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2	Engineering a dynamic, controllable infectivity switch in bacteriophage T7
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27 Abstract

28 Transcriptional repressors play an important role in regulating phage genomes. Here, 29 we examined how synthetic regulation based on repressors can be to create a dynamic. 30 controllable infectivity switch in bacteriophage T7. We engineered T7 by replacing a 31 large region of the early phage genome with combinations of ligand-responsive 32 promoters and ribosome binding sites (RBS) designed to control the phage RNA 33 polymerase. Phages with the engineered switch showed virulence comparable to 34 wildtype when not repressed, indicating the phage can be engineered without a loss of 35 fitness. When repressed, the most effective switch used a TetR promoter and a weak 36 RBS, resulting in a two-fold increase in latent period (time to lyse host) and change in 37 phage titer. Further, phage activity could be tuned by varying inducer concentrations. 38 Our study provides a proof of concept for a simple circuit for user control over phage 39 infectivity.

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41 Keywords

42 engineered bacteriophage, phage switch, ligand-responsive phages, phage genome43 editing

44 Introduction

45 Bacteriophages (or 'phages') are obligate parasites which require a bacterial host to complete their life cycle¹. Once a phage infects its host, a choreographed cascade of 46 47 phage genes is expressed to regulate subsequent steps in the phage's life cycle. Gene regulation in phages is frequently controlled by transcription repressors². Transcription 48 repressors act as switches that determine lifestyle decisions (e.g.: lytic vs. lysogenic) by 49 50 silencing or activating different sets of genes. This is best exemplified in the transcription repressor-based genetic switch in phage lambda and other temperate 51 phages with similar mechanisms (P22³, 434⁴, ϕ C31⁵). Phage lambda persists in 52 53 lysogenic state when transcription repressor CI represses early-stage lambda 54 promoters, halting the transcriptional cascade of lytic genes. To activate lytic genes and 55 release the prophage, a different repressor Cro, counteracts CI through differential 56 binding at the same promoters. Transcription repressors also control activation and 57 inactivation of non-integrating, episomal phage genomes, called pseudolysogens, in a nutrient dependent manner⁶. Transcription repressors are valuable tools for phages 58 59 because their mechanism of gene regulation simply relies on steric obstruction of the RNA polymerase making it largely host-independent⁷. Further, a transcription repressor 60 61 with strong affinity for its promoter can exert tight regulation over multiple open reading 62 frames commonly found in long phage operons. While natural transcription regulation in 63 phages is well studied, the engineering rules of introducing synthetic regulation into 64 phages has not been developed.

65 Bacteria have long served as a popular chassis for exploring and prototyping 66 synthetic regulation. This choice was largely driven by the availability of tools for

bacterial genome engineering and the application goal of bacterial biomanufacturing⁸. In 67 68 recent years, a strong impetus has emerged for engineering phages, driven similarly by 69 the availability of new tools for phage genome engineering and their potential applications in medicine and biotechnology^{9–11}. Phage genomes can be now edited with 70 71 high precision from single base to kilobase resolution using yeast cloning, homologydirected repair, and enzymatic recombination^{12,13}. Phages engineered using these 72 73 approaches could be powerful tools for killing antibiotic resistant bacteria and precisely 74 manipulating microbiomes with applications in agriculture, livestock, medicine, and 75 environment. Engineered phages have many advantages over their natural counterparts 76 including superior efficacy, greater programmability, higher compositional stability, and easier scalability of production¹⁴. Despite this promise, even basic rules of engineering 77 78 new regulation into phages have not been developed. This is in stark contrast to the 79 wealth of research on engineering promoters, switches, circuits, and pathways in 80 bacteria. Rules for engineering bacteria are not directly transferrable to phages due to 81 substantial differences in genome compactness, compartmentalization of regulation and 82 kinetics of growth and replication. Thus, rules for engineering new regulation for phages 83 must be developed by systematic design-build-test-learn analysis.

With this goal in mind, we examined how bacterial transcription repressors could be used to engineer synthetic genetic regulation in phages. We sought to engineer a repressor-based infectivity switch that can dynamically control the activity of an obligate lytic phage in a ligand-dependent manner. Such a system would provide direct user control of phage activity which would otherwise undergo unregulated exponential amplification upon infecting a host¹⁵. Natural phages such as lambda are not ligand90 inducible but instead rely on stochastic differences in the concentration of CI vs. Cro to 91 determine lysogenic vs. lytic choice¹⁶. A dynamic controllable infectivity switch would be 92 a valuable tool for activating a phage at user-defined times to carry out microbiome 93 editing in a natural or synthetic communities. Furthermore, inducible phage would be an 94 effective form of biocontainment, as the phages only remain active while the ligand is 95 provided but otherwise remain inert.

96 Here, we engineered an obligate lytic T7 phage with a synthetic regulatory switch 97 to control its infection cycle in a ligand-dependent manner. To introduce synthetic 98 regulation, we removed a large tract of native regulatory sequence from the phage 99 genome and replaced this region with a short ligand-responsive bacterial promoter that 100 regulates the T7 RNA polymerase gene. We saw no appreciable loss of fitness in the 101 synthetic phage relative to wildtype T7 (T7_WT). We tested different ligand-responsive 102 bacterial promoters and ribosome-binding sites (RBS) representing a range of 103 expression levels to characterize how these variables affect phage activity. We 104 measured our ability to attenuate phage replication by measuring the phage latent 105 period and change in phage titer and found that the strongest attenuation of phage 106 replication, or the 'OFF' state, occurred with a Tet-regulated promoter and a very weak 107 RBS, which resulted in a 2-fold increase in latent period and an approximately 2-fold 108 decrease in a change in phage titer. Ligand induced activation to the 'ON' state restored 109 activity of synthetic phages to levels comparable to unregulated T7 WT phage. Our 110 study provides a basic proof-of-concept for recoding a phage genome with synthetic 111 regulation, paving the way of engineering more sophisticated circuitry to enable phages 112 to carry out complex, user-defined tasks.

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114 Results

115 **Refactored synthetic phages retain wildtype infectivity**

116 We sought to create a simple, dynamically controllable infectivity switch for 117 phages using ligand-responsive transcriptional repressors naturally utilized for gene 118 regulation in bacteria. To test our synthetic switch, we chose bacteriophage T7, a well 119 characterized, prototypical obligate lytic phage that infects *Escherichia coli*. Bacterial 120 systems are routinely engineered with inducible gene expression by placing ligandresponsive promoters upstream of gene of interest¹⁷. Bacterial genomes are generally 121 122 tolerant to promoter substitutions as most substitutions have marginal effects on bacterial fitness¹⁸. In contrast, phage genomes have evolved to maintain compact 123 genomes containing mostly essential genes with overlapping regulation¹⁹⁻²¹. This 124 125 makes it challenging to identify a suitable T7 gene whose promoter can be replaced by 126 ligand-responsive bacterial promoter.

127 The T7 phage genome is approximately 40 kb long and is roughly partitioned into 128 early, middle and late-stage genes denoting their roles and when they are expressed at different stages of T7 lifecycle²². Transcription of phage genes occurs in two stages 129 130 after infecting its host. First, the host E. coli RNA polymerase transcribes T7 RNA polymerase and other early phage genes^{23–26}. After early genes have been expressed, 131 132 the host RNA polymerase is inhibited and T7 RNA polymerase assumes responsibility for transcribing middle and late-stage genes^{27,28}. T7 RNA polymerase is thus a "lynchpin" 133 134 gene and we reasoned that a ligand-responsive promoter regulating this gene (qp1) 135 would be an optimal site to exert systemic control. The region upstream of *qp1* gene is

comprised of several parts (Fig. 1A). The beginning of the genome contains elements
used for replication, including terminal repeats, promoter A0 recognized by the host
RNA polymerase, and origin of replication ØOL²⁹. Downstream of these components lies
early promoters A1, A2 and A3, all of which are recognized by the host RNA
polymerase, followed by early phage genes *gp0.3*, *gp0.4*, *gp0.5*, *gp0.6*, and *gp0.7*.
These genes are involved in host suppression and inhibition of host defenses³⁰.

