

Antagonistic interactions between phage and host factors control arbitrium lysis–lysogeny decision

Received: 26 May 2023

Accepted: 7 November 2023

Published online: 4 January 2024

 Check for updates

Sara Zamora-Caballero^{1,6}, Cora Chmielowska^{1,6}, Nuria Quiles-Puchalt^{1,2,3,6}, Aisling Brady^{1,2,4}, Francisca Gallego del Sol¹, Javier Mancheño-Bonillo¹, Alonso Felipe-Ruiz¹, Wilfried J. J. Meijer⁵, José R. Penades²✉ & Alberto Marina¹✉

Phages can use a small-molecule communication arbitrium system to coordinate lysis–lysogeny decisions, but the underlying mechanism remains unknown. Here we determined that the arbitrium system in *Bacillus subtilis* phage phi3T modulates the bacterial toxin–antitoxin system MazE–MazF to regulate the phage life cycle. We show that phi3T expresses AimX and YosL, which bind to and inactivate MazF. AimX also inhibits the function of phi3T_93, a protein that promotes lysogeny by binding to MazE and releasing MazF. Overall, these mutually exclusive interactions promote the lytic cycle of the phage. After several rounds of infection, the phage-encoded AimP peptide accumulates intracellularly and inactivates the phage antiterminator AimR, a process that eliminates *aimX* expression from the *aimP* promoter. Therefore, when AimP increases, MazF activity promotes reversion back to lysogeny, since AimX is absent. Altogether, our study reveals the evolutionary strategy used by arbitrium to control lysis–lysogeny by domesticating and fine-tuning a phage-defence mechanism.

Bacterial viruses (phages) parasitize their host’s machinery to replicate and produce progeny, a process resulting in lysis of the cell. Temperate phages possess an alternative stage to the lytic cycle, termed lysogeny, where the viral DNA integrates into the host chromosome and becomes dormant. The genetic determinants that govern the lysis–lysogeny decision are well understood in the *Escherichia coli* model temperate phage Lambda¹, but less is known for Gram-positive infecting phages². Recently, it was discovered that some *Bacillus*-infecting phages encode a peptide-based communication system, named arbitrium, which allows the virus to ‘talk’ to their kin and coordinate the lysis–lysogeny decision by incorporating the population-sensing factor into this process³. Although originally described in phages of the SPbeta group

(phi3T and SPβ), the arbitrium system is widespread in Bacillota and can also be found in conjugative elements and plasmids⁴.

In the phi3T phage, the arbitrium system consists of three main components, all encoded in the phage genome³. AimP, the small signalling peptide, is secreted into the medium after phage infection and is internalized by surrounding bacteria. The sensor, AimR, is a transcription factor that in the absence of AimP promotes the expression of *aimX*, which exerts a negative regulatory effect on lysogeny by an unknown mechanism. When the host:phage ratio is high during initial stages of the phage infection, AimP peptide concentration is low, *aimX* is expressed and the lytic cycle is favoured³. Accumulation of the peptide allows the phage to preferentially switch to lysogeny

¹Instituto de Biomedicina de Valencia (IBV)-CSIC and CIBER de Enfermedades Raras (CIBERER)-ISCIII, Valencia, Spain. ²Centre for Bacterial Resistance Biology, Imperial College London, London, UK. ³Department of Biomedical Sciences, Faculty of Health Sciences, Universidad CEU Cardenal Herrera, CEU Universities, Alfara del Patriarca, Spain. ⁴Institute of Infection, Immunity and Inflammation, University of Glasgow, Glasgow, UK. ⁵Centro de Biología Molecular “Severo Ochoa” (CSIC-UAM), Universidad Autónoma, Cantoblanco, Madrid, Spain. ⁶These authors contributed equally: Sara Zamora-Caballero, Cora Chmielowska, Nuria Quiles-Puchalt. ✉e-mail: j.penades@imperial.ac.uk; amarina@ibv.csic.es

when the susceptible bacterial population is reduced, which allows safe persistence of both the integrated prophage and the bacterial host². AimP binds to AimR⁵, inactivating its function and therefore inhibiting the expression of *aimX*, which results in promoting lysogeny. However, the molecular details of this mechanism remain elusive.

We initiated this study trying to understand how *aimX* controls the lysis–lysogeny decision in phage phi3T. To our surprise, our results not only redefine roles for *aimX* and AimR, as well as propose an additional function for *aimP*, but also bring forward new phage- and host-encoded players. The mutually exclusive interactions between these players are essential for the functionality of the arbitrium system and provide important insights into how the arbitrium system controls lysis–lysogeny decisions in phage communication.

Results

aimX expression is regulated by a transcriptional terminator

Our previous study identified two additional key components of the lysis–lysogeny decision system in phage SPβ, *yopR* and *yopN*, which are part of a six-gene operon (*yopM–yopR*), conserved in the SPbeta group of phages. YopR is the master repressor of SPβ, while YopN expression favours lysogenization by an unknown mechanism⁶. This operon in phi3T comprises genes *phi3T_92* to *phi3T_97* (ref. 6) (Supplementary Fig. 1). We proposed that genes *phi3T_93* and *phi3T_97* encode the SPβ YopN and YopR functional homologues, respectively. Indeed, deleting *phi3T_97* generated a lytic phage (Extended Data Fig. 1a,b), confirming its role as phi3T's master repressor. Additionally, and in agreement with a recent report⁷, overexpression of *phi3T_93* or *phi3T_97* in a recipient strain blocked phage reproduction and increased lysogenization (Extended Data Figs. 1c–e and 2c–e), similar to YopN and YopR in SPβ⁶. However, the deletion of *phi3T_93* did not impact the phage titre or number of lysogens after infection (Extended Data Fig. 2a,b).

In SPβ, *yopN* and *yopR* were originally identified by evolving the SPβ Δ *aimR* mutant, which is deficient in titre and produces cloudy plaques⁶. We repeated this strategy for the phi3T Δ *aimR* mutant phage. For six evolved phages, from independent experiments, we observed a recovery of phage titre following mitomycin C (MC) induction (Extended Data Fig. 3a). These evolved phages generated lysogens at a significantly reduced level and formed sharp plaques (Extended Data Fig. 3b,c). Surprisingly, none of these evolved phages carried mutations in *phi3T_93* or *phi3T_97*. Instead, they carried the same mutation in a non-coding region between *aimP* and *aimX* (Fig. 1a and Supplementary Table 1), where we identified a putative transcriptional terminator (TT). The compensatory mutation in the evolved phages was predicted (ARNold^{8,9} and RNAfold¹⁰ software) to impact the stem of the TT and alter its secondary structure (Fig. 1b).

We confirmed that the TT was functional by cloning the region containing the putative TT (or the evolved mutated TT*) with the *aimX* gene transcriptionally fused to the β -galactosidase (β -gal) reporter gene in a plasmid with an isopropyl β -D-1-thiogalactopyranoside (IPTG)-inducible P_{Spa}nk promoter (Fig. 1a). In the presence of IPTG, the wild-type (wt) TT blocked β -gal (*aimX*) expression but the mutated TT* did not (Fig. 1a). This suggests that the evolved phages restored their function just by disrupting this TT, enabling *aimX* expression from an uncharacterized promoter located upstream of *aimX*.

AimR is a transcriptional antiterminator

It has been postulated that AimR activates *aimX* expression by binding to the AimR operator located upstream of *aimX*³. This region has been characterized, and the structure of AimR bound to DNA has been solved^{5,11}. However, we found no putative promoter when analysing (phiSITE tool PromoterHunter^{12,13}) the region downstream of the AimR binding site. To test this, a two-plasmid system was used, with one plasmid overexpressing AimR^{phi3T} and a second reporter plasmid carrying the cloned region from the TT (including the AimR binding site) to *aimX* fused to the β -gal reporter. Surprisingly, both in the presence

or absence of AimR^{phi3T}, no β -gal expression was observed. Introducing the prophage did not result in reporter expression, which excluded the possibility that other phage-encoded factors are required. These results suggested that there is no promoter present for *aimX* in this region and AimR does not induce *aimX* expression as a traditional transcriptional activator. Therefore, we hypothesized that (1) AimR is not a transcriptional activator but an antiterminator and (2) the upstream *aimP* promoter produces the transcript that terminates at the TT in the absence of AimR.

To test this, we cloned *aimP–aimX* regions of different lengths that included either the wt or mutated TT (see Fig. 1c for details). In the absence of AimR^{phi3T}, no expression of β -gal was observed in plasmids carrying the wt TT or lacking the *aimP* promoter, while expression was observed in the plasmid containing the *aimP* promoter with the mutated TT* (Fig. 1c). In the presence of AimR^{phi3T}, expression of the β -gal occurred only in those plasmids that contained the *aimP* promoter (Fig. 1c). Reverse transcription PCR analyses confirmed that *aimP* and *aimX* are part of the same bicistronic operon that initiates at the *aimP* promoter (Supplementary Fig. 2). In addition, mutation of the AimR^{phi3T} binding site in the reporter constructs abolished expression (Fig. 1c), confirming that AimR binding to its specific site is required for antitermination to occur. Altering the distance (increase of 69 bp or decrease of 8 bp) between the TT and the AimR binding site also abolished AimR function, supporting the function of AimR as an antiterminator instead of a transcriptional activator (Fig. 1c).

In a similar way, we confirmed that AimR from SPβ (AimR^{SPβ}) also functions as an antiterminator, preventing transcript termination of a TT identified upstream of the AimR^{SPβ} binding site (Extended Data Fig. 4a,b). We also obtained 8 additional SPβ Δ *aimR* evolved phages with restored capacity to form normal plaques. Three of them contained a single bp mutation in the TT upstream of the AimR^{SPβ} binding site (Supplementary Data Table 1), which inactivated its functionality (Extended Data Fig. 4c). Finally, by swapping the AimR binding sites of phi3T and SPβ in a reporter construct carrying the phi3T region from *aimP* promoter to *aimX* (see scheme in Fig. 1d), we confirmed that the mechanism of antitermination is conserved in both phages and requires the interaction of AimR with their cognate DNA-binding sites^{5,11} (Fig. 1d).

In summary, our results challenge the current understanding of how the arbitrium system works, suggesting that *aimP* and *aimX* form a bicistronic operon regulated by an AimR antitermination mechanism.

phi3T *aimX* encodes a small peptide

While it was assumed that *aimX* encodes a small antisense RNA (sRNA)^{3,4}, the fact that *aimX* is part of a bicistronic transcript suggests that it may encode a peptide, a possibility already considered in the original article reporting the arbitrium system³. In the phi3T genome, a 51-amino-acid-long open reading frame, phi3T_91, was annotated in the *aimX* locus, and our in silico analysis identified all the elements required for translation (Fig. 2a).

To test this, we established an approach to verify *aimX* functionality in vivo. Since the role of AimR seems to be to allow *aimX* expression by preventing termination, we expressed different versions of *aimX* from an ectopic chromosomal locus and checked whether they restored the severely affected induction of the Δ *aimR* mutant prophage^{6,14}. The wt *aimX* and a version with an altered DNA sequence (*aimX*^{mt}), but encoding the same protein, successfully complemented the Δ *aimR* mutant phage, both in the donor (restored efficient prophage induction) and recipient cells (restored plaque morphology). In contrast, a construct with a stop codon in the fifth amino acid (*aimX*^{stop}) did not complement the Δ *aimR* mutation (Fig. 2b and Supplementary Fig. 3a–c). We also expressed AimX heterologously in *E. coli* (fused to an N-terminal histidine tag) and purified the product, yielding a soluble monomeric protein in solution (Supplementary Fig. 4a,c). This supports the idea that *aimX* encodes a peptide and not a regulatory RNA in phi3T.

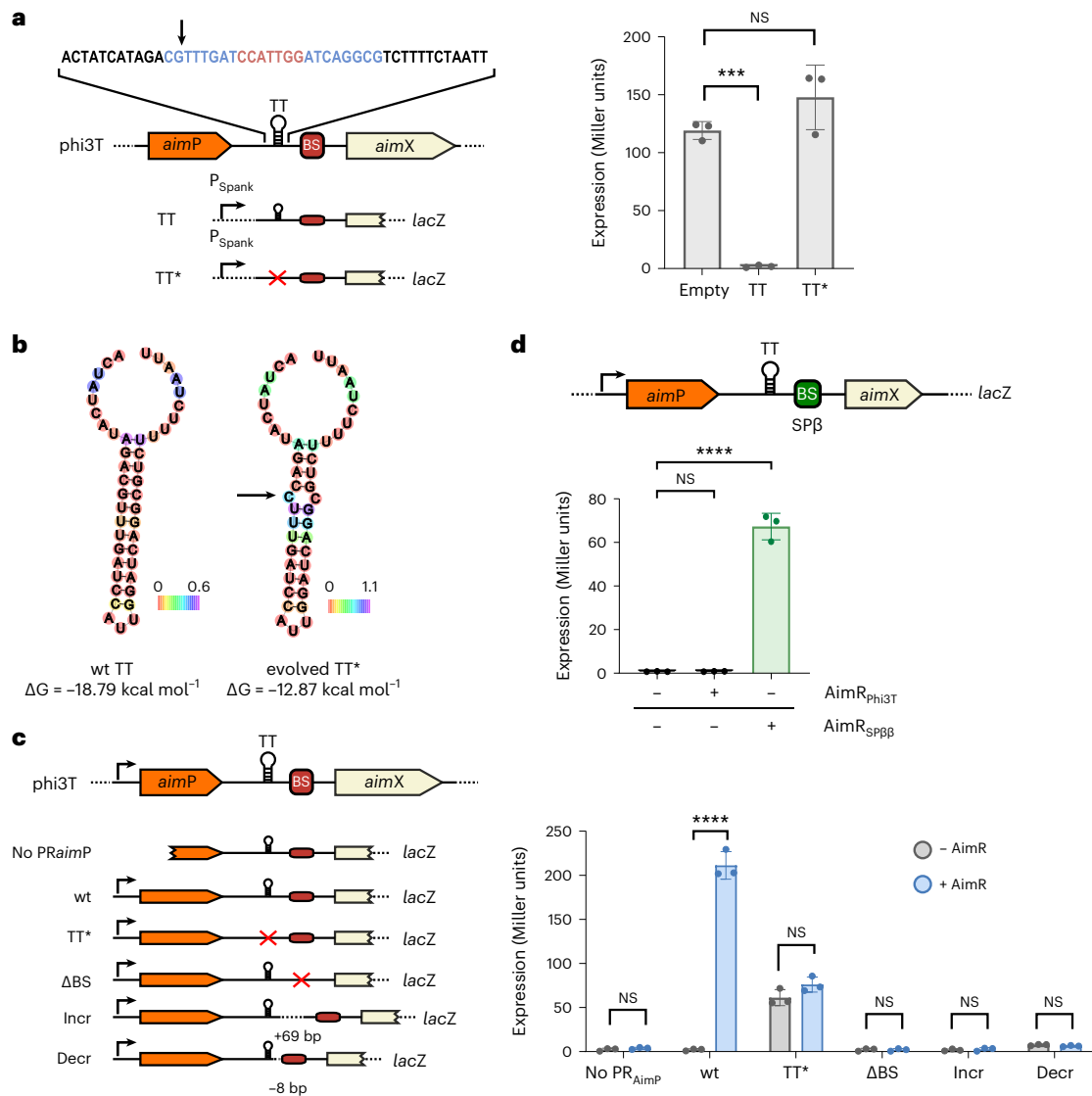


Fig. 1 | AimR functions as an antiterminator of a transcript that initiates at aimP. **a**, Left: schematic view of the regions cloned into the pDG1663+ P_{Spank} reporter plasmid and fused to the β -gal reporter gene. The DNA sequence (top) shows the predicted terminator and the regions corresponding to the stems (blue) and loop (red). Right: β -gal expression was measured in *B. subtilis* strains carrying the construction after IPTG induction of the P_{Spank} promoter. **b**, Predicted minimum free-energy secondary structures of the terminator for the wt and evolved sequence (RNAfold). Colours represent positional entropy in the structure. In **a** and **b**, the black arrow indicates the mutated base. **c**, Left: schematic view of relevant genetic features upstream from aimX and of the regions cloned into the reporter plasmid and fused with the β -gal gene. **d**, Top: schematic of the genetic features used to characterize cross-activity of AimR antitermination for SP β and phi3T where the binding site has been swapped.

c, Right and **d**, Bottom: β -gal expression measured in *B. subtilis* strains carrying the integrated reporter plasmid and pDR110 (empty or AimR_{phi3T}) or pDR110 (empty, AimR_{phi3T} or AimR_{SP β}) plasmid, respectively, after IPTG induction of the P_{Spank} promoter in pDR110. In the schematics: black arrows, promoters; black lollipop symbols, putative transcriptional terminators; TT, transcriptional terminator; TT*, evolved mutated TT; BS, AimR binding site (red, phi3T BS; green, SP β BS). Δ BS, mutated BS palindromes; Incr, increased distance between the TT and BS; Decr, decreased distance between the TT and BS. Data presented as mean \pm s.d. ($n = 3$). In **a** and **d**, an ordinary one-way analysis of variance (ANOVA), followed by Dunnett's multiple comparisons test, was performed; in **c**, two-tailed t -tests were performed. P values are indicated above each comparison: **** $P < 0.0001$; *** $P < 0.001$; NS, not significant.

