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## Prophage-encoded phage defense proteins with cognate self-immunity

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<b>Abstract:</b>	<p>Temperate phages are pervasive in bacterial genomes, existing as vertically-inherited islands termed prophages. Prophages are vulnerable to the predation of their host bacterium by exogenous phages. Here we identify BstA, a novel family of prophage-encoded phage defense proteins found in diverse Gram-negative bacteria. BstA drives potent suppression of phage epidemics through abortive infection. During lytic replication, the BstA-encoding prophage is not itself inhibited by BstA due to a self-immunity mechanism conferred by the anti-BstA (<i>aba</i>) element, a short stretch of DNA within the <i>bstA</i> locus. Inhibition of phage replication by distinct BstA proteins from <i>Salmonella</i>, <i>Klebsiella</i> and <i>Escherichia</i> prophages is generally interchangeable, but each possesses a cognate <i>aba</i> element. The specificity of the <i>aba</i> element ensures that immunity is exclusive to the replicating prophage, and cannot be exploited by heterologous BstA-encoding phages. BstA therefore allows prophages to defend host cells against exogenous phage attack, without sacrificing their own lytic replication.</p>
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# Prophage-encoded phage defense proteins with cognate self-immunity

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## 1 Summary

2  
3 Temperate phages are pervasive in bacterial genomes, existing as vertically-inherited islands termed  
4 prophages. Prophages are vulnerable to the predation of their host bacterium by exogenous phages.  
5 Here we identify BstA, a novel family of prophage-encoded phage defense proteins found in diverse  
6 Gram-negative bacteria. BstA drives potent suppression of phage epidemics through abortive infection.  
7 During lytic replication, the BstA-encoding prophage is not itself inhibited by BstA due to a self-immunity  
8 mechanism conferred by the anti-BstA (*aba*) element, a short stretch of DNA within the *bstA* locus.  
9 Inhibition of phage replication by distinct BstA proteins from *Salmonella*, *Klebsiella* and *Escherichia*  
10 prophages is generally interchangeable, but each possesses a cognate *aba* element. The specificity of  
11 the *aba* element ensures that immunity is exclusive to the replicating prophage, and cannot be exploited  
12 by heterologous BstA-encoding phages. BstA therefore allows prophages to defend host cells against  
13 exogenous phage attack, without sacrificing their own lytic replication.  
14

## 15 Introduction

16  
17 The eternal battle between bacteria and their viruses (phages) has driven the evolution of a diverse  
18 array of phage defense systems in bacteria (Bernheim and Sorek, 2020; Hampton et al., 2020; Houte  
19 et al., 2016; Rostøl and Marraffini, 2019). Conversely, it is increasingly recognised that phages have  
20 evolved mechanisms to subvert these defense systems (Maxwell, 2017; Samson et al., 2013;  
21 Trasanidou et al., 2019).  
22

23 Although the most intuitive form of phage defense involves the direct rescue of an infected cell, for  
24 example by targeted degradation of phage nucleic acids by CRISPR-Cas, or RM systems, many phage-  
25 defense systems function solely at the population level. In a mechanism conceptually analogous to the  
26 pathogen-stimulated programmed cell death driven by the innate immune systems of higher organisms  
27 (Abedon, 2012), phage infection can be prevented from sweeping across populations, at the cost of the  
28 lives of infected cells. These population-level phage defense systems are often grouped together under  
29 the umbrella term “abortive infection” (Abi) (Labrie et al., 2010; Lopatina et al., 2020) but actually  
30 represent diverse mechanisms to prevent phage replication and induce cell death (Bingham et al., 2000;  
31 Cohen et al., 2019; Fineran et al., 2009; Meeske et al., 2019; Pecota and Wood, 1996; Watson et al.,  
32 2019). Such mechanistic diversity and prevalence of abortive infection systems in nature emphasises  
33 the selective advantage the Abi strategy imparts in the battle against phages (Benler and Koonin, 2020).  
34

35 However, an important sub-plot in the bacteria-phage conflict is the widespread existence of so-called  
36 “temperate” or “lysogenic” phages *within* bacterial genomes. Temperate phages stably exist within the  
37 bacterial chromosome as latent, vertically-inherited islands known as a prophages. Crucially, to find new  
38 hosts, prophages must escape the bacterial genome and return to the lytic life-cycle.  
39

40 The prophage-state imposes unique existential pressures because the fitness of the phage is indefinitely  
41 dependent on that of the host bacterium. To favour their own fitness, prophages frequently encode  
42 “moron” or “accessory” loci that modulate the biology of their host bacteria (Bondy-Denomy and

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43 Davidson, 2014; Cumby et al., 2012; Fortier and Sekulovic, 2013; Howard-Varona et al., 2017), a  
44 phenomenon likened to altruism (Shub, 1994).

45

46 An important trait conferred by prophages that can significantly increase bacterial fitness is resistance  
47 against bacteriophage attack. Indeed, recent work has suggested that prophage accessory genes may  
48 represent an underexplored reservoir of phage-defense systems (Bondy-Denomy and Davidson, 2014;  
49 Dedrick et al., 2017; Snyder, 1995).

50 Here, we report a novel phage defense system driven by the BstA protein, that is encoded by prophages  
51 of diverse Gram-negative bacteria. When a bacterium harbours a BstA-encoding prophage, the BstA  
52 protein confers effective population-level defense against exogenous phage via abortive infection. The  
53 *bstA* locus includes an anti-BstA element, which suppresses the activity of BstA protein to allow the  
54 native prophage to switch to a lytic lifestyle. We propose that this self-immunity mechanism has evolved  
55 to allow prophages to both defend host cells from predatory phages, without compromising their own  
56 lytic replication cycle.

57

## 58 Results

59

### 60 The BstA protein encoded by prophage BTP1 mediates phage defense

61

62 *Salmonella enterica* subsp. *enterica* serovar Typhimurium (hereafter *S. Typhimurium*) strain D23580  
63 encodes the ~40 kb prophage BTP1 (figure 1A) (Owen et al., 2017). An operon within BTP1, the *gtr*  
64 locus (*gtrAC*<sup>BTP1</sup>), confers resistance against phage P22 by chemically modifying the cellular  
65 lipopolysaccharide (LPS), the receptor for phage P22 (Kintz et al., 2015). Unsurprisingly therefore,  
66 deleting the BTP1 prophage from strain D23580 (D23580 ΔBTP1) made the strain highly susceptible to  
67 infection by phage P22, confirming that resistance to phage P22 is conferred by BTP1 (figure 1B).  
68 However, inactivation of the *gtr* locus of prophage BTP1 (D23580 Δ*tsp-gtrAC*<sup>BTP1</sup>) did not restore  
69 sensitivity to phage P22 to the level of D23580 ΔBTP1 (figure 1B), suggesting the existence of a second  
70 BTP1-encoded phage resistance system.

71

72 Previously, we used transcriptomics to discover that the *bstA* gene was highly-expressed from prophage  
73 BTP1 during lysogeny, making it a candidate novel phage accessory gene (Owen et al., 2020). The  
74 *bstA* gene, encoded downstream of the prophage *cl* repressor locus, has been implicated phenotypically  
75 in both virulence and anti-virulence of *Salmonella* isolates, but no functional mechanism has been  
76 proposed (Herrero-Fresno et al., 2014, 2018; Spiegelhauer et al., 2020), and the BstA protein has not  
77 been characterized. We hypothesised that *bstA* was the second element in the BTP1 prophage that  
78 conferred defense against phage P22.

79

80 Consistent with this hypothesis, removal of the *bstA* gene from prophage BTP1 (D23580 Δ*bstA*)  
81 dramatically increased susceptibility to phage P22 (figure 1B). To confirm that phage resistance was  
82 directly mediated by BstA protein, we introduced two stop codons into the beginning of the *bstA* coding  
83 sequence by exchanging 4 nucleotides (D23580 *bstA*<sup>STOP</sup>) (figure 1C). D23580 *bstA*<sup>STOP</sup> was highly  
84 susceptible to P22 phage, to the same level as D23580 Δ*bstA*, demonstrating that BstA protein mediates  
85 defense against phage P22. Simultaneous deletion of the *gtr* locus and inactivation of the BstA protein  
86 (D23580 Δ*tsp-gtrAC*<sup>BTP1</sup> *bstA*<sup>STOP</sup>) recapitulated the susceptibility to phage P22 achieved by deleting the  
87 entire BTP1 prophage (D23580 ΔBTP1), indicating that resistance to phage P22 was solely mediated  
88 by the *bstA* and *gtrAC* loci in prophage BTP1. The findings were reproduced by assaying the replication  
89 of phage P22 on the same strains in liquid culture, demonstrating quantitatively that reduction of plaque  
90 formation by BstA truly reflected suppression of phage replication (figure 1D).

91

92 To investigate whether the defense function of the BstA protein depended on other elements from the  
93 BTP1 prophage, we constructed an inducible expression system in *S. Typhimurium* strain LT2 which  
94 does not contain the BTP1 prophage. LT2 is the type strain of *S. Typhimurium*, and is natively  
95 susceptible to many phages, including P22 (McClelland et al., 2001). Expression of the BstA<sup>BTP1</sup> protein  
96 in *S. Typhimurium* LT2 from within a neutral position on the chromosome (LT2 *tetR-P<sub>tetA</sub>-bstA*) conferred  
97 resistance to P22 and other phages, including ES18 and 9NA (figure 1E; figure S1A).

99 Whilst induced expression of BstA<sup>BTP1</sup> completely eliminated plaque formation of sensitive phages, at  
100 very high phage concentrations ( $10^{9-10}$  PFU/mL) these phages still produced clearing of the bacterial  
101 lawn (figure S1B), consistent with an abortive infection mechanism of phage defense. Expression of the  
102 derivative containing two stop codons in the *bstA* coding sequence (*bstA*<sup>STOP</sup>) conferred no phage  
103 resistance, demonstrating again that defense is mediated by *bstA* at the protein level (figure S1C).  
104 However, BstA did not mediate resistance against all phages tested: Det7, Felix O1, and notably, phage  
105 BTP1 (which encodes the *bstA* gene) were unaffected by expression of BstA, both at the level of plaque  
106 assay and replication in liquid culture (figure 1E; figure S1A). Induction of *bstA* or *bstA*<sup>STOP</sup> expression  
107 in the absence of phage infection did not cause a detectable effect on cell growth rate, suggesting that  
108 overexpression of BstA<sup>BTP1</sup> does not cause toxicity (figure S1D). We were unable to detect any pattern  
109 in the characteristics or gene repertoire of phages that were sensitive or insensitive to BstA protein  
110 which could relate to the mechanistic action of BstA protein.

### 111 **BstA represents a novel family of prophage-encoded phage defense proteins in diverse Gram-** 112 **negative bacteria**

113 Having established that BstA functions as a prophage-encoded phage defense system, we sought to  
114 further characterise the evolutionary conservation of this protein. We identified BstA homologs in the  
115 genomes of diverse Gram-negative bacteria (table S1) and compiled a dataset of 72 homologs  
116 representative of phylogenetic diversity. The majority (79%) of BstA homologs co-occurred with phage  
117 genes, and were designated as putatively-prophage associated (figure 2A). No known phage-  
118 associated genes were found in the vicinity of 21% (15 of 72) of BstA homologs, which were defined as  
119 putatively prophage-independent. A small subset of BstA homologs were plasmid-encoded (figure 2A).  
120 Strikingly, in many cases, BstA homologs were located downstream of putative prophage repressor  
121 proteins, mirroring the genetic architecture of BstA<sup>BTP1</sup> (figure 2B). We conclude that the BstA protein is  
122 highly associated with prophages of Gram-negative bacteria.

123 Whilst the BstA protein does not exhibit sequence homology to any functionally-characterised proteins,  
124 remote homology detection methods revealed a Kila-N (-like) domain in the N-terminal region (residues  
125 32-147 of BstA<sup>BTP1</sup>) (figure 2C). Though poorly characterised, the Kila-N domain is found in proteins  
126 from phages and eukaryotic DNA viruses, and contains the helix-turn-helix motif characteristic of DNA  
127 binding proteins (Iyer et al., 2002; Medina et al., 2019).

128 Certain residues in the BstA protein are highly conserved amongst homologs from diverse members of  
129 the Alpha-, Beta-, and Gamma- Proteobacteria (figure 2C; figure S2A). A small number of BstA  
130 homologs only exhibited homology to the N-terminal, Kila-N (-like) domain. A second small group of  
131 homologs were only homologous to the C-terminal region of BstA (shown at the bottom on the alignment  
132 in figure 2C). Such bipartite protein homology suggests that the BstA protein is composed of two  
133 functional domains. This conclusion is independently supported by evolutionary covariance analysis  
134 (figure S2B) where the clear depletion of predicted residue contacts between the ranges 1 to ~155 and  
135 ~156-307 of BstA<sup>BTP1</sup> suggests that there is a domain boundary (Rigden, 2002) around position 155,  
136 with the two folded domains making few contacts.

137 We selected two diverse BstA homologs from *Klebsiella pneumoniae* (48.4% amino acid identity to  
138 BstA<sup>BTP1</sup>) and *E. coli* (41.7% amino acid identity) to investigate the phage-resistance function of the  
139 larger BstA protein family (the native genetic context of these homologs is illustrated in figure 2B, and  
140 their identity to BstA<sup>BTP1</sup> is highlighted in the alignment in figure 2C). We engineered inducible expression  
141 systems mirroring the expression construct previously validated for BstA<sup>BTP1</sup> (figure 3A,B; figure 1E).  
142 Expression of BstA<sup>Kp</sup> and BstA<sup>Ec</sup> in *S. Typhimurium* LT2 conferred resistance to *Salmonella* phages at a  
143 similar level to BstA<sup>BTP1</sup>, despite these BstA homologs only sharing around 40% identity at the amino  
144 acid level (figure 3A,B; figure S3A). Unlike BstA<sup>BTP1</sup>, BstA<sup>Kp</sup> and BstA<sup>Ec</sup> prevented replication of phage  
145 BTP1 (which encodes *bstA*<sup>BTP1</sup>).

146 Finally, we tested the function of BstA against well-characterised coliphages. Heterologous expression  
147 of BstA<sup>BTP1</sup> in *E. coli* strain MG1655 conferred resistance to phage  $\lambda$ ,  $\phi$ 80, P1 and T7, but did not affect  
148 phages T4 and T5 (figure S3B,C). Surprisingly we found BstA<sup>Ec</sup> was slightly less active against  
149

155 coliphages than BstA<sup>BTP1</sup> (figure S3B,C). Replication in liquid culture was a more reliable and  
156 reproducible measure of phage susceptibility than plaque assay, and frequently revealed stronger  
157 resistance phenotypes than by plaque assay (figure S3B,C).

158  
159 We conclude that BstA represents a novel family of phage-resistance proteins associated with the  
160 prophages of diverse Gram-negative bacteria.

### 161 162 **BstA mediates effective population-level phage defense through abortive infection**

163  
164 Phage resistance systems operate *via* diverse functional mechanisms (Hampton et al., 2020; Rostøl  
165 and Marraffini, 2019). We used a microscopy to dissect BstA-mediated phage-resistance. Virulent P22  
166 phages (P22  $\Delta c2$ ) were used to infect *Salmonella* cells with and without native BstA<sup>BTP1</sup>, at high  
167 multiplicity of infection (MOI) to ensure that most cells were infected. We were surprised to observe that  
168 independent of BstA<sup>BTP1</sup>, all cells lysed within the time course of 3 hours (figure 4A, Supplementary  
169 Video 1), and BstA<sup>BTP1</sup> did not appear to confer any direct protection from phage infection at the level of  
170 individual infected cells. We conducted the same experiment in liquid culture, measuring phage  
171 replication and the fraction of surviving cells *post* phage infection. In cells possessing functional BstA  
172 (D23580  $\Delta tsp-gtrAC$ ), phage P22  $\Delta c2$  completely failed to replicate (figure 4B). In the absence of BstA  
173 function (D23580  $\Delta tsp-gtrAC bstA^{STOP}$ ), the phage replicated >100-fold. However, despite preventing  
174 the replication of phage P22, BstA<sup>BTP1</sup> had no effect on cell survival: independent of BstA<sup>BTP1</sup> function  
175 only 1-2% of cells survived following P22 infection (figure 4C). We hypothesised that BstA does not  
176 protect single cells and instead mediates phage defense at the population level by sacrificing the life of  
177 the infected cell.