142 We assembled engineered T7 phage genomes in yeast without the genomic 143 segments covering early promoters A0, A1, A2, A3, origin of replication ØOL, and gene 144 products *qp0.3*, *qp0.4*, *qp0.5*, *qp0.6*, and *qp0.7*, thereby retaining the essential 145 replication elements but simplifying the early genome to create an effective regulatory 146 switch. This approximately 3 kb long genomic tract was replaced with 95-100 bp long 147 ligand-responsive promoter NaIC, PmeR, or TetR (T7_NaIC, T7_PmeR, and T7_TetR, 148 respectively) placed immediately upstream of gp1 (T7 RNA polymerase) (Fig. 1A). 149 Engineered phages were subsequently 'rebooted' by transforming their genome into an *E. coli* host³¹. Once an engineered phage infects a host expressing the repressor, the 150 151 repressor will bind to its cognate promoter and block the expression of *gp1*, attenuating 152 phage activity (Fig. 1B). In the presence of the inducer or in a host where cognate 153 repressor is absent, the repressor is unable to bind to the promoter, initiating 154 transcription of *qp1* and reactivating the phage (Fig. 1B).

Since phage amplification occurs rapidly within the host, we wanted to evaluate different repressor-promoter systems for their OFF-states. We chose to test repressorpromoter systems NaIC, PmeR, and TetR as these systems have been previously engineered with tight and inducible gene regulation^{32–34}. However, before comparing the 159 ability of different repressor-promoters to regulate the phage, we first sought to 160 determine if our engineered phages (T7_NalC, T7_PmeR and T7_TetR) were viable 161 after removing approximately 3 kb (~7%) of native genes and regulatory regions. The 162 engineered phages were viable despite the substantial genomic disruption. Efficiency of 163 plating (EOP) assays showed no significant difference (p-value > 0.05) in ability to 164 plaque between our engineered phages and T7_WT (Fig. 1C). Furthermore, the plaque 165 morphology of the engineered phages was indistinguishable from T7 WT in *E. coli* with 166 and without *qp1* complementation (Fig. 1D). Taken together, our engineered phages 167 were able to maintain viability and infectivity on susceptible bacterial host compared to T7 WT. 168

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170 Repressing *gp1* diminishes activity of synthetic phages

171 We applied T7_NalC, T7_PmeR, and T7_TetR on E. coli host expressing the 172 cognate repressor from a plasmid (Fig. 2A, E. coli_NalC, E. coli_PmeR and E. 173 coli_TetR, respectively) or control wildtype E. coli host without repressor (Fig. 2A, E. 174 coli_WT). For each engineered phage, we evaluated the estimated latent period (the 175 time required to complete one infection cycle) by comparing cell densities at two 176 different multiplicities of infection (MOI or the phage-to-bacteria ratio). The latent period 177 can be estimated from the inflection in the growth curves between two MOIs where the 178 difference is equal to the phage burst size (see Materials and Methods) (Fig. 2A). All 179 three synthetic phages showed a delayed latent period in host expressing the cognate 180 repressor compared to *E.coli* WT lacking the repressor (Fig. 2A and 2B). The TetR 181 repressor had the strongest attenuation of phage activity among the three repressors 182 based on estimated lysis time (Fig. 2A). In E.coli_WT, T7_PmeR and T7_TetR required 183 approximately 20 minutes to lyse the host, which was comparable to that of T7_WT (Fig. 184 S1). This is also consistent with similar EOPs for wildtype and engineered phages (Fig. 185 1B). T7 NalC required a longer time of 30 minutes to lyse the host, indicating that the 186 NalC promoter may be affecting phage fitness even in the absence of the repressor. 187 When engineered phages were applied on hosts expressing their corresponding 188 repressors, the estimated lysis time of hosts E. coli_PmeR, E. coli_NalC and 189 E.coli TetR was approximately 40, 30 and 40 minutes, respectively. As expected, the 190 repressor was able to delay the activity of engineered phages (Fig. 2B). We found that 191 the TetR system gave the longest delay (2-fold) in estimated lysis time after normalizing 192 for differences in basal activity of each engineered phage on *E.coli_*WT (Fig. 2C). We 193 also examined residual bacterial density (OD₆₀₀) as it is indicative of the fraction of host 194 cells that remain unlysed. After 6 hours of incubation with the engineered phages, 195 bacterial density was highest for E. coli TetR, followed by E.coli NalC and then 196 *E.coli*_PmeR (Fig. 2A). Therefore, synthetic switches engineered into the phages able 197 to reduce phage efficacy and delay host cell lysis. Since the TetR system gave the 198 longest delay, we chose *E.coli*_TetR and T7_TetR for further optimization.

199

200 Engineering ribosome binding sites further enhances phage control

Although T7_TetR performed better than T7_PmeR and T7_NalC, T7_TetR still showed relatively high activity in the OFF-state (Fig. 2C). We hypothesized this could be due to high basal expression of *gp1* from the TetR-regulated promoter. To address this, we investigated if modifications to the ribosome binding site (RBS) could further 205 attenuate phage activity in the OFF-state by reducing basal expression of gp1. We 206 engineered T7_TetR with three RBS variants of the TetR-regulated promoter. The 207 chosen RBS variants spanned a range of translational activities based on a large-scale experimental study of *E. coli* RBS variants^{32,35}. Relative to the strength of the original 208 209 RBS at 100% strength (T7 TetR RBS1), we engineered T7 TetR with RBS variants at 210 39% strength (T7_TetR_RBS2), 3% strength (T7_TetR_RBS3), and 1% strength 211 (T7_TetR_RBS4) (see Methods). On E.coli_WT without repressor, all T7_TetR RBS 212 variants required an estimated 15-20 minutes to lyse the host, comparable to T7 WT, 213 indicating that strength of RBS had no measurable impact on phage activity in the 214 absence of repression (Fig. 3A and 3B). In contrast, when the engineered phages were 215 applied on *E. coli*_TetR, there was substantial delay in time required to lyse the host 216 compared to *E. coli* WT (Fig. 3B). The time required to lyse *E. coli* TetR was inversely 217 related to the strength of the RBS. T7 TetR RBS4 (the weakest RBS) required 95 218 minutes to lyse E. coli TetR, the longest time and a 4.75-fold greater delay relative to E. 219 coli_WT (Fig. 3C). T7_TetR_RBS4 also retained a high residual bacterial density after 6 220 hours, suggesting lowering basal expression of *gp1* successfully attenuated T7 activity 221 (Fig. 3A). Weaker RBSs are thus able to provide enhance control of phage replication 222 by further restricting phage activity in the OFF-state.

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224 Repression significantly delays replication in engineered phages

To further characterize the effectiveness of our synthetic switch, we examined the number of progeny phages over one infection cycle using one-step growth curves for T7_WT and T7_TetR_RBS4 on *E. coli_*WT and *E.coli_*TetR (Fig 4A). By counting the number of phages produced at different times, we determined the actual latent period (the time required to complete one infection cycle), and compared the log change in phage titer over one infection cycle. In prior literature, the latent period for T7_WT is approximately 20 minutes with an expected 2-2.5 log increase in the total phage population after one infection cycle^{36,37}. Since the TetR promoter and RBS does not have an impact on T7 fitness, we expected similar values for T7_TetR_RBS4 on *E. coli_WT*.

235 Our one step assay confirmed that both T7 WT and T7 TetR RBS4 have a 20 236 minute latent period on E.coli WT and T7 WT has a 20 minute latent period on 237 *E.coli* TetR (Figure 4A). After one infection cycle, the average increase in total phages 238 for T7_WT on E. coli_WT and E. coli_TetR was approximately 2.5±0.3 and 2.1±0.2 log respectively, while T7_TetR_RBS4 saw a comparable increase in total phages of 239 240 2.7±0.5 log on *E. coli* WT. These results confirm that the TetR repression system had 241 no apparent effect on phage activity for T7 WT or for T7 TetR RBS4 in the absence of 242 repression. In contrast, T7_TetR_RBS4 infecting E. coli_TetR had greatly extended 243 latent period of 40 minutes with only a 1.4±0.4 log increase in total phages (Fig. 4A), 244 indicated a dramatic delay of phage activity with very gradual production of phage 245 progeny during each replication cycle. We next determined how the TetR_RBS4 246 repression system affects the ability of the phage to plaque on E. coli TetR compared 247 to E. coli WT using an EOP assay. T7 TetR RBS4 phages had an EOP of -0.2±0.1 248 while T7_WT phage had an EOP of 0.2±0.1. The plaque activity of T7_TetR_RBS4 and 249 T7_WT was slightly significantly different (p-value = 0.05) and the T7_TetR_RBS4 250 plaques were significantly smaller than T7_WT plaques after nineteen hours of incubation, indicative of much slower phage activity and consistent with our one-step results (Fig 4B). In summary, engineered T7_TetR_RBS4 phages have a significant delay in replication compared to wildtype when the synthetic repressor is present, and phage activity can be fully recovered if the synthetic repressor is not present or under induction.

