and $\phi 80$ (a lambdoid phage) have strikingly similar organization, allowing
310 for easy construction of recombinant phage (48, 49). One prominent difference between λ and
311 $\phi 80$ is their use of different outer and inner membrane receptors. λ uses LamB (outer
312 membrane) and ManYZ (inner membrane) for adsorption and DNA injection respectively (50–
313 54), while $\phi 80$ uses FhuA (outer membrane) and the TonB complex (inner membrane) (48, 55,
314 56). The phage genes encoding determinants for utilization of host outer and inner membrane
315 receptors are located in the host range region of lambdoid phage genomes. Our results so far
316 suggest that the DicB-dependent phage resistance phenotype is not due to an effect on
317 adsorption to the outer membrane receptor but might be mediated at another early step of
318 infection such as injection of the phage genome through the inner membrane receptor. To test
319 this idea, we measured phenotypes of control and *dicBF*-expressing cells challenged with
320 recombinant λ phage containing the host range region of $\phi 80$ ($\lambda h80$). The $\lambda h80$ phage carry
321 most of the wild-type λ genome but have an altered host range region specifying use of the $\phi 80$
322 outer and inner membrane receptors. To confirm this, we tested the plaquing ability of $\lambda h80$
323 phages on wild-type, $\Delta fhuA$, $\Delta tonB$ and $\Delta manXYZ$ *E. coli* strains. As expected, the $\lambda h80$
324 phages, like $\phi 80$, did not plaque on $\Delta fhuA$ and $\Delta tonB$ strains but formed normal plaques on the
325 $\Delta manXYZ$ strain (Table S3). Next, we carried out EOP assays using *λvir* and a panel of
326 recombinant phage with the host range of λ or $\phi 80$ (Table S1) on control and P_{lac} -*dicBF* cells
327 (Fig. 4). We hypothesized that if DicB mediates resistance to λ phage by impairing injection of
328 phage DNA across the cytoplasmic membrane, then phage with the host range of λ would

329 remain inhibited by DicB, whereas λ h80 phage with altered inner membrane receptor specificity
330 would not be impacted by DicB. Results of the EOP assays demonstrate that phage with λ host
331 range remain sensitive to DicB-mediated inhibition while λ h80 phages had a similar EOP on
332 $P_{lac-dicBF}$ and control cells (Fig. 4). Together with our previous results, this observation
333 suggests that DicB-mediated resistance acts at the level of the inner membrane receptor
334 ManYZ used for λ phage DNA injection into the cytoplasm of *E. coli*. We note that the panel of
335 phages that we tested in this experiment had other genetic differences aside from the different
336 host ranges (Table S1). Only the host range was correlated with susceptibility to DicB-mediated
337 resistance.

338 Phage 434 (53) is another phage that uses ManYZ for injection of DNA through the
339 cytoplasmic membrane (Fig. 5A). Previous studies have shown that *manXYZ* deletion mutants
340 (also known as *pel* mutants) were resistant to infection by λ and phage 434, but not ϕ 80 (53). To
341 further test our hypothesis that DicB inhibits phage infection at the level of DNA entry through
342 ManYZ, we tested the ability of λ vir, phage 434 and ϕ 80 to infect control and *dicBF*-expressing
343 cells in *manXYZ*⁺ and Δ *manXYZ* backgrounds. We verified that the λ , phage 434 and ϕ 80
344 phages plaque as expected on wild-type and strains with mutations in specific receptors (Table
345 S4). As shown above (Fig. 2), ϕ 80 plaquing efficiency is not impacted by expression of the
346 *dicBF* operon in a *manXYZ*⁺ background. The Δ *manXYZ* mutant host also supported wild-type
347 EOPs for ϕ 80 plaquing regardless of whether *dicBF* was expressed (Fig. 5B). For λ vir, the EOP
348 on $P_{lac-dicBF}$ cells was 4% relative to control in *manXYZ*⁺ cells, whereas the Δ *manXYZ* host did
349 not support λ vir growth (Fig. 5B). The pattern of growth for phage 434 was very similar to that of
350 λ vir, with a reduced EOP of ~10% on $P_{lac-dicBF}$ cells in the *manXYZ*⁺ host and no countable
351 plaques on the Δ *manXYZ* host (Fig. 5B). The results of this experiment are consistent with the
352 hypothesis that DicB inhibits the use of the mannose phosphotransferase system proteins,
353 ManYZ, as an inner membrane receptor for productive phage infection.

354 **Growth of *dicBF*-expressing cells is inhibited on plates with mannose as the C source.**

355 Our previous results point to the small protein DicB inhibiting the activity of mannose transporter
356 ManYZ proteins with regard to DNA uptake during phage infection. To test whether DicB inhibits
357 the function of these proteins more broadly, we checked the growth of *dicBF*-expressing cells on
358 mannose as the sole C source. For this experiment, we used control, *P_{lac}-dicBF*, and *P_{lac}-dicBF*
359 Δ *dicB* strains. The strains were streaked on M63 minimal plates with different sugars with or
360 without 0.025mM IPTG (to induce *dicBF* expression) and incubated for 44 hours at 37°C (Table
361 2, Fig. S2). In the absence of inducer, all the strains had near normal growth on the different
362 sugars used. When *dicBF* expression was induced using 0.025mM IPTG, we observed growth
363 inhibition of *P_{lac}-dicBF* cells on mannose and glucosamine, but not on glucose, fructose or N-
364 acetyl glucosamine. Deletion of *dicB* relieved the growth inhibition on mannose and
365 glucosamine. Both mannose and glucosamine sugars are transported via the ManXYZ
366 transporter in *E. coli* (57–59). These results demonstrate that DicB affects growth specifically on
367 substrates of ManYZ. Growth on sugars that are transported by other PTS proteins was
368 unaffected. These data suggest that DicB impacts at least two different functions of ManYZ –
369 uptake of phage DNA during infection and transport of sugar substrates.