AimX interacts with phage phi3T_93 and host MazF

To explore AimX function and identify potential targets, we performed a sequence similarity search. Apart from multiple AimX homologues, BLAST¹⁵ analyses showed two additional candidates sharing >50% identity with a region of AimX: (1) MazE, a host-encoded antitoxin that binds to its cognate MazF toxin inactivating its ribonuclease activity¹⁶ and (2) YosL, encoded by both SP β and phi3T. MazE and YosL have similar sizes (100–130 residues) and show sequence similarity with AimX in their C-terminal regions (Fig. 3a). Importantly, this region of MazE is involved in binding to the MazF toxin¹⁶. Therefore, while AimX might bind to and modulate the function of one of the known lysogeny-promoting

phi3T proteins (phi3T_93 or phi3T_97), the similarity of AimX to MazE suggested another possible target, the toxin MazF.

To test the potential binding partners of AimX, we used biolayer interferometry (BLI) assays, which showed that AimX does not bind to the master repressor phi3T_97 but binds to both MazF and phi3T_93. Interestingly, the affinity is more than twice as high for MazF than for phi3T_93 (Extended Data Table 1). In addition, MazF has five times more affinity for AimX than for MazE, its cellular partner (Extended Data Table 1). In vitro competition assays confirmed this preference, showing that AimX rapidly displaces MazE from its complex with MazF (Supplementary Fig. 5). The affinity differences of both complexes are

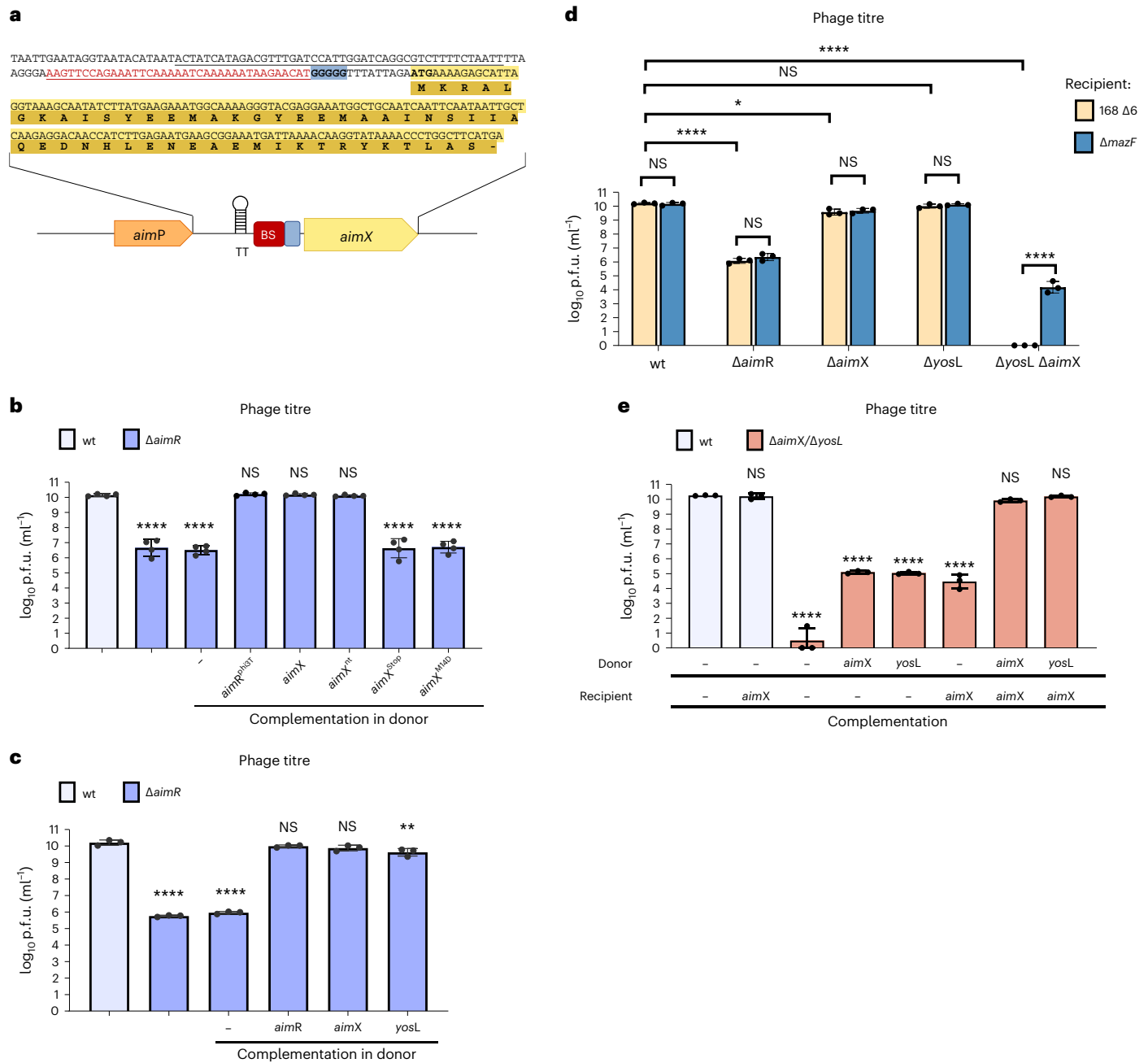


Fig. 2 | AimX is a peptide showing functional similarity to YosL.

a, Representation of the DNA region located between *aimP* and *aimX*. The TT is underlined in black. The AimR binding site (BS) is underlined in red. The Shine–Dalgarno region is highlighted in blue. The AimX coding sequence and the encoded protein sequence are highlighted in light and dark yellow, respectively, with the start codon in bold. **b, c**, Strains lysogenic for phages phi3T wt, Δ*aimR* and Δ*aimR* complemented (*amyE*::P_{spank}-) with *aimR*; different versions of the *aimX* (**b**) or *yosL* (**c**) gene were MC induced. Resulting phages were quantified using 168 Δ6 as the recipient strain. **d**, 168 Δ6 strains lysogenic for phi3T wt and mutant (Δ*aimR*, Δ*aimX*, Δ*yosL*, double Δ*aimX*/Δ*yosL*) were MC induced. Resulting phages were quantified using 168 Δ6 or 168 Δ6 Δ*mazF* mutant as recipient strains. **e**, Donor 168 Δ6 strains lysogenic for phages phi3T wt or

Δ*aimX*/Δ*yosL* and carrying empty vector (-) or expressing *aimX* or *yosL* were MC induced. Resulting phages were quantified using 168 Δ6 *amyE*::P_{spank}-empty (-) or 168 Δ6 *amyE*::P_{spank}-*aimX* as recipient strains. Results are presented as p.f.u.s ml⁻¹ and geometric mean and geometric s.d. are shown: **(b)** *n* = 4, **(c)** *n* = 3 **(d)** *n* = 3 and **(e)** *n* = 3. Values below the detection limit of 10 p.f.u.s ml⁻¹ were assigned a value of 1 and marked on the axis. An ordinary one-way ANOVA of log₁₀-transformed data, followed by Dunnett's multiple comparisons test, was performed to compare mean differences between titres in **b**, **c** and **e**. Comparisons were performed against wt phi3T. A two-way ANOVA of log₁₀-transformed data, followed by Tukey's multiple comparisons test, was performed to compare mean differences in titres in **d**. Adjusted *P* values: *****P* < 0.0001; ***P* < 0.005; **P* < 0.05.

mainly due to a decrease in dissociation constant (*k*_{off}), suggesting a more stable AimX–MazF complex with a potential regulatory function in the lysis–lysogeny decision (Extended Data Table 1).

Next, we analysed whether MazE and YosL^{phi3T} are also able to interact with phi3T₉₃ and MazF. In agreement with a recent study⁷,

MazE also binds to phi3T₉₃, which potentially results in an increased amount of active MazF^{7,16}. MazE has almost 15 times higher affinity for phi3T₉₃ than for its cellular target MazF (Extended Data Table 1). Surprisingly, while YosL^{phi3T} does not bind to the phage phi3T₉₃, it binds to the host MazF. MazF binds to YosL^{phi3T} with a higher affinity

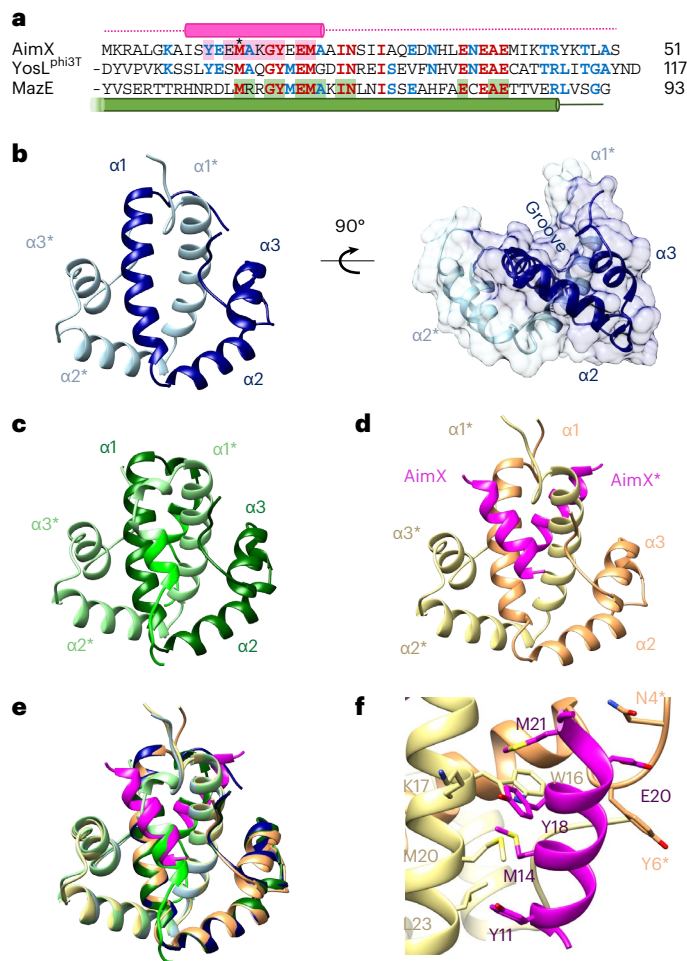


Fig. 3 | Molecular and structural characterization of arbitrium–MazE/F net of interactions. **a**, Sequence alignment for the homologous region of AimX, YosL^{phi3T} and MazE. Residues conserved in two or three of these sequences are highlighted in blue or red, respectively. Structural elements are shown for AimX and MazE in magenta and green, respectively. The AimX and MazE residues involved in contacts with their partners phi3T₉₃ and MazF are highlighted in pink and green background, respectively. The mutated methionine in AimX that abolishes phi3T₉₃ binding is pointed with a star. **b**, Ribbon representation of phi3T₉₃ (left) and side view of phi3T₉₃ surface (right). Each phi3T₉₃ monomer is depicted in a different blue shade and helices are numbered (* denotes second monomer). **c**, Ribbon representation of His-phi3T₉₃ structure. Each monomer is shown in a different shade of green. Residues belonging to the His-tag linker depicted in fluorescent green in both monomers. **d**, Ribbon representation of AimX-phi3T₉₃ complex with phi3T₉₃ monomers coloured in different shades of yellow and AimXs in magenta. **e**, Superimposition of AimX-phi3T₉₃ complex and His-phi3T₉₃ structures, following the same colour pattern as in **d** and **c**, respectively. **f**, Close-up view of the residues involved in phi3T₉₃ and AimX interaction. Interacting residues are labelled.

than with AimX or MazE, a similar affinity to that of the MazE–phi3T₉₃ complex (Extended Data Table 1). Similar results were obtained with YosL from phage SPβ (YosL^{SPβ}), confirming the YosL binding specificity of MazF (Extended Data Table 1). These results suggest that a common region of MazE, AimX and YosL is responsible for recognizing MazF, thus promoting AimX and YosL to putative antitoxins.

In vivo characterization of YosL

Our results suggested that the main role of AimR is regulating AimX expression, and overexpression of AimX from an ectopic chromosomal locus restored efficient $\Delta aimR$ prophage induction (Fig. 2b,c). However, induction of a $\Delta aimX$ mutant prophage resulted in a titre

similar to that of the wt phage, unlike the severely reduced titre of the $\Delta aimR$ prophage¹⁴ (Fig. 2b). One possible explanation is that AimR may control (directly or indirectly) the expression of another protein, whose function at least partially overlaps with the function of AimX. On the basis of our previous results, we hypothesized that YosL was a candidate.

Indeed, complementing the $\Delta aimR$ phi3T mutant by overexpressing *yosL* upon prophage induction restored high titre levels (Fig. 2c). We also attempted to complement the recipient strain during infection; however, we observed that overexpression of *yosL* prevented normal cell growth. During induction experiments, complementation was possible since we added IPTG and MC simultaneously, once the cells have already reached the desired density. Next, we generated a phi3T $\Delta yosL$ and a $\Delta aimX/\Delta yosL$ double mutant. No defect in phage titre was observed after the $\Delta yosL$ prophage induction (Fig. 2d); however, the $\Delta aimX/\Delta yosL$ prophage was severely affected in its ability to generate infective particles, even more than the $\Delta aimR$ mutant (Fig. 2d,e), and the cultures lysed badly upon induction. Simultaneous complementation of the $\Delta aimX/\Delta yosL$ mutant in the donor strain (with either *aimX* or *yosL*) and in the recipient (*aimX*) led to full titre recovery, while complementation in only recipient or donor resulted in partial titre recovery (Fig. 2e). Therefore, these proteins impact both the induction and infection processes and have, to some extent, an overlapping function.

MazF impacts the lysis–lysogeny outcome in phi3T

MazF is an RNA endoribonuclease¹⁷ during normal growth, forming a stable complex with its cognate antitoxin, MazE^{16,17}. However, if MazE is degraded or sequestered, MazF is released and cleaves cellular mRNAs to inhibit protein synthesis, leading to growth arrest¹⁶. The fact that both AimX and YosL bind to MazF suggested that this toxin may perform a central role in the life cycle of phi3T. The titre obtained when infecting the $\Delta mazF$ or $\Delta mazEF$ mutant strains with phi3T was identical to that obtained with the wt recipient strain (Fig. 4a); however, the number of lysogens was severely reduced in the absence of MazF (Fig. 4b) and the plaques produced were significantly sharper and larger (Fig. 4c). These observed differences were reverted by expression of the wt MazF in the recipient cells, but not after expression of the MazF T48A mutant which has no RNase activity¹⁶ (Fig. 4d). Taken together, these results indicated that MazF's activity impacts the lysis–lysogeny result by promoting lysogeny and/or inhibiting the lytic cycle.

A recent study suggested that the active MazF toxin suppresses phi3T phage's lytic propagation, possibly by an abortive infection mechanism⁷. Since AimX and YosL favour the lytic cycle of the phage, the formation of AimX–MazF or YosL–MazF complexes may block MazF activity. To test this, we introduced an IPTG-inducible *mazF* gene into the $\Delta mazEF$ strain, deprived of its natural antitoxin protein, MazE. Next, we lysogenized this strain with the wt phi3T phage or evolved phage carrying the double *aimR/TT* mutation, which constitutively expresses AimX. As predicted, toxicity of the IPTG-induced MazF was blocked in the presence of the evolved prophage expressing AimX, enabling bacterial growth (Fig. 4e).

We hypothesized that the $\Delta yosL/\Delta aimX$ double mutant phage produced a very low titre due to its inability to block MazF activity, which leads to suppression of lytic propagation. Indeed, the mutant phage formed more and sharper plaques in $\Delta mazF$ recipient strain. In addition, a $\Delta mazF$ strain lysogenized with the $\Delta aimX/\Delta yosL$ prophage lysed well after induction and produced an increased titre compared with the same prophage induced from the strain containing MazF. To fully restore the wt titre level, MazF had to be absent in both donor or recipient, indicating that both induction and infection are impacted by these interactions (Extended Data Fig. 5). Together, these results indicate that AimX or YosL needs to block MazF activity for the lytic cycle to proceed.