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257 Phage infectivity can be further controlled by tuning inducer concentration

258 To assess if phage activity could be dynamically controlled using a small 259 molecule inducer, we compared the activities of T7 TetR RBS4 with and without the inducer anhydrotetracycline (aTC)³⁸. Comparison of spot plates showed that repressed 260 261 T7_TetR_RBS4 (Fig. 5A, left) regained activity comparable to T7_WT when maximally 262 induced at 1 µM aTC (Fig 4A, right, Fig. S2). A control experiment confirmed that aTC 263 had no impact on activities of T7 TetR RBS4 on E. coli WT host (Fig. S3). However, 264 host cells grown over a gradient of aTC concentrations experienced a minor fitness 265 deficit at higher aTC concentrations suggesting inducer toxicity may have partially contributed to cell death³⁹. To evaluate if the activity of T7_TetR_RBS4 could be tuned 266 267 by the inducer in a dose-dependent manner without cell toxicity, we performed a time 268 course experiment over a range of aTC concentrations from 0 to 1 µM (Fig. 5B). 269 T7_TetR_RBS4 showed switch-like behavior from no activity to high activity with 270 modest dose-dependent activity over a narrow range of aTC concentration from 15-23 271 nM. Above 23 nM, T7 TetR RBS4 was fully switched 'ON' and activity was comparable 272 to activity at maximal induction at 1 µM (Fig. 5B). Below 15 nM, the activity of 273 T7 TetR RBS4 was nearly the same as the activity with no inducer. The tunable concentration range (15-23 nM) showed large variations which arise due to high stochasticity in the expression of *gp1* across the host population at these concentrations. Since T7 RNA polymerase can be recycled for phage gene expression⁴⁰, even stochastic bursts of expression would eventually lead to bacterial lysis. Altogether these results show that a moderate aTC concentration can provide a tunable level of control over phage activity.

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282

281 Discussion

283 In this study, we engineered synthetic gene regulation into T7 phage to create a 284 phage-bacterial system with ligand-regulated infectivity. We deleted a large tract of the 285 genome from T7 phage and inserted a suite of ligand-regulated promoters to control the 286 expression of *gp1*, the T7 RNA polymerase. The deletion of phage early promoters and 287 genes and replacement with ligand-regulated promoter resulted in a viable phage with 288 infectivity comparable to T7_WT. Our most optimized engineered phage. 289 T7 TetR RBS4, had a 2-fold delay in estimated latent period and an approximately 2-290 fold decrease in change in phage titer after one replication cycle compared to T7 WT. 291 The activity of the engineered phage could be fully recovered in dose-dependent 292 manner by adding the inducer ligand.

Our engineered phage sets the stage for several improvements to further refine and control phage infectivity. Because the infectivity of our engineered phage was determined solely by control over T7 RNAP, even basal levels of expression would allow the phage to escape repression proceed with infection. Adding multiple levels of repression throughout the phage genome could substantially improve our system and decrease the activity the phage in the OFF state^{41,42}. Critical genes such as late structural genes would make natural targets for improving control over the phage replication. Our study and other previous reports suggest that the phage genome may be tolerant to such changes without affecting viability^{36,43}.

302 An engineered phage with a dynamic controllable infectivity switch can be used as a potent tool for bacterial community control^{44,45}. Utilizing different repression 303 304 systems across different bacterial hosts, phages can be redirected to different bacteria in the same community⁴⁶. Phages could also be continuously propagated in a 305 306 community using a 'feeder' host whose susceptibility could be turned ON or OFF as needed. By tuning inducer concentrations^{47,48}, phages could be engineered to repress 307 308 but not eliminate specific bacterial hosts in a complex community. Alternatively, our 309 engineered phages can act as pseudo-lysogenic phages, maintaining a low burden on 310 the targeted microbe until being triggered to eliminate the host. This could be useful for 311 applications like starter cultures, where timed bacterial lysis is a critical consideration for cheese maturation⁴⁹. Our study presents a simple genetic regulatory technique that can 312 313 be further engineered to create a more controllable phage-bacterial system to precisely manipulate microbial communities^{50–52}. 314

315

Figure descriptions

316 Figure 1. Engineered phages retain wildtype infectivity

317 (A) Schematic illustration of the T7 bacteriophage genome with early genes shown in 318 an expanded closeup. The organization of the phage genome is shown in the middle. 319 wildtype phage is shown top, and the engineered phage is shown bottom. The portion of 320 the phage genome shown boxed in black is engineered with a replacement synthetic 321 promoter and ribosome binding site (RBS). (B) On top (1), an illustration of the phage 322 "OFF" state, where repressor (green) prevents RNA Polymerase (RNAP, blue) binding 323 and blocks *qp1* transcription, stopping phages from killing host. On bottom (2), a 324 schematic illustration of the phage "ON" state. Infection without the repressor present 325 allows RNAP to transcribe gp1 and phages replicate and kill the bacterial host. (C) 326 Ability of engineered T7 phages to infect E. coli by Efficiency of Plating (EOP) using E. 327 coli complementing qp1 as a reference host. Compared to wildtype T7 (T7 WT), T7 328 engineered with NaIC, PmeR, and TetR (T7 NaIC, T7 PmeR, and T7 TetR, 329 respectively) show no significant difference (n.s.) in ability to plague. Data is 330 represented as mean ± SD of biological triplicates. (D) Plaque morphology for wildtype 331 and engineered phages shown by plague assay on wildtype E. coli and E. coli with qp1 332 complemented host after a 19-hour incubation. Plaques for engineered phages retain 333 wildtype plaque morphology.

334

Figure 2. Repression of *gp1* **reduces activity in engineered phages**

(A) Bacterial cell density over time measured by absorbance (OD₆₀₀) of wildtype host (*E. coli_*WT; dotted lines) and host expressing NalC, PmeR or TetR repressor (*E. coli_*NalC,

338 E. coli PmeR, E. coli TetR, respectively; solid lines) after addition of T7 NaIC (red), 339 T7_PmeR (green) or T7_TetR (blue) repressors or T7_WT (black). Phages were applied at time 0 at a MOI of 10⁻³ and 10⁻⁵. Phages with synthetic promoters have 340 341 delayed lysis on hosts expressing the cognate repressor but have a comparable 342 estimated latent period in hosts without repressor. All data represented as mean OD₆₀₀ 343 ± SD in technical triplicates. (B) The estimated latent period of engineered phages on 344 wildtype host compared to host expressing the cognate repressor, as determined by 345 comparing the inflection in the growth curves between two MOIs where the difference is 346 equal to the phage burst size (C) The relative estimated latent period of engineered 347 phage on hosts expressing the cognate repressor compared to wildtype host. See also 348 Figure S1.

349

Figure 3. Engineered ribosome binding sites further enhances control over phage activity

352 (A) Bacterial cell density over time measured by absorbance (OD_{600}) for wildtype (E. 353 coli_WT; dotted lines) and host expressing TetR repressor (*E. coli_*TetR; solid lines) 354 after application of engineered T7 with an RBS at 100% strength (T7_TetR_RBS1, dark 355 blue), 39% strength (T7_TetR_RBS2, light blue), 3% strength (T7_TetR_RBS3, yellow) 356 and 1% strength (T7 TetR RBS4, orange). Phages were added at time 0 at a MOI of 10⁻³ and 10⁻⁵. The delay in estimated latent period increases as the RBS strength 357 358 decreases. All data represented as mean $OD_{600} \pm SD$ in biological triplicates. (B) The 359 estimated latent period of engineered phages on E. coli_WT compared to hosts 360 expressing the TetR repressor. (C) The relative estimated latent period of engineered

phage on host expressing the TetR receptor compared to wildtype host. Single asterisk
(*) represents significant difference (*p*-value < 0.05). Non-significant difference not
shown. See also Figure S2.