370 **MinC mutants that do not interact with DicB lose the phage resistance and sugar**

371 **phenotypes.** The only characterized activity of DicB is inhibition of cell division (20, 60). The
372 mechanism by which DicB impacts cell division requires a protein-protein interaction with MinC,
373 one of the proteins involved in controlling septal ring placement in *E. coli*. MinC is an inhibitor of
374 FtsZ polymerization, and normally MinC concentrations are highest at cell poles so that septum
375 formation is inhibited at polar sites and directed instead to mid-cell (61). Previous work
376 demonstrated that DicB interacts with MinC and brings it to mid-cell via an interaction with ZipA,
377 a septal protein (28). DicB-mediated localization of MinC to cell center inhibits FtsZ
378 polymerization and promotes filamentation (27, 28). To determine if the DicB-MinC interaction is
379 necessary for the DicB-dependent phenotypes we found in this study, we constructed strains

380 with *minC* mutant alleles that produce MinC proteins that are defective for interaction with DicB.
381 We used two different MinC mutants: MinC R172A, which interacts weakly with DicB, and MinC
382 E156A mutant, which does not interact with DicB (37). In strains expressing these *minC* alleles,
383 DicB has a modest (MinC R172A) or no (MinC E156A) impact on cell division, consistent with
384 their reduced binding to DicB. We used MinC E156A and R172A mutant hosts to test whether
385 the DicB-mediated phage resistance or sugar growth phenotypes required the DicB-MinC
386 interaction.

387 As observed previously, in the wild-type *minC*⁺ background, *dicBF*-expressing cells
388 showed reduced EOP for λ vir compared to control cells (Fig. 6). However, in the MinC R172A
389 (reduced binding to DicB) strain, the resistance phenotype was diminished – *dicBF* expression
390 in this host gave an EOP of 12% compared to the control strain. In the MinC E156A (abrogated
391 binding to DicB) background, the EOP of λ vir on *dicBF*-expressing cells was very similar to the
392 control strain (Fig. 6). These results suggested that the DicB-MinC interaction is required for the
393 DicB-mediated resistance to λ phage infection. The same strains were grown on M63 minimal
394 plates with different sugars without or with 0.025mM IPTG to induce the *dicBF* operon. As
395 shown above, in the wild-type *minC*⁺ background, expression of *dicBF* inhibited growth on
396 plates with mannose and glucosamine, but not on plates with glucose (Table 3). In contrast,
397 *dicBF* expression in *minC* mutant strains (E156A and R172A) did not inhibit growth on any of
398 the sugars tested (Table 3). Collectively, these data indicate that the new DicB-associated
399 phenotypes we have identified – phage resistance and inhibition of growth on sugars that are
400 transported by ManYZ – require the previously defined molecular mechanism of DicB interaction
401 with the host protein MinC.

402 **Discussion**

403 The existence of cryptic or defective prophages on bacterial chromosomes was
404 discovered long ago (4), but their potential beneficial functions for host cells are still coming to
405 light. In part, this is because we do not know the functions of the majority of genes encoded on

406 these prophages. In this study, we have identified a new functional role for the cryptic prophage-
407 encoded protein DicB in *E. coli* K12. We showed that induction of the *dicBF* operon makes cells
408 resistant to infection by phages that use the ManYZ PTS proteins as inner membrane receptors
409 for DNA injection (Figs. 1, 5). DicB, a 62-amino acid protein encoded by the *dicBF* operon, plays
410 the primary role in conferring this phage resistance phenotype (Fig. 1). Our results are
411 consistent with the model that DicB inhibits phage DNA injection through the mannose
412 transporter proteins ManYZ (Figs. 4, 5). The DicB effect on ManYZ also inhibits ManYZ-
413 dependent transport of sugar substrates (Tables 2, 3), suggesting that DicB affects the general
414 structure or function of these transport proteins. Previous work demonstrated that DicB inhibits
415 cell division by interacting with and affecting localization and activity of the cell division proteins
416 MinC and FtsZ (23, 26–28). In this study, we found that the DicB-associated phage resistance
417 and sugar utilization phenotypes are dependent on DicB-MinC interactions (Fig. 6).