178  
179 To investigate whether BstA protein mediated population-level phage defense, we conducted a second  
180 microscopy experiment, wherein approximately only 1 in every 1000 cells was infected with phage P22.  
181 Unlike culture in liquid media, our microscopy setup involved immobilisation of cells on agarose pads,  
182 which restricts the movement of phage particles. The spread of infection was tracked as primary infected  
183 cells lysed and produced secondary infections. To visualise these phage epidemics, we used a reporter  
184 phage engineered to encode the red fluorescent protein mCherry within the early lytic operon (P22  $\Delta c2$   
185 *P-mCherry*); the fluorescence signal indicated phage replication (figure 4D).

186  
187 In the population lacking functional BstA<sup>BTP1</sup> (D23580  $\Delta tsp-gtrAC bstA^{STOP}$ ), primary infected cells lysed  
188 after around 30 minutes (figure 4D, Supplementary Video 2). Subsequently, the red fluorescence signal  
189 was observed in neighbouring cells revealing secondary infection, followed by cell lysis, a cycle which  
190 repeated until all cells in the radius of the primary infected cell had lysed, reminiscent of plaque formation  
191 (figure 4C). The impact of the epidemic of phage infection upon bacterial cells lacking BstA<sup>BTP1</sup> can be  
192 visualised in Supplementary Video 2.

193  
194 In contrast, in the D23580  $\Delta tsp-gtrAC$  population no secondary infection was observed in neighbouring  
195 cells following the lysis of the primary infected cells. Instead, cells continued to divide normally,  
196 eventually forming a confluent lawn (figure 4D, Supplementary Video 2). The lack of subsequent rounds  
197 of infection after the primary cell lysis events shows that few or no infectious phage particles were  
198 generated.

199  
200 Taken together, these experiments demonstrate that BstA protein inhibits successful phage replication,  
201 but does not prevent the death of the infected cell. BstA therefore provides phage defense at the  
202 population-level and prevents the spread of phage epidemics. Accordingly, we propose that BstA is a  
203 novel abortive infection system.

### 204 205 **BstA protein responds dynamically to phage infection and co-localises with phage DNA**

206  
207 To explore the molecular activity of BstA during phage infection, we first constructed a functional  
208 translational fusion of the BstA<sup>BTP1</sup> protein to superfolder green fluorescent protein (<sub>sf</sub>GFP) (figure SS4).  
209 We then used time-lapse fluorescence microscopy to observe the dynamics of BstA protein inside  
210 individual cells during infection with two BstA-sensitive phages, P22 and 9NA. In the absence of phage

211 infection, BstA protein was distributed diffusely within the cytoplasm of the cells, suggesting no particular  
212 sub-cellular localisation (figure S5A, Supplementary Video 3). However, approximately 20 minutes after  
213 infection with phages P22 and 9NA, we consistently observed BstA protein aggregating into discrete  
214 foci towards the centre of infected cells (figure S5B, Supplementary Video 3). Cell lysis occurred  
215 approximately 40 minutes after the formation of BstA foci.

216  
217 We speculated that the dynamic establishment of foci by BstA in response to phage infection was likely  
218 to reflect the mechanistic activity of the protein. We noticed that the foci dynamics of BstA proteins  
219 during phage infection resembled phage replisomes (Cenens et al., 2013; Trinh et al., 2017). We  
220 therefore speculated that the focus of BstA protein in phage infected cells might correspond to the  
221 replicating phage DNA. To test this hypothesis, we used a ParB-*parS* system to track the sub-cellular  
222 localisation of phage DNA relative to BstA protein. We inserted a *parS* site into the P22 phage  
223 chromosome, and expressed a ParB-mCherry fusion protein inside cells already expressing BstA<sub>-sf</sub>GFP.  
224 ParB protein oligomerises onto DNA at *parS* sites, labelling *parS*-tagged DNA with ParB-mCherry foci.  
225 We conducted a microfluidic infection experiment to co-locate BstA foci and infecting P22 phage DNA,  
226 and observed that the position of ParB-mCherry foci (corresponding to phage P22 DNA) overlapped  
227 with foci formed by BstA<sub>-sf</sub>GFP (figure S5C, Supplementary Video 4). The microscopy data therefore  
228 suggest that BstA protein interacts with the replicating DNA of infecting phages. Consistent with the  
229 other microscopy data (figure 4A, figure S5B), cells proceeded to lysis after the formation of BstA/ParB-  
230 mCherry foci. We note that the strain used in this experiment (SVO251, table S2) is cured of all  
231 prophages, ruling out the possibility that cell lysis is caused by native prophage induction.

232  
233 In summary, our data are consistent with a model that involves the movement of BstA protein to sites of  
234 phage DNA replication inside infected cells, followed by prevention of phage replication.

### 235 236 **BstA phage resistance systems contain anti-BstA elements (*aba*) that suppress the activity of** 237 **BstA**

238  
239 When characterising the sensitivity of different phages to the activity of BstA<sup>BTP1</sup> (figure 1E), we observed  
240 that phage BTP1, (which itself encodes the *bstA*<sup>BTP1</sup> gene) was not affected by heterologous expression  
241 of BstA<sup>BTP1</sup> (schematised in figure 5A). We hypothesized that BTP1 carried an anti-BstA determinant: a  
242 self-immunity factor that allows phage BTP1 to replicate without being targeted by its own abortive  
243 infection protein. Consistent with this hypothesis, phage BTP1 became sensitive to BstA<sup>BTP1</sup> expression  
244 when the *bstA* coding sequence was deleted (BTP1  $\Delta$ *bstA*). (figure 5B). The self-immunity function of  
245 the *bstA* locus was not affected by the introduction of the double stop codon mutation into the beginning  
246 of the coding sequence (as described in figure 1C), indicating that self-immunity is not mediated by the  
247 BstA protein itself, but by an alternative genetic element encoded within the *bstA* locus (figure 5B). Here,  
248 and for the duration of this report, we define the *bstA* “locus” as the region including the *bstA* coding  
249 sequence with its 5’ upstream sequence.

250  
251 To identify the genetic basis of BstA self-immunity, we constructed a series of BTP1 mutant phages,  
252 carrying truncations of different lengths from the 3’ end of the *bstA* locus (figure S6A) and screened  
253 these phages for the ability to replicate in the presence of BstA<sup>BTP1</sup> expression. Self-immunity (i.e.  
254 insensitivity to BstA<sup>BTP1</sup> expression) was preserved in all mutant phages except the mutant with the  
255 longest *bstA* truncation (BTP1 *bstA* <sup>$\Delta$ 24</sup>), in which just the first 24 bp of the *bstA* reading frame were intact  
256 (figure S6A). A similar truncation mutant containing just the first 34 bp of *bstA* (BTP1 *bstA* <sup>$\Delta$ 34</sup>) retained  
257 immunity to BstA, suggesting that the first 34 bp of the *bstA* gene are essential for the activity of the  
258 anti-BstA determinant. Transfer of *bstA* <sup>$\Delta$ 34</sup> (the first 34 bp of *bstA*, along with the upstream sequence) to  
259 phage P22 (P22 *bstA* <sup>$\Delta$ 34</sup>), conferred BstA immunity (figure S6B). To identify the minimal sequence  
260 required for BstA self-immunity, we constructed further P22 *bstA* <sup>$\Delta$ 34</sup>-derived phages, successively  
261 truncating the transferred sequence from the 5’ end (P22 *bstA* <sup>$\Delta$ 34I</sup> - P22 *bstA* <sup>$\Delta$ 34V</sup>, figure S6B). We  
262 discovered that a 63 bp sequence  
263 (GCCCGCCACACTTTAACAAAGGAAAATCAAATGGTTAATCAGATAAGGTCCATATCACCCCGCC)  
264 spanning 29 bp of the upstream region and the first 34 bp of the *bstA* coding sequence (start codon  
265 underlined) was necessary and sufficient to confer the self-immunity (figure 5C). We designated this  
266 element ‘*aba*’, for anti-BstA. Supplying the 63 bp *aba* sequence on the high-copy number pUC18  
267 plasmid (pUC18-*aba*) rescued P22 phage replication in the presence of BstA protein, demonstrating



268 that the self-immunity effect of *aba* is retained in *trans* (when *aba* is not carried by the targeted phage,  
269 but is supplied on another replicative element) (figure S7A). The intracellular localisation of BstA protein  
270 following phage infection was unaffected by the presence of the pUC18-*aba* plasmid (figure S7B).

### 272 **The *aba* element appears to be DNA-based**

273  
274 In the native BTP1 prophage, the *aba* sequence overlaps the start of the *bstA* gene, preventing  
275 mutational disruption of the *aba* element without modification of the BstA protein sequence. We  
276 therefore used the plasmid *trans*-complementation system (wherein the BstA protein and the *aba*  
277 sequence are independently encoded) to probe the function of the *aba* sequence (Supplementary  
278 figures 7A). A notable feature of the *aba* sequence is the presence of a direct “CCCGCC” repeat at the  
279 terminal ends, which we hypothesised was functionally important. Single nucleotide exchange of the  
280 CCCGCC→CCCTCC in the first and second repeat (*aba*<sup>mut1</sup> and *aba*<sup>mut2</sup>, respectively) abolished the  
281 self-immunity function of the *aba* element, both when located on a phage (figure 5B) and from a plasmid  
282 *in trans* (figure S8A), showing that the *aba* terminal direct repeats are required for *aba* function. Plasmid-  
283 borne expression of BstA efficiently suppressed plaque formation of P22 and BTP1 phages lacking a  
284 functional *aba* sequence (P22 WT, BTP1  $\Delta$ *bstA* or BTP1 *aba*<sup>mut1</sup>) but had no effect on BTP1 WT which  
285 natively encodes *aba* (figure S8B).

286  
287 The *aba* plasmid *trans*-complementation system additionally allowed us to interrogate the genetic nature  
288 of the *aba* element, which we hypothesised was either DNA, RNA or peptide-based. Though three short  
289 open reading frames exist within the *aba* sequence, non-synonymous mutation of the reading frames  
290 did not ablate *aba* function (figure S9A), suggesting the *aba*-driven immunity is not mediated by a small  
291 peptide.

292  
293 Secondly, we investigated whether the *aba* element is DNA or RNA based by assessing whether  
294 transcription of *aba* is necessary for suppression of BstA. The *aba* sequence was cloned into the high-  
295 copy pUC18 vector with no promoter, and flanked by terminators to abrogate transcription (figure S9B).  
296 This created a scenario with high copy *aba* DNA and minimal *aba* transcription. In parallel, we inserted  
297 the *aba* sequence into the *Salmonella* chromosome, downstream of the arabinose-inducible  $P_{BAD}$   
298 promoter (D23580  $\Delta\Phi$  *tetR-bstA*<sup>BTP1</sup>  $P_{BAD}$ -*aba-gfp*; figure S9B). In this scenario, *aba* exists as a single  
299 copy of DNA but is highly transcribed. In both plasmid and chromosomal constructs a *gfp* gene was  
300 transcriptionally fused to the *aba* sequence to report the level of transcription. Our chromosomal  $P_{BAD}$ -  
301 *aba-gfp* construct generated a high level of green fluorescence in our assay conditions, whereas  
302 fluorescence was barely detectable for our plasmid-based *aba* constructs (figure S9B), demonstrating  
303 that much more *aba* RNA is transcribed from the single copy chromosomal construct than the high copy  
304 plasmid construct. We assessed the activity of BstA in both scenarios by challenging the cells against  
305 phages P22 and 9NA. *aba* only functioned to suppress the activity of BstA (i.e. allow plaquing of P22  
306 and 9NA) in the high copy DNA, low transcription scenario, suggesting that the *aba* element is DNA-  
307 based. However, a single chromosomal copy of *aba* did not confer self-immunity (figure S9B),  
308 suggesting that *aba* can only suppress BstA when supplied on high-copy replicative elements. Further  
309 mutational disruption of the *aba* sequence revealed that the self-immunity function was sensitive to  
310 mutation at multiple sites in the 63 bp sequence (figure S9C).

311  
312 Collectively, our data suggest that *aba*-driven suppression of BstA is neither peptide or transcript-  
313 mediated, and supports a model where BstA suppression is mediated by *aba* DNA.

### 315 **The *aba* element prevents the *bstA*-encoding prophage from aborting its own lytic replication**

316  
317 Unlike most mechanistically-characterised abortive infection systems, a unique feature of the BstA  
318 system is its frequent occurrence on prophages (figure 2A). Prophages must be able to switch to lytic  
319 replication, or else the prophage-state becomes an evolutionary dead-end for the phage.

320  
321 We hypothesised that the primary biological role of the *aba* element is to allow the endogenous *bstA*-  
322 encoding phage to escape BstA-mediated inhibition upon induction from the prophage-state. To test  
323 this, we measured the level of induction of prophage P22 in the presence of heterologously-expressed  
324 BstA<sup>BTP1</sup> protein (figure 5D). In the absence of BstA<sup>BTP1</sup> expression, the P22 prophage generated a titer

325 of  $\sim 4 \times 10^9$  PFU/mL after 5 hours growth with inducing agent (mitomycin C, MitC). However, with BstA<sup>BTP1</sup>  
326 expression, the MitC-induced titer of P22 dropped >300-fold to  $\sim 1 \times 10^7$  PFU/mL, showing that BstA  
327 inhibited P22 phage replication. Transfer of the *aba* sequence to prophage P22 (P22 *aba*) significantly  
328 increased the induced titer in the presence of BstA<sup>BTP1</sup> to  $\sim 1.5 \times 10^9$ , restoring it to the level seen in the  
329 absence of BstA, and showing that the *aba* element rescues prophage induction *via* suppression of  
330 BstA.  
331

332 Finally, we validated the importance of the *aba* element in the context of native BTP1 prophage  
333 induction. The presence of additional copies of the *aba* sequence *in trans* on the pUC18 plasmid did not  
334 affect the titer of BTP1 phage generated after 5 hours growth with inducing agent (MitC), suggesting  
335 that native levels of BstA protein do not constrain BTP1 prophage induction in the presence of the native,  
336 functional *aba* element (figure 5E). However, when the *aba*<sup>mut1</sup> mutation (exchange of a single  
337 functionally important nucleotide in the terminal direct repeat) was introduced into the BTP1 prophage,  
338 the MitC-induced titer of phage BTP1 was reduced  $\sim 40$ -fold in the presence of native BstA expression.  
339 This reduction was almost entirely rescued when the *aba* sequence was supplied *in trans* on the pUC18  
340 plasmid, confirming that the *aba*<sup>mut1</sup> mutation ablates the function of the *aba* element. When the *aba*<sup>mut1</sup>  
341 mutation was introduced into the BTP1 prophage in the absence of native BstA protein expression  
342 (D23580  $\Delta\Phi$  [BTP1 *aba*<sup>mut1</sup> *bstA*<sup>STOP</sup>]) there was no effect on prophage induction (figure S10), confirming  
343 that the effect of the *aba*<sup>mut1</sup> mutation is dependent on the presence of BstA.  
344

345 These experiments demonstrate that a functional *aba* element is required for the *bstA*-encoding  
346 prophage to switch from a lysogenic to lytic lifestyle. In the absence of *aba*, the *bstA*-encoding prophage  
347 suffers replication inhibition by endogenous BstA protein (self-targeting), presumably by the same  
348 abortive infection mechanism that inhibits exogenous phage infection.  
349

### 350 **Distinct BstA proteins are associated with cognate *aba* elements**

351  
352 Finally, we determined whether the *aba* sequence from *bstA*<sup>BTP1</sup> could suppress the activity of the BstA  
353 proteins of other bacteria. We challenged the P22 *bstA*<sup>BTP1</sup> phage (immune to expression of BstA<sup>BTP1</sup>  
354 due to the presence of *aba*) against expression of BstA<sup>Ec</sup> or BstA<sup>Kp</sup>. The *bstA*<sup>BTP1</sup> locus did not protect  
355 P22 from the heterologous BstA proteins, suggesting that the *aba* element from *bstA*<sup>BTP1</sup> only confers  
356 immunity against BstA<sup>BTP1</sup>, and therefore that heterologous BstA proteins have cognate *aba* elements  
357 (figure 5F). To test this hypothesis, we engineered P22 phages to encode either *bstA*<sup>Ec</sup> or *bstA*<sup>Kp</sup>  
358 (including the respective upstream sequence). Consistent with a cognate BstA-*aba* interaction, P22  
359 *bstA*<sup>Ec</sup> became specifically immune to expression of BstA<sup>Ec</sup>, and P22 *bstA*<sup>Kp</sup> gained specific immunity to  
360 BstA<sup>Kp</sup> expression (figure 5F).  
361

362 We conclude that whilst BstA proteins are broadly functionally interchangeable in terms of their phage-  
363 defense activity, each *bstA* locus contains a cognate *aba* element that is inactive against heterologous  
364 BstA proteins. The specificity of the *aba* self-immunity element means that heterologous *bstA*-encoding  
365 phages are unable to bypass BstA-mediated abortive infection, making *aba*-mediated suppression of  
366 BstA exclusive to the induced *bstA*-encoding prophage.  
367

### 368 **BstA protein does not affect phage lysogenic development but inhibits DNA replication during** 369 **lytic development**

370  
371 To interrogate how the BstA protein interacts with infecting bacteriophages, we determined whether  
372 lysogenic phage development, where the infecting phage integrates into the genome of the bacterium,  
373 was affected by BstA expression. We used an antibiotic-tagged derivative of P22 (P22  $\Delta pid::aph$ ) to  
374 determine the frequency of lysogeny with and without BstA expression. We found that the frequency of  
375 lysogeny was approximately 6% (figure 6A) regardless of the presence of BstA, suggesting that BstA  
376 expression does not affect phage lysogenic development. This finding suggests that BstA activity is  
377 triggered by, or targets, an aspect of phage lytic replication not shared by lysogenic development.  
378 Further, it implies that BstA has no effect on initial stages of phages infection that occur prior to  
379 lysogenic development i.e. adsorption and DNA translocation.  
380

381 Sequence-based analysis of BstA protein homologs suggested that the N-terminal domain may bind  
382 DNA (figure 2C), and fluorescence microscopy showed BstA protein co-localizing with phage DNA  
383 (figure S5C). The replication of DNA is crucial for phage morphogenesis, as a new copy of the phage  
384 chromosome is required for packaging into the capsid of each new virion. To test whether BstA protein  
385 inhibits phage DNA replication during lytic development in a manner that can be suppressed by *aba*, we  
386 conducted Southern blot experiments to monitor levels of phage DNA during infection. Using our  
387 prophage-negative, inducible BstA-expression strain (D23580  $\Delta\Phi$  *tetR-P<sub>tetA</sub>-bstA<sup>BTP1</sup>*), we first tested the  
388 replication of the BstA-sensitive virulent phage, 9NA. In the absence of BstA expression, the level of  
389 phage 9NA DNA gradually increased over a 50 minute infection time course, reflecting successful phage  
390 replication (figure 6B). However, no accumulation of phage 9NA DNA was observed in the presence of  
391 BstA<sup>BTP1</sup>, suggesting that BstA protein strongly inhibited the replication of phage DNA.

