A functional selection reveals previously undetected anti-phage defence systems in the E. coli pangenome

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The ancient, ongoing coevolutionary battle between bacteria and their viruses, bacteriophages, has given rise to sophisticated immune systems including restriction-modification and CRISPR-Cas. Many additional anti-phage systems have been identified using computational approaches based on genomic co-location within defence islands, but these screens may not be exhaustive. Here we developed an experimental selection scheme agnostic to genomic context to identify defence systems in 71 diverse E. coli strains. Our results unveil 21 conserved defence systems, none of which were previously detected as enriched in defence islands. Additionally, our work indicates that intact prophages and mobile genetic elements are primary reservoirs and distributors of defence systems in E. coli, with defence systems typically carried in specific locations or hotspots. These hotspots encode dozens of additional uncharacterized defence system candidates. Our findings reveal an extended landscape of antiviral immunity in E. coli and provide an approach for mapping defence systems in other species.

Bacteriophages (or simply, phages) are an extraordinarily diverse and ubiquitous class of viruses that pose a nearly constant threat to bacteria. Phages are the most abundant biological entity on the planet, with estimates of $10^{31}$ particles that drive the daily turnover of ~20% of all bacteria in some environments. Bacteria and their viral predators are locked in a perpetual coevolutionary battle, leading to the emergence of sophisticated mechanisms by which phage manipulate and exploit their hosts, and an equally diverse set of bacterial immune mechanisms collectively referred to as anti-phage defence systems. These immunity systems include both innate mechanisms, such as restriction-modification systems, and adaptive mechanisms, such as CRISPR-Cas. Recent studies have begun to identify many new defence systems, but the full inventory probably extends well beyond what is currently defined.

Identifying additional anti-phage defence systems promises to provide new insight into the ancient coevolutionary conflict between viruses and their hosts. Recent work has found that many defence systems have homologues with similar function in eukaryotic innate immunity, indicating a potentially ancient origin and cross-kingdom conservation of many immune systems. Additionally, previous studies of anti-phage defence have produced precision molecular tools such as CRISPR and restriction enzymes, so the discovery of new immune mechanisms may enable new tools for manipulating cells and genomes. Finally, there is growing interest in using phages to treat antibiotic-resistant bacterial infections and to manipulate microbiomes. A more complete understanding of the diverse mechanisms by which bacteria defend themselves may be critical for these endeavours.

Multiple groups have previously used computational methods to identify uncharacterized defence systems on the basis of the observation that anti-phage defence systems often cluster in bacterial genomes in high density, forming so-called ‘defence islands’. However, not all defence systems may be detectable in these guilt-by-association approaches. For systems that are rare or not widely conserved, it may be difficult to detect enrichment within defence islands, and not all defence systems necessarily associate with defence islands. Additionally, candidates identified computationally must be expressed in a model laboratory organism and then tested against a panel of phages. Some systems may not work in a heterologous host or protect against the phages examined, and demonstrating that a given system provides defence in its native context is typically not tested or even possible. We reasoned that an experimental selection scheme to uncover antiviral proteins may reveal new insights into bacterial immunity, including identifying defence systems that remain uncharacterized and revealing the relative frequency of the different genomic contexts of these bacterial immune systems (that is, defence islands, mobile genetic elements (MGEs) or other contexts). To this end, we took a functional metagenomic approach to map the range of defence systems in the Escherichia coli pangenome (Fig. 1). In addition to revealing previously uncharacterized phage defence systems in E. coli, our work also provides a robust screening methodology that can now be adapted to systematically identify phage defence mechanisms in virtually any bacterial genome or metagenomic sample.

Results

Identification of novel anti-phage defence systems. To sample the immune landscape of E. coli, we collected a diverse set of wild isolate strains from the ECOR collection as well as 19 clinical isolates (71 strains in total), all with available draft genome sequences (Extended Data Fig. 1). The ECOR collection is a set of strains curated to span the phylogenetic diversity of the species. Together, the 71 strains collected encode 21,149 unique gene clusters, of which >10,000 exist in only one or two strains (Extended Data Fig. 1). From genomic DNA, we constructed a 100x-coverage library...
of fosmids, each harbouring a ~40 kb genome fragment, in EPI300, a derivative of E. coli K12. We used large-insert fosmids to minimize the size of the library, to include potentially large defence systems, and because the copy number is maintained around one, minimizing false positives due to overexpression.