418 Prior to this work, the only known function of DicB was inhibition of cell division. DicB
419 directly interacts with MinC of the Min system, which consists of the proteins MinC, MinD and
420 MinE, which play a role in spatial positioning of the FtsZ ring at mid-cell for cell division. MinC is
421 a negative regulator of FtsZ polymerization, and in *E. coli* MinC oscillates between the two cell
422 poles (driven by MinD and MinE) in order to inhibit Z ring assembly at the poles (62, 63).
423 However, when DicB is expressed, a DicB-MinC complex is formed which interacts with the
424 septal ring component ZipA and stimulates Z ring depolymerization at mid-cell leading to cell
425 filamentation (28). Both activities of DicB, cell division inhibition and the ManYZ inhibition
426 phenotypes reported in this study, involve interaction with MinC. These results imply a
427 previously unsuspected link between the Min system or other components of the cell division
428 machinery and the mannose PTS. A few studies have examined localization of various PTS
429 proteins. The general PTS proteins, EI and HPr, were found to localize primarily to cell poles
430 (64, 65). Localization of EII sugar permeases has been less studied, but there is recent
431 evidence that these proteins cluster together around the cell membrane (66). It will be

432 interesting in future work to explore the subcellular localization of ManYZ and examine if or how
433 it is impacted by MinC and DicB.

434 It is clear that active prophages can protect their hosts from superinfection by other
435 phages (67, 68). One of the earliest identified phage resistance mechanisms is called
436 superinfection exclusion, which can occur by a number of different mechanisms (67, 68). One
437 mechanism of superinfection exclusion is mediated by prophage-encoded proteins that block
438 entry of a superinfecting phage's DNA by mechanisms that are not well defined, but may involve
439 modifying the activity or function of inner membrane receptors (67). For example, protein gp15
440 of HK97 prophage inhibits superinfection by HK97 phage by preventing phage DNA entry
441 through the PtsG inner membrane receptor (47, 69). SieA is an inner membrane protein
442 encoded by the P22 prophage of *Salmonella typhimurium* that prevents infection by P22, L,
443 MG178, and MG40 phages by inhibiting DNA entry into the cytoplasm (70, 71). The P1
444 prophage-encoded Sim protein was also shown to confer superinfection immunity against P1
445 and few other phages by an unknown mechanism that occurs after adsorption of the infecting
446 phage (72). While these examples illustrate superinfection exclusion by active prophages, our
447 results suggest that in at least some cases defective prophages can play similar roles in
448 protecting their hosts from phage infection.

449 *E. coli* strains have several defective prophages residing on their chromosomes, many of
450 which have undergone extensive deletions since their original integration (16). There is
451 evidence that many of the genes remaining on the cryptic prophages, including the core phage
452 genes which code for tail and lysis proteins, are under strong purifying selection, implying that
453 these genes now perform functions beneficial to the host (16). *E. coli* K12 has 9 defective
454 prophages, 8 of which do not respond to conditions that usually induce an active prophage (19).
455 Some of these prophages have lost genes important for excision from the host chromosome.
456 For example, CP4-44 prophage entirely lacks an integrase, Qin prophage has an inactive
457 integrase and DLP12 prophage has lost part of the excisionase gene (19, 73). Pathogenic *E.*

458 *coli* O157:H7 strain Sakai harbors 18 prophages and 17 of these were found to have defects in
459 the prophage genes required for excision and morphogenesis (15). Though not fully functional,
460 three of these 18 defective prophages carry virulence factors and were capable of packaging
461 and transferring their DNA to *E. coli* K12 (15). These observations demonstrate that even
462 though defective prophages have undergone significant genetic changes through their life
463 history with the host, some retain functions that allow them to continue acting as gene-transfer
464 agents.

465 It has been speculated that another beneficial role of defective prophages could be
466 encoding functions that are important for host cell adaptation to stress conditions. In a study by
467 Wang *et al.* (19), defective prophages of *E. coli* K12 were shown to increase resistance to
468 environmental stresses like oxidative stress, osmotic stress, and to certain antibiotics like
469 quinolones and beta-lactams. This study reported that Δqin derivatives of the parent strain were
470 more sensitive to beta-lactam antibiotics and that $\Delta dicB$ strains showed greater sensitivity to
471 azlocillin and nalidixic acid (19). We constructed Δqin and $\Delta dicB$ strains and examined
472 sensitivity to nalidixic acid and ampicillin, and we found no differences in sensitivity between
473 parent strains and mutants (data not shown). It is possible that phenotypes vary with strain
474 background – our strains are MG1655 derivatives and Wang, *et al.*, used BW25113 (19).

475 Identifying the signals or conditions that induce prophage genes will be key to
476 understanding their physiological roles in host cells. We have exposed our strains to various
477 conditions that are known to induce prophage gene expression, including DNA damage,
478 starvation, and exposure to antibiotics, and have not yet identified conditions that substantially
479 induce transcription from the native *dicBF* promoter (data not shown). Another study (74),
480 reported that *E. coli* K12 MG1655 cells undergo DicF-dependent filamentation under
481 anaerobic conditions (growth in large volume anaerobic fermenters). The authors of this study
482 suggest that stability of DicF is differentially regulated such that it is more stable under
483 anaerobic growth conditions and degraded faster under aerobic conditions. A very recent

484 study found that four DicF orthologs encoded by different prophages in *E. coli* O157:H7 are
485 produced under microaerobic growth conditions (31). These DicF sRNAs promote low oxygen-
486 responsive virulence gene expression via base pairing-mediated regulation of a key virulence
487 transcription factor. These studies suggest that in at least some *E. coli* strain backgrounds,
488 oxygen is an important signal for modulation of *dicBF* operon transcription or DicF mRNA
489 stability. However, we have not observed any DicF- or DicB-mediated filamentation of
490 MG1655 cells grown in small volume LB liquid cultures in an anaerobic chamber (data not
491 shown), so we speculate that additional signals or conditions might contribute to *dicBF* operon
492 expression in our host strain background.