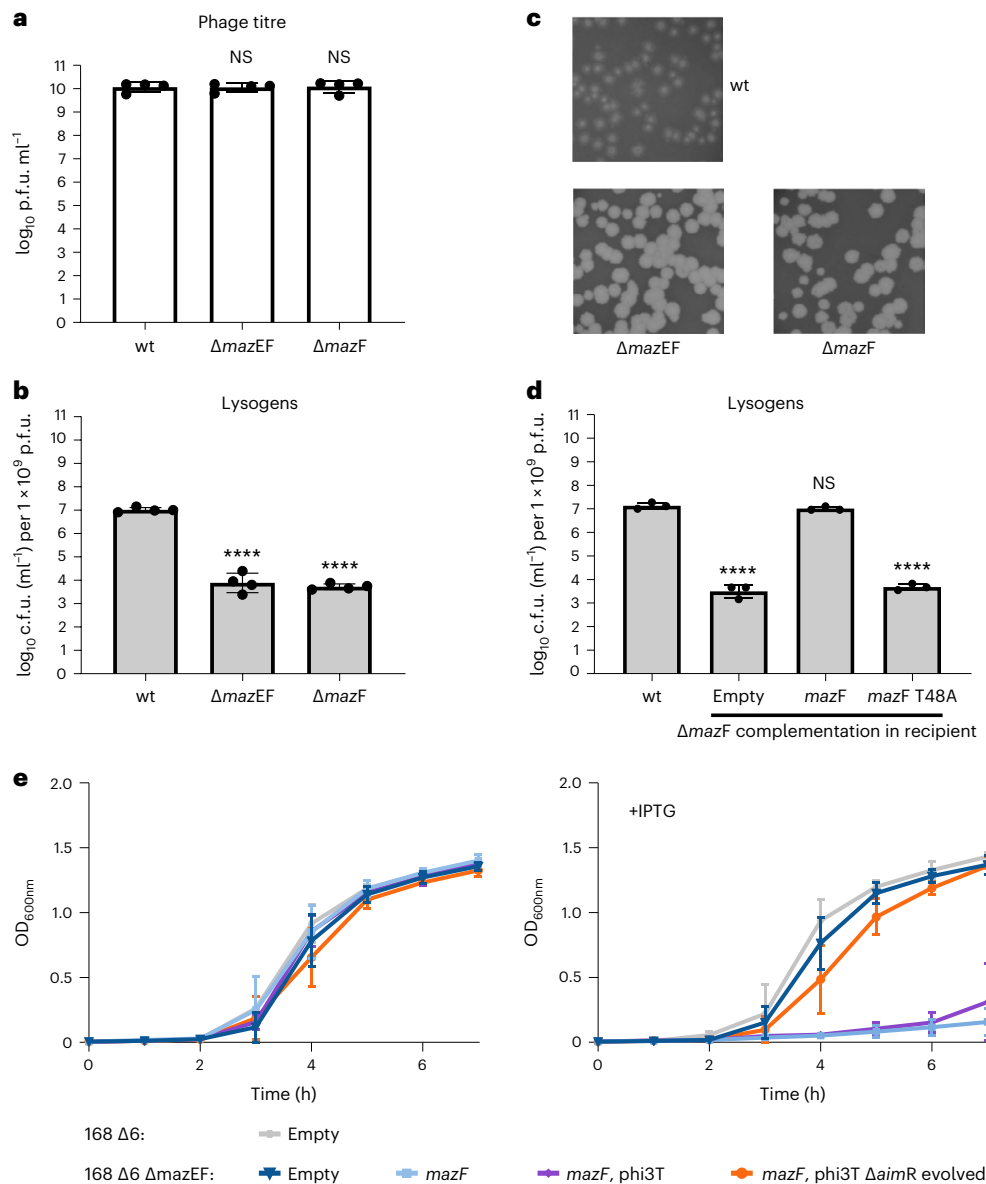


Fig. 4 | MazF activity is important for efficient lysogeny of phi3T phage. 168 $\Delta 6$ strain lysogenic for phi3T was MC induced. **a, b**, The number of resulting phages (**a**) and lysogens (**b**) were quantified using 168 $\Delta 6$ (wt) or mutated 168 $\Delta 6$ ($\Delta mazEF$, $\Delta mazF$) recipient strains. Data represent the geometric mean and geometric s.d. ($n = 4$). **c**, The resulting plaque morphologies were photographed. **d**, To test for complementation of the 168 $\Delta 6$ $\Delta mazF$ strain, phi3T was used to infect 168 $\Delta 6$ (wt) or 168 $\Delta 6$ $\Delta mazF$ -expressing amyE::P_{Spank}-empty/*mazF*/*mazF*^{T48A} (mutant of *mazF* with no RNase activity) as the recipient strains. Data represent geometric mean and geometric s.d. ($n = 3$). In **a**, results are shown as p.f.u.s ml⁻¹, and in **b** and **d** as c.f.u.s ml⁻¹ normalized by p.f.u.s ml⁻¹ and represented as the c.f.u. of an average phage titre (1×10^9 p.f.u.s ml⁻¹).

e, Growth curves (OD₆₀₀ readings) for strains carrying different IPTG-inducible expression vectors (amyE::P_{Spank}) and lysogenic phages that were grown without inducer (left) or with IPTG (right). Mean \pm s.d. ($n = 3$) are presented. Strains used: wt 168 $\Delta 6$ with empty vector (grey); 168 $\Delta 6$ $\Delta mazEF$ with empty vector (dark blue), expressing *mazF* (light blue), expressing *mazF* and lysogenic for wt phi3T prophage (purple), expressing *mazF* and lysogenic for evolved $\Delta aimR$ phi3T prophage (orange). An ordinary one-way ANOVA of log₁₀-transformed data, followed by Dunnett's multiple comparisons test, was performed to compare mean differences in titre numbers in **a**, **b** and **d**. Comparisons were performed against wt phi3T infecting 168 $\Delta 6$. Adjusted P values: **** $P < 0.0001$.

Structural characterization of the AimX–phi3T₉₃ complex

On the basis of these results, we propose that AimX, YosL, phi3T₉₃, MazE and MazF are involved in a network of mutually exclusive interactions essential for the lysis–lysogeny decision. Therefore, we attempted the structural characterization of its components. The structures of MazF in complex with MazE (PDB 4ME7) or with a target RNA (PDB 4MDX) have already been reported¹⁶. We obtained the structure of phi3T₉₃ alone (wt and its N-terminal His-tagged version; His-phi3T₉₃) and in complex with AimX (Extended Data Table 2). In all structures, phi3T₉₃ shows an identical all-helices fold composed of a longer

N-terminal helix ($\alpha 1$), followed by two shorter helices ($\alpha 2$ and $\alpha 3$) connected by a long unstructured loop (Fig. 3b–d) forming a V-like shape. The $\alpha 1$ helices of two phi3T₉₃ monomers interact, resulting in a W-shaped tight and globular dimer that displays two shallow grooves, one across each side of the dimer molecule (Fig. 3b). Size exclusion chromatography coupled to multi-angle light scattering (SEC–MALS) analysis confirmed the phi3T₉₃ dimeric organization in solution (Supplementary Fig 4b,c).

The structure of the AimX–phi3T₉₃ complex showed that a phi3T₉₃ dimer binds two independent AimX molecules that are

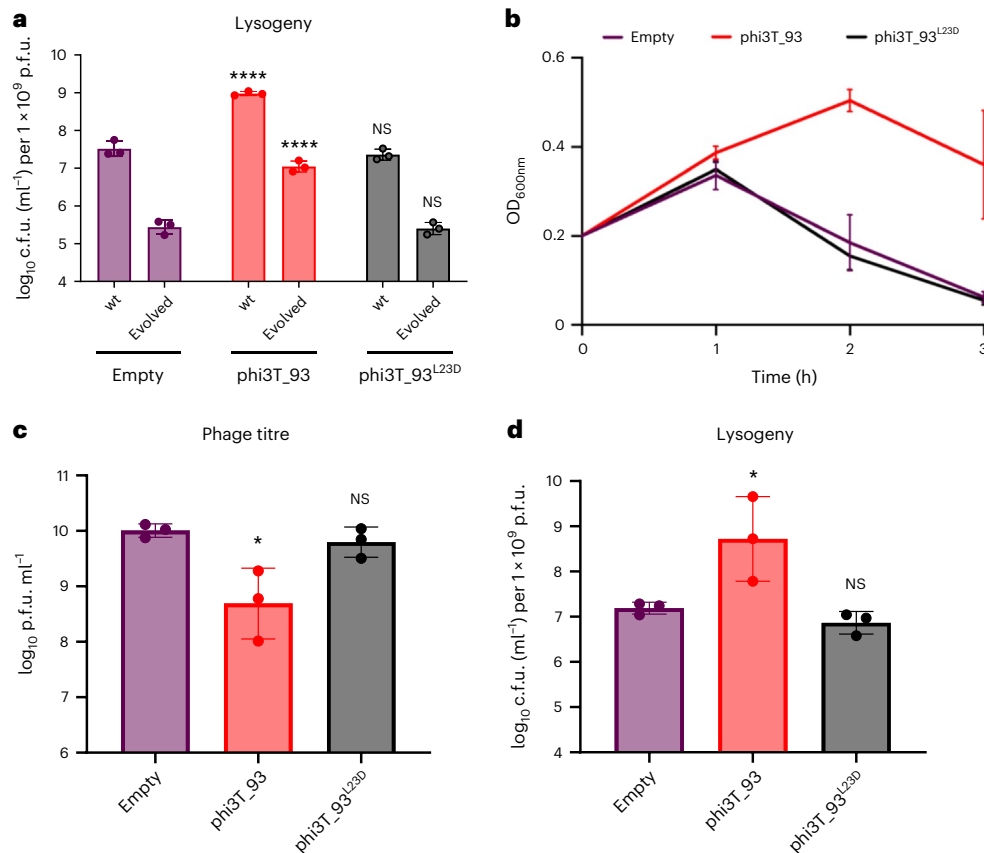


Fig. 5 | phi3T_93 promotes lysogeny. **a**, Strains lysogenic for phi3T wt and evolved $\Delta aimR$ phages were MC induced. The number of lysogens were quantified using 168 $\Delta amyE::P_{Spank}$ -empty or 168 $\Delta amyE::P_{Spank}$ -phi3T_93 (wt/L23D) as the recipient strains. **b–d**, *B. subtilis* strains either devoid of (purple) or expressing wt phi3T_93 (red) or phi3T_93^{L23D} (grey) were infected with phage phi3T wt at an MOI of 1:10 (phage:bacteria). For all infection experiments, growth (OD₆₀₀) readings (**b**), phage titres (**c**) and the relative number of lysogens (**d**) were determined after 3 h. Results are presented as p.f.u.s ml⁻¹ in **c**, or c.f.u.s ml⁻¹

normalized by p.f.u.s ml⁻¹ and presented as the c.f.u. of an average phage titre (1×10^9 p.f.u. ml⁻¹) in **a** and **d**. In **a**, **c** and **d**, geometric mean and geometric s.d. are shown ($n = 3$), and in **b**, mean \pm s.d. are shown ($n = 3$). An ordinary one-way ANOVA of log₁₀-transformed data, followed by Dunnett's multiple comparisons test, was performed to compare mean differences in titre or relative lysogen numbers. Comparisons were performed against results for the same phage (wt phi3T or evolved) infecting 168 $\Delta 6$ or 168 $\Delta amyE::P_{Spank}$ -empty strain. Adjusted *P* values: *****P* < 0.0001; **P* < 0.05.

accommodated in the shallow grooves (Fig. 3d,e). Only a short fragment of AimX (residues 8–22; 30% of the protein), which folds as a 3-turn helix, is visible in the AimX–phi3T_93 crystal (Fig. 3a,d). The main contacts between AimX and phi3T_93 are nucleated by the projection of AimX M14 and M21 towards the hydrophobic bottom of the phi3T_93 groove generated by the side chains of K13, K17, W16, M20 and L23, pointing to these hydrophobic interactions as key for phi3T_93–AimX recognition (Fig. 3f and Supplementary Table 2). Additional polar and hydrophobic interactions stabilize the complex (Supplementary Table 2). Interestingly, in the His-phi3T_93 structure, the N-terminal His-tag folds as helices and are also accommodated in the dimer grooves replicating the AimX contacts (Fig. 3c,e, Extended Data Fig. 6b and Supplementary Table 2), supporting the idea that the phi3T_93 grooves are specifically designed to accommodate a short helix. The portion of AimX visible in the structure corresponds to an area of high sequence similarity with MazE and YosL (Fig. 3a), explaining why all these three proteins bind to MazF. In MazE, this region also folds as a helix and AlphaFold also proposed a similar fold for YosL^{phi3T} (Extended Data Fig. 7a,b). Despite the structural differences between phi3T_93 (all alpha fold) and MazF (mainly beta fold), both AimX and MazE use an identical recognition mechanism by introducing a helix in the groove in the dimer's interface (Fig. 3b and Extended Data Fig. 7a). Structural alignment showed identical residues in the helix of AimX (M14, Y18, E20 and M21 for AimX) and MazE involved in the interactions with the

corresponding partners, and YosL^{phi3T} also presents identical residues in these positions (Fig. 3a and Extended Data Fig. 7a–c). Models of the putative AimX–MazF, YosL–MazF, MazE–phi3T_93 and YosL–phi3T_93 complexes showed that these key conserved residues occupy identical spatial arrangements allowing them to mimic interactions with their alternative partners (Extended Data Fig. 7d–g). In agreement with the BLI results (Extended Data Table 1), no steric hindrance was observed in the AimX–MazF, YosL–MazF and MazE–phi3T_93 models; however, for the YosL–phi3T_93 complex, steric clashes were observed between the YosL N-terminal domain and the phi3T_93 dimer (Extended Data Fig. 7d–i).

In vitro and in vivo validation of the interaction network

Our in silico complexes indicate that the AimX M14 residue plays an essential role in phi3T_93 and MazF recognition and substituting for a charged residue in that position should prevent AimX function (Fig. 3a,f and Extended Data Fig. 7c). Indeed, while the AimX^{M14D} mutant showed similar yield and behaviour in solution as the wt protein (Supplementary Fig. 4a,c), BLI analysis showed no binding of AimX^{M14D} to either phi3T_93 or MazF (Extended Data Table 1). As expected, the expression of the AimX^{M14D} mutant, unlike that of the wt AimX, did not complement the defective induction of the phi3T $\Delta aimR$ prophage (Fig. 2b).

Next, we produced phi3T_93^{L23D} by replacing a hydrophobic residue that interacts with the AimX M14 for a negatively charged one.

The new charged residue had no impact on the protein integrity and quaternary organization as confirmed by solving its crystal structure and SEC analyses (Extended Data Fig. 6c and Table 2, and Supplementary Fig. 4b,c). BLI analysis showed no binding of ϕ 3T₉₃^{L23D} to AimX and a strong reduction (two orders of magnitude) in its affinity for MazE (Extended Data Table 1). We hypothesize that ϕ 3T₉₃ promotes lysogeny by binding AimX and MazE, reducing the amount of antitoxins that can block MazF activity. Therefore, we used the evolved ϕ 3T Δ aimR TT-mutant which expresses AimX constitutively and as a result produces less lysogens (Extended Data Fig. 3b). Overexpression in the recipient cell of ϕ 3T₉₃, but not of the ϕ 3T₉₃^{L23D}, restored the ability of the evolved phage to produce lysogens (Fig. 5a). This confirmed that ϕ 3T₉₃–AimX interactions are important for the lysis–lysogeny decision. In addition, strains expressing ϕ 3T₉₃, unlike those expressing ϕ 3T₉₃^{L23D}, blocked the lytic cycle and promoted lysogeny of the wt ϕ 3T (measured by plaque and colony counts 3 h after infection; Fig. 5b–d). These results validate both the structural complexes and the common mechanism of recognition observed for AimX– ϕ 3T₉₃ and MazE–MazF, which are driven mainly by the insertion of a hydrophobic helix into the grooves of the dimers of ϕ 3T₉₃ and MazF. Given the conservation of residues in the proposed interacting helix of YosL, this model could also be extended to complexes of this putative antitoxin with MazF.

Collectively, our in vitro and in vivo results confirm the intricate network of mutually exclusive, and to some extent functionally redundant, interactions involving the phage arbitrium system and host MazEF toxin–antitoxin systems in the lysis–lysogeny decision.