364

Figure 4. Repression significantly delays replication in engineered phages

366 (A) One step growth assay showing phage titer (PFU/mI) of wildtype (T7_WT) and 367 engineered (T7 TetR RBS4) phage on wildtype (E. coli WT) and engineered (E. 368 coli TetR) host. Inset table shows the latent period (minutes) and an estimated log 369 change in phage titer produced from the first replication cycle. All data shown as mean 370 ± SD in biological triplicates. (B) Plaque assay of wildtype (T7_WT) and T7 with TetR 371 and a 1% strength ribosome binding site (T7_TetR_RBS4) phages on wildtype host (E. 372 coli WT) and host expressing TetR repressor (E. coli TetR) host after 19 hours of 373 incubation.

374

Figure 5. Phage infectivity can be tuned by changing inducer concentration

(A) Spot plate assay measuring the ability of wildtype T7 (T7_WT) and T7 with TetR and a 1% strength ribosome binding site (T7_TetR_RBS4) to infect *E. coli* expressing the TetR repressor without inducer (Repressed) or with 1 μ M anhydrotetracycline (aTC) inducer (Induced). Phages are spotted from 10⁻³ to 10⁻⁶ dilution and spots are imaged after 4 hours of incubation. Induction restores the phage ability to spot comparable to wildtype. (B) Bacterial cell density over time measured by absorbance (OD₆₀₀) for *E. coli* expressing TetR repressor. T7_TetR_RBS4 phage is added at timepoint 0 at a MOI 383 of 10^{-5} , and each culture contains aTC inducer ranging from a concentration of 0 to 384 1000 nM (color gradient). See also Figure S3.

385 Supporting Information

Figure S1. Expression of repressors has no effect on wildtype T7

Bacterial cell density over time measured by absorbance (OD_{600}) of wildtype T7 (T7_WT) on wildtype host (*E. coli*_WT; black) and host expressing NalC (*E. coli*_NalC, red), PmeR (*E. coli*_PmeR, green) or TetR (*E. coli*_TetR, blue) repressor. Phages were applied at time 0 at a MOI of 10⁻³ and 10⁻⁵. There is no delay of estimated latent period when any repressor is expressed compared to wildtype host. All data represented as mean $OD_{600} \pm SD$ in technical triplicates.

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386

395 Figure S2. Activity of engineered phages is rescued by aTC inducer

396 Bacterial cell density over time measured by absorbance (OD₆₀₀) for *E. coli* expressing 397 TetR repressor after application of no phages (T7_None, grey), wildtype phage (T7_WT, 398 black), engineered T7 with an RBS at 100% strength (T7 TetR RBS1, dark blue), 39% 399 strength (T7 TetR RBS2, light blue), 3% strength (T7 TetR RBS3, yellow) and 1% 400 strength (T7_TetR_RBS4, orange). Phages and 1 µM aTC was added at timepoint. 401 Addition of aTC rescues phage function and results in comparable time to lysis for 402 engineered phages and T7_WT. All data represented as mean $OD_{600} \pm SD$ in biological 403 triplicates.

404

405 Figure S3. Inducer has no effect on bacterial growth or phage activity

406 Bacterial cell density over time measured by absorbance (OD₆₀₀) for (A) E. coli 407 expressing TetR (E. coli_TetR) without phage (T7_None), (B) wildtype E. coli (E. 408 coli WT) and engineered phage with a 1% strength RBS (T7 TetR RBS4), and (C) E. 409 coli expressing TetR (E. coli TetR) and wildtype phage (T7 WT). Anhydrotetracycline 410 (aTC) is added at concentrations from 0-1000 nM (color gradient) at time zero. E. coli is 411 able to grow productively under all aTC concentrations and application of phages 412 results in a reduction of cell density at the same time for all aTC concentrations. All data 413 represented as mean $OD_{600} \pm SD$ in biological triplicates.

- 414
- 415 **Table S1: Primers and Plasmids used in this study**
- 416 **Table S2: Promoter and Ribosome Binding Site sequences**

417 Materials and Methods

418 Microbes and Culture Conditions

419 Escherichia coli (E. coli) 10G, a highly competent derivative of DH10β was obtained

420 from Lucigen (60107-1)⁵³, T7 bacteriophage was obtained from ATCC (ATCC[®] BAA-

421 1025-B2), and *Saccharomyces cerevisiae* BY4741⁵⁴ is a laboratory stock.

422

Bacterial cultures were grown in LB (Luria-Bertani) media (1% Tryptone, 0.5% Yeast Extract, 1% NaCl). For plating, LB agar contains 1.5% agar, while top agar contains 0.5% agar (Teknova). Spectinomycin (115 µg/ml final concentration, GoldBio[®]) was added to media for selections of pSC101_NalC, pSC101_PmeR and pSC101_TetR. All incubations of bacterial and bacteriophage cultures were performed at 37°C, with the liquid culture shaking at 200-250 rpm consistently, otherwise specified.

429

T7 bacteriophage was grown and propagated using *E. coli* 10G in LB media. Phage
stocks were tittered using plaque assay and stored in LB at 4°C.

432

S. cerevisiae BY4741 was grown in YPD (2% Peptone, 1% Yeast Extract, 2% Glucose)
media prior to transformation. Yeast transformants were selectively grown on SD-Leu
(0.17% Yeast Nitrogen Base, 0.5% Ammonium Sulfate, 0.162% Amino Acid-Leucine,
[Sigma Y1376], 2% Glucose). YPD and SD-Leu plates contain 2.4% and 2% agar
additionally, respectively. Yeast incubation was performed at 30°C, with liquid culture
shaking at 200-250 rpm.

440 Short-term storage of liquid culture and plates were performed at 4°C and long-term 441 storages of bacterial and yeast stock culture were performed at -80°C in screw-capped 442 cryotubes, with 25% glycerol added as a cryoprotectant.

443

Bacterial and phage transformants were recovered in SOC (2% Tryptone, 0.5% Yeast
Extract, 0.2% 5 M NaCl, 0.25% 1 M KCl, 1% 1 M MgCl₂, 1% 1 M MgSO₄, 2% Glucose)
liquid media.

447

448 General Cloning Procedure

PCR and cloning were adapted and performed using standard laboratory procedures¹². 449 450 Briefly, PCR amplification was performed using KAPA HiFi (Roche KK2101) for all amplifications with plasmid or phage templates. KAPA2G Robust PCR kits (Roche 451 452 KK5005) were used to perform colony PCR and multiplex PCR for a screening of Yeast Artificial Chromosomes (YACs). All primer oligos were obtained from IDT[™]. Golden 453 454 Gate assembly was performed using New England Biosciences (NEB) Golden Gate 455 Assembly Kit (Bsal-HFv2, E1601L). DNA purification was performed using EZNA Cycle Pure Kits (Omega Bio-tek D6492-01) using centrifugation protocol. YAC extraction was 456 457 performed using YeaStar Genomic DNA Extraction Kits (Zymo Research D2002). 458 Gibson Assembly mixture was made in the laboratory (final concentration 100 mM Tris-459 HCL pH 7.5, 20 mM MgCl₂, 0.2 mM dATP, 0.2 mM dCTP, 0.2 mM dGTP, 10 mM dTT, 5% PEG-8000, 1 mM NAD⁺, 4 U/ml T5 exonuclease, 4 U/µl Tag DNA Ligase, 25 U/mL 460 461 Phusion polymerase). PCR product visualization was performed using agarose gel

462 electrophoresis with appropriate agarose concentration and SYBR[®] Safe DNA Gel Stain
463 (Invitrogen).

464

465 PCR amplification using plasmid templates was performed with 0.1 ng DNA template. 466 Phage fragment amplification was performed using 1 µl phage crude lysis treated at 467 65°C for 10 minutes as a template. Deletions and insertions of the T7 genome were 468 performed using PCR primers skipping or adding desired sequences, respectively. All 469 plasmid-template PCR products were treated with DpnI (NEB) following standard 470 protocol. Briefly, purified PCR product was combined with 5 µl 10x CutSmart Buffer, 1 µl 471 Dpnl, and dH₂O to 50 µl reaction. Digestion was performed at 37°C for at least 2 hours, 472 followed by heat inactivation at 80°C, 20 minutes. PCR purification was performed 473 afterward. All PCR products were quantified using NanoDrop 2000 Spectrophotometer 474 (Thermo Scientific).