493 Previous studies from our lab characterized the mRNA target regulon of DicF (25). In
494 addition to the previously discovered DicF target *ftsZ* mRNA, we found that DicF base pairs with
495 and represses translation of *xyIR*, *pykA* and *manXYZ* mRNAs, encoding the xylose repressor,
496 pyruvate kinase, and mannose PTS components, respectively (25, 75). Thus, the *dicBF* operon
497 encodes a base pairing-dependent sRNA regulator (DicF) and a small protein (DicB) that act at
498 different levels to inhibit the synthesis and activity of a PTS sugar transporter (ManXYZ). This is
499 strikingly similar to the regulation of the glucose PTS (*ptsG*, enzyme IICB^{Glc}) by the dual-
500 function sRNA SgrS and the small protein it encodes, SgrT. SgrS base pairs with and
501 represses translation of *ptsG* mRNA (76, 77), while SgrT inhibits PtsG activity at a post-
502 translational level (78–80). Perhaps regulation of PTS enzyme synthesis and activity by
503 sRNAs and small proteins is a common mechanism for post-transcriptional control of these
504 systems. Future studies on the multitude of sRNAs and small proteins encoded on prophages
505 and bacterial chromosomes promise to reveal more surprising connections between phages
506 and their hosts.

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733 regulatory peptide SgrT and the EIICB^{Glc} of the glucose-phosphotransferase system of *E.*
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	Control	$P_{lac}-dicBF$	$P_{lac}-dicBF \Delta dicB$	$P_{lac}-dicBF \Delta dicF$	$P_{lac}-dicBF \Delta dicF \Delta dicB$
% adsorption	99	99	99	99	99
Standard error (+/-)	0.22	0.25	0.69	0.23	0.15

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743 **Table 1. The *dicBF* operon does not affect phage adsorption to host cells.** The strains
744 used in this experiment are control (DJ480), $P_{lac}-dicBF$ (DB240), $P_{lac}-dicBF \Delta dicB$ (DB243), $P_{lac}-$
745 $dicBF \Delta dicF$ (DB247) and $P_{lac}-dicBF \Delta dicF \Delta dicB$ (DB248). Cells were infected at an MOI of 0.1
746 with λvir and allowed to adsorb for 10 mins. at 37°C. After adsorption, the samples were
747 centrifuged and the supernatant containing the unadsorbed phages were titered on phage
748 sensitive cells for quantification (residual titer). Control titer was calculated by carrying out the
749 assay with TM buffer, without bacterial cells. The percentage of adsorption was calculated as
750 $(\text{control titer} - \text{residual titer}) * 100 / \text{control titer}$. Standard error was calculated as standard
751 deviation of values from three biological replicates.

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Carbon Source	Control		P_{lac} - <i>dicBF</i>		P_{lac} - <i>dicBF</i> Δ <i>dicB</i>	
	Uninduced	Induced	Uninduced*	Induced	Uninduced*	Induced
Mannose	+++	+++	++	+	+++	+++
Glucose	+++	+++	+++	+++	+++	+++
Glucosamine	+++	+++	++	+	+++	+++
Fructose	+++	+++	+++	+++	+++	+++
N-acetyl glucosamine	+++	+++	+++	+++	+++	+++

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762 **Table 2. Growth of *dicBF* expressing cells is inhibited on plates with mannose as the C**

763 **source.** The strains were streaked on M63 minimal medium with 0.2% sugars as C source

764 without or with 0.025mM IPTG and incubated for 44 hours at 37°C. The strains used were

765 control (DJ624), P_{lac} -*dicBF* (DB240) and P_{lac} -*dicBF* Δ *dicB* (PR165). +++ indicates normal

766 growth, ++ and + indicates decremental growth. * denotes that the P_{lac} promoter is leaky and we

767 suspect low level expression of the *dicBF* operon even at 0mM IPTG.

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	Glucose		Mannose		Glucosamine	
	Uninduced	Induced	Uninduced	Induced	Uninduced	Induced
Control	+++	+++	+++	+++	+++	+++
P_{lac} - <i>dicBF</i>	+++	+++	++	+	++	+
Control <i>minC</i> R172A	+++	+++	+++	+++	+++	+++
P_{lac} - <i>dicBF</i> <i>minC</i> R172A	+++	+++	+++	+++	+++	+++
Control <i>minC</i> E156A	+++	+++	+++	+++	+++	+++
P_{lac} - <i>dicBF</i> <i>minC</i> E156A	+++	+++	+++	+++	+++	+++

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783 **Table 3. MinC mutants that do not interact with DicB regain ability to grow on Mannose**

784 **and Glucosamine.** The strains were streaked on M63 minimal medium plates with 0.2% sugars

785 and 0.025mM IPTG to induce *dicBF* operon. The plates were incubated for 44 hours at 37°C.