392  
393 Consistent with the self-immunity function of *aba*, BTP1 phage DNA replication was not affected by the  
394 expression of BstA<sup>BTP1</sup>, unless the *aba* element was non-functional (BTP1 *aba<sup>mut1</sup>*) (figure 6C). Likewise,  
395 successful replication of phage P22 DNA in the presence of BstA<sup>BTP1</sup> only occurred when the phage  
396 possessed a functional *aba* element (figure 5B).

397  
398 To confirm that BstA protein inhibits DNA replication, we constructed small phage-derived plasmids  
399 ('phagemids') based on the phage P22 replication module (pP22) (figure 6D), and a P22 phagemid that  
400 included the 63 bp *aba* sequence (pP22-*aba*).

401  
402 *Salmonella* cells were transformed with the phagemids in the presence or absence of BstA<sup>BTP1</sup> protein  
403 expression. In the absence of BstA, the stable replication of both P22 phagemids in *Salmonella* cells  
404 generated >10<sup>6</sup> transformants/ng phagemid. However, expression of BstA<sup>BTP1</sup> reduced the  
405 transformation efficiency of pP22 (lacking the *aba* sequence) to around 10 transformants/ng. Addition  
406 of the *aba* sequence to the phagemid (pP22-*aba*) restored the transformation efficiency of the phagemid  
407 in the presence of BstA to BstA-negative levels (figure 6D).

408  
409 We conclude that phage DNA replication is strongly suppressed by BstA, but replication can be rescued  
410 by the *aba* element, presumably by suppression of BstA protein activity. As replicated phage DNA is an  
411 essential substrate for packaging into phage capsids, the inhibition of DNA replication is likely to prevent  
412 the production of infectious progeny phages, consistent with the observation that infectious phages are  
413 not released from BstA-expressing cells following cell lysis (figure 4). We propose that BstA protein  
414 suppresses phage DNA replication, a process that can be blocked by the native prophage carrying the  
415 *aba* self-immunity element.

## 416 Discussion

417  
418 Here, we have discovered a novel family of prophage-encoded abortive infection proteins (BstA) which  
419 efficiently defend bacterial populations from phage epidemics. BstA protein is constitutively expressed  
420 inside cells that carry the prophage, and provides effective population-level phage defense through  
421 abortive infection, inhibiting phage replication at the cost of the viability of individual infected cells.  
422 Possession of such innate phage defense systems by active prophages imposes an obvious challenge:  
423 the prophage must avoid self-targeting by its own defense system when switching to lytic replication.

424  
425 The BstA system solves this problem with the *aba* element (anti-BstA), a co-encoded short DNA  
426 sequence that specifically suppresses the activity of BstA protein upon prophage induction, giving the  
427 induced prophage self-immunity against endogenous BstA protein. Theoretically, such a system might  
428 leave BstA-expressing cells vulnerable to infection by heterologous BstA-encoding phages, which could  
429 use their own *aba* element to bypass native BstA. This problem is avoided by cognate BstA-*aba* pairs,  
430 as each BstA protein is suppressed only by the cognate *aba* element, ensuring that BstA suppression  
431 is specific to the native BstA-encoding prophage.

432  
433  
434 Though we present a high-level overview of the BstA phage defense system, and the corresponding  
435 anti-BstA *aba* element, we are left with two major questions regarding the activity of the BstA protein.  
436 Firstly, what are the phage determinants for BstA sensitivity? Though BstA was active against  
437 approximately 50% of the phages tested, we did not detect similarities between BstA-targeted and non-

438 targeted phages that could reflect the molecular determinants of sensitivity. It remains possible that  
439 rather than responding to a physical phage stimulus, such as phage DNA or protein, BstA protein  
440 responds to a cellular stimulus produced by the infection of specific types of phages, for example the  
441 recruitment of DNA replication machinery.  
442

443 Secondly, what is the molecular mechanism by which BstA protein inhibits phage DNA replication? Our  
444 data suggest that phage DNA does not replicate in the presence of BstA. Though numerous Abi systems  
445 in *Lactococcus* have been proposed to interfere with phage DNA replicative functions (Chopin et al.,  
446 2005), the molecular mechanisms have not been well characterised. The existence of a putative DNA-  
447 binding domain in BstA proteins, and microscopic observation of BstA co-localisation with phage DNA  
448 make it tempting to speculate that BstA interacts physically with phage DNA to prevent replication, for  
449 example by occlusion of a replication initiation site.  
450

451 Alongside the mechanistic details of the BstA protein that remain to be established, little is known about  
452 the interaction of BstA with the *aba* element. Our data show that *aba* interacts with BstA in DNA form,  
453 but the mechanism by which *aba* DNA suppresses BstA protein is unclear. Our findings indicate that  
454 multiple copies of the *aba* element are required to suppress BstA protein in *trans*. However, copy-  
455 number cannot be the only factor affecting *aba* functionality, because a prophage is evidently able to  
456 suppress BstA protein right from the initial stages of prophage induction, when *aba* is present as just a  
457 single copy on the chromosome. Whilst we did not observe a loss of BstA focus formation during  
458 suppression with high-copy *aba* DNA, we are cautious to interpret this as evidence against a direct  
459 interaction between *aba* DNA and BstA protein. It remains possible that the phage-DNA colocalization  
460 behaviour and abortive activity of BstA proteins are mechanistically uncoupled (indeed BstA is  
461 predicated to contain two domains), or that reduction in focus formation is beyond the sensitivity of our  
462 microscopy methods. Further study of the BstA-*aba* system is required to resolve the precise molecular  
463 mechanisms by which BstA-encoding prophages, such as BTP1, achieve self-immunity.  
464

465 We consistently observed that phage-infected cells that contained BstA protein underwent lysis,  
466 probably in the absence of infectious progeny phage release. However we cannot be certain whether  
467 BstA protein acts actively or passively to cause cell lysis. Abi systems have frequently been termed  
468 “altruistic suicide” systems, which mediate “programmed cell death” in response to phage infection  
469 (Abedon, 2012; Shub, 1994). Whilst perhaps a useful conceptual analogy for the strictly population-level  
470 effect of Abi systems, this narrative implies that Abi systems actively cause cell death. Though this may  
471 often be the case, such as in the CBASS system (Cohen et al., 2019), Abi can also be achieved by  
472 simple disruption of the phage replication pathway. Because phage lysis is generally a temporally  
473 programmed event that occurs independently of successful virion morphogenesis (Cahill & Young,  
474 2018), phage-mediated cell lysis can occur in the absence of virion assembly. For example many  
475 *Lactococcus* Abi systems target aspects of phage replication, such as AbiZ which is thought to interact  
476 with phage holin proteins, to stimulate premature cell-lysis before virion assembly is completed (Durmaz  
477 and Klaenhammer, 2007).  
478

479 It is possible that BstA protein simply inhibits viable phage particle formation, whilst allowing the phage  
480 lytic pathway to proceed unperturbed to cell lysis. However, inhibition of phage DNA replication would  
481 dramatically reduce substrates for transcription and translation of phage lysis gene products, yet we did  
482 not observe a difference in the timing of cell lysis for phage infected cells in the presence or absence of  
483 BstA during microscopy studies. The exact mechanism of cell lysis during BstA Abi activity will require  
484 further study.  
485

486 An intriguing feature of the BstA phage defense system is its tight association with prophages, and  
487 specifically, with the prophage repressor locus. Though we found homologs in diverse Gram-negative  
488 bacteria, the genetic architecture of the *bstA* locus (i.e. lying downstream of, and presumably sharing  
489 the promoter of the prophage repressor) was strikingly conserved. The region between the repressor  
490 (*cI*) and *n* gene of lambdoid phages has previously been identified as a hotspot of mosaic diversity  
491 (Degnan et al., 2007). In fact, the corresponding site in phage Lambda harbours the *rexAB* genes,  
492 perhaps the most widely studied prophage-encoded abortive infection system (Snyder, 1995). Despite  
493 >60 years of research, the molecular mechanisms of RexAB activity are poorly understood. RexB is  
494 reported to be an ion channel, which triggers loss of cell membrane potential upon activation by the

495 intracellular sensor RexA (Labrie et al., 2010; Snyder, 1995). Whilst not mechanistically comparable to  
496 BstA, perhaps the shared synteny of the BstA and RexAB abortive infection systems points to a  
497 functional significance of this genomic region, as the *cl* repressor gene is one of the most highly  
498 transcribed prophage promoters during lysogeny.

499  
500 Though somewhat functionally analogous to toxin-antitoxin systems, to our knowledge no other example  
501 of self-immunity mechanisms have been described within prophage-encoded abortive infection  
502 systems. However, some evidence supports the widespread existence of such mechanisms. For  
503 example, the activity of Lambda RexB protein can be suppressed by overexpression of the *rexB* gene  
504 relative to *rexA*. It has been speculated, but not shown experimentally, that high levels of RexB might  
505 allow phage Lambda to replicate lytically in the presence of RexAB (Parma et al., 1992) i.e. giving the  
506 Lambda prophage self-immunity against its own Abi proteins.

507  
508 In conclusion, the discovery of the BstA-*aba* system opens unexplored avenues of research into the  
509 mechanisms used by prophages to suppress their own phage-defense activities. We anticipate that  
510 similar strategies may be widespread and commonplace, perhaps existing within known prophage-  
511 encoded phage defense systems. Given the huge mosaic diversity of temperate phages, and high  
512 prevalence of uncharacterised accessory genes, the reservoir of prophage-encoded phage defense and  
513 self-immunity systems is likely to be vast and largely unexplored.

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533 Conceptualization, S.V.O., N.W., and J.C.D.H.; Methodology, S.V.O., N.W., C.L.D., N.Q., and D.J.R.;  
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535 Writing – Original Draft, S.V.O. and N.W.; Writing – Review & Editing, S.V.O., N.W., C.L.D., M.B., and  
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537 J.C.D.H.

## 538 539 **Declaration of Interests**

540 The authors declare no competing interests.

## 541 Figure Legends

542

### 543 **figure 1: The *bstA* gene of prophage BTP1 confers phage defense**

544 (A) Genomic architecture of prophage BTP1 of *S. Typhimurium* D23580, according to Owen *et al.*, 2020:  
545 the LPS modification genes *gtrAC*<sup>BTP1</sup> (Kintz *et al.*, 2015) and the immunity region carrying *bstA*  
546 (downstream of the *cl* repressor gene) are detailed. Bent arrows represent promoters. For reference  
547 purposes the locus tags of important genes in this study in the D23580 reference genome (accession:  
548 FN424405) are shown. (B) Removal of prophage BTP1 from strain D23580 results in enhanced  
549 sensitivity to phage P22. Two BTP1 genes confer resistance to P22; *gtrAC* and *bstA*. Plaque assays  
550 were performed with phage P22 HT 105/1 *int-201* (P22 HT) applied to lawns of *S. Typhimurium* D23580  
551 WT or  $\Delta$ BTP1,  $\Delta$ *bstA*, *bstA*<sup>STOP</sup>,  $\Delta$ *tsp-gtrAC*<sup>BTP1</sup> and  $\Delta$ *tsp-gtrAC*<sup>BTP1</sup> *bstA*<sup>STOP</sup> mutants (strains JH3877,  
552 SSO-204, SSO-78, JH4287 and SNW431, respectively). The requirement for the inactivation of *tsp* is  
553 describes in the Methods. (C) The 4 nucleotide substitutions leading to two nonsense mutations in the  
554 *bstA*<sup>STOP</sup> strain are indicated. SD: putative Shine-Dalgarno sequence of the *bstA* gene. The beginning  
555 of the *bstA* open reading frame is highlighted in blue. (D) Phage replication assays in liquid culture using  
556 P22 HT and the same D23580 derivative strains shown in the plaque assay in B. Replication was  
557 measured 3 hours post-infection and phages were enumerated on lawns of D23580  $\Delta$ *tsp-gtrAC* *bstA*<sup>STOP</sup>  
558 (SNW431). Phage replication is presented as the mean of biological triplicates  $\pm$  SD. (E) BstA protein  
559 confers phage defense in *S. Typhimurium* LT2. Phages P22, ES18, P22HT and 9NA are inhibited by  
560 BstA. Phages Det7, Felix O1 and BTP1 are not affected by BstA expression. Plaque assays were carried  
561 out with the indicated *Salmonella* phages applied to lawns of LT2 *tetR-P<sub>tetA</sub>-bstA*<sup>BTP1</sup> (JH4400) in the  
562 absence (BstA-) or the presence of the inducer anhydrotetracycline (AHT, BstA++). The *tetR-P<sub>tetA</sub>-bstA*  
563 insertion replacing a part of the *STM1553* pseudogene of strain JH4400 is schematized above: *tetR*  
564 encodes the tetracycline repressor that represses the *P<sub>tetA</sub>* promoter in the absence of AHT induction,  
565 “*frr*” denotes the 84 nt scar sequence of pKD4 and the hairpin represents the native *bstA* Rho-  
566 independent terminator (*term*).

567

### 568 **figure 2: BstA homologs are found in diverse bacterial taxa and are frequently associated with**

569 **prophages**

570 (A) A dataset of 72 BstA homologs representative of taxonomic diversity were manually curated and  
571 analyzed for prophage association based on the co-occurrence of phage-related Pfam-domains in the  
572 20 kb either side of each homologs (yielding a total 40 kb window) (table S1). Homologs without co-  
573 occurring phage-related protein domains were assigned to be “Putatively prophage-independent”. A  
574 further subset of the BstA homologs were encoded on plasmids. The top ten most commonly co-  
575 occurring Pfam domains with prophage-associated and putatively prophage-independent BstA  
576 homologs are shown as bar graphs. (B) Gene maps showing the genetic context of a selection of 6  
577 prophage-associated and 6 putatively prophage-independent BstA homologs (homologs indicated by  
578 the grey rectangle). Putative prophage repressor genes are highlighted in red. The top three BstA  
579 proteins from BTP1 (BstA<sup>BTP1</sup>, blue), *K. pneumoniae* 52.145 (BstA<sup>Kp</sup>, green) and *E. coli* NCTC10963  
580 (BstA<sup>Ec</sup>, orange) are studied experimentally in later stages of this work, and so are highlighted. Open  
581 reading frames associated with functional annotations are shown as solid black arrows, and functional  
582 gene name or Pfam domains are annotated. (C) An alignment of the 72 BstA protein homologs to  
583 BstA<sup>BTP1</sup>, with colors indicating amino acid conservation (Clustal colour scheme). Alignment columns  
584 containing gaps relative to the reference sequence (BstA<sup>BTP1</sup>) have been collapsed and are indicated  
585 with blue lines and triangles at the base of the alignment (an expanded alignment can be found in figure  
586 S2). The position of BstA<sup>Kp</sup> and BstA<sup>Ec</sup> within the alignment is highlighted. The position of the Kila-N (-  
587 like) domain (BstA<sup>BTP1</sup> residues 32-147) is indicated by a grey box. Heatmaps on the left of the alignment  
588 indicate the prophage and plasmid association of each homolog (lanes 1 & 2), and the taxonomic group  
589 each homolog derives from (lane 3). Prophage association was split into high and low confidence based  
590 on gene co-occurrence criteria (see Methods).