475

476 E. coli 10G competent cells were made by mixing 192 mL SOC with 8-ml overnight 477 culture and incubating at 21°C and shaking at 200 rpm until OD₆₀₀~0.4 was reached, determined by the Agilent Cary 60 UV-Vis Spectrometer using manufacturer 478 479 documentation. Cells were centrifuged at 4°C, 800 xg to1000 xg for 20 minutes, the 480 supernatant was discarded, and cells were resuspended in 50 ml of pre-cooled 10% 481 glycerol. Centrifugation and washing were repeated three times. Cells were 482 resuspended in a final volume of ~1 ml 10% glycerol and were aliquoted and stored at -483 80°C. Cells made by this protocol are competent for both plasmid and YAC 484 transformation using electroporation.

485

Electroporation was performed using 40 µl 10G competent cell for both plasmids and YACs using a Biorad MicroPulser (165-2100), EC2 setting with 2-mm cuvette, 2.5 kV, single pulse. All cuvettes and Eppendorf tubes were chilled prior to the electroporation. After electroporation, recovery was performed by adding 950 µl pre-warmed SOC and incubated at 37°C for 1 hour and plated on relevant selective media.

491

492 Detailed protocols for cloning are available on request. All primers and plasmids used in 493 this study are listed in supporting document (Table S1).

494

495 **Phage Engineering**

496 A large segment (2961 bp) in the left end region of the wild type (WT) T7 phage 497 genome, including promoters A0, A1, A2, A3 and gp0.3-0.7 genes, was deleted. Two 498 identical operator (repressor binding) sites were inserted, one upstream of -35 and the 499 other in between the -35 and -10 consensus sequences replacing the spacer sequence 500 of promoter A1 driving the expression of *gp1* gene. The three operator sites are NalC1, 501 PmeR2, and TetO correspond to repressors, NalC, PmeR and TetR, respectively, in 502 bacterial system. The Bujard RBS (Ec-TTL-R111) in TetO engineered phage was 503 replaced by BBa_J61106 (Ec-TTL-R065), BBa_J61133 (Ec-TTL-R003), or DeadRBS 504 (Ec-TTL-R001) with 39%, 3%, and 1% translational strength relative to the Bujard RBS as 100% strength³⁵, respectively. The sequences of all consensus sequences, operator 505 506 sites, RBS, and repressors used are included in supporting document (Table S2).

508 Engineered phage genomes were assembled using yeast assembly^{4,5}, which requires 509 yeast transformation of relevant DNA segments. A prs315 yeast centromere plasmid 510 was split into three segments by PCR, separating the centromere and leucine selection 511 marker, which has been shown to improve assembly efficiency by limiting 512 recircularization events⁶. T7 genomic segments were made by PCR using WT T7 as a 513 template.

514

515 Relevant DNA fragments were mixed (0.1 pmol/fragment) and transformed into S. 516 *cerevisiae* BY4741 using the high efficiency yeast transformation protocol⁷. Successfully 517 assembled YACs were selected using SD-Leu media. After 2-3 days of incubation at 518 30°C, colonies were picked and directly assayed by multiplex colony PCR to screen for 519 colonies with correctly assembled YACs. Multiplex PCR was an effective way of 520 distinguishing correctly assembled YACs by interrogating junctions in the YACs. 521 Correctly assembled YACs were purified and transformed into E. coli 10G cells, and 522 after 1-hour recovery, 400 µl was inoculated in 4.6 ml LB. This culture was incubated 523 until a complete lysis, engineered phages were purified, sequence-confirmed, and 524 stored at 4°C in LB.

525

526 Plasmid Descriptions

527

528 pSC101_NalC, pSC101_PmeR and pSC101_TetR contain a pSC101 backbone, 529 spectinomycin resistance cassette, and repressors NalC, PmeR and TetR under 530 constitutive promoter apFAB61, respectively. 531

532 pHT7_gp1 contains a pBR backbone, kanamycin resistance cassette, mCherry 533 fluorescence marker, and the T7 RNAP gene, *gp1*. Both mCherry marker and *gp1* are 534 under constitutive expression.

535

536 Bacterial Methods and Phage Titer Quantification

Bacterial concentration was determined using a drop plate with 10-fold serial dilution of bacterial culture. Ten-microliters of bacterial dilution were dropped on LB plate in triplicate and counted after overnight incubation as colony-forming unit per milliliter (CFU/ml). Optical density (OD) measurement for all microplate-reader experiments were performed using microplate reader (Synergy HTX) at 600 nm wavelength, except the preparation of electrocompetent cell and the determination of log phase of bacterial culture that were performed using an Agilent Cary 60 UV-Vis Spectrometer.

544

545 Bacterial culture used for bacteria-phage interaction experiment and phage 546 quantification was obtained at log phase when the initial phage was added, unless 547 otherwise specified. Briefly, overnight bacterial culture was diluted 1:20 in liquid LB 548 media. Bacterial culture was collected when the OD₆₀₀ reaches 0.4-0.6 as determined 549 using an Agilent Cary 60 UV-Vis Spectrometer.

550

Phage stock was produced by a complete infection of phage in *E. coli* 10G. Bacterial
lysate was centrifuged at 4,400 xg for 5 minutes and supernatant was filtered through a
0.22 µm filter (Celltreat 151205-051). Phage titer was determined by plaque assay with

554 a 10-fold serial dilution of phage lysate. Typically, 10-20 µl of phage dilution is mixed 555 thoroughly with 250 µl log-phase bacterial culture and 3.5-ml 0.5% top agar and plated on 37°C pre-warmed LB plate. Plague number was counted after overnight incubation 556 557 as plaque-forming unit per milliliter (PFU/ml). Spot assay was considered as a 558 preliminary quantification of phage titer, which was performed occasionally prior to 559 whole-plate plaque assay. Log-phase bacterial culture was mixed with 3.5-ml top agar 560 and plated on pre-warmed LB plate. One-and-a-half microliters of phage dilution from 10⁰ to 10⁻⁸ were spotted in triplicate on set top agar. The number of PFU was 561 562 approximated determined at 4-6 hours of incubation and the appropriate dilution was 563 chosen to perform whole-plate plaque assay.

564

565 Multiplicity of Infection (MOI) was determined by dividing phage titer by bacterial 566 concentration. MOI used in this study was approximately 10^{-3} and 10^{-5} , except for one-567 step growth curve which was performed with MOI ~ 0.1.

568

Efficiency of Plating (EOP) was calculated using wildtype *E. coli* 10G as a reference host. After performing a plaque assay using desired bacterial hosts and phages, EOP was determined by dividing the experimental phage titer to the phage titer on the reference bacterial host. Typically, a 10-fold serial dilution was performed with phage stock in triplicate and 10 µl appropriate phage dilutions were applied for plaque assay as described in desired bacterial host.

575

576 Infection time course curves

577 Preliminary change in phage activity of engineered phages was determined by 578 evaluating bacterial growth after addition of different phages. Seven microliters of 579 overnight bacterial culture were inoculated in 133 µl of relevant media as a 1:20 dilution 580 with corresponding antibiotics if necessary or 1 µM aTC to the total volume of 140 µl in a 96-well plate. Ten microliters of phages were added in triplicate at an MOI of 10⁻³ and 581 582 10^{-5} . The culture was continuously incubated in microplate reader (Synergy HTX) at 583 37°C with OD₆₀₀ measured every 10 minutes for 6 hours. The bacterial growth curve 584 was constructed by averaging OD_{600} .

585

586 Phage latent period estimation with infection time courses

With the assumption that T7 phage produces an average of 100 progeny phage at 37°C when infecting $10G^{37}$, time to lysis was estimated by comparing the inflection point of bacterial growth (or the point at which the culture begins to lyse) for two cultures after addition of two phage titers spanning 2 orders of magnitude. An MOI of 10^{-3} and 10^{-5} were chosen for this estimation. The estimated latent period was calculated as the difference between the last time point immediately before the inflection point in OD_{600} between the two cultures.