786 The strains used in this experiment are control (DJ624), P_{lac} -*dicBF* (DB240), control *minC*

787 R172A (PR181), P_{lac} -*dicBF minC* R172A (PR183), control *minC* E156A (PR180) and P_{lac} -*dicBF*

788 *minC* E156A (PR182). +++ indicates normal growth, ++ and + indicates decremental growth.

789 The P_{lac} promoter is leaky and we suspect low level expression of the *dicBF* operon even at

790 0mM IPTG.

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798 **Figure 1. Transient induction of the *dicBF* operon protects against λ vir infection.** (A) The
799 *dicBF* locus on Qin prophage of *E. coli* K12. The red arrow indicates where P_{lac} is inserted on
800 the chromosome, replacing the native *dicBp* promoter. (B) Efficiency of Plaquing (EOP) is
801 defined as (λ vir titer on the test strain)/ (λ vir titer on the control strain). Strains used in this
802 experiment were: control (DJ480), P_{lac} -*dicBF* (DB240), P_{lac} -*dicBF* Δ *dicB* (PR165), P_{lac} -*dicBF*
803 Δ *dicF* (DB247) and P_{lac} -*dicBF* Δ *dicF* Δ *dicB* (DB248). All strains were grown to the same state of
804 growth with the *dicBF* operon induced with 0.5 mM IPTG for 60 minutes. The inducer was
805 washed off, the strains were resuspended in TM buffer, infected with λ vir and plated to calculate
806 titer. Error bars were calculated as standard deviation of values from three biological replicates.
807 (C) Efficiency of center of infection (ECOI) is calculated as (number of infectious centers/ml from
808 test strain)* 100/(number of infectious centers/ml from control strain). The strains used in this
809 experiment are control (DJ480), P_{lac} -*dicBF* (DB240), P_{lac} -*dicBF* Δ *dicB* (DB243), P_{lac} -*dicBF* Δ *dicF*
810 (DB247) and P_{lac} -*dicBF* Δ *dicF* Δ *dicB* (DB248). The cells were grown with induction of the *dicBF*
811 operon with 0.5 mM IPTG and infected with λ vir at an MOI of 0.1. The unadsorbed phages were
812 removed, the phage-host complex was added to phage sensitive cells (DJ480) and plated onto
813 LB agar for counting plaques (infectious centers). Error bars were calculated as standard
814 deviation of values from three biological replicates.

815 **Figure 2. The *dicBF* operon confers resistance against λ phage but not other phages.** For
816 each of the nine phages, titer in terms of pfu/ml was calculated by infection of control (DJ480)
817 and P_{lac} -*dicBF* (DB240) cells. The cells were prepared for infection and the EOP was calculated
818 for each phage as described for Fig.1B. Error bars were calculated as standard deviation of
819 values from three biological replicates.

820 **Figure 3. One step growth curve of λ vir control and P_{lac} -*dicBF* cells.** The cells were grown
821 with induction of the *dicBF* operon and infection was carried out an MOI of 0.1, similar to the
822 center of infection assay. After removing unadsorbed phages, the cells were diluted in LB
823 medium (with IPTG to induce the *dicBF* operon) and incubated at 37°C for the entire duration of

824 the growth curve. At each time point starting at 30 minutes from the start of infection, samples
825 were removed and added to the phage-sensitive strain (DJ480), and plated to count plaques.
826 Burst size was calculated as (phage titer at 100 minutes - initial titer at 30 minutes)/initial titer at
827 30 minutes. Latent period was calculated as the time at the mid-point of the exponential phase
828 of the curve. Error bars were calculated as standard deviation of values from three biological
829 replicates.

830 **Figure 4. λ phage with the host range of $\phi 80$ is not affected by DicB.** Recombinant λ
831 phages with either the λ or $\phi 80$ host range were plaqued on control (DJ480) or $P_{lac-dicBF}$
832 (DB240) cells. Phage names from left to right are: 185, 148, 169, 173, 158, 138 and λvir (see
833 Table S1 for phage genotypes). The cells were prepared for infection and the EOP was
834 calculated for each phage as described for Fig. 1B. Error bars were calculated as standard
835 deviation of values from three biological replicates.

836 **Figure 5. Phage 434 plaquing on $ManYZ^+$ strains is inhibited by the *dicBF* operon.** (A)
837 Outer and inner membrane receptor specificity of phages λvir , 434 and $\phi 80$. (B) EOP assay was
838 carried out by preparing cells and calculating titer of phages on the different strains as described
839 in Fig.1B. The strains used in this experiment are control (DJ624), $P_{lac-dicBF}$ (DB240), control
840 $\Delta manXYZ::kan$ (PR187) and $P_{lac-dicBF} \Delta manXYZ::kan$ (PR191). EOP of phages for each strain
841 is calculated with respect to the control strain in the same background. Error bars were
842 calculated as standard deviation of values from three biological replicates. * denotes strains for
843 which plaques could not be counted.