591

### 592 **figure 3: BstA homologs from *Salmonella*, *E. coli* and *K. pneumoniae* confer phage defense**

593 Heterologous expression of *bstA* homologs from (A) *E. coli* NCTC10963 (*bstA*<sup>Ec</sup>) and (B) *K. pneumoniae*  
594 Kp52.145 (*bstA*<sup>Kp</sup>) in *Salmonella* strain LT2 confers phage defense at similar levels to *bstA*<sup>BTP1</sup>, but show  
595 additional activity against phage BTP1. (C) BstA<sup>BTP1</sup> confers defense against coliphages in *E. coli*  
596 MG1655. Plaque assays were carried out with the indicated phages applied on mock-induced (BstA-)  
597 or AHT-induced (BstA++) lawns of LT2 *tetR-P<sub>tetA</sub>-bstA*<sup>Ec</sup> (JH4408), LT2 *tetR-P<sub>tetA</sub>-bstA*<sup>Kp</sup> (JH4404) or

598 MG1655 *tetR-P<sub>tetA</sub>-bstA<sup>BTP1</sup>* (JH4410). The genetic context for the *tetR-P<sub>tetA</sub>-bstA* insertions within the  
599 *SM1553* pseudogene of LT2 or in the *glmS-pstS* intergenic region of MG1655 are depicted above.

600

#### 601 **figure 4: BstA mediates population-level phage defense through abortive infection**

602 BstA protein does not protect individual cells from phage infection. (A) Cells natively expressing BstA  
603 (D23580  $\Delta$ *tsp-gtrAC*, JH4287) or possessing a mutated BstA locus (D23580  $\Delta$ *tsp-gtrAC bstA<sup>STOP</sup>*,  
604 SNW431) were infected with the obligately virulent P22-derivate phage, P22  $\Delta$ c2, at an MOI of 5 to  
605 increase the likelihood of infecting all cells. Infected cells were imaged on agarose pads and the images  
606 represent a time series. Regardless of BstA function, almost all cells were observed to lyse (indicated  
607 by loss of defined cell shape and phase contrast). Videos of the time series are presented in  
608 Supplementary Video 1. (B) A phage replication assay showed that P22  $\Delta$ c2 phage failed to replicate  
609 after three hours growth on the BstA+ strain (D23580  $\Delta$ *tsp-gtrAC*), but replicated ~136-fold when BstA  
610 was inactivated (D23580  $\Delta$ *tsp-gtrAC bstA<sup>STOP</sup>*). (C) Survival assay of the same strains after infection by  
611 phage P22  $\Delta$ c2, at an MOI of 5. Consistent with the microscopy data in (A), BstA function did not affect  
612 cell survival from phage infection. D23580  $\Delta$  $\Phi$  [P22] (SSO-128), a phage P22 lysogen (and therefore  
613 natively resistant), was used as a negative control. (D) A fluorescent reporter module for phage  
614 replication was added to P22  $\Delta$ c2 (P22  $\Delta$ c2 *P-mcherry*) so that phage replication yielded red  
615 fluorescence (*mcherry* was inserted into the replicative genes of the phage). A similar experiment to (A)  
616 was conducted, but P22  $\Delta$ c2 *P-mcherry* infected cells were mixed 1:1000 with uninfected cells. In the  
617 BstA+ cells, primary infected cells lysed, but did not stimulate secondary infections of neighbouring cells,  
618 and eventually formed a confluent lawn. In BstA- cells, primary lysis events caused secondary infections  
619 (neighbouring cells showing red fluorescence and subsequent lysis) causing an epidemic of phage  
620 infection reminiscent of plaque formation. Cartoons schematise the outcomes of these experiments in  
621 the two strain backgrounds. Videos of the time series are presented in Supplementary Video 2. All  
622 experiments were carried out in liquid or solid M9 Glu<sup>+</sup> media.

623

#### 624 **figure 5: BstA systems include cognate self-immunity elements, *aba*, which are required for 625 successful prophage induction**

626 (A) Cartoon summarising the data from figure 1E. The BTP1 phage, which encodes the *bstA* locus, is  
627 not affected by heterologous expression of BstA<sup>BTP1</sup>, whilst the replication of phage P22 is inhibited.  
628 (B) Schematic of the BTP1-derived phages used and the corresponding effect on sensitivity to  
629 BstA<sup>BTP1</sup> expression (plaque assay). (C) Schematic of the P22-derived phages used. In all cases,  
630 introduced sequences (*bstA* homologs or fragments) were inserted downstream of the *c2* repressor  
631 gene of P22 and are linked to the *frt* sequence. Hairpins represent Rho-independent terminators.  
632 Insensitivity of the BTP1 phage to BstA<sup>BTP1</sup> is dependent on the *bstA* locus on the phage chromosome.  
633 However, only the first 34 bp of the *bstA* gene are required, along with 29 bp upstream (in total a 63  
634 bp sequence). The *aba* sequence (native in BTP1, or engineered into P22) counteracts the BstA-  
635 driven phage resistance. The G→T mutation (*aba<sup>mut1</sup>*) causes loss of *aba* function, and suppresses the  
636 anti-*bstA* interference. (D) BstA represses P22 prophage induction in the absence of *aba*. P22  
637 induction was measured in D23580  $\Delta$  $\Phi$  *tetR-P<sub>tetA</sub>-bstA<sup>BTP1</sup>* lysogenized with prophages P22 WT or P22  
638 *aba* (strain SNW583 and SNW585, respectively). The induced phage titer was measured 5 hours post  
639 induction with Mitomycin C (MitC). (E) Endogenous BstA represses BTP1 prophage induction in the  
640 presence of the *aba<sup>mut1</sup>* mutation, but replication can be rescued by supply of a functional *aba* in *trans*.  
641 Prophage induction was measured in strain D23580  $\Delta$  $\Phi$   $\Delta$ Tn21 (Ap<sup>S</sup>) lysogenized with BTP1 WT  
642 (*aba<sup>WT</sup>*) or BTP1 *aba<sup>mut1</sup>* (strain SNW597 and SNW598, respectively). Lysogens were transformed with  
643 pUC18 (vector) or pUC18-*aba* (pNAW203, +*aba*) and prophage induction was measured 5 hours post-  
644 induction with Mitomycin C. Data in D & E are presented as the mean of biological triplicates  $\pm$  SD.  
645 Groups were compared using unpaired two-tailed Student *t*-test and P values and significance is  
646 indicated by \*, \*\*, \*\*\*, \*\*\*\* or ns (not significant). (F) Each *bstA* locus encodes a homolog-specific anti-  
647 *bstA* element (*aba*) that suppresses BstA-mediated phage defense. Transfer of each *bstA* locus to  
648 phage P22 only confers immunity against the cognate BstA protein. Plaque assays were carried out  
649 with the indicated phages, applied on mock-induced (BstA-) or AHT-induced (BstA++) lawns of the  
650 indicated strain: SNW576 for panels B & C and strains JH4400 (BstA<sup>BTP1</sup>++), JH4404 (BstA<sup>Kp</sup>++) or  
651 JH4408 (BstA<sup>Ec</sup>++) for panel F.

652

#### 653 **figure 6: BstA protein does not affect phage lysogeny but inhibits phage DNA replication in the 654 absence of *aba***

655 (A) Frequency of lysogeny of the P22  $\Delta pid::aph$  phage in mock-induced (BstA-) or AHT-induced  
656 (BstA++) D23580  $\Delta\Phi tetR-P_{tetA}-bstA^{BTP1}$  (SNW576). DNA replication of 9NA phage (B), BTP1 and  
657 P22-derived phages (C) in the absence or presence of BstA expression. Phage DNA was detected by  
658 Southern blotting with total DNA extracted from mock-induced (BstA-) or AHT-induced (BstA++) host  
659 strain D23580  $\Delta\Phi tetR-P_{tetA}-bstA^{BTP1}$  (SNW576), infected by the indicated phage at MOI=5. Before the  
660 transfer procedures, total stained DNA was visualized from gels under UV light and the resulting  
661 pictures served as loading control. Min PI = minutes post infection. Non-infected SNW576 DNA was  
662 used as negative control to check the DNA probe specificity. (D) *aba* dramatically increases the  
663 transformation efficiency of P22 derived phagemids in BstA expressing *Salmonella*. The Km<sup>R</sup>  
664 phagemids pP22 (pNAW229) and pP22-*aba* (pNAW230) are schematized, and the efficiency of  
665 transformation for each phagemid was measured in mock- or AHT-induced competent bacteria of  
666 strain D23580  $\Delta\Phi tetR-P_{tetA}-bstA^{BTP1}$  (SNW576). Data are presented as the mean of biological  
667 triplicates  $\pm$  SD.



668 **STAR Methods**

669

670 **RESOURCE AVAILABILITY**

671 Further information and requests for bacterial and bacteriophage strains should be addressed to the  
672 corresponding author.

673

674 **EXPERIMENTAL MODEL AND SUBJECT DETAILS**

675 **Bacteria and bacteriophages**

676 The full list of bacterial strains used and constructed is available in table S2. All the *Salmonella* strains  
677 were derived from the African *S. Typhimurium* ST313 strain D23580 (GenBank: FN424405.1) (Kingsley  
678 et al., 2009) or the model *S. Typhimurium* strain LT2 (GenBank: AE006468.2)(McClelland et al., 2001b;  
679 Zinder and Lederberg, 1952). All the *Escherichia coli* strains constructed were derived from *E. coli* strain  
680 K-12 substrain MG1655 (GenBank: NC\_000913.3) (Riley et al., 2006). The *bstA* homolog genes were  
681 cloned from *E. coli* NCTC10963 (GenBank: NZ\_CAADJH010000002.1) or from *K. pneumoniae*  
682 Kp52.145 (GenBank: FO834906.1) (Bialek-Davenet et al., 2014). Bacteriophages (phages), including  
683 the temperate phages P22 (GenBank: NC\_002371.2) (Pedulla et al., 2003) and BTP1 (GenBank;  
684 NC\_042346.1) (Owen et al., 2017) and their derivatives, are described in table S2. The genomic  
685 coordinates and gene identifiers indicated below refer to the GenBank accession numbers mentioned  
686 above.

687

688 **METHOD DETAILS**

689 **Growth conditions and transformation**

690 All suppliers of chemical and reagents are specified in the Key Resources Table. Unless stated  
691 otherwise, bacteria were grown at 37°C in autoclaved Lennox Broth (LB: 10 g/L Bacto Tryptone, 5 g/L  
692 Bacto Yeast Extract, 5 g/L NaCl) with aeration (shaking 220 rpm) or on LB agar plates, solidified with  
693 1.5% Agar. The salt-free LBO media contained 10 g/L Bacto Tryptone, 5 g/L Bacto Yeast Extract. Pre-  
694 cultures were inoculated with isolated colonies from agar plates and grown to stationary phase (for at  
695 least 6 hours) in 5 mL LB in 30 mL universal glass tubes or in 50 mL plastic tubes (Greiner).

696

697 Cultures were typically prepared by diluting the pre-cultures (1:100) or (1:1000) in LB, and bacteria were  
698 grown in conical flasks containing 10% of their capacity of medium (*i.e.* 25 mL LB in a 250 mL conical  
699 flask) with aeration. For fluorescent microscopy experiments, bacteria were grown in M9 minimal  
700 medium (Sambrook and Russell, 2001) supplemented with 0.4% glucose and 0.1% Bacto Casamino  
701 Acids Technical (M9 Glu<sup>+</sup>).

702

703 When required, antibiotics were added to the media: 50 µg/mL kanamycin monosulfate (Km), 100 µg/mL  
704 Ampicillin sodium (Ap), 25 µg/mL tetracycline hydrochloride (Tc), 20 µg/mL gentamicin sulfate (Gm), 20  
705 µg/mL chloramphenicol (Cm). Bacteria carrying inducible constructs with genes under the control of the  
706 *P<sub>BAD</sub>* or *P<sub>m</sub>* promoters were induced by adding 0.2 % (w/v) L-(+)-arabinose or 1 mM *m*-toluate,  
707 respectively. For the strains carrying *tetR-P<sub>tetA</sub>* modules, *P<sub>tetA</sub>* induction was triggered by adding 500  
708 ng/mL of anhydrotetracycline hydrochloride (AHT, stock solubilized in methanol). For these constructs,  
709 the same volume of methanol was added to the non-induced cultures (mock treatment). Chemically-  
710 competent *E. coli* were prepared with RbCl-based solutions and were transformed by heat shock (Green  
711 and Rogers, 2014).

712

713 For the preparation of electro-competent cells, bacteria were grown in the salt-free medium LBO to an  
714 Optical Density at 600 nm (OD<sub>600</sub>) of 0.4-0.5. The bacteria were washed twice with cold sterile Milli-Q  
715 water (same volume as the culture volume) and were concentrated 100 times in cold 10% glycerol, prior  
716 to storage at -80°C. When ultra-competent *Salmonella* cells were required, the bacteria were grown in  
717 LBO at 45°C to OD<sub>600</sub> 0.4-0.5, because growth at high temperature inactivates the *Salmonella* restriction  
718 systems (Edwards et al., 1999). Competent cells (10-50 µL) were mixed with 10-5000 ng of DNA in  
719 electroporation cuvettes (2 mm gap) and the reactions were electroporated (2.5 kV) using a MicroPulser  
720 electroporator (Bio-Rad). Bacteria were re-suspended in 0.5-1 mL LB and incubated for recovery at  
721 37°C (30°C for temperature sensitive plasmids) with aeration, for at least one hour. Finally, the  
722 transformed bacteria were spread on selective LB agar plates and transformant colonies were obtained  
723 after at least 12 hours incubation at 30-37°C.

724

725 For assessment of strain growth kinetics with BstA expression, a FLUOstar Omega plate reader (BMG  
726 LABTECH) was used as follows: bacteria were inoculated at an initial OD<sub>600</sub> of 0.01 (six replicates) in  
727 200 µL of LB or LB + AHT in 96-well plates (Greiner). Bacteria were grown at 37°C with aeration (500  
728 rpm, orbital shaking) and the OD<sub>600</sub> was monitored every 15 min for 15 hours. Uninoculated LB medium  
729 was used as blank.

730

### 731 Cloning procedures

732 All the plasmids and DNA oligonucleotides (primers) are listed in table S2. DNA manipulation and  
733 cloning procedures were carried out according to the enzyme and kit supplier recommendations and to  
734 standard procedures (Sambrook and Russell, 2001). DNA purity and concentration were measured with  
735 a DeNovix DS-11 FX spectrophotometer/fluorometer and using the Qubit dsDNA HS assay Kit.

736

737 For all the cloning procedures, Polymerase Chain Reactions (PCRs) were performed with the Phusion  
738 High Fidelity DNA polymerase, purified template DNA and primers in the presence of 3 % Dimethyl  
739 Sulfoxide and 1 M betaine, when required. Prior to Sanger sequencing of the constructs, PCR reactions  
740 were carried out directly from bacteria or phages with MyTaq Red Mix 2X. PCR fragments were analysed  
741 by electrophoresis, purified and finally sequenced with the appropriate primers (Lightrun service,  
742 Eurofins Genomics) (table S2).