594

595 Phage Growth Quantification: Latent Period and Phage Titer change estimation

596 One-step phage growth assays were performed to construct one-step growth curves 597 using an adaptation of a standard protocol^{55,56}. Briefly, Seven-hundred-fifty microliters of 598 an overnight bacterial were added into 15-ml LB media with antibiotics if necessary. The 599 culture was incubated shaking at 250 rpm, 37°C in a 50-ml flask. Bacterial culture was 600 collected when the OD₆₀₀ ~ 0.25 was reached in 15-ml tube and concentrated into 1.5 601 ml by centrifugation at 4,400 xg for 5 minutes. Bacterial culture was transferred into a 2-602 ml Eppendorf tube, phage was added into the culture at MOI ~ 0.1. Phage-bacteria 603 culture was incubated for absorption without shaking at 37°C for 5 minutes. Culture was 604 washed four times by centrifugation at 10,000 xg for 30 seconds, discarding the 605 supernatant, and resuspending with 1-ml LB media, with antibiotic if necessary. In 606 between washes, culture was resuspended by vortexing. After the final wash, 1-ml 607 media was added and transferred into 14-ml pre-warmed media in a 50-ml flask. Eight-608 hundred microliters of culture was collected immediately and centrifuged at 10,000 xg 609 for 30 seconds. Seven-hundred-microliter supernatant was filtered through 0.22 µm 610 (Celltreat 151205-051) into a new Eppendorf tube. Fifteen-milliliter culture was 611 incubated shaking at 250 rpm at 37°C. The culture sample was collected every 5 or 10 612 minutes, depending on the type of phage culture for 60 or 120 minutes. Phage titer was 613 quantified by plaque assay using wildtype E. coli 10G. Plated culture was incubated at 614 37°C for 12 hours and the numbers of plaques were counted and calculated. The 615 dilution of the phage lysate was made if necessary. The growth curve of each bacteria-616 phage combination was constructed in biological triplicates.

617

The latent period of each phage-bacteria combination was quantified by the first phage replication cycle⁵⁶ using one-step phage growth curve. The mid-timepoint at exponential-phase of replication cycle in between plateau phage titer was considered as a latent period.

The change in total phage titer was estimated using the phage growth curve by calculating the log difference of number of phage progenies at the plateau after the first replication cycle and at the initial plateau where phage titer added into the experimental culture.

627

628 Inducer dependency assay

629 The activity of phage in response to a range of aTC inducer concentrations was 630 determined by quantifying bacterial growth using OD₆₀₀. Briefly, 7 µl overnight bacterial 631 culture was inoculated in 133 µl of relevant medium as 1:20 dilution to the total volume 632 of 140 µl in a 96-well plate. Media used was prepared with a range of aTC concentration 633 from 0 to 1 µM. Culture was incubated in a microplate reader (Synergy HTX), and OD 634 was measured at 600 nm wavelength. When bacterial $OD_{600} \sim 0.25$ was reached, 10 µl of desired phage or LB as control was added into the culture for MOI ~ 10⁻⁵. Culture was 635 636 continuously incubated for 7 hours using vertical shaking mode, and OD₆₀₀ was 637 measured every 10 minutes. The experiment was performed in biological triplicates.

638

639 Statistical Analysis

All data analyses were performed in Microsoft Excel 2020 and R v4.0.4⁵⁷. Bonferroni pairwise *t*-test^{58,59} was used to detect the difference amongst conditions in all experiments. Statistical analyses were considered significant at p < 0.05.

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- 667 performed the experiment, curated the data, performed the analysis, wrote the original
- manuscript, and designed the figures. S.R., C.C., C.L., P.H., and M.F. reviewed and
- 669 edited the manuscript. All authors discussed the results and provided critical feedback

on the manuscript.

671

647

- 672 **Notes**
- 673
- 674 S.R. is on the scientific advisory board of MAP/PATH LLC. All other authors declare no
- 675 competing financial interest.

676 Acknowledgements

677 This work was supported by National Institute for Allergy and Infectious Disease grant

678 1R21AI156785-01 (to S.R). C.C was supported by a graduate training scholarship from

679 the Anandamahidol Foundation (Thailand).

680

- 681 References
- 682

(1) Azam, A. H., and Tanji, Y. Bacteriophage-host arm race: an update on the

- 684 mechanism of phage resistance in bacteria and revenge of the phage with the 685 perspective for phage therapy.
- 686 (2) Yang, H. H., Ma, Y., Wang, Y., Yang, H. H., Shen, W., and Chen, X. (2014, July 23)

Transcription regulation mechanisms of bacteriophages: Recent advances and future
 prospects. *Bioeng. Bugs.* Landes Bioscience.

- (3) Prell, H. H. (1975) Regulation of gene expression in Salmonella phage P22 II.
- 690 Regulation of expression of late functions. *MGG Mol. Gen. Genet.* 136, 351–360.
- 691 (4) Bushman, F. D., and Ptashne, M. (1986) Activation of transcription by the
- 692 bacteriophage 434 repressor (in vitro transcription/RNA polymerase/ethylation
- 693 interference/protein-protein interaction). *Proc. Nati. Acad. Sci. USA*.
- 694 (5) Kuhstoss, S., and Rao, R. N. (1991) Analysis of the integration function of the 695 streptomycete bacteriophage φ C31. *J. Mol. Biol.* 222, 897–908.
- 696 (6) Court, D. L., Oppenheim, A. B., and Adhya, S. L. (2007, January 15) A new look at
- bacteriophage λ genetic networks. *J. Bacteriol.* American Society for Microbiology Journals.
- 699 (7) Rojo, F. (1999) Repression of Transcription Initiation in Bacteria. *J. Bacteriol.* 181, 700 2987.
- 701 (8) Guzmán-Trampe, S., Ceapa, C. D., Manzo-Ruiz, M., and Sánchez, S. (2017, June
- 15) Synthetic biology era: Improving antibiotic's world. *Biochem. Pharmacol.* ElsevierInc.
- (9) Lemire, S., Yehl, K. M., and Lu, T. K. Phage-Based Applications in Synthetic Biology
 5, 453–476.
- (10) Brown, R., Lengeling, A., and Wang, B. (2017) Phage engineering: how advances
- in molecular biology and synthetic biology are being utilized to enhance the therapeutic
 potential of bacteriophages. *Quant. Biol.* 5, 42–54.
- (11) Huss, P., and Raman, S. (2020) Engineered bacteriophages as programmable
 biocontrol agents. *Curr. Opin. Biotechnol.*, pp 116–121. Elsevier Ltd.
- (12) Huss, P., Meger, A., Leander, M., Nishikawa, K., and Raman, S. (2021) Mapping
- the functional landscape of the receptor binding domain of t7 bacteriophage by deep
- 713 mutational scanning. *Elife 10*.
- (13) Kilcher, S., and Loessner, M. J. (2019) Engineering Bacteriophages as Versatile
- 715 Biologics. *Trends Microbiol.* 27, 355–367.

(14) Pires, D. P., Cleto, S., Sillankorva, S., Azeredo, J., and Lu, T. K. (2016) Genetically