844 **Figure 6. MinC mutants that do not interact with DicB lose the phage resistance effect.**
845 The cells were grown with induction of the *dicBF* operon with 0.5mM IPTG, infected with λvir
846 and the EOP calculated as described in Fig.1B. EOP of λvir for each strain is calculated with
847 respect to the control strain in the same background. The strains used in this experiment are
848 control (DJ624), $P_{lac-dicBF}$ (DB240), control *minC* R172A (PR181), $P_{lac-dicBF} minC$ R172A

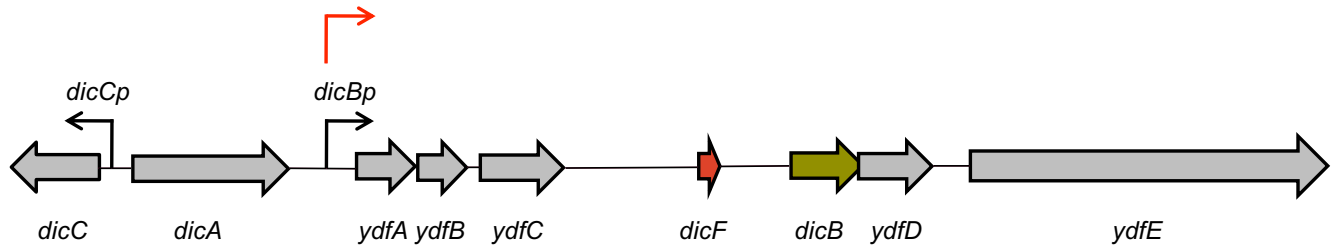
849 (PR183), control *minC* E156A (PR180) and $P_{lac-dicBF}$ *minC* E156A (PR182). Error bars were
850 calculated as standard deviation of values from three biological replicates.

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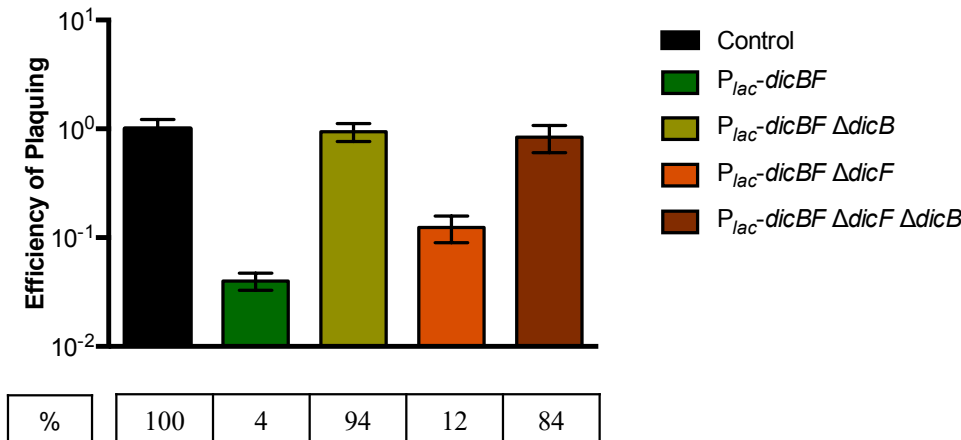
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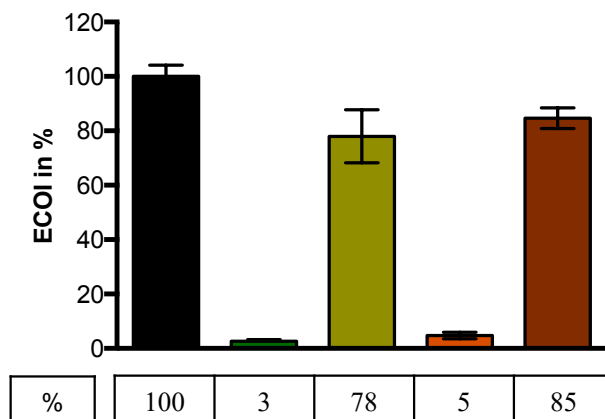


Figure 1. Transient induction of the *dicBF* operon protects against *λvir* infection. (A) The *dicBF* locus on Qin prophage of *E. coli* K12. The red arrow indicates where P_{lac} is inserted on the chromosome, replacing the native *dicBp* promoter. (B) Efficiency of Plaquing (EOP) is defined as (*λvir* titer on the test strain) / (*λvir* titer on the control strain). Strains used in this experiment were: control (DJ480), P_{lac}-*dicBF* (DB240), P_{lac}-*dicBF* Δ*dicB* (PR165), P_{lac}-*dicBF* Δ*dicF* (DB247) and P_{lac}-*dicBF* Δ*dicF* Δ*dicB* (DB248). All strains were grown to the same state of growth with the *dicBF* operon induced with 0.5 mM IPTG for 60 minutes. The inducer was washed off, the strains were resuspended in TM buffer, infected with *λvir* and plated to calculate titer. Error bars were calculated as standard deviation of values from three biological replicates. (C) Efficiency of center of infection (ECOI) is calculated as (number of infectious centers/ml from test strain) * 100 / (number of infectious centers/ml from control strain). The strains used in this experiment are control (DJ480), P_{lac}-*dicBF* (DB240), P_{lac}-*dicBF* Δ*dicB* (DB243), P_{lac}-*dicBF* Δ*dicF* (DB247) and P_{lac}-*dicBF* Δ*dicF* Δ*dicB* (DB248). The cells were grown with induction of the *dicBF* operon with 0.5 mM IPTG and infected with *λvir* at an MOI of 0.1. The unadsorbed phages were removed, the phage-host complex was added to phage sensitive cells (DJ480) and plated onto LB agar for counting plaques (infectious centers). Error bars were calculated as standard deviation of values from three biological replicates.