743

744 All the plasmids were constructed as detailed in the table S2 and were verified by Sanger sequencing.  
745 Insertions of DNA fragments into plasmids were performed by digestion/ligation procedures, using  
746 restriction enzymes and the T4 DNA ligase. In addition, PCR-driven restriction-free cloning techniques  
747 were used: overlap extension PCRs (Heckman and Pease, 2007) and plasmid assembly by PCR cloning  
748 (Van Den Ent and Löwe, 2006) were performed with chimeric primers, purified template DNA and  
749 Phusion DNA polymerase, as described previously (Owen et al., 2020). Cloning reactions were  
750 transformed by heat shock into *E. coli* Top10 (Invitrogen) or S17-1  $\lambda$ pir (Simon et al., 1983). New  
751 template plasmids were constructed to insert fluorescent protein encoding genes into *Salmonella* or *E.*  
752 *coli* chromosomes, as reported previously (Gerlach et al., 2007). These plasmids carry the *oriR6K*  $\gamma$   
753 origin of replication of pEMG, the *frt-aph-frt* (Km<sup>R</sup>) module of pKD4 linked to *gfp*<sup>+</sup> (pNAW52), *s<sub>gfp</sub>*  
754 (encoding for superfolder GFP, pNAW62) or *mcherry* (pNAW73), amplified respectively from plasmids  
755 pZEP09 (Hautefort et al., 2003), pXG10-SF (Corcoran et al., 2012) and pFCcGi (Figueira et al., 2013).  
756 A similar template plasmid, carrying the *frt-aph-frt-tetR-P<sub>tetA</sub>* module (pNAW55) was constructed and  
757 was used to insert the *tetR* repressor and the AHT-inducible promoter *P<sub>tetA</sub>* upstream of genes of interest,  
758 as reported earlier (Schulte et al., 2019). For the construction of gentamicin resistant plasmids, the  
759 *aacC1* resistance gene was obtained from plasmid pME4510 (Rist and Kertesz, 1998).

760

761 The high copy number plasmid pUC18 was used to clone the different versions of the anti-*bstA* (*aba*)  
762 fragment: the *aba* fragments (*aba1-aba14* alleles) were amplified by PCR, digested with EcoRI and  
763 BamHI and ligated into the corresponding sites of pUC18. For cloning of the *aba* fragments fused to  
764 *gfp*<sup>+</sup> and flanked by terminators, 20nt overlapping DNA fragments were amplified with Q5 high fidelity  
765 polymerase, pooled and digested with DpnI prior to four piece isothermal assembly using the NEBuilder  
766 HiFi DNA Assembly Cloning Kit.

767

768 Phagemids based on the phage P22 replication module were constructed by EcoRI/KpnI digestion and  
769 ligation, as follows: the *P<sub>R</sub>* promoter and the *cro-c1-orf48-O-P* genes of P22 (coordinates 31648-34683)  
770 were amplified and circularized by ligation with the *aph* Km<sup>R</sup> cassette of pKD4 or with the *aba-aph*  
771 modules, amplified from strain SNW617. The ligation reactions were purified and electroporated into  
772 ultra-competent SNW555, a prophage-free and plasmid-free derivative of *S. Typhimurium* D23580. The  
773 resulting phagemids pNAW229 (pP22-*aph*), pNAW230 (pP22-*aba-aph*) were obtained after selection  
774 on Km medium.

775

776

777 Phage DNA was extracted from high titer lysates in LBO: nine volume of the phage lysates were mixed  
778 with one volume of 10 X DNase buffer (100 mM Tris-HCl, 25 mM MgCl<sub>2</sub>, pH 7.5) supplemented with  
779 RNase A (40 µg/mL final) and DNase I (400 µg/mL final). After 1 hour incubation at 37°C, DNase I was  
780 heat-inactivated at 75°C for 10 min and phage DNA was extracted from 500 µL of the nuclease-treated

781 lysates with the Norgen Phage DNA Isolation after Proteinase K treatment, as specified by the  
782 manufacturer.

783

### 784 **Genome editing techniques**

785 Strain constructions are detailed in table S2. For chromosomal insertions and deletions,  $\lambda$  *red*  
786 recombination was carried out with the arabinose-inducible plasmid pKD46 (for *E. coli*) or with the heat  
787 inducible plasmid pSIM5-*tet* (for *Salmonella*), both expressing the  $\lambda$  *red* genes. Bacteria were grown to  
788 exponential phase in LBO, according to the resistance and induction condition of the respective  $\lambda$  *red*  
789 plasmid (Datsenko and Wanner, 2000; Hammarlöf et al., 2018; Koskiniemi et al., 2011) and electro-  
790 competent cells were prepared as mentioned above. PCR fragments carrying a resistance cassette  
791 were constructed by overlap extension PCR or were directly obtained by PCR from the appropriate  
792 plasmid or strain. Electro-competent cells (40-50  $\mu$ L) were transformed with 500-5000 ng of the PCR  
793 fragments and the recombinants were selected on selective LB agar plates.

794

795 Mutations or insertions linked to selective markers were transduced into *Salmonella* strains using the  
796 P22 HT 105/1 *int-201* (P22 HT) transducing phage (Owen et al., 2017; Schmieger, 1972). For *E. coli*,  
797 the transducing phage P1 *vir* was used (Ikeda and Tomizawa, 1965; Tiruvadi Krishnan et al., 2015).  
798 Transductants were grown on selective LB agar plates supplemented with 10 mM EGTA. After two  
799 passages, clearance of the transducing phages was confirmed by diagnostic PCR using primer pairs  
800 NW\_62/NW\_63 for P22 HT or NW\_392/NW\_393 for P1 *vir* and by a passage on Green Agar medium  
801 (Maloy, 1990). To remove the antibiotic cassettes, flanked by FLP recognition target sites (*frt*), the FLP  
802 recombinase expressing plasmids pCP20, pCP20-TcR and pCP20-Gm were used, as previously  
803 reported (Cherepanov and Wackernagel, 1995; Doublet et al., 2008; Hammarlöf et al., 2018; Kintz et  
804 al., 2015). The inducible *tetR-P<sub>tetA</sub>-bstA* modules were constructed by fusing the *frt-aph-frt-tetR-P<sub>tetA</sub>*  
805 module of pNAW55 to the *bstA* gene of D23580 (*bstA<sup>BTP1</sup>*, *STMMW\_03531*), *E. coli* NCTC10963 (*bstA<sup>Ec</sup>*,  
806 *E4V89\_RS07420*) or *K. pneumoniae* Kp52.145 (*bstA<sup>Kp</sup>*, *BN49\_1470*). Each construct carries the native  
807 *bstA* ribosome binding site and Rho-independent terminator. The *tetR-P<sub>tetA</sub>-bstA* modules were inserted  
808 by  $\lambda$  *red* recombination into the *STM1553* pseudogene of *S. Typhimurium* LT2 (between coordinates  
809 1629109-1629311), corresponding to *STMMW\_15481* in D23580 (coordinates 1621832-3). Previously  
810 we have shown that the *STM1553* and *STMMW\_15481* genes are not expressed at the transcriptional  
811 level (Canals et al., 2019).

812

813 In *E. coli* MG1655, the *bstA* modules were inserted into the *glmS-pstS* intergenic region (coordinates  
814 3911773-4). To generate Ap and Cm sensitive D23580 strains, the pSLT-BT plasmid-encoded Tn21-  
815 like element, that carries the resistance genes (Kingsley et al., 2009), was replaced by the Km<sup>R</sup> cassette  
816 of pDK4 by  $\lambda$  *red* recombination (deletion coordinates 34307 to 57061, GenBank: NC\_013437.1). The  
817 resulting large single-copy plasmid pSLT-BT  $\Delta$ Tn21::*aph* was extracted (Heringa et al., 2007) and  
818 electroporated into the strains of interest. After selection on Km medium, the Ap and Cm sensitivity was  
819 confirmed and the Km<sup>R</sup> cassette was flipped out using pCP20-Gm. For scarless genome editing, the  
820 pEMG plasmid-based allelic exchange system was used (Martínez-García and de Lorenzo, 2011). The  
821 pEMG derivative suicide plasmids were constructed as specified in table S2 and were replicated in *E.*  
822 *coli* S17-1  $\Delta$ *pir*. Conjugation of the resulting plasmids into *Salmonella* and subsequent merodiploid  
823 resolution with plasmid pSW-2 were carried out as previously described (Canals et al., 2019; Owen et  
824 al., 2017). Some key strains and phages (indicated in table S2) used in this study were whole-genome  
825 sequenced (Illumina) at MicrobesNG (Birmingham, UK).

826

### 827 **Plasmid deletion in *S. Typhimurium* D23580**

828 The pSLT-BT, pBT1, pBT2 and pBT3 plasmids (Kingsley et al., 2009) were cured from strain D23580,  
829 using the CRISPR-Cas9-based methodology (Lauritsen et al., 2017). A novel CRISPR-Cas9 Km  
830 resistant plasmid (pNAW136) was obtained by ligating the CRISPR-Cas9 module of plasmid pCas9  
831 (Jiang et al., 2013) with the unstable origin of replication *oriRK2*, the *trfA* replication gene and the *aph*  
832 Km<sup>R</sup> gene. Anti-plasmid protospacers (30 bp) were generated by the annealing of 5'-phosphorylated  
833 primer pairs that targeted the pSLT-BT, pBT1, pBT2 and pBT3 plasmids, designed according to the  
834 Marraffini Lab protocol (Jiang et al., 2013). The protospacers were ligated into *Bsal*-digested pNAW136  
835 with T4 DNA ligase and the resulting plasmids were checked by Sanger sequencing, using primer  
836 NW\_658.

837

838 The resulting plasmids pNAW168 (anti-pSLT-BT) and pNAW169 (anti-pBT1), pNAW139 (anti-pBT2)  
839 and pNAW191 (anti-pBT3) were electroporated into D23580-derived strains and transformants were  
840 selected on Km plates. After two passages on Km, the loss of the pSLT-BT, pBT1, pBT2 or pBT3  
841 plasmids was confirmed by diagnostic PCR. The absence of the unstable pNAW136-derived plasmids  
842 was confirmed by the Km sensitive phenotype of colonies after two passages on non-selective medium.

### 843 **Phage stock preparation and plaque assays**

844 All phage stocks were prepared in LB or LBO. For *Salmonella* phages, the prophage-free strain *S.*  
845 *Typhimurium* D23580  $\Delta\Phi$  (JH3949) was used as host (Owen et al., 2017). Exponential phase cultures  
846 of D23580  $\Delta\Phi$  were infected with  $\sim 10^5$  Plaque Forming Units (PFU) and infected cultures were incubated  
847 for at least 3 hours at 37°C (with aeration). Phages lysates were spun down (4,000 X *g*, 15 min) and  
848 supernatants were filter-sterilized (0.22  $\mu\text{m}$ , StarLab syringe filters). The resulting phage lysates were  
849 stored at 4°C in the presence 1% chloroform to prevent bacterial contamination.

850  
851 Coliphage lysates were prepared similarly with *E. coli* MG1655 as host. When required, maltose (0.2%),  
852  $\text{CaCl}_2$  (10 mM) and  $\text{MgSO}_4$  (10 mM) were added during the infection ( $\lambda$ , P1 *vir* and  $\Phi 80\rho\text{SU}3^+$ ). For  
853  $\Phi 80$ -derived phages, the infection temperature was reduced to 30°C (Rotman et al., 2010).

854  
855 Phage lysates were serial-diluted (decimal dilutions) with LB and virion enumeration was performed by  
856 double-layer overlay plaque assay (Kropinski et al., 2009), as follows. Bacterial lawns were prepared  
857 with stationary phase cultures of the reporter strains, diluted 40 times with warm Top Agar (0.5 % agar  
858 in LB, 50°C). The seeded Top Agar was poured on LB 1.5% agar bottom layer: 4 mL for 8.6 cm diameter  
859 petri dishes or 8 mL for 12 x 12 cm square plates.

860  
861 When inducible  $P_{tetA}$  or  $P_{BAD}$  constructs were present in the reporter bacteria, 500 ng/mL of AHT or 0.2  
862 % arabinose were added in the Top Agar. When required, antibiotics were added in the Top Agar layer.  
863 The bacterial lawns were incubated for 30 min at room temperature with the appropriate inducer, to  
864 allow solidification and the expression of the inducible genes. Finally, phages suspensions (5-20  $\mu\text{L}$ )  
865 were applied on the Top Agar surface and pictures of the resulting plaques were taken with an  
866 ImageQuant Las 4000 imager (GE Healthcare) after 16-20 hours incubation at 30 or 37°C.

### 868 **Construction of P22 virulent phages**

869 For the generation of obligately virulent P22 phages, a 633 bp in-frame deletion (coordinates 31028-  
870 31660) was introduced in the *c2* repressor gene by  $\lambda$  *red* recombination in a P22 lysogen as follows.  
871 Two fragments of  $\sim 500$  bp, flanking *c2*, were amplified with primers pairs NW\_818 / NW\_819 and  
872 NW\_820 / NW\_821. The two amplicons were fused by overlap extension PCR and 1000-3000 ng of the  
873 resulting  $\Delta c2$  fragment were electroporated into P22 lysogens (in the prophage-free D23580  $\Delta\Phi$   
874 background) carrying the  $\lambda$  *red* recombination plasmid pSIM5-*tet*, as described above. The  
875 transformation reactions were re-suspended in 5 mL LB and incubated for 2 hours at 37°C with aeration.  
876 The culture supernatants were filter sterilized and serial-diluted to  $10^{-2}$ . Ten microliters of each dilution  
877 were mixed with 100  $\mu\text{L}$  of a D23580  $\Delta\Phi$  stationary phase culture and with 4 mL of warm Top Agar. The  
878 mixtures were poured on LB agar plates and the plates were incubated for  $\sim 16$  hours at 37°C. P22  $\Delta c2$   
879 recombinants were identified by the clear morphology of their plaques, compared to the turbid plaques  
880 of WT P22. The  $\Delta c2$  deletion was confirmed by PCR and Sanger sequencing with primers NW\_406 and  
881 NW\_805.

### 883 **Use of the $\Delta tsp$ -*gtrAC* genetic background**

884 Where possible, experiments were carried out with native BstA expression (from its natural locus within  
885 the BTP1 prophage), to best recapitulate the natural biological activity of the protein. However, as the  
886 *gtr* locus of phage BTP1 blocks attachment of many phages including P22 and BTP1, to achieve efficient  
887 phage infections we consistently used a strain background where the *gtr* locus has been inactivated  
888 ( $\Delta tsp$ -*gtrAC*). The BTP1 prophage spontaneously induces to a titer of  $\sim 10^9$  PFU/mL in liquid culture  
889 (Owen et al., 2017), and in the absence of *gtr* activity in surrounding cells, free BTP1 phages mediate  
890 cleavage of the O-antigen *via* the putative enzymatic activity of the tailspike protein (Kintz et al., 2015).  
891 Consequently, to avoid an unnatural, short LPS phenotype as a result of *gtr* inactivation in a BTP1  
892 lysogen, we additionally inactivated the upstream gene encoding for the BTP1 tailspike (*tsp*) (D23580  
893  $\Delta tsp$ -*gtrAC*, JH4287). Full details of the construction of this strain can be found in the table S2.

894

### 895 **Phage replication assay**

896 Stationary phase cultures of the reporter bacteria were diluted to OD<sub>600</sub> 0.4 with LB. Aliquots (0.2 mL)  
897 were prepared in 1.5 mL tubes and phage stock suspensions were added to a final phage titer of 100-  
898 1000 PFU/mL. The infections were carried out at 37°C (30°C for Φ80pSU3<sup>+</sup>) with shaking for 2-4 hours  
899 and were stopped by the addition of 20 μL of chloroform. After a 10 sec vortex, the lysates were  
900 centrifuged (20,000 X g, 5 min) and serial diluted. When M9 Glu<sup>+</sup> was used, *Salmonella* strains were  
901 grown to OD<sub>600</sub> ~ 0.5 in this medium prior to phage infection.

902

903 Phage titer was determined by plaque assay: 10 μL of the dilutions were applied to bacterial lawns of  
904 the appropriate reporter strain in technical triplicates. Plaques were enumerated after 16-20 hours of  
905 incubation and phage titers (PFU/mL) were calculated for each lysate. To measure the phage input at  
906 time 0 (T<sup>0</sup>), the same volume of stock phage suspension was added to 0.2 mL of bacteria-free LB and  
907 the titer was determined as described above. The fold-replication for each phage was calculated as the  
908 phage titer of the lysate post infection divided by the input phage titer at T<sup>0</sup>. When the phage titer in the  
909 lysate was lower than the phage input, the replication was considered to be null (<1-fold). When AHT  
910 inducible *tetR-P<sub>tetA</sub>-bstA* strains were used, AHT (500 ng/mL) or methanol (mock) were added to the  
911 diluted bacterial suspension and phages were added after 15 min of incubation at 37°C with aeration.