717 Engineered Phages: a Review of Advances over the Last Decade. *Microbiol. Mol. Biol.*

- 718 *Rev. 80*, 523–543.
- (15) Kulczyk, A. W., and Richardson, C. C. (2016) The Replication System of
- 720 Bacteriophage T7, in *Enzymes*, pp 89–136. Academic Press.
- (16) Casjens, S. R., and Hendrix, R. W. (2015, May 1) Bacteriophage lambda: Early
- pioneer and still relevant. *Virology*. Academic Press Inc.
- 723 (17) Chen, Y., Ho, J. M. L., Shis, D. L., Gupta, C., Long, J., Wagner, D. S., Ott, W.,
- Josić, K., and Bennett, M. R. (2018) Tuning the dynamic range of bacterial promoters
- regulated by ligand-inducible transcription factors. *Nat. Commun.* 2017 91 9, 1–8.
- (18) Gruber, T. M., and Gross, C. A. (2003) Multiple Sigma Subunits And The
- 727 Partitioning Of Bacterial Transcription Space. Annu. Rev. Microbiol 57, 441–66.
- (19) Pachl, C. A., and Young, E. T. (1976) Detection of polycistronic and overlapping
- bacteriophage T7 late transcripts by in vitro translation. *Proc. Natl. Acad. Sci. U. S. A.*
- 730 **73**, **312**.
- (20) Teif, V. B. (2010) Predicting Gene-Regulation Functions: Lessons from Temperate
 Bacteriophages. *Biophys. J.* 98, 1247.
- 733 (21) Jack, B. R., Boutz, D. R., Paff, M. L., Smith, B. L., and Wilke, C. O. (2019)
- Transcript degradation and codon usage regulate gene expression in a lytic phage.
- 735 *Virus Evol.* 5.
- (22) Dunn, J. J., and Studier, F. W. Complete Nucleotide Sequence of Bacteriophage
 T7 DNA and the Locations of T7 Genetic Elements.
- (23) Kemp, P., Gupta, M., and Molineux, I. J. (2004) Bacteriophage T7 DNA ejection
- into cells is initiated by an enzyme-like mechanism. *Mol. Microbiol.* 53, 1251–1265.
- (24) P, K., LR, G., and IJ, M. (2005) Changes in bacteriophage T7 virion structure at the
 initiation of infection. *Virology 340*, 307–317.
- 742 (25) García, L. R., and Molineux, I. J. (1995) Rate of translocation of bacteriophage T7
- 743 DNA across the membranes of Escherichia coli. J. Bacteriol. 177, 4066.
- 744 (26) Savalia, D., Robins, W., Nechaev, S., Molineux, I., and Severinov, K. (2010) The
- Role of the T7 Gp2 Inhibitor of Host RNA Polymerase in Phage Development. *J. Mol. Biol.* 402, 118–126.
- (27) McAllister, W. T., and Wu, H. L. (1978) Regulation of transcription of the late genes
 of bacteriophage T7. *Proc. Natl. Acad. Sci. U. S. A.* 75, 804.
- (28) FW, S., and JJ, D. (1983) Organization and expression of bacteriophage T7 DNA.
- 750 Cold Spring Harb. Symp. Quant. Biol. 47 Pt 2, 999–1007.
- (29) J;, K. T., and A.;, T. C. An Intermediate in the Replication of Bacteriophage T7 DNA
 Molecules.
- (30) McAllister, W. T., and Barrett, C. L. (1977) Roles of the Early Genes of
- 754 Bacteriophage T7 in Shutoff of Host Macromolecular Synthesis. J. Virol. 23, 543.
- (31) Ando, H., Lemire, S., Pires, D. P., and Lu, T. K. (2015) Engineering Modular Viral
- 756 Scaffolds for Targeted Bacterial Population Editing. *Cell Syst.* 1, 187.
- 757 (32) Liu, X., Gupta, S. T. P. P., Bhimsaria, D., Reed, J. L., Jos', J., Rodríguez-Martínez,
- J. A., Ansari, A. Z., Raman, S., Jos´, J., Rodríguez-Martínez, J. A., Ansari, A. Z., and
- Raman, S. (2019) De novo design of programmable inducible promoters 47, 10452–10463.
- 761 (33) Ramos, J. L., Martínez-Bueno, M., Molina-Henares, A. J., Terán, W., Watanabe, K.,

- Zhang, X., Gallegos, M. T., Brennan, R., and Tobes, R. (2005) The TetR Family of
 Transcriptional Repressors. *Microbiol. Mol. Biol. Rev.* 69, 326–356.
- (34) Lutz, R., and Bujard, H. (1997) Independent and tight regulation of transcriptional
- vinits in Escherichia coli via the LacR/O, the TetR/O and AraC/I 1-I 2 regulatory
- 766 elements. Oxford University Press.
- 767 (35) Kosuri, S., Goodman, D. B., Cambray, G., Mutalik, V. K., Gao, Y., Arkin, A. P.,
- Endy, D., and Church, G. M. (2013) Composability of regulatory sequences controlling
- transcription and translation in Escherichia coli. *Proc. Natl. Acad. Sci. U. S. A. 110*,
 14024–14029.
- (36) Nguyen, H. M., and Kang, C. Lysis Delay and Burst Shrinkage of Coliphage T7 by
 Deletion of Terminator Tφ Reversed by Deletion of Early Genes 88, 2107–2115.
- (37) You, L., Suthers, P. F., and Yin, J. (2002) Effects of Escherichia coli Physiology on
- Growth of Phage T7 In Vivo and In Silico. J. Bacteriol. 184, 1888–1894.
- (38) Gossen, M., and Bujard, H. (1993) Anhydrotetracycline, a novel effector for
- tetracycline controlled gene expression systems in eukaryotic cells. Nucleic Acids Res.
- (39) Halling-Sørensen, B., Sengeløv, G., and Tjørnelund, J. (2002) Toxicity of
- 778 Tetracyclines and Tetracycline Degradation Products to Environmentally Relevant 770 Bacteria Including Selected Tetracycline Posistant Bacteria
- 779 Bacteria, Including Selected Tetracycline-Resistant Bacteria.
- 780 (40) R, F., C, R., and G, D. (2004) Transcription reinitiation properties of bacteriophage
- T7 RNA polymerase. *Biochem. Biophys. Res. Commun.* 315, 376–380.
- 782 (41) Ponta, H., Rahmsdorf, H. J., Pai, S. H., Hirsch-Kauffmann, M., Herrlich, P., and
- Schweiger, M. (1974) Control of gene expression in bacteriophage T7: Transcriptional
 controls. *MGG Mol. Gen. Genet.* 134, 281–287.
- (42) Endy, D., Kong, D., and Yin, J. (1997) Intracellular Kinetics of a Growing Virus: A
- Genetically Structured Simulation for Bacteriophage T7. *Biotechnol Bioeng*. John Wiley
 & Sons, Inc.
- (43) Chan, L. Y., Kosuri, S., and Endy, D. (2005) Refactoring bacteriophage T7. *Mol.* Syst *Biol* 1 2005 0018
- 789 Syst. Biol. 1, 2005.0018.
- (44) Hu, Y. O. O., Hugerth, L. W., Bengtsson, C., Alisjahbana, A., Seifert, M., Kamal, A.,
- Sjöling, Å., Midtvedt, T., Norin, E., Du, J., and Engstrand, L. Bacteriophages Synergize
- with the Gut Microbial Community To Combat Salmonella 3, e00119--18.
- (45) Ganeshan, S. D., and Hosseinidoust, Z. antibiotics Phage Therapy with a Focus onthe Human Microbiota.
- (46) Hennig, S., Rödel, G., and Ostermann, K. (2015) Artificial cell-cell communication
- as an emerging tool in synthetic biology applications. *J. Biol. Eng.* 2015 91 9, 1–12.
- 797 (47) Bertram, R., and Hillen, W. (2007) The application of Tet repressor in prokaryotic 798 gene regulation and expression. *Microb. Biotechnol.*
- 799 (48) Schmitter, S., Fieseler, L., Klumpp, J., Bertram, R., and Loessner, M. J. (2017)
- 800 TetR-dependent gene regulation in intracellular Listeria monocytogenes demonstrates 801 the spatiotemporal surface distribution of ActA. *Mol. Microbiol.* 105, 413–425.
- 802 (49) Lortal, S., and Chapot-Chartier, M. P. (2005) Role, mechanisms and control of
- lactic acid bacteria lysis in cheese, in *International Dairy Journal*, pp 857–871. Elsevier.
- (50) Vrancken, G., Gregory, A. C., Huys, G. R. B., Faust, K., and Raes, J. (2019,
- 805 December 1) Synthetic ecology of the human gut microbiota. *Nat. Rev. Microbiol.*
- 806 Nature Publishing Group.
- 807 (51) Mabwi, H. A., Kim, E., Song, D. G., Yoon, H. S., Pan, C. H., Komba, E. V. G., Ko,

- G. P., and Cha, K. H. (2021, January 1) Synthetic gut microbiome: Advances and
- 809 challenges. *Comput. Struct. Biotechnol. J.* Elsevier B.V.
- (52) de Souza, R. S. C., Armanhi, J. S. L., and Arruda, P. (2020) From Microbiome to
- 811 Traits: Designing Synthetic Microbial Communities for Improved Crop Resiliency. Front.
- 812 Plant Sci. 11, 1179.
- 813 (53) Durfee, T., Nelson, R., Baldwin, S., Plunkett, G., Burland, V., Mau, B., Petrosino, J.
- F., Qin, X., Muzny, D. M., Ayele, M., Gibbs, R. A., Csörgo, B., Pósfai, G., Weinstock, G.
- M., and Blattner, F. R. (2008) The complete genome sequence of Escherichia coli
- B16 DH10B: Insights into the biology of a laboratory workhorse. *J. Bacteriol.* 190, 2597–
- 817 **2606**.
- 818 (54) Baker Brachmann, C., Davies, A., Cost, G. J., Caputo, E., Li, J., Hieter, P., and
- 819 Boeke, J. D. (1998) Designer Deletion Strains derived from Saccharomyces cerevisiae
- S288C: a Useful set of Strains and Plasmids for PCR-mediated Gene Disruption and
 Other Applications. *Yeast*.
- 822 (55) Yu, Y.-P., Gong, T., Jost, unter, Liu, W.-H., Ye, D.-Z., and Luo, Z.-H. Isolation and
- characterization of five lytic bacteriophages infecting a Vibrio strain closely related to
- 824 Vibrio owensii.
- 825 (56) Chaudhry, W. N., Haq, I. U., Andleeb, S., and Qadri, I. (2014) Characterization of a
- virulent bacteriophage LK1 specific for Citrobacter freundii isolated from sewage water. *J. Basic Microbiol.* 54, 531–541.
- 82/ J. Basic Microbiol. 54, 531–541.
- (57) (2021), R. C. T. R: A language and environment for statistical computing. *R Found. Stat. Comput. Vienna, Austria.*
- (58) Goeman, J. J., and Solari, A. (2014) Multiple hypothesis testing in genomics. *Stat. Med.* 33, 1946–1978.
- 832 (59) Bland, j. M., and Altman, D. G. (1995) Multiple significance tests: The Bonferroni
- 833 method. *BMJ 310*, 170.
- 834