Discussion

Arbitrium communication systems are highly diverse and are present in numerous phages and other mobile genetic elements⁴. Despite this diversity, it has been widely assumed that they share a conserved mode of action: AimR is a transcriptional regulator controlling the expression of *aimX*, which then directly inhibits the phage master repressor using an uncharacterized *cis*-antisense mechanism³. Our study challenges the roles associated with the arbitrium players and introduces additional ones, both for the phage (ϕ 3T₉₃, YosL) and for the host (MazEF). We have demonstrated that MazF is key in determining whether ϕ 3T will enter the lytic or the lysogenic cycle, and that the major function of the arbitrium system in ϕ 3T is to control MazF activity. Our model assumes that blocking MazF activity is essential for the lytic cycle to proceed. MazF has been described as a defence mechanism that blocks phage infection, so we expect that upon infection and in response to cellular damage, MazE is degraded and MazF released^{18,19}. Therefore, at early stages of infection, AimR inhibits the function of a TT located between *aimP* and *aimX*, allowing simultaneous expression of the *aimP*–*aimX* transcript (Extended Data Fig. 8). AimX, as well as another phage protein, YosL, bind to MazF and inactivate its function. AimX can also bind to ϕ 3T₉₃, preventing it from binding to MazE which would release MazF. Once AimP concentration increases in later infection stages, it will bind to AimR, inactivating its function. AimX will not be produced, allowing ϕ 3T₉₃ to bind MazE, thus liberating MazF (Extended Data Fig. 8). MazF then promotes lysogeny by an uncharacterized mechanism, possibly involving preferential cleaving of transcripts involved in the lytic cycle²⁰. Intriguingly, deletion of ϕ 3T₉₃ did not visibly impact phage titre or lysogen numbers. This, in agreement with a recent paper²⁰, indicates that there might be functional redundancy not only between AimX and YosL, but also between another protein and ϕ 3T₉₃, suggesting an even more complex regulatory network. This network of mutually exclusive interactions seems to be based on two main concepts: redundancy to ensure a fine-tuning control of the process and molecular mimicry which uses a conserved mechanism of recognition of the MazEF toxin–antitoxin system that the phage mimics with AimX, YosL and ϕ 3T₉₃. The precise regulation of the lysis–lysogeny decision appears to rely on a complex balance among all of these components, encompassing differences in

their binding strength, expression levels and potentially the timing of their expression.

Interestingly, ϕ 3T not only suppresses the host's defence systems, but possibly exploits it for its own benefit. Since toxin–antitoxin systems can be indicators of cellular stress²¹, involving them in the lysis–lysogeny decision may help the phage sense the physiological state of the host, beyond the classical SOS response required for induction. The connection between arbitrium and SOS pathways remains unknown in ϕ 3T and may involve other genes encoded in the *phi3T_92*–*phi3T_97* operon^{3,4,22}.

Another question is why *aimX* expression depends on an antiterminator and not a transcriptional activator? Potentially, this could accelerate the lysis–lysogeny switch by linking, via AimR, signalling (AimP) and effector (AimX) molecules with antagonistic functions. Interestingly, many SPbeta phages seem to encode an sRNA, not an AimX protein^{3,4,22}. Possibly, this sRNA may act as a preferential substrate of MazF, blocking the activity of this toxin on other targets by competition and thus enabling the lytic cycle. In addition, we have previously demonstrated that AimR binds to different binding sites across the bacterial genome⁵. Therefore, a potential AimR antiterminator regulon might control the lysis–lysogeny decision in arbitrium phages.

The arbitrium system was the first report of a viral communication system regulating lysis–lysogeny decisions. However, it is now clear that communication between phages and/or with their host for decision making is a more generalized feature as observed in *Vibrio* phages with quorum-sensing receptors that regulate lysis–lysogeny in response to bacterial host inducers, or in examples of bacterial quorum-sensing molecules inducing the lytic cycle in different phages^{23,24}. Our findings reveal a set of additional regulatory layers related to the lysis–lysogeny decision pathway, which must integrate different signals from the phage, host and the mutual environment to optimize the phage's survival. Since arbitrium is present not only in phages but also in other mobile genetic elements that infect both pathogenic and non-pathogenic bacteria, unravelling these complex networks of interactions is crucial to understanding how these phages, mobile genetic elements and bacteria evolve.

Finally, we would like to mention that in a parallel study we completed the characterization of the six-gene operon that contains the *yopMNOPQR* genes in phage SP β and the *phi3T_92* to *phi3T_97* genes in phage ϕ 3T²². Based on its function, we have renamed this operon as the 'SPbeta phages repressor operon' (*sro*) and proposed that the genes in the operon now be referred to as *sroABCDEF*^{SP β} or *sroABCDEF* ^{ϕ 3T}.

Methods

Bacterial strains and growth conditions

All bacterial strains used in this study are listed in Supplementary Table 3. *B. subtilis* and *E. coli* strains were routinely grown at 37 °C in liquid broth (LB Miller or LB Lennox, respectively) with shaking at 200 r.p.m. and 180 r.p.m., or plated into corresponding LB media with 1.5% (w/v) bacteriological agar plates. When required, antibiotics were utilized at the following concentrations: erythromycin (1 μ g ml⁻¹ or 5 μ g ml⁻¹), kanamycin (10 μ g ml⁻¹), ampicillin (100 μ g ml⁻¹), tetracycline (10 μ g ml⁻¹) or spectinomycin (100 μ g ml⁻¹).

Strain construction

Phages SP β (NC_001884) and ϕ 3T (KY030782) were used in this study. To generate the conditional *phi3T_97* deletion mutant in phage ϕ 3T, we first introduced into the Δ 6 ϕ 3T *phi3T_5::kan* strain a copy of the *phi3T_97* gene under the control of the P_{spank} promoter, adding 1 mM IPTG to ensure expression (see further information in 'Plasmids and cloning'). We generated overlapping PCRs containing the erythromycin marker (including *lox* sites) and 1 kb of flanking region for the *phi3T_97* gene. After transformation of the PCRs into Δ 6 ϕ 3T strain, insertion of the erythromycin cassette was confirmed by PCR and sequencing, followed by removal of the antibiotic resistance cassette²⁵.

Briefly, plasmid pDR244 was transformed into strains harbouring the *loxP*-flanked antibiotic resistance cassette with selection for spectinomycin resistance at 30 °C to allow for Cre/*lox*-mediated loop-out of the cassette. Transformant colonies were streaked onto LB plates with 1 mM IPTG and incubated overnight at 42 °C for removal of the temperature-sensitive plasmid. Strains were screened for plasmid curing (loss of spectinomycin resistance) and the antibiotic resistance cassette (loss of erythromycin resistance). Clean mutants were confirmed by PCR and sequencing. The additional mutants used in this study listed in Supplementary Table 3 were generated by overlapping PCRs as described above without complementation in *trans*. To generate mutants in *mazF*, we used strain BKE04660 as a template for the PCR.

Plasmids and cloning

All plasmids, primers and reagents are listed in Supplementary Tables 4, 5 and 6, respectively. Competent cell preparation and transformation was performed as follows: *B. subtilis* cells were grown in GM1 minimal media (SBase 1x (15 mM ammonium sulfate, 61 mM K₂HPO₄, 44 mM KH₂PO₄, 3.4 mM sodium citrate), 0.5% glucose, 0.1% yeast extract, 0.02% casein hydrolysate, 0.8 mM MgSO₄, 0.025% D/L tryptophan, 0.02% L-methionine) to early stationary phase to induce natural competence. Then, 1 µg of DNA (PCR or plasmid) was added to a subculture of 500 µl cells grown in 5 ml GM2 (GM1 supplemented with 3.3 mM MgSO₄ and 0.5 mM CaCl₂), followed by incubation at 37 °C for 1 h with shaking at 210 r.p.m.²⁶. Cultures were centrifuged at 6,000 g for 1 min, 800 µl of the supernatant removed, and the pellet resuspended and plated onto the relevant antibiotic plates. Plates were incubated at 37 °C overnight. For overexpression in *B. subtilis*, the genes were cloned into the *amyE* integration vector pDR110 under the control of the IPTG-inducible P_{spank} promoter²⁷. The *phi3T_97* gene was cloned into a pDR110 where the spectinomycin resistance marker (*spec*) was replaced with a tetracycline (*tetL*) resistance marker. The pDR110 was digested with EcoRI and AsiSI restriction enzymes, and a 6,544-bp band was gel-extracted and ligated to a PCR amplifying the *tetL* resistance cassette from pDG1515. The reporter plasmid pDG1663 (ref. 28) was modified to introduce a new multicloning site with EcoRI-NotI-HindIII-SpeI-NheI-SphI-BamHI. Then, we cloned the P_{spank} promoter into the modified version of pDG1663, generating pDG1663+ P_{spank}.

β-galactosidase reporter assays

Strains were grown at 37 °C overnight in LB containing the appropriate antibiotics. Cultures were diluted 1:100 into fresh LB with 0.1 mM MnCl₂ and 5 mM MgCl₂, and grown at 37 °C with 210-r.p.m. shaking until an optical density (OD)₆₀₀ of 0.2. When needed, 1 mM IPTG was added and cultures were grown for an additional hour before 1 ml of sample was taken and pelleted. OD₅₉₅ values were measured for each sample. To measure β-galactosidase levels, the Miller method was used²⁹. Briefly, cells were resuspended in Z buffer (60 mM Na₂HPO₄·7H₂O, 40 mM NaH₂PO₄, 10 mM KCl, 1 mM MgSO₄, 50 mM β-mercaptoethanol) and permeabilized with lysozyme. Following incubation with ONPG solution (4.0 mg ml⁻¹ *o*-nitrophenyl-β-D-galactoside in Z buffer), the time taken for samples to change colour was noted and absorbance (A)₄₂₀ readings were taken. Miller units were calculated as (1,000 × A₄₂₀)/(reaction time (min) × OD₅₉₅).

Bacteriophage induction assay

For induction, overnight cultures were diluted 1:100 in LB with 0.1 mM MnCl₂ and 5 mM MgCl₂, and grown at 37 °C with 210-r.p.m. shaking until an OD₆₀₀ of 0.2. This step was repeated twice to ensure the cells were in exponential growth. After the second growth, MC at 0.5 µg ml⁻¹ was added. When experiments were performed to test the complementation of the mutants, 1 mM of IPTG was added simultaneously with MC. The induced cultures were incubated at 30 °C with 80-r.p.m. shaking for 4 h and then left overnight at room temperature. Following lysis, samples were filtered (0.2 µm) and lysates were stored at 4 °C until use.

Total RNA extraction

An induction experiment was performed using wild-type *phi3T*, *ΔaimR* and *ΔaimR* evolved. After the second growth and before adding MC, 5 ml of sample was taken (time 0'), the rest of the culture was divided into two flasks with 10 ml of sample in each, adding MC (0.5 µg ml⁻¹) to one flask. After 1 h incubation at 80 r.p.m. at 32 °C, 5 ml of each sample (time 60' with or without MC treatment) were taken. For RNA extraction, each sample was mixed with two volumes of RNAprotect bacteria reagent (Qiagen) and incubated at room temperature for 5 min, then centrifuged. Pellets were snap frozen on dry ice and resuspended in 1 ml TRIzol reagent (Ambion) and lysed in a FastPrep-24 homogenizer (MP Biomedicals Lysing Matrix B tubes) using two cycles of 60 s at 6.5 m s⁻² interrupted by 5 min incubation on ice. Total RNA was extracted using the TRIzol Plus RNA purification kit following manufacturer instructions. Genomic DNA was removed using an on-column DNase digestion step with an RNase-free DNase kit (Qiagen). Residual DNA was removed by a second DNase treatment using RQ1 DNase (Promega). Total RNA was measured using Nanodrop 2000/2000c (Thermo Scientific).

Reverse transcription-PCR

The High-Capacity cDNA Reverse Transcription kit (Applied Biosystems) was used to convert RNA into complementary DNA. To verify the absence of genomic DNA in every RNA sample, the reverse transcription (RT) reaction was performed in the presence and absence of MultiScribe reverse transcriptase. In each reaction, 0.5 µg total RNA was subjected to RT following manufacturer instructions. The cDNA obtained was purified using QIAquick PCR purification kit (Qiagen) and 1 µl was used for PCR using Platinum *Taq* DNA polymerase HiFi (Life Technologies).

Bacteriophage titring assay

Overnight cultures of the recipient strains (*B. subtilis* Δ6 or with the corresponding integration vector) were diluted 1/100 in LB with 0.1 mM MnCl₂ and 5 mM MgCl₂, and then grown at 37 °C with 210-r.p.m. shaking until reaching an OD₆₀₀ of 0.2. If needed, 1 mM IPTG was added. Then, 100 µl of recipient bacteria was infected with 100 µl of serial dilutions of phage lysate in phage buffer (PhB; 1 mM NaCl, 0.05 M Tris pH 7.8, 0.1 mM MnCl₂, 5 mM MgCl₂) at room temperature for 10 min. Then, 3 ml of phage top agar (LB with 0.1 mM MnCl₂, 5 mM MgCl₂ and 0.7% agar) at 55 °C was added to the culture-phage mix and immediately poured over phage base agar plates (LB with 0.1 mM MnCl₂, 5 mM MgCl₂ and 1.5% agar). Plaques were counted after overnight growth at 37 °C and photographed.

Lysogenization assays

Lysogens were quantified by growing a recipient strain to OD₆₀₀ 0.2. Lysates containing a kanamycin marker were serially diluted in PhB and 100 µl was added to 1 ml of the recipient bacteria in 12 ml tubes. After incubating at 37 °C for 30 min to allow the phage to infect bacteria, the mixture was then transferred to 1.5 ml Eppendorf tubes and centrifuged at 3,750 g for 1 min. The supernatant was removed and the bacterial pellet was resuspended in fresh LB broth before plating onto selective antibiotic LB agar plates. Plates were incubated overnight at 37 °C. The number of colony-forming units (c.f.u.) was calculated.

Infection experiments

Cultures of the indicated strains were grown to OD₆₀₀ 0.2 corresponding to ~3 × 10⁷ c.f.u. ml⁻¹. If needed, 1 mM IPTG was added. Then, 15 ml of this culture was infected with the defined phage lysates at an MOI of 1:10 (phage:bacteria) and incubated at 30 °C and 80 r.p.m. At the indicated timepoints, samples were taken to assess the OD₆₀₀. At the end timepoint, samples were taken to assess the number of lysogens and phage titres. For evaluating the phage titres, 1 ml of the co-culture was filtered (0.2 µm) to generate phage lysates. The number of phage

particles was quantified by a titring assay against 168 Δ6. To quantify the number of lysogens, 1 ml of the co-culture was centrifuged at 3,750 *g* for 1 min and resuspended in 200 μl of fresh LB. The cells were then serially diluted in PBS and spotted onto LB kanamycin plates. All plates were incubated overnight at 37 °C and the number of plaque forming units (p.f.u.s) and c.f.u.s were calculated.

Growth experiments

Overnight cultures of the relevant strains were diluted to OD₆₀₀ 0.05 in LB with 0.1 mM MnCl₂ and 5 mM MgCl₂ and, if indicated, 1 mM IPTG, then grown at 37 °C with 120-r.p.m. shaking for 7 h. At the marked timepoints, samples were taken to assess the OD₆₀₀.

Statistical analysis

Statistical analyses were performed as indicated in the figure legends using GraphPad Prism 9 software. The *P* values represented in each figure are shown in the figure legends and detailed statistical test data, including exact *P* values, are provided in the source data file. In all figure descriptions, the number *n* represents biologically independent experiment results.

Recombinant protein expression and purification

The *aimX*, *phi3T_93*, *phi3T_97* and *yosL^{phi3T}* genes were amplified using genomic DNA from *B. subtilis* strain 1L1. The *yosL^{SPβ}*, *mazE* and *mazF* genes were amplified using genomic DNA from *B. subtilis* strain 168. In both cases, additional overhangs necessary for cloning were added (see primers in Supplementary Table 5). PCRs were cloned into the pLIC-SGC1 plasmid using NEBuilder HiFi DNA Assembly master mix (New England Biolabs). After sequence confirmation, plasmids were transformed into *E. coli* strain BL21 (DE3) RIL (Agilent) for protein overexpression.

For protein production, strains carrying the expression plasmid were grown overnight at 37 °C in 50 ml of LB with 50 μl ampicillin (100 μg ml⁻¹) and 50 μl chloramphenicol (33 μg ml⁻¹). The culture was used to inoculate 1 l of LB containing ampicillin and chloramphenicol, and was grown at 37 °C until OD₆₀₀ 0.6–0.8. Then, the temperature was set to 20 °C and protein expression was induced with 0.4 mM IPTG. The culture was incubated at 20 °C for an additional 16 h. Cells were collected by centrifugation, washed with PBS and frozen at –20 °C until use.