912

913 For replication assays of the coliphages λ, P1 *vir* and Φ80pSU3<sup>+</sup>, *E. coli* strains were grown to  
914 exponential phase (OD<sub>600</sub> 0.4) in LB and phages were added as mentioned above. To stimulate infection  
915 by these phages, maltose (0.2%), CaCl<sub>2</sub> (10 mM) and MgSO<sub>4</sub> (10 mM) were added during the infection  
916 and in the lawns of the reporter *E. coli* MG1655. All the phage replication experiments presented were  
917 carried out at least twice with biological triplicates.

918

### 919 **Induction of P22 and BTP1 prophages**

920 D23580 ΔΦ-derived lysogens that carried the different versions of P22 and BTP1 were constructed as  
921 detailed in the table S2. For complementation with the pUC18-derived plasmids (Ap<sup>R</sup>), Ap sensitive  
922 lysogens were constructed by the inactivation of the Tn21-like element, as described above. The  
923 resulting lysogens were grown to stationary phase in LB and the pre-cultures were diluted 100-1000  
924 times in fresh LB and grown to OD<sub>600</sub> 0.4-0.5, prior addition of Mitomycin C (MitC, 2 μg/mL). The induced  
925 cultures were incubated for 3-5 hours at 37°C with aeration and cultures were filter sterilized and serial  
926 diluted. The phage titer was measured by plaque assay on the appropriate host strain lawn with technical  
927 replicates, as described above. All the prophage induction experiments were carried out at least twice  
928 with biological triplicates.

929

### 930 **Survival assays**

931 For the survival assays, D23580 Δ*tsp-gtrAC* (JH4287), D23580 Δ*tsp-gtrAC bstA*<sup>STOP</sup> (SNW431) or  
932 D23580 ΔΦ [P22] (SSO-128) were grown in M9 Glu<sup>+</sup> to OD<sub>600</sub> ~0.5 and two 0.5 mL subcultures were  
933 prepared for each culture. The use of D23580 ΔΦ [P22] in these experiments controlled for the effect of  
934 lysis from without due to use of high multiplicity of infection (MOI). The strain is a lysogen for WT P22  
935 phage, and therefore is highly resistant to infection by P22-derived phages. P22 Δ*c2* was added at an  
936 MOI of 5. The same volume of LB was added to the two remaining subcultures (non-infected controls).  
937 Samples were incubated for 15 min at 37°C to allow phage attachment. To stop phage development,  
938 the cultures were chilled on ice and bacteria were washed with 0.5 mL of cold PBS. All the samples  
939 were serial-diluted in PBS to 10<sup>-6</sup> and kept on ice. For the measure of survival post-infection, 10 μL of  
940 diluted infected or non-infected cultures were applied in technical triplicates on LB agar supplemented  
941 with 10 mM EGTA (EGTA was used to minimize secondary infection by free phages). Colony forming  
942 Units (CFU) were enumerated and the survival rate, was calculated as the ratio of CFUs in infected  
943 cultures divided by the CFUs obtained from non-infected cultures (in %). All the survival experiments  
944 were carried out at least twice with biological triplicates.

945

### 946 **Frequency of lysogeny assays**

947 For the frequency of lysogeny assays, a derivative of phage P22 was used that has the *pid* locus replaced  
948 with an *aph* cassette yielding kanamycin resistant lysogens (P22 Δ*pid::aph*, SNW490). The *pid* locus  
949 has previously been shown to be non-essential in phage P22 and does not establishment of lysogeny  
950 (Cenens et al., 2013). D23580 ΔΦ *tetR-P<sub>tetA</sub>-bstA* (SNW576) cells were grown in 3 mL of LB to OD<sub>600</sub>

951 ~0.35. Methanol (mock) or AHT (500 ng/mL, inducer) were added to the cultures and bacteria were  
952 incubated to induce BstA for 1 hour at 37°C. 200 µL samples of the bacteria were mixed in triplicate with  
953 P22  $\Delta$ *pid::aph* phage to achieve a MOI of 0.1, and incubated at 37°C for 20 minutes to allow adsorption  
954 and ejection of nucleic acids. Cells were pelleted and resuspended in LB media supplemented with 10  
955 mM EGTA to minimize secondary infection by any free phages (along with methanol or AHT) and  
956 incubated at 37°C for a further 20 minutes to allow integration and expression of the kanamycin  
957 resistance determinant. Colony forming Units (CFU) were enumerated on LB kanamycin. Frequency of  
958 lysogeny was determined as the kanamycin resistant CFU/mL divided by the PFU/mL of input phage.  
959

### 960 **Phage DNA detection by Southern Blotting**

961 D23580  $\Delta\Phi$  *tetR-P<sub>tetA</sub>-bstA* (SNW576) was grown in 50 mL LB to OD<sub>600</sub> ~0.35. The culture was split in  
962 two 20 mL sub-cultures and methanol (mock) or AHT (500 ng/mL, inducer) were added to each  
963 subculture. Bacteria were incubated to induce BstA for 20 min at 37°C and the phage of interest was  
964 added at an MOI of 5. Infections were carried out at 37°C with aeration and total DNA was extracted  
965 (Quick-DNA™ Universal Kit Zymo) from 1.5 mL of culture at 0, 10, 20, 30, 35, 40 and 50 minutes post  
966 infection. Total DNA (100 ng, according to QuBit quantification) was size-separated (2 hours at 100 V  
967 in TAE 1X) on a 0.8 % agarose-TAE gel containing Midori Green DNA staining (4 µL for 100 mL gel).  
968 One hundred nanograms of none-infected D23580  $\Delta\Phi$  *tetR-P<sub>tetA</sub>-bstA* genomic DNA were used as a  
969 negative control. DNA was fragmented by exposing the agarose gel to UV light for 5 min on a UV-  
970 transilluminator. DNA was denatured by soaking the gel in the Denaturation Solution (0.5 M NaOH, 1.5  
971 M NaCl) for 30 min and then in the Neutralization Solution (1.5 M NaCl, 1 M Tris-HCl, pH 7.6) for 30  
972 min. DNA was transferred on a positively-charged Nylon membrane using the capillary blotting method.  
973 Phage DNA was detected with DIG labelled dsDNA probes generated by PCR amplification with MyTaq  
974 DNA polymerase (Bioline), buffer, phage DNA and primers (0.4 µM each), in the presence of 0.2 mM  
975 dATP, 0.2 mM dCTP, 0.2 mM dGTP, 0.13 mM dTTP and 0.07 mM DIG-11-dUTP. For the 9NA probe a  
976 588 bp PCR fragment was generated with primer pair NW\_602 / NW\_603 and for the P22/BTP1 probe  
977 a 725 bp PCR fragment was generated with primer pair SO-22 / SO-23. The DNA probes were heat-  
978 denatured at 95°C for 15 min and the DNA-DNA hybridizations were carried out at 45°C for 16 hours in  
979 DIG-Easy Hyb buffer. The washing and immunodetection procedures were carried out, as specified in  
980 the DIG Application Manual for Filter Hybridization (Roche) and the chemiluminescence signal was  
981 detected using an ImageQuant LAS 4000 imager (GE Healthcare). Prior to DNA transfer onto the  
982 membrane, the Midori green-stained DNA was visualized under UV and the resulting image was used  
983 as a loading control.  
984

### 985 **Phagemid efficiency of transformation**

986 To avoid a reduction in transformation caused by interference interspecies DNA  
987 modification/restriction interference between *E. coli* and *Salmonella*, all the P22-derived  
988 phagemids were replicated and extracted from *S. Typhimurium* SNW555.  
989

990 *Salmonella* strains carrying the *tetR-P<sub>tetA</sub>-bstA* module were grown in 50 mL LBO culture. When  
991 OD<sub>600</sub> ~0.4 was reached, each culture was split into two 25 mL sub-cultures and methanol  
992 (mock) or AHT (inducer) were added to each subculture. Bacteria were incubated for BstA  
993 induction during 15 min at 37°C. The cultures were incubated on ice for 5 min and bacteria  
994 were washed twice with cold water (25 mL) and were concentrated in 0.1 mL of ice-cold sterile  
995 10% glycerol. The OD<sub>600</sub> of each electro-competent cell sample was measured by diluting 10  
996 µL of competent cells with 990 µL of 10% glycerol. Cell concentration was adjusted with 10%  
997 glycerol for each sample, according to the sample with the lowest OD<sub>600</sub>. The competent cells  
998 (20 µL) were mixed with 10 ng (estimated by Qubit) of the P22 phagemids, pP22 (pNAW229)  
999 or pP22-*aba* (pNAW230) and the mixture was incubated on ice until electroporation (2.5 KV).  
1000 Transformation reactions were re-suspended in 1 mL LB or 1 mL LB + AHT (for the *bstA*-  
1001 induced bacteria) and were incubated for 60 min at 37°C, for recovery. The transformations  
1002 were diluted (decimal dilution to 10<sup>-5</sup>) in LB or LB+AHT and 100 µL of each dilution (including  
1003 the non-diluted sample) were spread on LB agar Km or LB agar Km+AHT plates. After  
1004 incubation at 37°C, the number of Km<sup>R</sup> transformants was enumerated for each transformation  
1005 and efficiency of transformation was defined as the number of transformants obtained per ng

of phagemid. This experiment was performed with biological triplicates and was repeated twice with LT2 *tetR-P<sub>tetA</sub>-bstA* (SNW389) and once with D23580  $\Delta\Phi$  *tetR-P<sub>tetA</sub>-bstA* (SNW576), giving similar results.

## Microscopy- general

For all imaging experiments, bacteria were sub-cultured in liquid M9 Glu<sup>+</sup> media. All images were collected with a wide field Nikon Eclipse Ti-E inverted microscope equipped with an Okolab Cage Incubator warmed to 37°C with Cargille Type 37 immersion oil. A Nikon CFI Plan Apo DM Lambda 100X 1.45 NA Oil objective and a Nikon CFI Plan Apo DM Lambda 20X .75 NA objective were used with Perfect Focus System for maintenance of focus over time. Superfolder GFP, mCherry and SYTOX Orange Nucleic Acid Stain (ThermoFisher) were excited with a Lumencor Spectra X light engine with Chroma FITC (470/24) and mCherry (575/25) filter sets, respectively and collected with a Spectra Sedat Quad filter cube ET435/26M-25 ET515/30M-25 ET595/40M-25 ET705/72M-25 and a Spectra CFP/YFP/mCherry filter cube ET475/20M-25 ET540/21M-25 ET632/60M-25. Images were acquired with an Andor Zyla 4.2 sCMOS controlled with NIS Elements software. For time-lapse experiments, images were collected every 3 minutes (unless specified otherwise) *via* ND acquisition using an exposure time of 100 ms and 50% or 100% illumination power for fluorescence. Multiple stage positions (fields) were collected using the default engine Ti Z. Fields best representing the overall experimental trend with the least technical artefacts were chosen for publication. Gamma, brightness, and contrast were adjusted (identically for compared image sets) using FIJI (Schindelin et al., 2012). The FIJI plugins Stack Contrast (Capek et al., 2006) and StackReg (Thevenaz et al., 1998) were used for brightness matching and registering image stacks.

## Microscopy- agarose pads

Agarose pads were prepared with 2% agarose and M9 Glu<sup>+</sup> media, and mounted on MatTek dishes (No. 1.5 coverslip, 50 mm, 30 mm glass diameter, uncoated). Cells (D23580 $\Delta$ *tsp-gtrAC* (JH4287) or D23580 $\Delta$ *tsp-gtrAC bstA<sup>STOP</sup>* (SNW431) were grown to log phase (OD<sub>600</sub> ~ 0.4) in M9 Glu<sup>+</sup> at 37°C with shaking (220 RPM), and where required, diluted in fresh M9 Glu<sup>+</sup> to achieve the desired cell density on the agarose pad. For experiments where all cells were infected (figure 4A), phage P22  $\Delta$ c2 was added at an MOI of 5. Phage adsorption and initial infection was facilitated by incubation at 37°C with shaking for 10 minutes. Subsequently, infected cells were pelleted at 5000 x g and resuspended in ice-cold PBS to pause phage development. Two microliters of chilled, infected cells were spotted onto opposite sides of an agarose pad (two strains were imaged on the same pad) and inverted onto the MatTek imaging dish. Experimental MOIs were immediately confirmed by CFU and PFU /mL measurement of the cell and phage preparations. Phase-contrast images using the 100X objective were collected every 3 minutes for 3 hours.

Procedures for experiments involving a subset of infected cells (figure 4C) were identical, except cells infected with P22  $\Delta$ c2 *P-mcherry* were washed an additional 4 times in ice-cold PBS to reduce the concentration of un-adsorbed, free phage. In parallel, uninfected cells were washed once in ice-cold PBS. Infected cells were mixed at a ratio of 1:1000 with uninfected cells of the same genotype before being spotted onto the agarose pad. This ratio of uninfected to infected cells was optimised such that in randomly chosen microscopy fields (without prior knowledge of which cells in the field were infected) there was likely to be at least 1 infected cell. Infected cells were retrospectively identified during image analysis by their synchronised lysis within a 10-minute window at the beginning of the microscopy timelapse. For these experiments, phase-contrast and fluorescence images (mCherry) using the 20X objective were collected every 3 minutes for 3 hours.

## Microscopy- microfluidic infection

The CellASIC ONIX2 system from EMD Millipore with B04A plates was used for microfluidic imaging experiments (figure S5). Phages used in microfluidic infection experiments shown in 5B (P22 HT or 9NA) were stained with SYTOX Orange Nucleic Acid Stain according to the protocol previously described (Valen et al., 2012). Stained phages washed 4 times in 15 mL M9 Glu<sup>+</sup> media using Amicon Ultra-15 centrifugal filter units. After staining, the titer and viability of phages were immediately assessed by plaque assay, and once stained, phages were used for no longer than 2 weeks. For use in the

1061 microfluidic experiments, SYTOX Orange strained phages were normalised to a titer of approximately  
1062  $10^{10}$  PFU/mL. Cells (D23580 *bstA-sf::gfp*, SNW403) were grown to early exponential phase ( $OD_{600} \sim 0.1$ )  
1063 in M9 Glu<sup>+</sup> at 37°C with shaking (220 RPM) before being loaded into CellASIC B04A plates using the  
1064 pressure-driven method according to the manufacturer protocol for bacterial cells. The slanted chamber  
1065 of the plate immobilises the cells, but allows media to flow continuously. Firstly, cells were equilibrated  
1066 with constant M9 Glu<sup>+</sup> media flow for approximately 30 minutes. Secondly, stained phages suspended  
1067 in M9 Glu<sup>+</sup> media were flowed over the cells until the majority of cells were infected (typically 10-30  
1068 minutes). In the case of P22 HT phage (which exhibits inefficient adsorption to D23580 *bstA-sf::gfp* due  
1069 to the *gtr* locus of prophage BTP), phages were continuously flowed. Finally, M9 Glu<sup>+</sup> media was flowed  
1070 over the cells for the duration of the experiment. Microfluidic experiments typically lasted 5 hours, after  
1071 which time uninfected cells outgrew the chamber. Phase-contrast and fluorescence images were  
1072 collected every 1.5 minutes for the experiments in figure S5B.

1074 For the microfluidic imaging experiments shown in figure S5C, strain SVO251 (*S. Typhimurium* D23580  
1075  $\Delta\Phi$  STM1553::(*P*<sub>tetA</sub>-*bstA-sf::gfp-frt*)  $\Delta$ pSLT-BT  $\Delta$ pBT1 pAW61 (*P*<sub>BAD</sub>-*parB-mcherry*) was used. This strain  
1076 contains the *bstA-sf::gfp* fusion contrast under the control of the *P*<sub>tetA</sub> promoter. However, this strain lacks  
1077 the *tetR* gene, and therefore expression of *bstA-sf::gfp* is constitutive (not inducible). Additionally, this  
1078 strain is cured of two natural plasmids that contain native partitioning systems (pSLT-BT and pBT1),  
1079 and there for might interfere with the correct function of the ParB-*parS* system used for phage DNA  
1080 localization. The ParB-mCherry fusion protein is expressed from the pAW61 plasmid (Ap<sup>R</sup>) under the  
1081 control of the *P*<sub>BAD</sub> promoter (induced by L-arabinose). Strain SVO251 was grown in M9 Glu<sup>+</sup>  
1082 supplemented with 100  $\mu$ g/mL ampicillin to maintain the pAW61 plasmid and 0.2% L-arabinose to induce  
1083 expression of ParB-mCherry. The same supplemented media was used in the microfluidic chamber.  
1084 Cells were grown to  $\sim OD_{600}$  0.1 before loading into the CellASIC B04A plate as described above. After  
1085 15 minutes growth, phage P22  $\Delta$ *pid*::(*parS-aph*) [which contains one *parS* site along with a kanamycin  
1086 resistance locus, *aph*, in place of the non-essential *pid* locus (Cenens et al., 2013)] diluted to a  
1087 concentration of  $10^8$  PFU/mL (in M9 Glu<sup>+</sup> amp100 0.2% L-ara) was flowed into the chamber. Phase  
1088 contrast and red and green fluorescence images were collected every 2 minutes for 4 hours.