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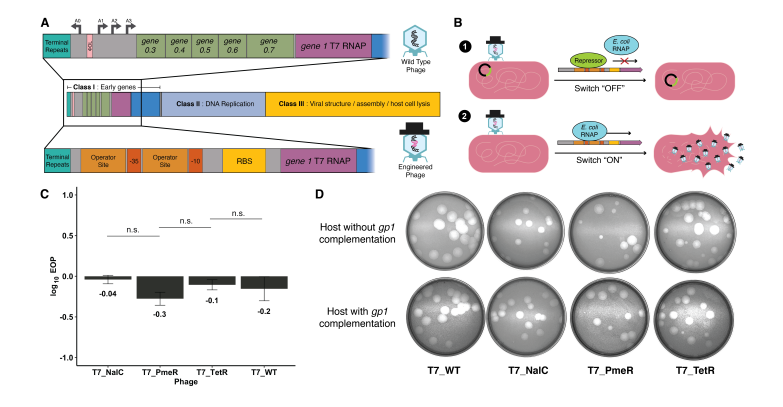


Figure 1. Engineered phages retain wildtype infectivity

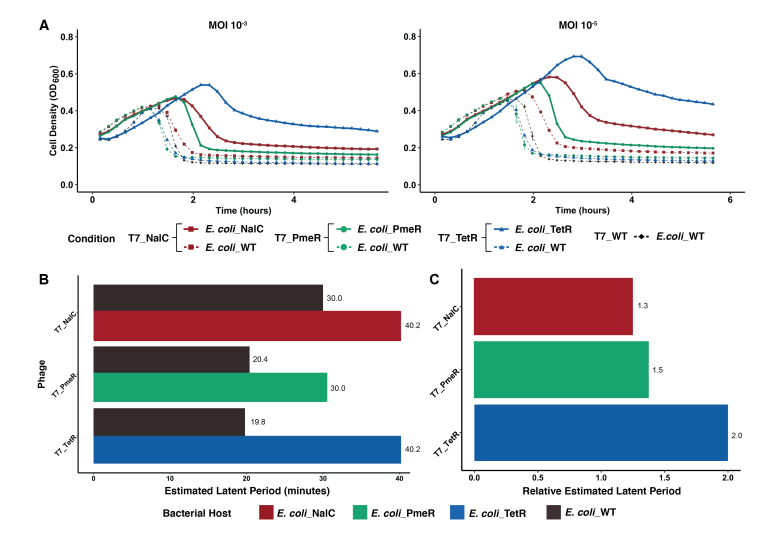


Figure 2. Repression of gp1 reduces activity in engineered phages

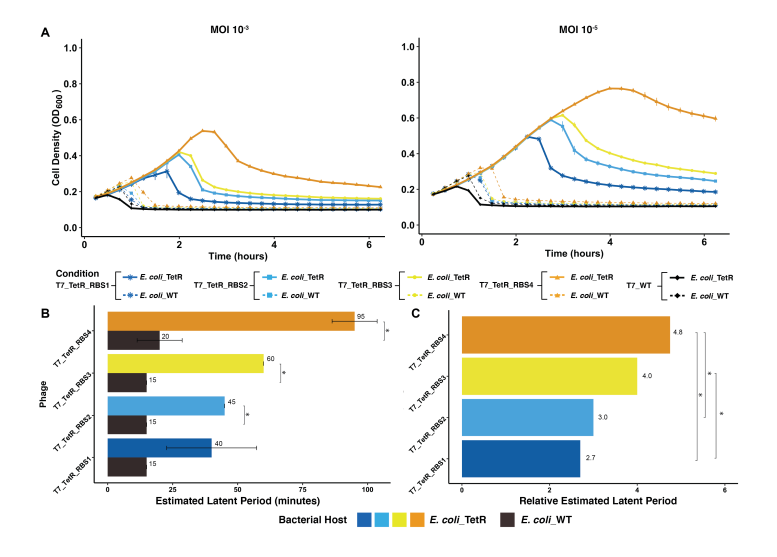


Figure 3. Engineered ribosome binding sites further enhances control over phage activity

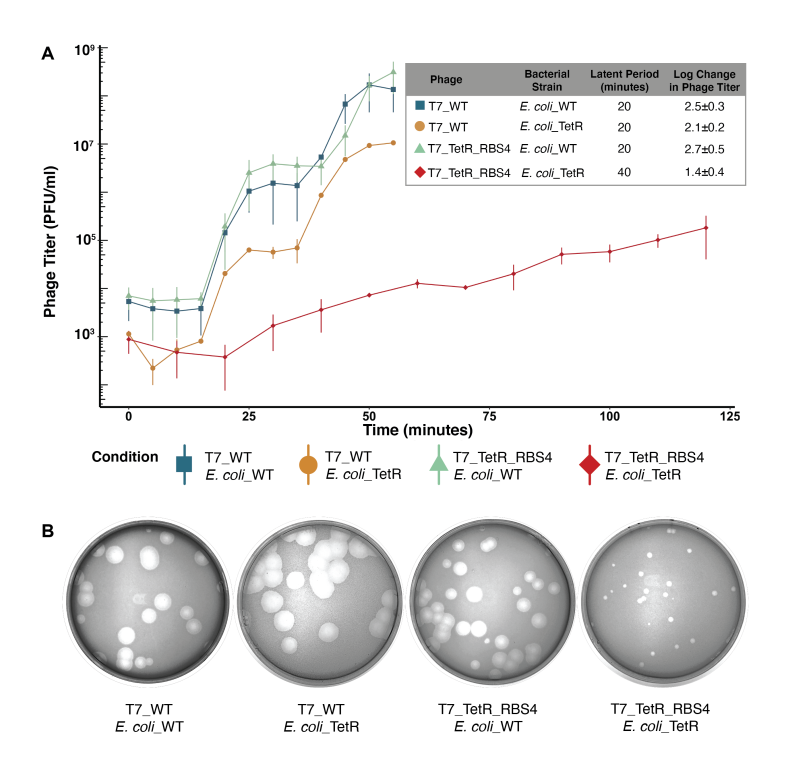
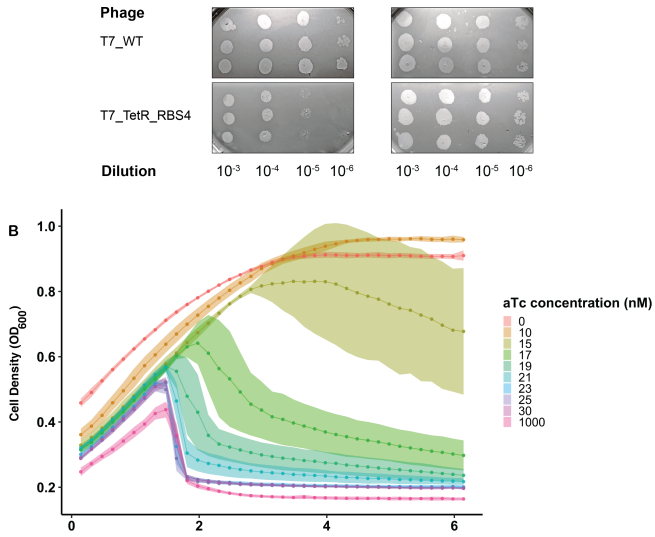


Figure 4. Repression significantly delays replication in engineered phages

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Repressed

Induced



Time (hours)

Figure 5. Phage infectivity can be tuned by changing inducer concentration

Α