For protein purification, cell pellets were resuspended in lysis buffer (50 mM Tris pH 8, 500 mM NaCl) and sonicated for 10 min on ice. Cell debris was separated by centrifugation at 10,000 *g* for 1 h. The supernatant was filtered (0.45 μm) and loaded onto a 1 ml TALON (Cytiva) affinity column equilibrated with buffer A (50 mM Tris pH 8, 250 mM NaCl), washed with 10 column volumes of buffer A supplemented with 25 mM imidazole and stepwise eluted with buffer B (buffer A with 500 mM imidazole). Fractions containing the purest protein as confirmed by SDS–PAGE were pooled and digested, if needed, with TEV protease (50:1 molar ratio protein:TEV) for 16 h at 4 °C while being dialysed against buffer A supplemented with 1 mM EDTA and 1 mM β-mercaptoethanol. Digested and dialysed samples were reloaded onto the TALON column pre-equilibrated with buffer A to remove the His-tag and the protease. The non-retained protein was concentrated by centrifugal filtration in a 3-kDa-cut-off Amicon Ultra system (Millipore) and loaded onto a Superdex 75 increase 10/300 (GE Healthcare) gel filtration column equilibrated in buffer A. After an isocratic elution with buffer A, the purest fractions were pooled, concentrated by centrifugal filtration, flash frozen in liquid nitrogen and stored at –80 °C until use. In all cases, protein purity levels were superior to 90% according to SDS–PAGE. AimX, phi3T_93, phi3T_97, YosL^{phi3T}, YosL^{SPβ} and MazE^{T78A} yielded 5–15 mg of pure protein per litre of LB. MazE yielded ~2 mg of pure protein per litre of LB in a soluble and folded form as confirmed by circular dichroism (Supplementary Fig. 6).

Protein crystallization

All crystals were obtained in the IBV-CSIC crystallogenes facility at 21 °C using the sitting-drop vapour-diffusion method. His-phi3T_93 crystals were grown by mixing 0.3 μl protein solution at 10 mg ml⁻¹ and 0.3 μl reservoir solution (0.1 M Tris pH 8, 5% glycerol and 0.6 M Na/K tartrate). phi3T_93 crystals were obtained by mixing 0.3 μl protein solution at 10 mg ml⁻¹ and 0.3 μl reservoir solution (10% PEG 8000, 50 mM magnesium acetate and 100 mM sodium acetate). phi3T_93^{L23D} crystals were grown by mixing 0.3 μl protein solution at 10 mg ml⁻¹ and 0.3 μl reservoir solution (18% w/v PEG 8000, 100 mM HEPES sodium salt pH 7.5, 200 mM calcium acetate). Crystals of AimX–phi3T_93 complex were obtained by co-crystallization. Samples were prepared by mixing 65 μl of phi3T_93 at 2 mM and 387 μl of AimX at 336 μM (1:1 molar ratio) and concentrating the mixture to 100 μl. Crystals were obtained by mixing 0.3 μl of the protein mixture and 0.3 μl reservoir solution (0.1 M HEPES pH 7.5, 10% PEG 8000 and 0.2 M calcium acetate).

Data collection, structure solution and refinement

Before data collection, crystals were flash frozen in liquid nitrogen. For all phi3T_93 crystals, the corresponding crystallization solution was supplemented with 10% glycerol as cryoprotectant. AimX–phi3T_93 complex crystals were frozen in paratone. All diffraction experiments were carried out at 100 K.

His-phi3T_93, phi3T_93 and phi3T_93^{L23D} (PDB codes 8ANT, 8ANU and 8C8E, respectively) diffraction data were collected at Xaloc beamline, ALBA synchrotron in Barcelona using a wavelength of 0.97925 Å for phi3T_93 and 0.97918 Å for His-phi3T_93 and phi3T_93^{L23D}. Data reduction was carried out in CCP4. Phases for His-phi3T_93 were obtained by ab initio phasing using ARCIMBOLDO LITE³⁰ as implemented in CCP4i2 (ref. 31), whereas phi3T_93 and phi3T_93^{L23D} phases were obtained by molecular replacement using Phaser as implemented in Phenix³² and His-phi3T_93 or phi3T_93 as respective models. AimX–phi3T_93 (PDB code 8ANV) crystals were collected at beamline I04 of Diamond Light Source synchrotron (Oxfordshire, UK) using a wavelength of 0.9795 Å. Data reduction was carried out by the beamline software pipeline. The structure of the complex was solved at 2.2 Å resolution by molecular replacement using the phi3T_93 structure as a model for Phaser³³ in CCP4 suite. Following phase determination, model refinement was carried out combining manual building with Coot³⁴ (v.0.9.8.8) and computational refinement using phenix.refine³⁵. Extended Data Table 2 details all the X-ray data collection and refinement statistics.

Structure prediction

The predicted structures of YosL from phi3T and SPβ phages were determined with AlphaFold2 run online in Google ColabFold v.1.5.2 using default (auto) settings³⁶. Proteins with structural similarity to the structures determined experimentally or predicted by Alphafold were searched using the DALI server³⁷. Structures were visualized and figures prepared with PyMOL 2.1 (Schrodinger) and Chimera (<https://www.cgl.ucsf.edu/chimera/>).

Binding kinetics analysis

Binding kinetics were measured by biolayer interferometry using the BLITZ system (FortéBio, PALL) and Ni-NTA biosensors (Sartorius) at room temperature. All proteins were diluted in binding buffer (25 mM Tris pH 8, 150 mM NaCl, 3 mM imidazole and 0.005% Tween 20). Measurements were carried out by immobilizing the protein bait at 0.3 mg ml⁻¹ for 120 s to the biosensors through its N-terminal His-tag. Association and dissociation were monitored for 60 s each using the protein prey at concentrations from 1.25 to 0.07813 μM, obtained by 1:2 serial dilution from the highest concentration. Data analysis was performed using BLITZ pro software (FortéBio).

SEC–MALS experiments

For SEC–MALS assays, 20 μl of phi3T_93 (wt and the different mutants) or AimX constructs at 1 mg ml^{-1} were injected onto a Shodex KW-402.5-4F column, equilibrated with 100 mM HEPES pH 7.5 and 250 mM NaCl using a Shimadzu HPLC equipped with a manual injector at a flow rate of 0.3 ml min^{-1} . MALS analysis was performed using a miniDAWN Treos MALS detector followed by an optilab T-REX and a DLS detector (Wyatt). Data analysis was performed using Astra software (Wyatt).

Circular dichroism

Data collection was performed between wavelengths 200–275 nm in a J-1500 spectrometer with quartz cuvettes, using 150 μl 27 μM MazE diluted in PBS (20 mM) at pH 7.0. Millidegrees were used for secondary structure determination. The composition of secondary structures was estimated from the data using the software BeStSel³⁸.

Native-PAGE

A modified version of Native-PAGE analysis³⁹ was used to study the dynamics of complex formation. Briefly, a MazE–MazF complex was initially obtained by mixing MazE (30 μM) and MazF (60 μM) (ratio 1:2) in buffer A (200 mM NaCl, 50 mM Tris-HCl pH 8.0) and incubating at room temperature for 15 min. After complex formation, AimX was added to the MazE–MazF mixture to a final concentration of 30 μM at 20 min, 10 min and 5 min before loading the sample into the gel. The samples were separated in 8% acrylamide native gel (100 V, 4 °C, 2 h) and visualized by Coomassie blue staining.

Reporting summary

Further information on research design is available in the Nature Portfolio Reporting Summary linked to this article.

Data availability

The atomic coordinates of the phi3T_93, His-phi3T_93, phi3T_93^{L23D} and AimX-phi3T_93 complex have been deposited at <http://www.pdb.org>, with PDB codes 8ANT, 8ANU, 8CSE and 8ANV, respectively. The previously determined structures used in this study are available from the PDB (<http://www.pdb.org>) under the accession codes indicated. The rest of the data are available in the main text, supplementary materials and auxiliary files. Plasmids and bacterial strains generated during this work are listed in Supplementary Tables 3 and 4 and are available upon request. Source data are provided with this paper.

References

- Court, D. L., Oppenheim, A. B. & Adhya, S. L. A new look at bacteriophage λ genetic networks. *J. Bacteriol.* **189**, 298–304 (2007).
- Brady, A. et al. Molecular basis of lysis–lysogeny decisions in Gram-positive phages. *Annu. Rev. Microbiol.* **75**, 563–581 (2021).
- Erez, Z. et al. Communication between viruses guides lysis–lysogeny decisions. *Nature* **541**, 488–493 (2017).
- Stokar-Avihail, A., Tal, N., Erez, Z., Lopatina, A. & Sorek, R. Widespread utilization of peptide communication in phages infecting soil and pathogenic bacteria. *Cell Host Microbe* **25**, 746–755.e5 (2019).
- Gallego del Sol, F., Penadés, J. R. & Marina, A. Deciphering the molecular mechanism underpinning phage arbitrium communication systems. *Mol. Cell* **74**, 59–72.e3 (2019).
- Brady, A. et al. The arbitrium system controls prophage induction. *Curr. Biol.* **31**, 5037–5045.e3 (2021).
- Cui, Y. et al. Bacterial MazF/MazE toxin-antitoxin suppresses lytic propagation of arbitrium-containing phages. *Cell Rep.* **41**, 111752 (2022).
- Macke, T. et al. RNAMotif: a new RNA secondary structure definition and discovery algorithm. *Nucleic Acids Res.* **29**, 4724–4735 (2001).
- Gautheret, D. & Lambert, A. Direct RNA motif definition and identification from multiple sequence alignments using secondary structure profiles. *J. Mol. Biol.* **313**, 1003–1011 (2001).
- Hofacker, I. L. Vienna RNA secondary structure server. *Nucleic Acids Res.* **31**, 3429–3431 (2003).
- Gallego del Sol, F., Quiles-Puchalt, N., Brady, A., Penadés, J. R. & Marina, A. Insights into the mechanism of action of the arbitrium communication system in SPbeta phages. *Nat. Commun.* **13**, 3627 (2022).
- Klucar, L., Stano, M. & Hajduk, M. PhiSITE: database of gene regulation in bacteriophages. *Nucleic Acids Res.* **38**, D366–D370 (2009).
- Stano, M. & Klucar, L. PhiGENOME: an integrative navigation throughout bacteriophage genomes. *Genomics* **98**, 376–380 (2011).
- Bruce, J. B., Lion, S., Buckling, A., Westra, E. R. & Gandon, S. Regulation of prophage induction and lysogenization by phage communication systems. *Curr. Biol.* **31**, 5046–5051.e7 (2021).
- Boratyn, G. M. et al. BLAST: a more efficient report with usability improvements. *Nucleic Acids Res.* **41**, W29–W33 (2013).
- Simanshu, D. K., Yamaguchi, Y., Park, J.-H., Inouye, M. & Patel, D. J. Structural basis of mRNA recognition and cleavage by toxin MazF and its regulation by antitoxin MazE in *Bacillus subtilis*. *Mol. Cell* **52**, 447–458 (2013).
- Park, J. H., Yamaguchi, Y. & Inouye, M. *Bacillus subtilis* MazF-bs (EndoA) is a UACAU-specific mRNA interferase. *FEBS Lett.* **585**, 2526–2532 (2011).
- Alawneh, A. M., Qi, D., Yonesaki, T. & Otsuka, Y. An ADP-ribosyltransferase Alt of bacteriophage T4 negatively regulates the *Escherichia coli* MazF toxin of a toxin-antitoxin module. *Mol. Microbiol.* **99**, 188–198 (2016).
- Hazan, R. & Engelberg-Kulka, H. *Escherichia coli* mazEF-mediated cell death as a defense mechanism that inhibits the spread of phage P1. *Mol. Genet. Genomics* **272**, 227–234 (2004).
- Guler, P. et al. Arbitrium communication controls phage lysogeny through non-lethal modulation of a host toxin–antitoxin defense system. *Nat. Microbiol.* <https://doi.org/10.1038/s41564-023-01551-3> (2024).
- Yamaguchi, Y., Park, J.-H. & Inouye, M. Toxin-antitoxin systems in Bacteria and Archaea. *Annu. Rev. Genet.* **45**, 61–79 (2011).
- Brady, A. et al. Characterization of a unique repression system present in arbitrium phages of the SPbeta family. *Cell Host Microbe* <https://doi.org/10.1016/j.chom.2023.11.003> (2023).
- Duddy, O. P. & Bassler, B. L. Quorum sensing across bacterial and viral domains. *PLoS Pathog.* **17**, e1009074 (2021).
- Silpe, J. E. & Bassler, B. L. A host-produced quorum-sensing autoinducer controls a phage lysis-lysogeny decision. *Cell* **176**, 268–280.e13 (2019).
- Koo, B.-M. et al. Construction and analysis of two genome-scale deletion libraries for *Bacillus subtilis*. *Cell Syst.* **4**, 291–305.e7 (2017).
- Serrano, E. & Carrasco, B. Measurement of the length of the integrated donor DNA during *Bacillus subtilis* natural chromosomal transformation. *Bio Protoc.* **9**, e3338 (2019).
- Carniol, K., Ben-Yehuda, S., King, N. & Losick, R. Genetic dissection of the sporulation protein SpoIIE and its role in asymmetric division in *Bacillus subtilis*. *J. Bacteriol.* **187**, 3511–3520 (2005).
- Guérout-Fleury, A. M., Frandsen, N. & Stragier, P. Plasmids for ectopic integration in *Bacillus subtilis*. *Gene* **180**, 57–61 (1996).
- Harwood, C. R. & Cutting, S. M. *Molecular Biological Methods for Bacillus* (John Wiley, 1990).
- Sammito, M. et al. ARCIMBOLDO-LITE: single-workstation implementation and use. *Acta Crystallogr. D* **71**, 1921–1930 (2015).
- Potterton, L. et al. CCP 4 i 2: the new graphical user interface to the CCP 4 program suite. *Acta Crystallogr. D* **74**, 67–84 (2018).

32. Adams, P. D. et al. PHENIX: a comprehensive Python-based system for macromolecular structure solution. *Acta Crystallogr. D* **66**, 213–221 (2010).
33. McCoy, A. J. et al. Phaser crystallographic software. *J. Appl. Crystallogr.* **40**, 658–674 (2007).
34. Emsley, P. & Cowtan, K. Coot: model-building tools for molecular graphics. *Acta Crystallogr. D* **60**, 2126–2132 (2004).
35. Afonine, P. V. et al. Towards automated crystallographic structure refinement with phenix.refine. *Acta Crystallogr. D* **68**, 352–367 (2012).
36. Mirdita, M. et al. ColabFold: making protein folding accessible to all. *Nat. Methods* **19**, 679–682 (2022).
37. Holm, L. Dali server: structural unification of protein families. *Nucleic Acids Res.* **50**, W210–W215 (2022).
38. Micsonai, A. et al. BeStSel: webserver for secondary structure and fold prediction for protein CD spectroscopy. *Nucleic Acids Res.* **50**, W90–W98 (2022).
39. Gallego del Sol, F. & Marina, A. Structural basis of rap phosphatase inhibition by Phr peptides. *PLoS Biol.* **11**, e1001511 (2013).

Acknowledgements

We thank the IBV-CSIC Crystallogenes Facility for protein crystallization screenings. The structural results reported in this Article derive from measurements made at the synchrotron DLS (Didcot, UK), ALBA (Cerdanyola del Valles, Spain) and ESRF (Grenoble, France). Data collection experiments for the best crystals were carried out at XALOC and IO4 beamlines at ALBA and DLS Synchrotrons, respectively. X-ray diffraction data collection was supported by block allocation group (BAG) DLS Proposal MX28394, ALBA Proposal 2020074406 and ESRF proposal MX-2452. We acknowledge the ESRF, ALBA and DLS synchrotrons for provision of beam time and the beamline staff for assistance. Extended Data Figs. 4a and 8 were created with BioRender.com. This work was supported by grants PID2019-108541GB-I00 and PID2022-137201NB-I00 from the Spanish Government (Ministerio de Ciencia e Innovación), PROMETEO/2020/012 by the Valencian Government and the European Commission NextGenerationEU fund (EU 2020/2094), through CSIC's Global Health Platform (PTI Salud Global) to A.M., and grants MR/M003876/1, MR/V000772/1 and MR/S00940X/1 from the Medical Research Council (UK), BB/N002873/1, BB/V002376/1 and BB/S003835/1 from the Biotechnology and Biological Sciences Research Council (BBSRC, UK), ERC-ADG-2014 Proposal no. 670932 Dut-signal (from EU), and Wellcome Trust 201531/Z/16/Z to J.R.P. A.F.-R. received an FPU predoctoral fellowship from the Spanish Ministry of Universities, reference FPU19/00433.