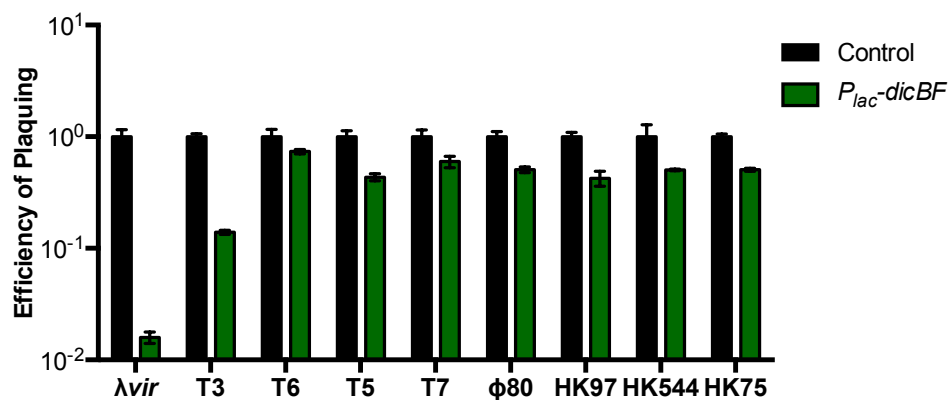


Figure 2. The *dicBF* operon confers resistance against λ phage but not other phages. For each of the nine phages, titer in terms of pfu/ml was calculated by infection of control (DJ480) and $P_{lac-dicBF}$ (DB240) cells. The cells were prepared for infection and the EOP was calculated for each phage as described for Fig.1B. Error bars were calculated as standard deviation of values from three biological replicates.

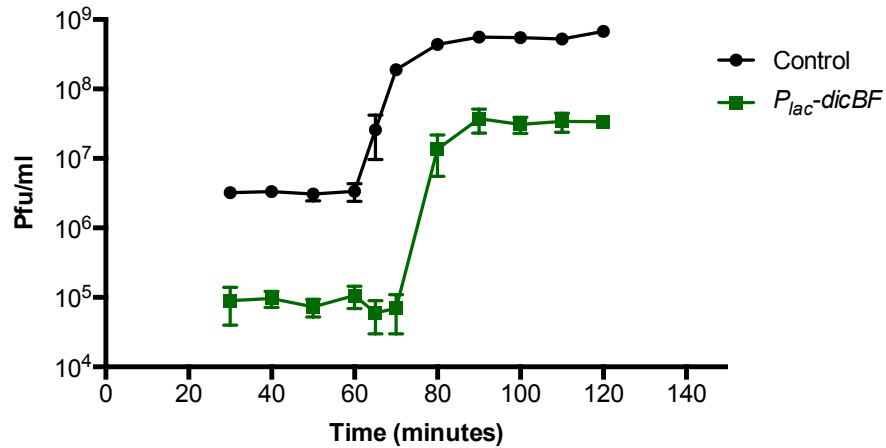


Figure 3. One step growth curve of λvir control and $P_{lac-dicBF}$ cells. The cells were grown with induction of the *dicBF* operon and infection was carried out an MOI of 0.1, similar to the center of infection assay. After removing unadsorbed phages, the cells were diluted in LB medium (with IPTG to induce the *dicBF* operon) and incubated at 37°C for the entire duration of the growth curve. At each time point starting at 30 minutes from the start of infection, samples were removed and added to the phage-sensitive strain (DJ480), and plated to count plaques. Burst size was calculated as (phage titer at 100 minutes - initial titer at 30 minutes)/initial titer at 30 minutes. Latent period was calculated as the time at the mid-point of the exponential phase of the curve. Error bars were calculated as standard deviation of values from three biological replicates.

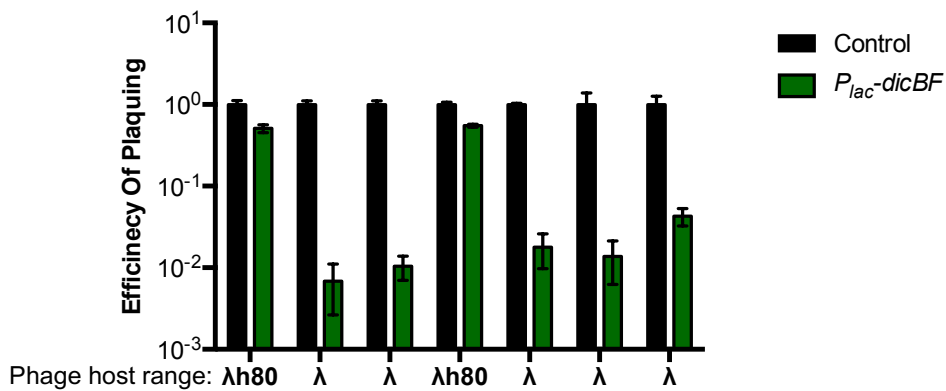


Figure 4. λ phage with the host range of $\phi 80$ is not affected by DicB. Recombinant λ phages with either the λ or $\phi 80$ host range were plaqued on control (DJ480) or $P_{lac-dicBF}$ (DB240) cells. Phage names from left to right are: 185, 148, 169, 173, 158, 138 and λvir (see Table S1 for phage genotypes). The cells were prepared for infection and the EOP was calculated for each phage as described for Fig. 1B. Error bars were calculated as standard deviation of values from three biological replicates.