### 1090 BstA protein homolog analysis

1091 BstA protein homologs were identified using tblastn (database: non-redundant nucleotide collection)  
1092 and the HMMER webserver (Potter et al., 2018) (database: Reference Proteomes). The dataset of BstA  
1093 protein homologs was manually curated to reflect the diversity of taxonomic background harbouring  
1094 homologs. Evolutionary covariance analysis was done using DeepMetaPSICOV (Buchan and Jones,  
1095 2019) at the PSI-PRED server (Kandathil et al., 2019). To analyse the genetic context of BstA homologs,  
1096 the sequence region 20 Kb either side of the homolog (40 kb total) was extracted (BstA 40 kb  
1097 neighbourhoods). To produce homogenous and comparable annotations, each region was re-annotated  
1098 using Prokka 1.13 (Seemann, 2014). Additionally, the resulting annotated amino acid sequences were  
1099 queried against our custom BstA profile-hmm and the Pfam 31.0 database (El-Gebali et al., 2019) with  
1100 hmmercan (Eddy, 1998), and the highest scoring significant hit per ORF was considered for the results  
1101 shown in figure 2. All the code is available in [https://github.com/baymLab/2020\\_Owen-BstA](https://github.com/baymLab/2020_Owen-BstA).

1103 Pairwise identity of homologs in figure 2B to BstA<sup>BTP1</sup> was computed using the EMBOSS Needle  
1104 webserver (Needleman and Wunsch, 1970). BstA homologs were designated “putatively-prophage  
1105 associated” if annotated genes in the 40 kb neighbourhood contained any instance or the word “phage”  
1106 or “terminase”. For categorisation in figure 2C, homologs were classed as having “high confidence  
1107 association” if instances of gene annotations including the aforementioned key words occurred both  
1108 before, and after, the BstA gene within the 40 Kb neighbourhood (i.e., to account for the possibility that  
1109 a prophage-independent homolog could co-occur next to a prophage region by chance). Homologs  
1110 classed as having “low confidence association” had at least one instance of genes whose annotations  
1111 included “phage” or “terminase” either in the upstream or downstream 20Kb, but not both. Plasmid status  
1112 was determined from information in the sequence records. The HHpred webserver was used to annotate  
1113 the putative Kila-N domain (Zimmermann et al., 2018). All homolog neighbourhoods, homolog  
1114 alignments and sequences is available to download [https://github.com/baymLab/2020\\_Owen-BstA](https://github.com/baymLab/2020_Owen-BstA).



## QUANTIFICATION AND STATISTICAL ANALYSES

The phage replication, survival rate, efficiencies of transformation and of lysogeny were calculated as mentioned above. The numerical data were plotted and analyzed using GraphPad Prism 8.4.1. Unless stated otherwise in the figure legends, data are presented as the mean of biological triplicates  $\pm$  standard deviation. The unpaired *t*-test was used to compare the groups and statistical significance is indicated on the figures. P values are reported using the following criteria:  $< 0.0001 = ****$ ,  $0.0001$  to  $0.001 = ***$ ,  $0.001$  to  $0.01 = **$ ,  $0.01$  to  $0.05 = *$ ,  $\geq 0.05 = ns$ .

## KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
<b>Antibodies</b>		
Anti-Digoxigenin-AP, Fab fragments (Roche)	MilliporeSigma	Cat#000000011093274910
<b>Bacterial and Virus Strains</b>		
All the bacterial strains and bacteriophages are listed and described in table S2		
<b>Chemicals, Peptides, and Recombinant Proteins</b>		
L-(+)-arabinose	Melford	Cat#A51000-100.0
Betaine	MilliporeSigma	Cat#B2629
Bacto Agar (BD)	Appleton Woods	Cat#MN663
Bacto Casamino Acids Technical (BD)	Appleton Woods	Cat#223110
EGTA	MilliporeSigma	Cat#E3889
M9 Salts 5X	MilliporeSigma	Cat#M6030
Maltose	MilliporeSigma	Cat#M5885
<i>m</i> -toluic acid	MilliporeSigma	Cat#T36609
Tryptone (BD)	Appleton Woods	Cat#MN649
Yeast Extract (BD)	Appleton Woods	Cat#DM832
Sodium Chloride	MilliporeSigma	Cat#S3014
Ampicillin Sodium	Melford	Cat#A40040-25.0
Anhydrotetracycline hydrochloride (AHT)	MilliporeSigma	Cat#37919
Chloramphenicol	MilliporeSigma	Cat#C0378
Gentamicin sulfate	Melford	Cat#G38000-25.0
Kanamycin monosulfate	Melford	Cat#K22000-25.0
Mitomycin C	MilliporeSigma	Cat#M0503
Tetracycline hydrochloride	MilliporeSigma	Cat#T7660
EcoRI	ThermoFisher Scientific	Cat#ER0271
BamHI	ThermoFisher Scientific	Cat#ER0051
Dpnl	New England Biolabs	Cat#R0176S
Kpnl-HF	New England Biolabs	Cat#R3142S
Bsal-HF	New England Biolabs	Cat#R3535S
SmaI	ThermoFisher Scientific	Cat#ER0661
Sall	ThermoFisher Scientific	Cat#ER0641
XbaI	New England Biolabs	Cat#R0145S
T4 Polynucleotide Kinase	ThermoFisher Scientific	Cat#EK0031
T4 DNA ligase	ThermoFisher Scientific	Cat#EL0014
CutSmart buffer	New England Biolabs	Cat#B7204S
Tango Buffer 10X	ThermoFisher Scientific	Cat#BY5
T4 DNA ligase Reaction Buffer	New England Biolabs	Cat#B0202S
Phusion High Fidelity DNA polymerase	New England Biolabs	Cat#M0530S
MyTaq Red PCR mix 2X	Bioline	Cat#BIO-25043
Taq DNA polymerase	Bioline	Cat#BIO-21105
NEBuilder® HiFi DNA Assembly Cloning Kit	New England Biolabs	Cat#E5520S
DNase I	MilliporeSigma	Cat#DN25
RNase A	MilliporeSigma	Cat#R6513

Proteinase K	Bioline	Cat#BIO-37037
dNTP mix	Bioline	Cat#BIO-39025
dATP	Bioline	Cat#BIO-39036
dGTP	Bioline	Cat#BIO-39037
dCTP	Bioline	Cat#BIO-39038
dTTP	Bioline	Cat#BIO-39039
DIG-11-dUTP, alkali-stable (Roche)	MilliporeSigma	Cat#11093088910
Midori Green DNA/RNA staining	Nippon Genetics	Cat#MG06
Electroporation cuvettes	GeneFlow	Cat#E6-0060
Nylon membrane, positively charged (Roche)	MilliporeSigma	Cat#11417240001
Blocking Reagent (Roche)	MilliporeSigma	Cat#11096176001
DIG Easy Hyb™ Granules (Roche)	MilliporeSigma	Cat#11796895001
<b>Critical Commercial Assays</b>		
ISOLATE II Plasmid Mini Kit	Bioline	Cat#BIO-52057
ISOLATE II PCR and Gel Kit	Bioline	Cat#BIO-52060
Norgen Phage DNA Isolation Kit (46850)	GeneFlow	Cat#P4-0134
Quick-DNA™ Universal Kit	Zymo	Cat#D4069
Invitrogen Qubit dsDNA HS Assay Kit	ThermoFisher Scientific	Cat#Q32851
SYTOX Orange Nucleic Acid Stain – 5 mM in DMSO	ThermoFisher Scientific	Cat#S11368
Amicon Ultra-15 centrifugal filter units	MilliporeSigma	Cat#UFC910024
<b>Oligonucleotides</b>		
All the DNA oligonucleotides are listed in table S2		
<b>Recombinant DNA</b>		
All the plasmids are listed and described in table S2		
<b>Software and Algorithms</b>		
GraphPad Prism 8.4.1		
HMMER webserver	<a href="https://www.ebi.ac.uk/Tools/hmmer/">https://www.ebi.ac.uk/Tools/hmmer/</a>	
Prokka 1.13		
HHPred webserver	<a href="https://toolkit.tuebingen.mpg.de/tools/hhpred">https://toolkit.tuebingen.mpg.de/tools/hhpred</a>	
EMBOSS Needle webserver	<a href="https://www.ebi.ac.uk/Tools/psa/emboss_needle/">https://www.ebi.ac.uk/Tools/psa/emboss_needle/</a>	