Author contributions

J.R.P. and A.M. conceived the study; S.Z.-C., C.C., A.B., N.Q.-P., F.G.d.S., J.M.-B. and A.F.-R. performed the experiments; S.Z.-C., C.C., A.B., N.Q.-P., F.G.S., J.M.-B., A.F.-R., W.J.J.M., J.R.P. and A.M. processed data; J.R.P. and A.M. wrote the manuscript with inputs from all authors.

Competing interests

The authors declare no competing interests.

Additional information

Extended data is available for this paper at <https://doi.org/10.1038/s41564-023-01550-4>.

Supplementary information The online version contains supplementary material available at <https://doi.org/10.1038/s41564-023-01550-4>.

Correspondence and requests for materials should be addressed to José R. Penadés or Alberto Marina.

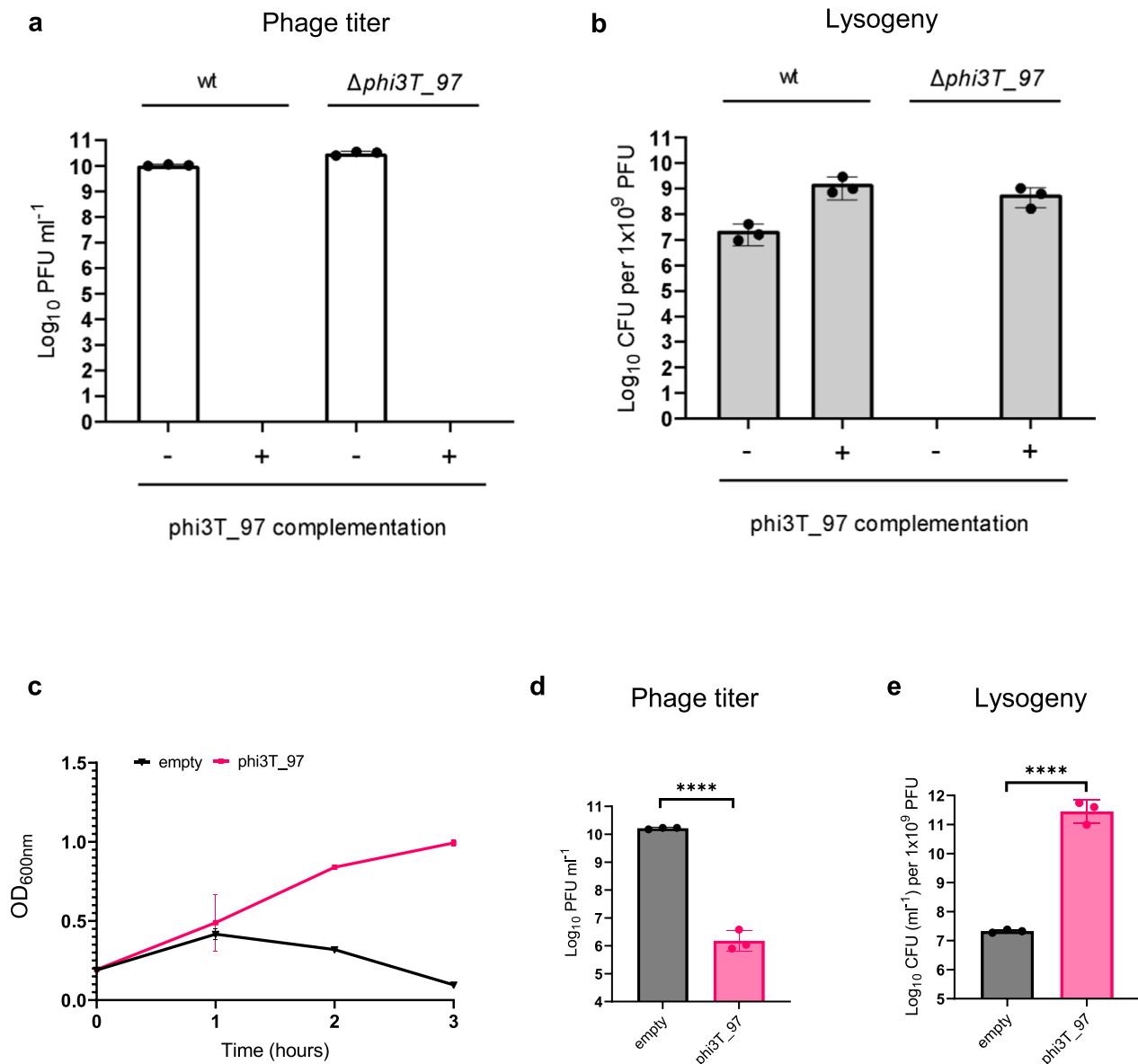
Peer review information *Nature Microbiology* thanks Aude Bernheim and the other, anonymous, reviewer(s) for their contribution to the peer review of this work.

Reprints and permissions information is available at www.nature.com/reprints.

Publisher's note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

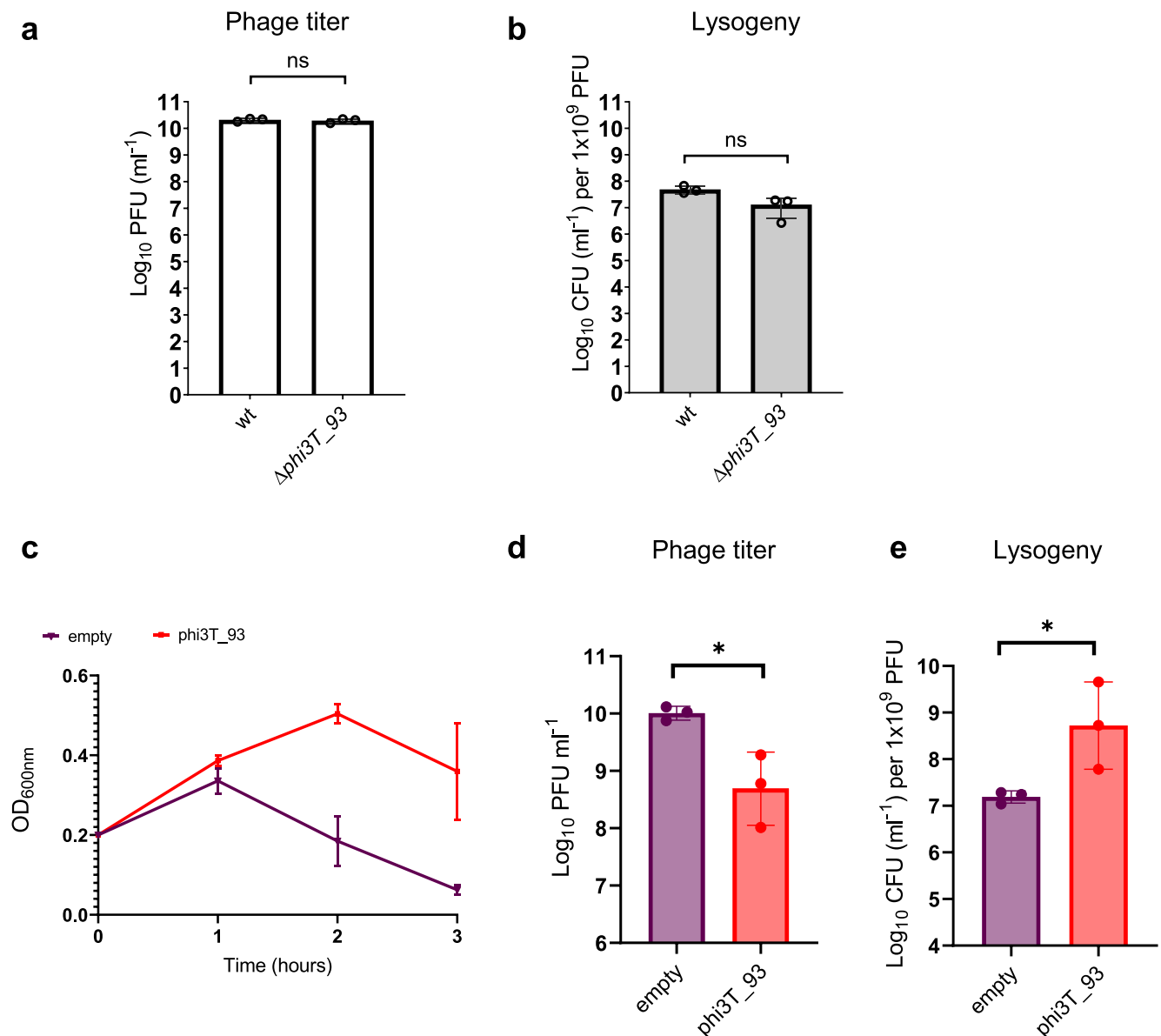
Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The images or other third party material in this article are included in the article's Creative Commons license, unless indicated otherwise in a credit line to the material. If material is not included in the article's Creative Commons license and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this license, visit <http://creativecommons.org/licenses/by/4.0/>.

© The Author(s) 2024

**Extended Data Fig. 1 | Characterisation of phi3T master repressor.**

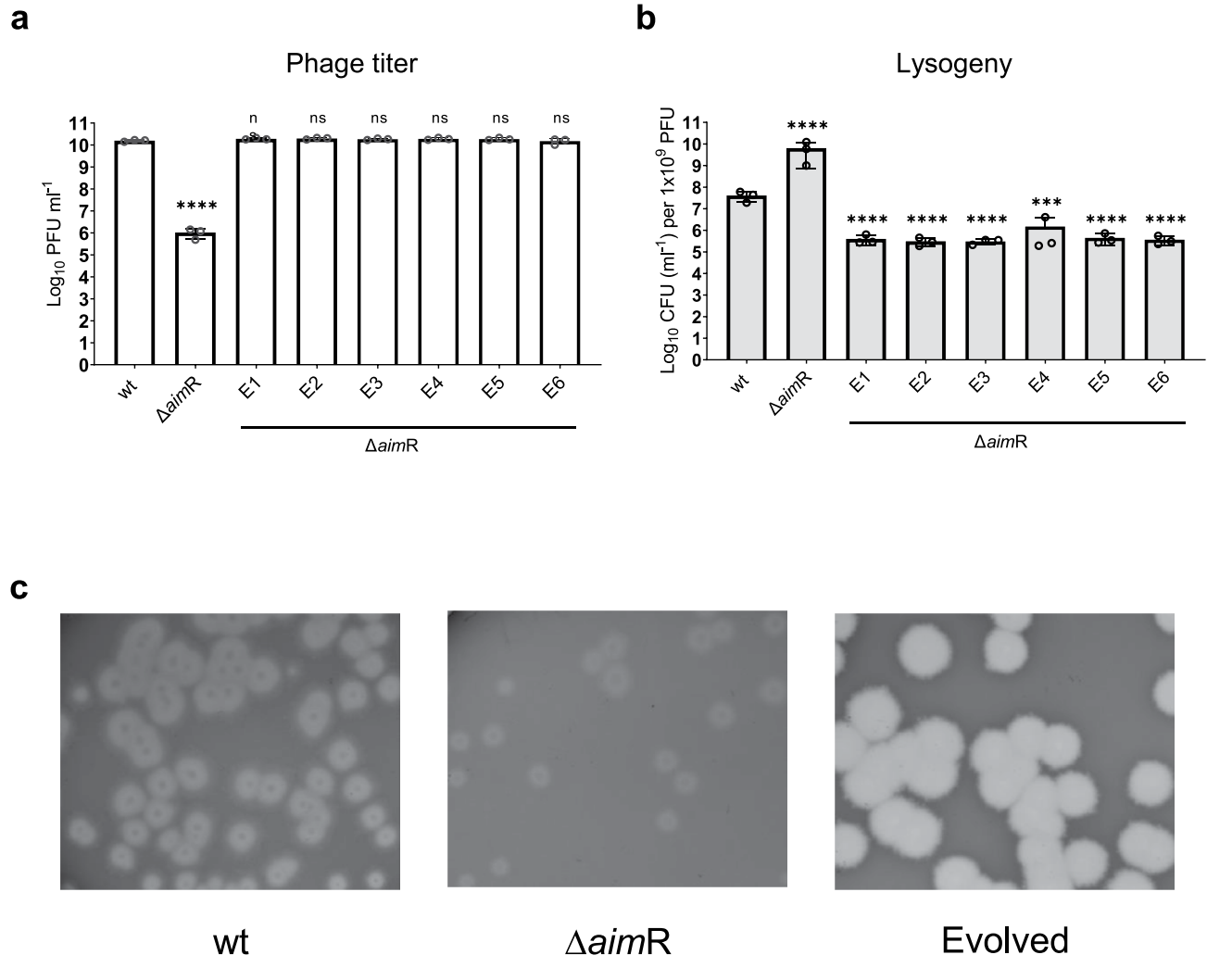
(a–b) Lysates containing phages phi3T wt or phi3T $\Delta phi3T_{97}$ were used to infect strains 168 $\Delta 6$ or 168 $\Delta 6 amyE::P_{spank}$ -phi3T₉₇, which overexpresses phi3T₉₇. The number of resulting plaques and lysogens were then quantified. The geometric mean and SD are presented (n = 3). (c) *B. subtilis* strains either devoid of (black) or expressing phi3T₉₇ (pink) were infected with phage phi3T wt at an MOI of 1:10 (phage:bacteria). The mean and SD of OD₆₀₀ readings (n = 3) are

presented. (d–e) After 3 h, the number of resulting plaques and relative number of lysogens were quantified. The geometric mean and SD are presented (n = 3). The results are shown as PFUs ml⁻¹ or as CFUs ml⁻¹ normalised by PFUs ml⁻¹ and represented as the CFU of an average phage titer (1x10⁹ PFU ml⁻¹). Values below detection limit (1x10¹ PFUs ml⁻¹ or CFUs ml⁻¹) are not marked. In panel (d) and (e) a two-tailed t-test was performed to compare mean differences; p values indicated were as follows: ****p < 0.0001.



Extended Data Fig. 2 | Characterisation of phi3T_93. (a-b) 168 $\Delta 6$ strains lysogenic for wt phi3T and phi3T mutant $\Delta phi3T_{93}$ were MC induced. The number of resulting phages were quantified using 168 $\Delta 6$ as recipient strain. The number of resulting plaques (PFUs ml⁻¹) and lysogens (CFUs ml⁻¹) are presented and geometric mean and SD (n = 3) are marked. A two-tailed t-test of log₁₀-transformed data was performed to compare mean differences. p values indicated are as follows: ns (not significant). (c) *B. subtilis* strains either devoid of (purple) or expressing phi3T_93 (pink) were infected with phage phi3T wt at an