A

	λ	434	$\phi 80$
Outer membrane	LamB	OmpC	FhuA
Inner membrane	ManYZ	ManYZ	TonB

B

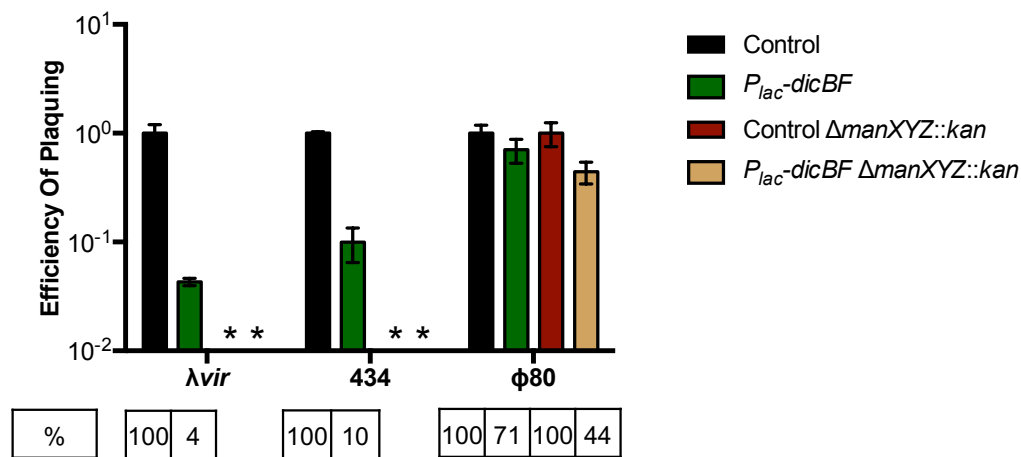


Figure 5. Phage 434 plaquing on ManYZ⁺ strains is inhibited by the *dicBF* operon. (A) Outer and inner membrane receptor specificity of phages λvir , 434 and $\phi 80$. (B) EOP assay was carried out by preparing cells and calculating titer of phages on the different strains as described in Fig.1B. The strains used in this experiment are control (DJ624), *P_{lac}-dicBF* (DB240), control $\Delta manXYZ::kan$ (PR187) and *P_{lac}-dicBF* $\Delta manXYZ::kan$ (PR191). EOP of phages for each strain is calculated with respect to the control strain in the same background. Error bars were calculated as standard deviation of values from three biological replicates. * denotes strains for which plaques could not be counted.

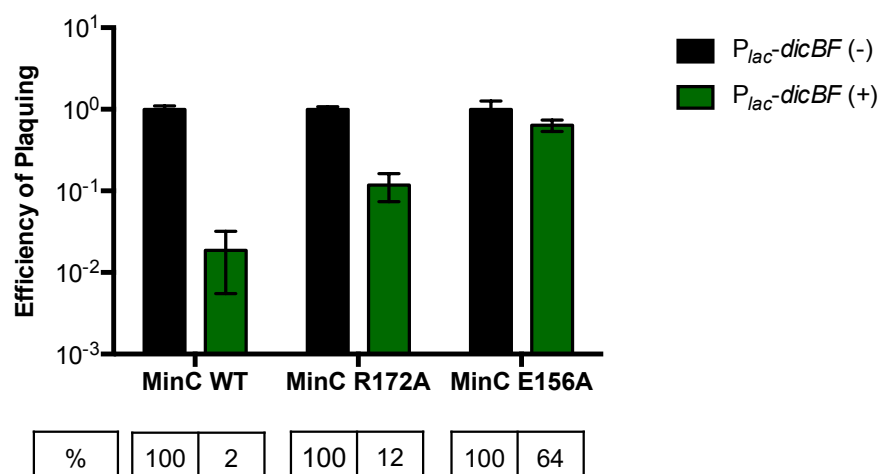


Figure 6. MinC mutants that do not interact with DicB lose the phage resistance effect. The cells were grown with induction of the *dicBF* operon with 0.5mM IPTG, infected with λvir and the EOP calculated as described in Fig.1B. EOP of λvir for each strain is calculated with respect to the control strain in the same background. The strains used in this experiment are control (DJ624), $P_{lac-dicBF}$ (DB240), control *minC* R172A (PR181), $P_{lac-dicBF}$ *minC* R172A (PR183), control *minC* E156A (PR180) and $P_{lac-dicBF}$ *minC* E156A (PR182). Error bars were calculated as standard deviation of values from three biological replicates.