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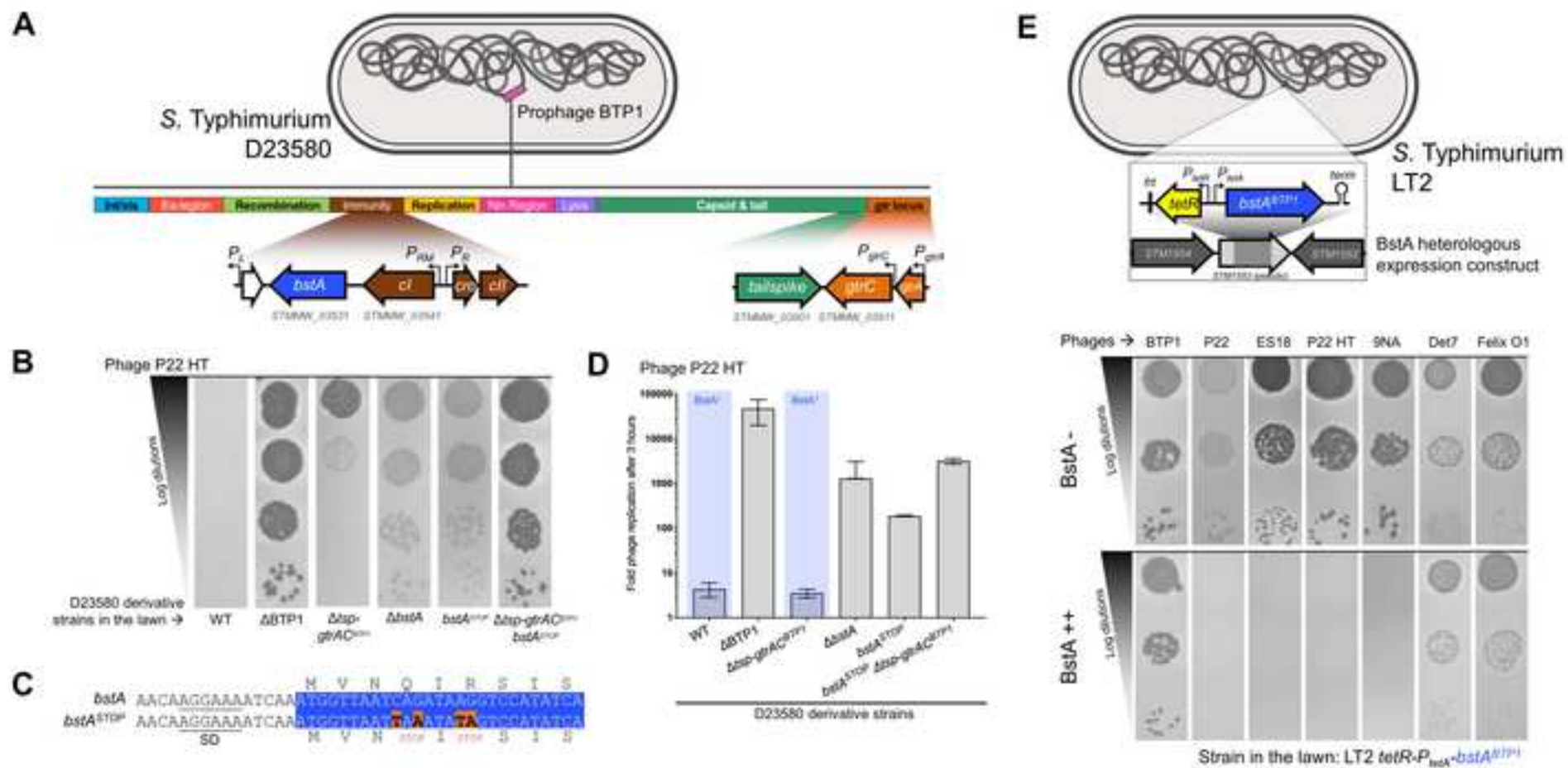
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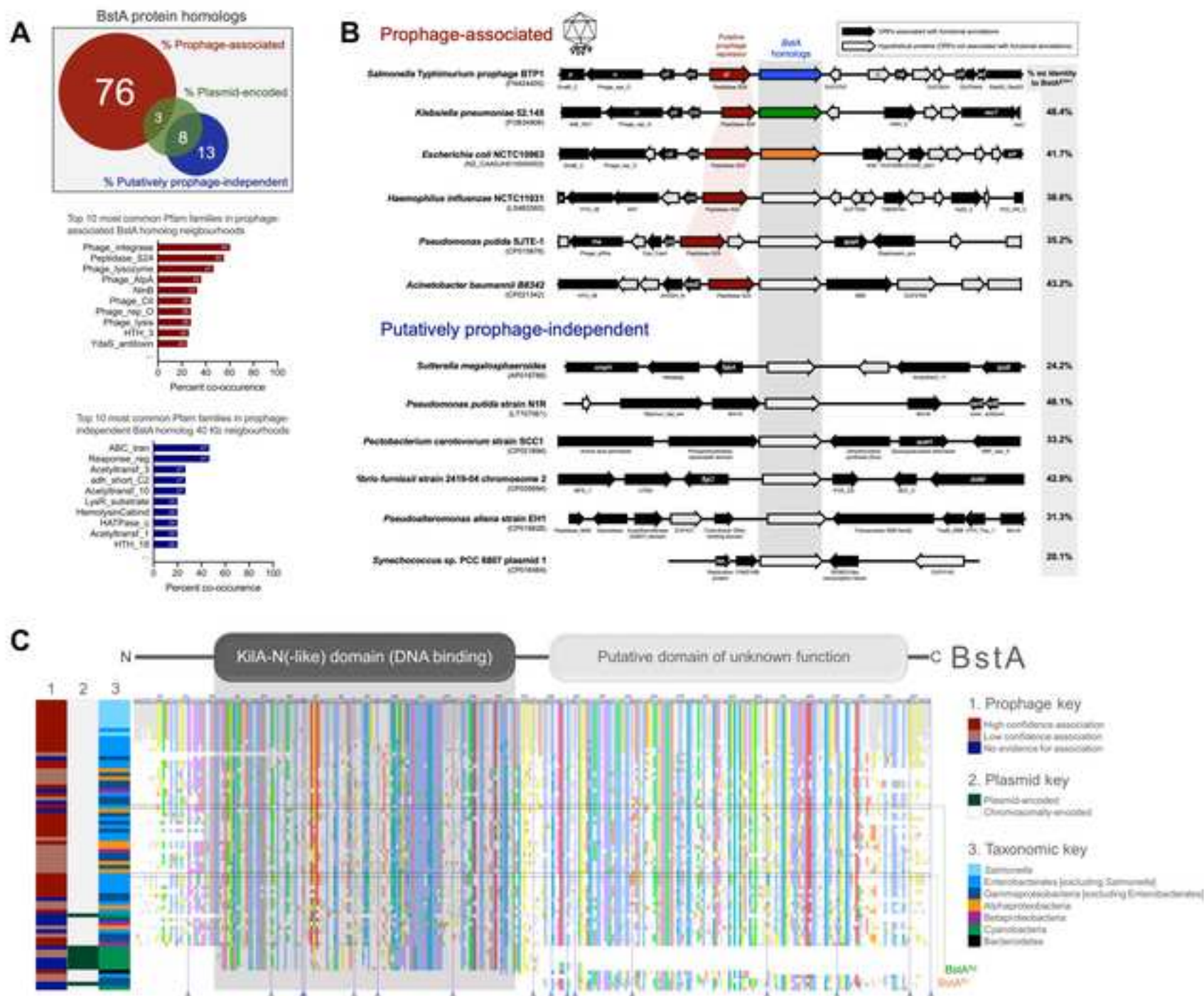
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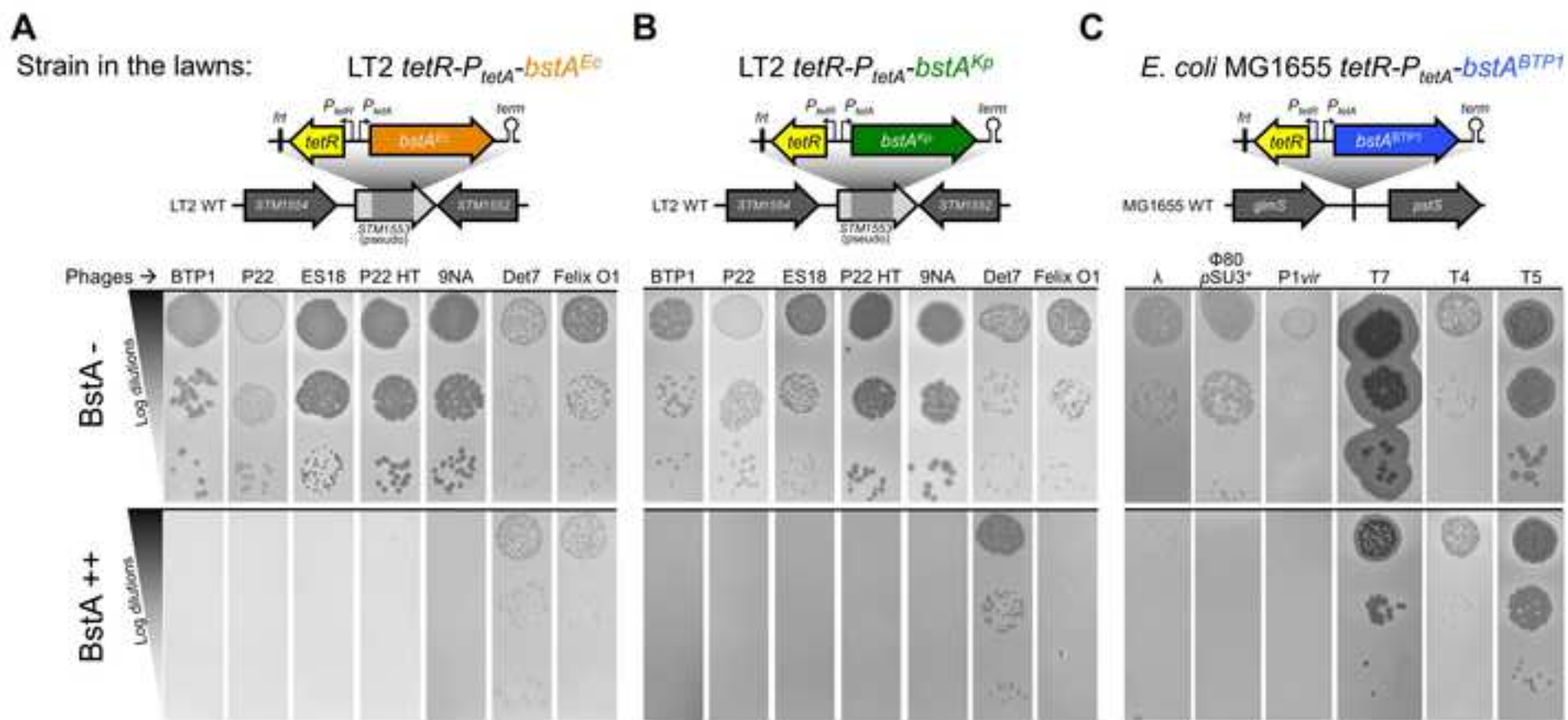
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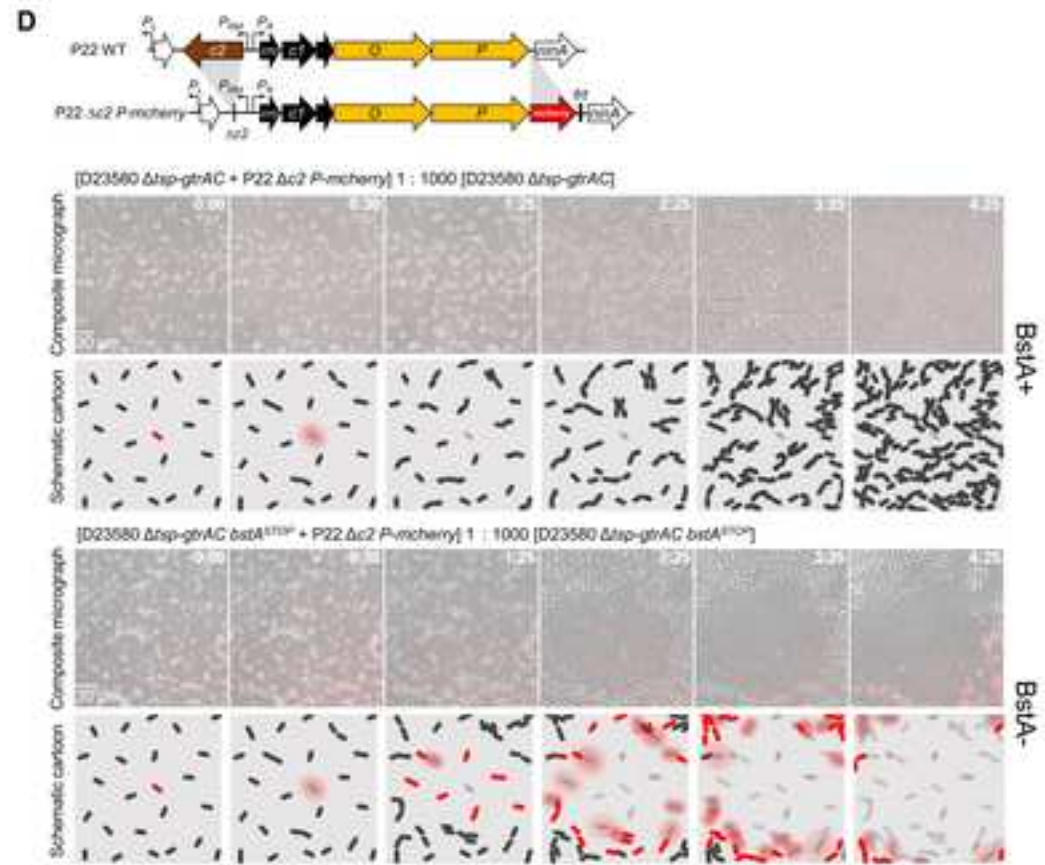
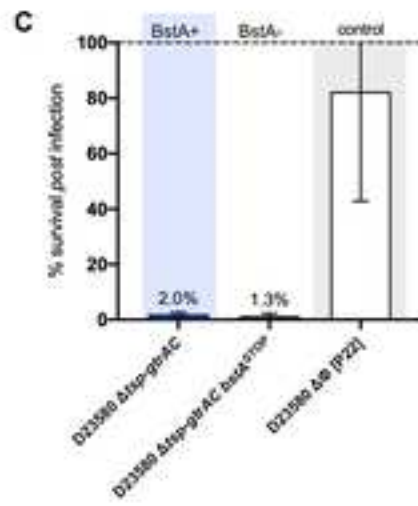
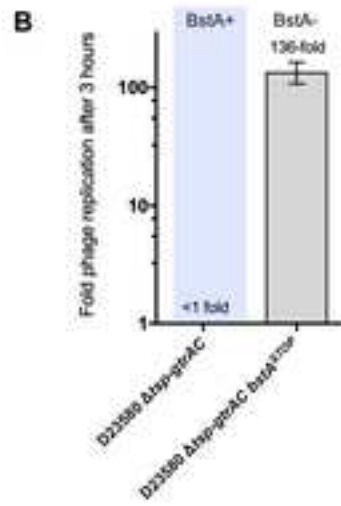
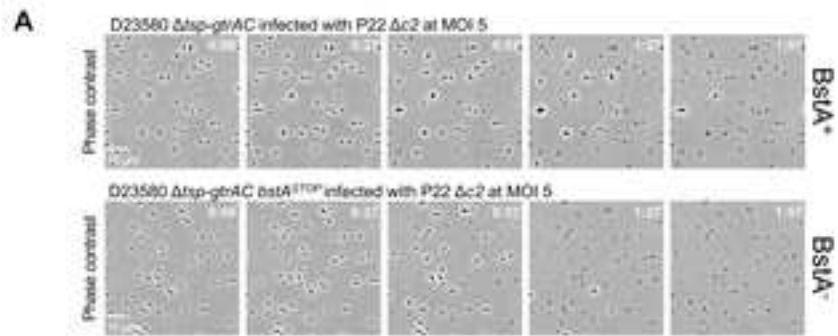
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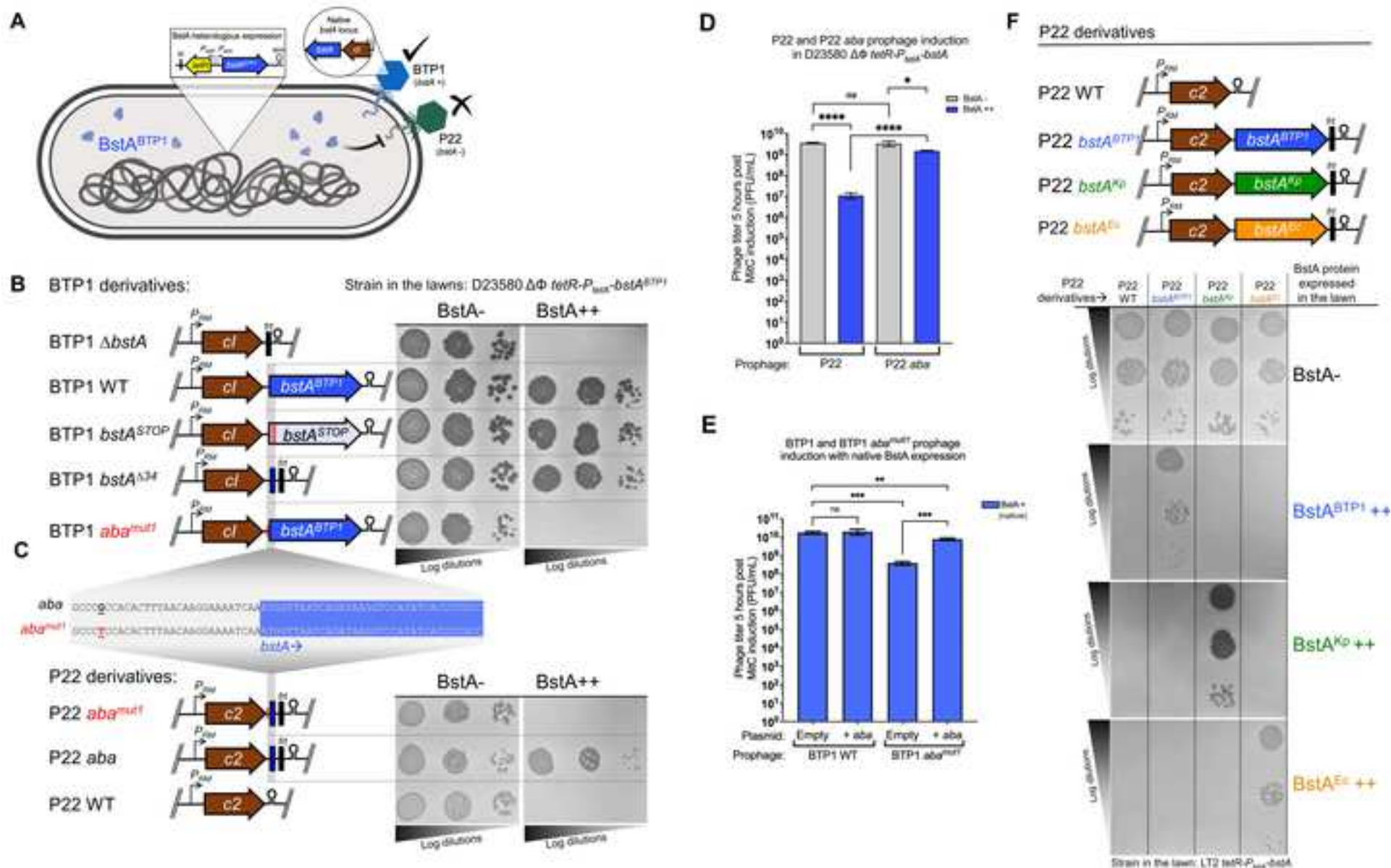


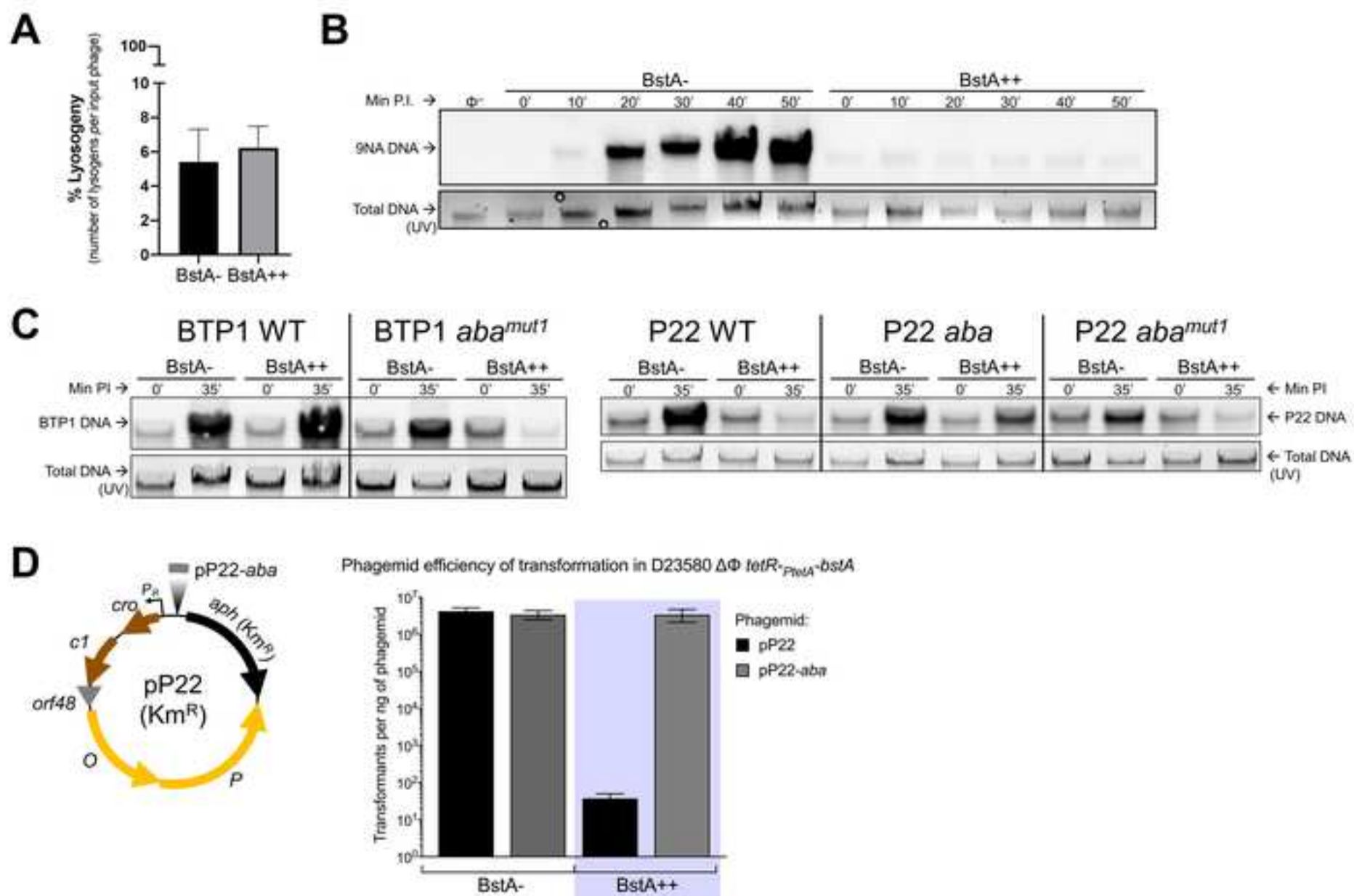








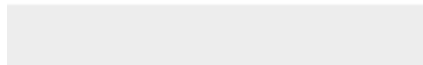






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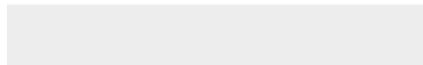
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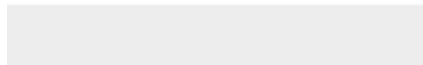
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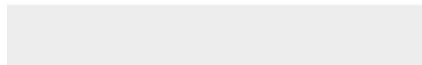




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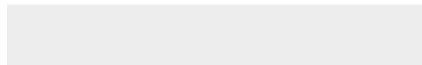




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