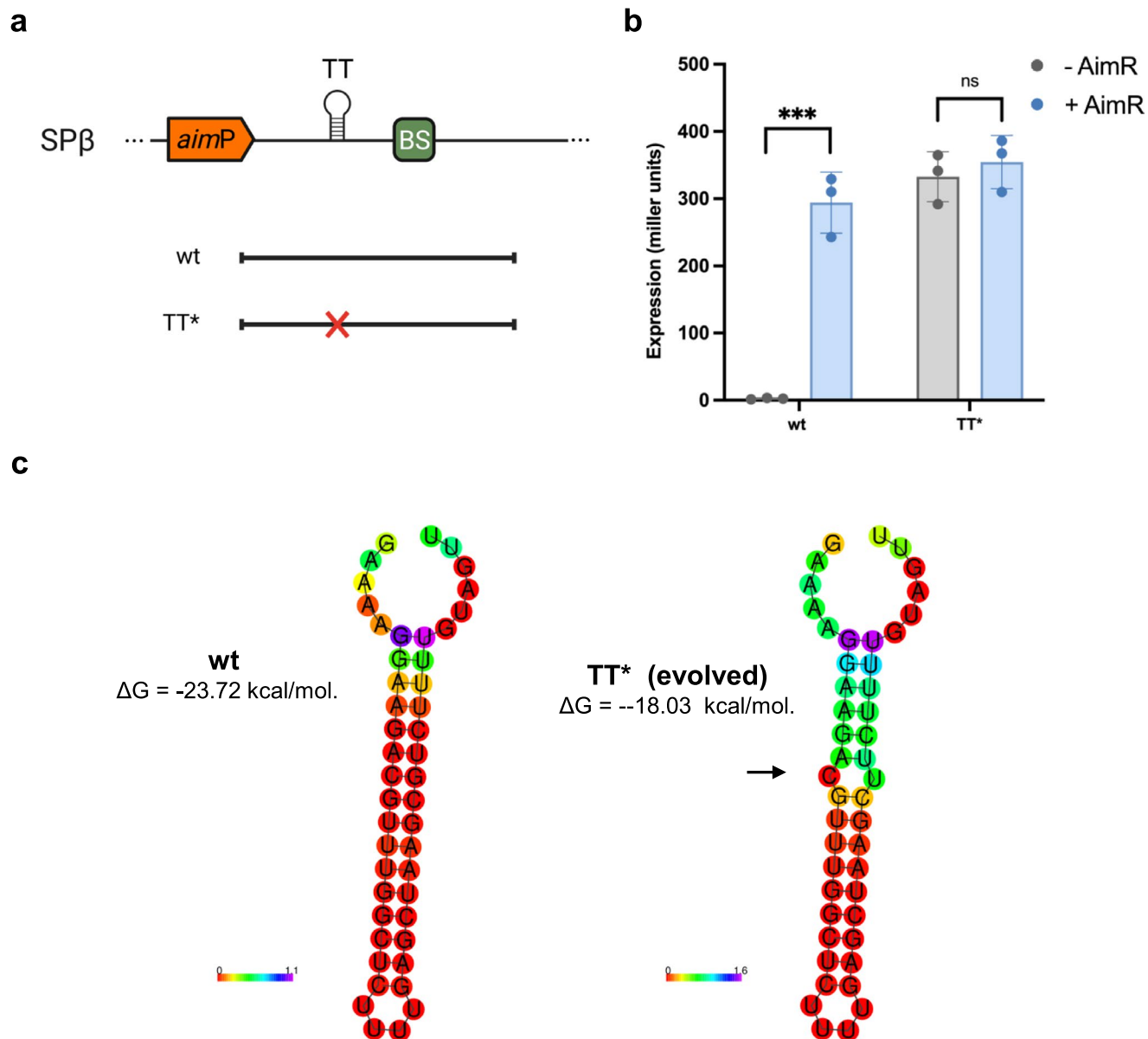
MOI of 1:10 (phage:bacteria). The mean and SD of OD₆₀₀ readings are presented (n = 3). (d-e) After 3 h the number of resulting plaques and relative number of lysogens were quantified. Geometric mean and SD are presented in panels (d) and (e). The results are shown as PFUs ml⁻¹ or as CFUs ml⁻¹ normalised by PFUs ml⁻¹ and represented as the CFU of an average phage titer (1 × 10⁹ PFU ml⁻¹). Two-tailed t-tests were performed to compare mean differences; p values indicated were as follows: *p < 0.05. Results presented in panels (c-e) are a subset of a more extended experiment, presented in full in Fig. 5.



phi3T

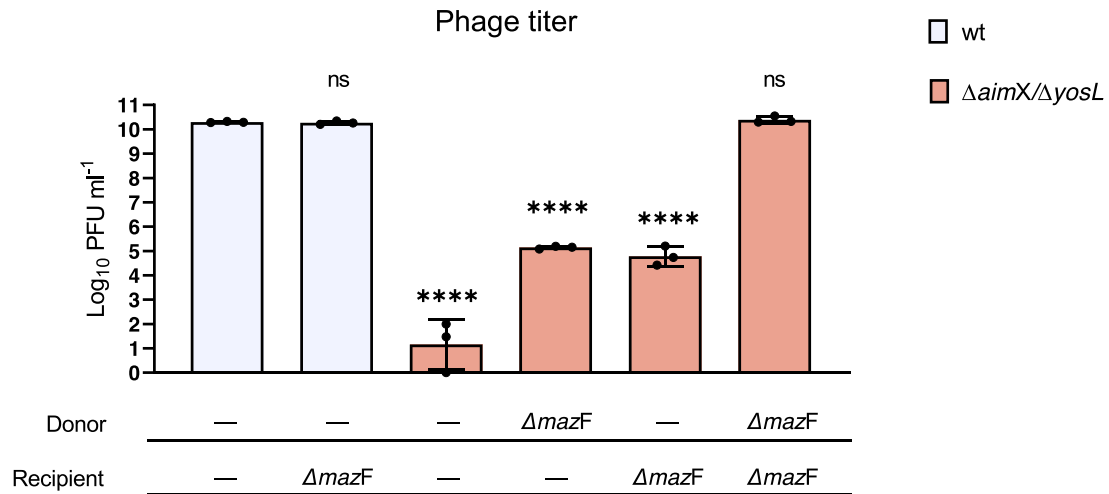
Extended Data Fig. 3 | Evolved phi3T ΔaimR phages have reduced lysogenisation. Strains lysogenic for phages phi3T wt, ΔaimR, and evolved ΔaimR phages were MC induced. The number of resulting plaques (**a**) and lysogens (**b**) were quantified using 168 Δ6 as the recipient strain. The results are represented as PFUs ml⁻¹ or CFUs ml⁻¹ normalised by PFUs ml⁻¹ and represented as the CFU of an average phage titer (1x10⁹ PFU ml⁻¹). Geometric mean and SD

are marked (n = 3). An ordinary one-way ANOVA of log₁₀-transformed data followed by Dunnett's multiple comparisons test was performed to compare mean differences. Comparisons were performed against wt phi3T infecting 168 Δ6. Adjusted p values were as follows: ***p < 0.001; ****p < 0.0001; ns (not significant). (**c**) The resulting plaque morphologies were photographed.



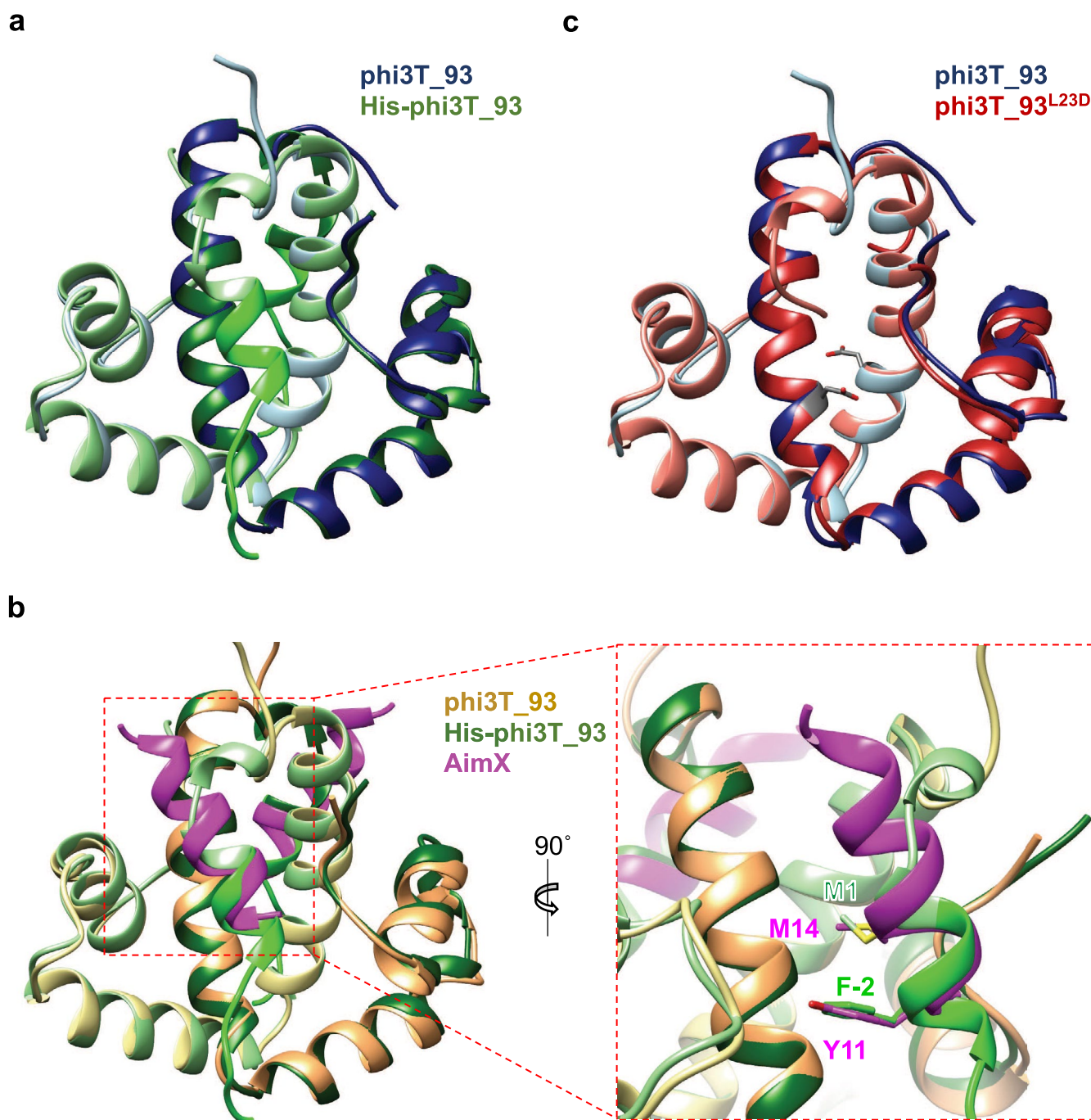
Extended Data Fig. 4 | AimR is an antiterminator in phage SP β . (a) Schematic view of the regions of SP β DNA cloned into the pDG1663+ P_{Sp β} reporter plasmid. TT*: mutated transcriptional terminator. (b) *B. subtilis* strains carrying pDR110 (empty or AimR^{SP β}) and the reporter plasmids were grown to OD₆₀₀ 0.2. IPTG was added for induction of the P_{Sp β} promoter in the pDR110 plasmid (empty or AimR). β -gal expression was measured using the Miller method. Data presents

the mean and SD (n = 3). Two-tailed t-tests were performed to compare mean differences; p values indicated were as follows: ***p = 0.0004; ns (not significant). replicates. (c) Predicted minimum free energy secondary structures of the terminator for the wt and TT* (evolved) sequence according to RNAfold. Colours are representative of positional entropy in the structure. Black arrow indicates the mutated bp.



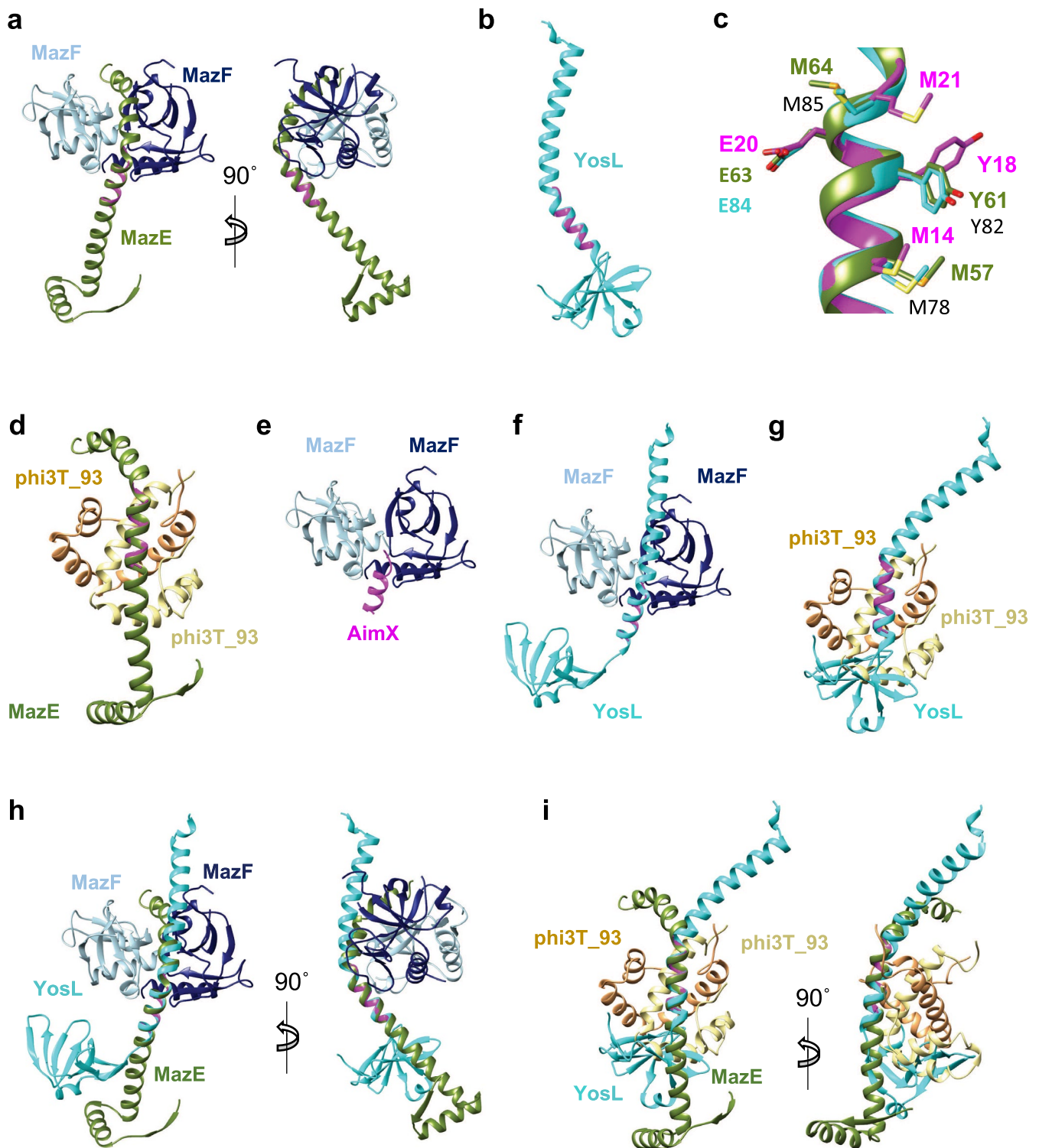
Extended Data Fig. 5 | When MazF is present in donor or recipient strains, there is a reduced number of plaques resulting from a phi3T *ΔaimX ΔyosL* mutant infection. Strains lysogenic for phi3T wt or phi3T mutant *ΔaimX/ΔyosL*, were MC induced from different donor strains: 168 $\Delta 6$ (indicated as -) and 168 $\Delta 6 \Delta mazF$. The number of resulting plaques was quantified using 168 $\Delta 6$ or 168 $\Delta 6 \Delta mazF$ mutants as recipient strains. Geometric mean and SD ($n = 3$) of the

number of resulting plaques (PFUs ml⁻¹) are presented. An ordinary one-way ANOVA of log₁₀-transformed data followed by a Dunnett's multiple comparisons test was performed to compare mean differences. Comparisons were performed against wt phi3T infecting 168 $\Delta 6$. Adjusted p values indicated are as follows: ****p < 0.0001; ns (not significant). Values below the detection limit of 10 PFUs ml⁻¹ were assigned value 1 and marked on the axis.



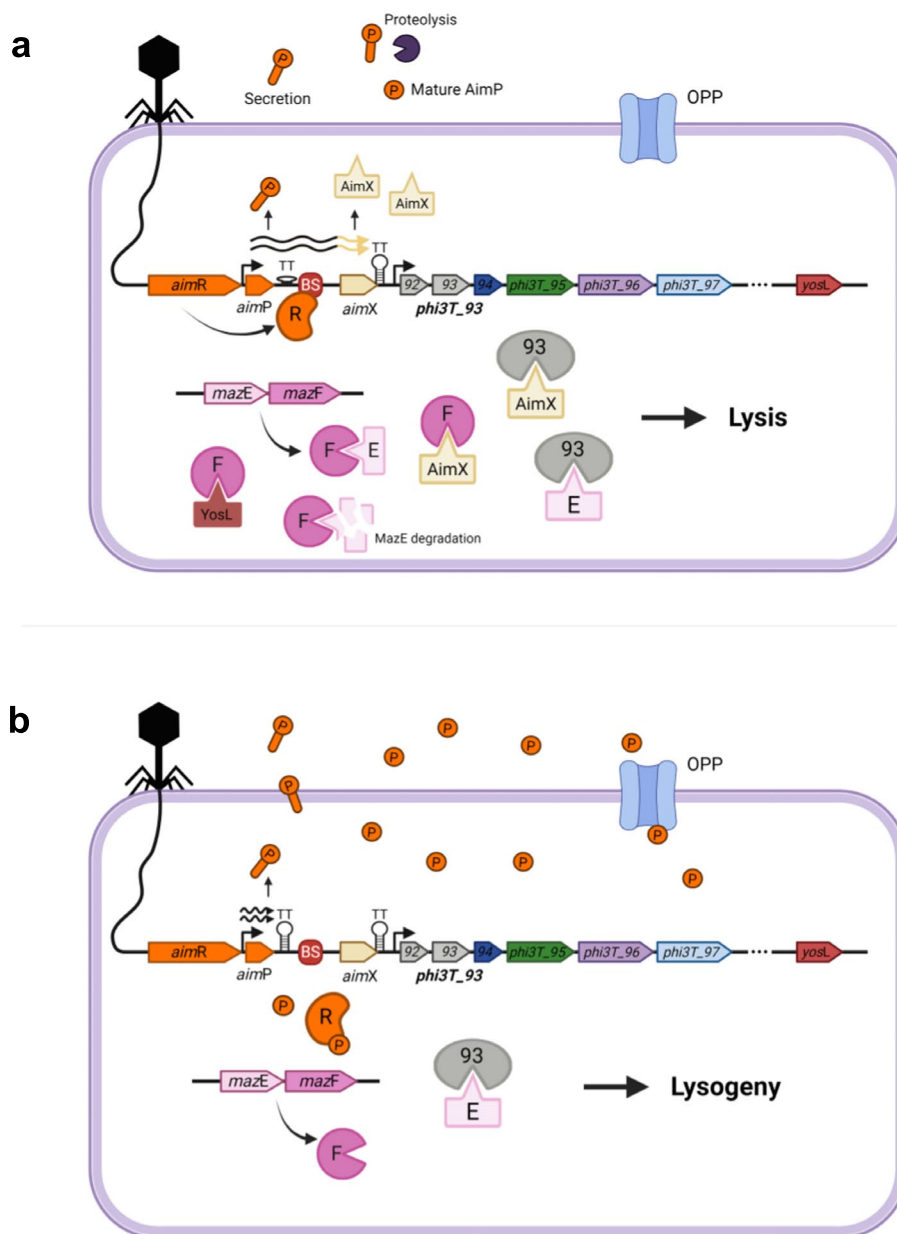
Extended Data Fig. 6 | phi3T_93 structures superimposition. Superimposition of the ribbon represented crystal structures of phi3T_93 (shades of blue) and His-phi3T_93 (shades of green) (a), phi3T_93, His-phi3T_93 and AimX-phi3T_93 complex (b) and phi3T_93 and phi3T_93^{L23D} (c). Note that phi3T_93 and His-phi3T_93 dimers are coloured in shades of blue and green respectively. Residues

belonging to the linker with the His-tag are coloured in fluorescent green for both monomers. AimX is coloured in magenta whereas phi3T_93 bound to AimX is coloured in shades of yellow. phi3T_93^{L23D} dimer is coloured in shades of red, whereas the mutated residue is displayed in dark grey.



Extended Data Fig. 7 | Experimental and *in silico* models for the net of mutually exclusive interactions. (a) Experimental complex (PDB 4ME7) of MazF (blue shades) and MazE (green) superimposed to the homologous region of AimX (magenta). (b) AlphaFold model for YosL^{phi3T} (cyan) superimposed to the homologous region of AimX (magenta). (c) Close view of the homologous region of AimX, MazE and YosL^{phi3T} showing the conserved residues involved in partner recognition in sticks and labelled. Models of the complexes (d) MazE-phi3T_93, (e) MazF-AimX, (f) YosL-MazF and (g) phi3T_93-YosL generated by the

superimposition of the homologous alpha helix. (h) Two orthogonal views of the superimposition of YosL to the MazF-MazE complex show no steric problems for YosL-MazF complex formation. (i) Two orthogonal views for the superimposition of potential phi3T_93-YosL and phi3T_93-MazE complexes revealing steric hindrances for the N-terminal domain of YosL with phi3T_93 dimer but not for MazE. Note that in all the models AimX is depicted in magenta to show the relative position in the complex against the rest of the components.



Extended Data Fig. 8 | Model for the arbitrium system. (a) In early stages of the infection process or after induction of the SOS response, when the concentration of AimP is low, AimR binds to its cognate DNA-binding site (BS), acting as an antiterminator that allows the expression of *aimX* as part of a bicistronic operon starting from the *aimP* promoter. The AimX peptide binds to the toxin MazF and to *phi3T_93*, preventing its binding to the antitoxin MazE, freeing MazE to bind to MazF. Additionally, YosL binds to MazF. All these interactions result in blocking

the function of MazF and promote the phage lytic cycle. **(b)** During late stages of the infection, AimP accumulates inside the bacteria and binds to AimR, inhibiting its function as an antiterminator. When this occurs, the mRNA transcript that starts at *aimP* will stop at the transcriptional terminator (TT) localised upstream *aimX*, preventing its expression. In absence of AimX, *phi3T_93* will bind to MazE, liberating MazF whose activity promotes lysogeny by an unknown mechanism. Model created with BioRender.com.

Extended Data Table 1 | BLI calculated binding affinities

Immobilised protein	Protein in solution	K _D (nM)	k _{on} (M ⁻¹ s ⁻¹)	k _{off} (s ⁻¹)
AimX	phi3T_97	NBD ^a	NBD	NBD
AimX	phi3T_93	184 ± 10	2.7 ± 0.8 × 10 ⁵	4.5 ± 0.9 × 10 ⁻²
AimX	MazF	81.3 ± 5	8.0 ± 0.4 × 10 ⁴	6.6 ± 3.4 × 10 ⁻³
AimX ^{M14D}	phi3T_93	NBD	NBD	NBD
AimX ^{M14D}	MazF	NBD	NBD	NBD
AimX	phi3T_93 ^{L23D}	NBD	NBD	NBD
MazE	MazF	404 ± 34	3.6 ± 2.6 × 10 ⁴	1.0 ± 0.2 × 10 ⁻²
MazE	phi3T_93	27.3 ± 21	7.0 ± 4.4 × 10 ⁴	1.3 ± 1.0 × 10 ⁻³
MazE	phi3T_93 ^{L23D}	3770 ± 150	2.6 ± 1.8 × 10 ³	8.3 ± 3.1 × 10 ⁻³
MazF	YosL ^{SPβ}	15 ± 2	5.3 ± 0.5 × 10 ⁴	8.5 ± 0.1 × 10 ⁻⁴
MazF	YosL ^{phi3T}	20.2 ± 5	1.6 ± 0.1 × 10 ⁴	3.3 ± 0.7 × 10 ⁻⁴
YosL ^{SPβ}	phi3T_93	NBD	NBD	NBD
YosL ^{phi3T}	phi3T_93	NBD	NBD	NBD

^aNBD: No binding detected
Results are given as mean ± SE.

Extended Data Table 2 | Data collection and refinement statistics

	His-phi3T	phi3T_93	AimX-phi3T_93	phi3T_93 ^{L23D}
Data collection				
Space group	P2 ₁	I2	P3 ₂ 21	P321
Cell dimensions				
<i>a</i> , <i>b</i> , <i>c</i> (Å)	37.5, 50.8, 55.8	46.5, 49.9, 70.5	66.4, 66.4, 85.1	71.9, 71.9, 68.3
α , β , γ (°)	90, 102.0, 90	90, 107.7, 90	90, 90, 120	90, 90, 120
Resolution (Å)	50.8-1.9	40.1-2.3	57.6-2.2	62.7-2.2
	(1.94-1.9)	(2.44-2.3)	(2.32-2.2)	(2.28-2.2)
<i>R</i> _{merge}	0.08 (0.38)	0.10 (0.21)	0.03 (0.48)	0.12 (1.12)
<i>I</i> / σ <i>I</i>	8.0 (1.7)	8.5 (4.4)	7.4 (1.5)	15 (3.1)
Completeness (%)	99.7 (98.8)	97.1 (96)	99.7 (93.2)	100 (100)
Redundancy	6.0 (5.2)	4.9 (5.1)	18.5 (18.4)	16.7 (14.4)
Refinement				
Resolution (Å)	1.9	2.3	2.2	2.2
No. reflections	16124	13240	11412	10705
<i>R</i> _{work} / <i>R</i> _{free}	0.21/0.25	0.21/0.26	0.22/0.23	0.21/0.24
No. atoms	1419	1173	1370	1157
Protein	1375	1134	1330	1128
Ligand/ion	-	2	2	1
Water	44	37	38	28
<i>B</i> -factors				
Protein	50.10	38.4	63.99	62.3
Ligand/ion	-	45	63.21	70.9
Water	52.20	31	63.05	62.1
R.m.s. deviations				
Bond lengths (Å)	0.017	0.010	0.009	0.008
Bond angles (°)	1.63	1.16	1.02	0.89
PDB ID	8ANU	8ANT	8ANV	8C8E

Each data set was collected from a single crystal. Values in parentheses correspond to the highest resolution shell.

Reporting Summary

Nature Portfolio wishes to improve the reproducibility of the work that we publish. This form provides structure for consistency and transparency in reporting. For further information on Nature Portfolio policies, see our [Editorial Policies](#) and the [Editorial Policy Checklist](#).

Statistics

For all statistical analyses, confirm that the following items are present in the figure legend, table legend, main text, or Methods section.

- | n/a | Confirmed |
|-------------------------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| <input type="checkbox"/> | <input checked="" type="checkbox"/> The exact sample size (n) for each experimental group/condition, given as a discrete number and unit of measurement |
| <input type="checkbox"/> | <input checked="" type="checkbox"/> A statement on whether measurements were taken from distinct samples or whether the same sample was measured repeatedly |
| <input type="checkbox"/> | <input checked="" type="checkbox"/> The statistical test(s) used AND whether they are one- or two-sided
<i>Only common tests should be described solely by name; describe more complex techniques in the Methods section.</i> |
| <input type="checkbox"/> | <input checked="" type="checkbox"/> A description of all covariates tested |
| <input type="checkbox"/> | <input checked="" type="checkbox"/> A description of any assumptions or corrections, such as tests of normality and adjustment for multiple comparisons |
| <input type="checkbox"/> | <input checked="" type="checkbox"/> A full description of the statistical parameters including central tendency (e.g. means) or other basic estimates (e.g. regression coefficient) AND variation (e.g. standard deviation) or associated estimates of uncertainty (e.g. confidence intervals) |
| <input type="checkbox"/> | <input checked="" type="checkbox"/> For null hypothesis testing, the test statistic (e.g. F , t , r) with confidence intervals, effect sizes, degrees of freedom and P value noted
<i>Give P values as exact values whenever suitable.</i> |
| <input checked="" type="checkbox"/> | <input type="checkbox"/> For Bayesian analysis, information on the choice of priors and Markov chain Monte Carlo settings |
| <input checked="" type="checkbox"/> | <input type="checkbox"/> For hierarchical and complex designs, identification of the appropriate level for tests and full reporting of outcomes |
| <input checked="" type="checkbox"/> | <input type="checkbox"/> Estimates of effect sizes (e.g. Cohen's d , Pearson's r), indicating how they were calculated |

Our web collection on [statistics for biologists](#) contains articles on many of the points above.

Software and code

Policy information about [availability of computer code](#)

Data collection	Synchrotron Alba (Cerdanyola del Vallès, Spain), Diamond Light source (Didcot, UK) and ESRF (Grenoble, France) data collection software provided by the beamline
Data analysis	<p>Size exclusion chromatogram figures and Statistical Analyses were carried out with the scientific software GraphPad Prism 9. Open source software (https://www.graphpad.com/scientific-software/prism/)</p> <p>Crystallographic processing was carried out with programs from CCP4 (http://www.ccp4.ac.uk/), CCP4i2 (http://ccp4i2.fg.oisin.rc-harwell.ac.uk) and Phenix (https://www.phenix-online.org/), free and open source suites.</p> <p>Model building was carried out with Coot 0.9.8.8 (https://www2.mrc-lmb.cam.ac.uk/personal/pemsley/coot/).</p> <p>Size Exclusion Chromatography with Multi-Angle Light Scattering (SEC-MALS) data acquisition and analysis was carried out with Astra 7.1.2 software (https://www.wyatt.com/products/software/astra.html) from Wyatt.</p> <p>Structure prediction was carried out with AlphaFold2 run online in Google ColabFold v1.5.2 (https://colab.research.google.com/github/sokrypton/ColabFold/blob/main/beta/AlphaFold2_advanced.ipynb#scrollTo=pc5-mbsX9PZC).</p> <p>Proteins with structural similarity were identified with DALI server (http://ekhidna2.biocenter.helsinki.fi/dali/).</p> <p>Protein sequence homology search was carried out with BLASTp suite at the NCBI BLAST homepage (https://blast.ncbi.nlm.nih.gov).</p>

Biolayer Interferometry (BLI) data acquisition and analysis was carried out with BLItz Pro 1.2 software (www.fortebio.com) from Fortebio

Secondary structure determination and fold recognition from protein circular dichroism spectra was carried out with BestSel (<https://bestsel.elte.hu/index.php>)

Figures for three-dimensional structures were generated with Pymol 2.1 (<https://pymol.org/2/>) or Chimera v1.15 (<https://www.cgl.ucsf.edu/chimera/>).

Some figures in this manuscript have been created with Biorender ([Biorender.com](https://biorender.com))

All this information has been included in the manuscript.

For manuscripts utilizing custom algorithms or software that are central to the research but not yet described in published literature, software must be made available to editors and reviewers. We strongly encourage code deposition in a community repository (e.g. GitHub). See the Nature Portfolio [guidelines for submitting code & software](#) for further information.

Data

Policy information about [availability of data](#)

All manuscripts must include a [data availability statement](#). This statement should provide the following information, where applicable:

- Accession codes, unique identifiers, or web links for publicly available datasets
- A description of any restrictions on data availability
- For clinical datasets or third party data, please ensure that the statement adheres to our [policy](#)

The atomic coordinates of the phi3T_93, His-phi3T_93, phi3T_93L23D and AimX-phi3T_93 complex has been deposited with PDB codes 8ANT, 8ANU, 8C8E and 8ANV, respectively, at <http://www.pdb.org>. The previously determined structures used in this study are available from the PDB (<http://www.pdb.org>) under the accession codes indicated. The rest of the data are available in the main text, supplementary materials and auxiliary files. Plasmids and bacterial strains generated during this work are listed in Supplementary Tables S3 and S4 and are available upon request. Correspondence and requests for materials can be addressed to A.M. (amarina@ibv.csic.es) or J.R.P. (j.penades@imperial.ac.uk). Source data are provided with this paper.

Research involving human participants, their data, or biological material

Policy information about studies with [human participants or human data](#). See also policy information about [sex, gender \(identity/presentation\), and sexual orientation](#) and [race, ethnicity and racism](#).

Reporting on sex and gender

Reporting on race, ethnicity, or other socially relevant groupings

Population characteristics

Recruitment

Ethics oversight

Note that full information on the approval of the study protocol must also be provided in the manuscript.

Field-specific reporting

Please select the one below that is the best fit for your research. If you are not sure, read the appropriate sections before making your selection.

Life sciences Behavioural & social sciences Ecological, evolutionary & environmental sciences

For a reference copy of the document with all sections, see [nature.com/documents/nr-reporting-summary-flat.pdf](https://www.nature.com/documents/nr-reporting-summary-flat.pdf)

Life sciences study design

All studies must disclose on these points even when the disclosure is negative.

Sample size

Data exclusions

Replication

Binding affinities were measured by BLI three times for each sample. No discrepancies superior of 10 % were observed between samples. SEC-MALS assays were carried two times for each sample. No discrepancies superior of 5 % were observed between samples.

Randomization Not relevant for our study. Samples not allocated in sample groups. Independent bacterial cultures for each bacterial strain were cultured.

Blinding The results reported in this study are not related to a clinical research study, thus blinding was not necessary

Reporting for specific materials, systems and methods

We require information from authors about some types of materials, experimental systems and methods used in many studies. Here, indicate whether each material, system or method listed is relevant to your study. If you are not sure if a list item applies to your research, read the appropriate section before selecting a response.

Materials & experimental systems

n/a	Involvement in the study
<input checked="" type="checkbox"/>	<input type="checkbox"/> Antibodies
<input checked="" type="checkbox"/>	<input type="checkbox"/> Eukaryotic cell lines
<input checked="" type="checkbox"/>	<input type="checkbox"/> Palaeontology and archaeology
<input checked="" type="checkbox"/>	<input type="checkbox"/> Animals and other organisms
<input checked="" type="checkbox"/>	<input type="checkbox"/> Clinical data
<input checked="" type="checkbox"/>	<input type="checkbox"/> Dual use research of concern
<input checked="" type="checkbox"/>	<input type="checkbox"/> Plants

Methods

n/a	Involvement in the study
<input checked="" type="checkbox"/>	<input type="checkbox"/> ChIP-seq
<input checked="" type="checkbox"/>	<input type="checkbox"/> Flow cytometry
<input checked="" type="checkbox"/>	<input type="checkbox"/> MRI-based neuroimaging