Many anti-phage defence systems work by an abortive infection (Abi) mechanism in which an infected cell sacrifices its viability to prevent phage replication and thereby protect uninfected cells in a population16. Thus, it is impossible to directly select for clones containing an Abi-based defence system because the infected cell dies. Instead, we used a selection strategy, historically known as the ‘tab’ (T4 abortive) method17. In brief, cells harbouring the gDNA library are mixed with phage in a structured medium (soft agar) at varying concentrations of phage (Fig. 2a). At intermediate phage concentrations, individual clones from the library can grow and form small populations before encountering a phage particle. Any such micro-colonies that harbour an Abi defence system can be infected, but the initially infected cell will die without producing progeny phage, enabling the rest of the population to survive and produce a colony. Thus, our screening approach allows the identification of both Abi and conventional defence systems.

Using this general strategy, we challenged cells harbouring the fosmid library with three lytic phages: T4, λ, and T7, each representing a major class of Caudovirales, the tailed bacteriophages (Fig. 1). From each selection, we isolated approximately 90 surviving colonies and then sequenced the ends of the vector insert in each clone to identify the genomic region and strain of origin of each fragment. For each positive clone, we then measured the efficiency of plaquing (EOP)—the ratio of plaques formed by a phage plated on the positive clone to plaques formed by a phage-sensitive control strain (Fig. 2b). In some cases, these EOP assays revealed nearly complete protection even at phage titres of >10^9 p.f.u. ml^-1. Upon initial sequencing, representative clones with this phenotype consistently encoded lipopolysaccharide (LPS) or capsule biosynthesis genes. This phenotype is consistent with a loss of adsorption that results from modification of the core surface properties of the cell (Fig. 2b and Extended Data Fig. 2a). Other positive clones produced protection accompanied by a high rate of escape plaques, a phenotype seen with many restriction-modification (RM) systems (Fig. 2b). Escape plaques can arise with many defence systems but are well known to arise at high frequencies for RM systems due to epigenetic escape18. Upon sequencing, the clones exhibiting high-frequency escape plaques indeed encoded RM systems. Thus, clones with these two phenotypes were excluded from further analyses (Methods). From 257 initial clones, 117 and 9 were eliminated as probably resulting from changes in cell surface properties affecting adsorption or RM-mediated defence, respectively. After also accounting for redundancy in the remaining 131 clones, we had 43 clones that we hypothesized to encode novel defence systems.

To pinpoint possible defence systems, we pooled the three sets of remaining fosmids corresponding to each phage and generated random 6–12 kb fragments. These fragments were then sub-cloned into a low-copy plasmid vector to create three high-coverage sub-libraries. Cells harbouring these high-coverage sub-libraries were then selected for resistance to their respective phage, and plasmids from positive clones were sequenced by Nanopore long-read sequencing. The sub-library clones that survived selection each contained the originally selected defence system, flanked by random lengths of adjacent DNA from the original fosmid insert. Thus, when reads were mapped back to the genome fragment in a given fosmid, the coverage maxima typically delineated the boundaries of each candidate defence system (Fig. 2c). In some cases, this identified previously characterized systems, including type III and IV RM systems and an Old-family endonuclease, or non-defence genes that account for phage resistance, such as malI, a regulator of the lambda receptor LamB (Supplementary Table 1). After discarding such cases, there were 21 unique candidate defence systems, with 10, 6 and 5 systems from the selections for T4, λ, and T7, respectively.
Validation of candidate defence systems. To validate these novel defence systems, we cloned each candidate open reading frame (ORF) or operon into a low-copy vector under the control of its native promoter in wild-type MG1655 (Supplementary Table 2). We confirmed that each system did not affect phage adsorption (Extended Data Fig. 2b) and then challenged each with a panel of 10 diverse phages (Fig. 3a). Each candidate system was confirmed to substantially reduce the EOP for the phage originally used to select the system, and often others. In some cases, a given defence system did not change the EOP of a phage but instead produced smaller plaques. Although most defence systems were relatively specific, protecting against only a few phages, some systems provided relatively broad protection, such as PD-\(\lambda\)-5, which affected EOP or plaque size for all but one of the phages tested. The 10 systems selected to defend against T4 also generally protected against the other, related T-even phages, T2 and T6. Most systems protected most strongly against the phage originally used to select it, but with some exceptions. For instance, PD-\(\lambda\)-5, PD-T7-1 and PD-T7-3 protected more strongly against the T-even phages than against the \(\lambda\) used to identify them. The fact that these systems provided robust defence against T4 but were not identified in the T4 selection indicates that our screen was not saturating and that the systems identified represent only a subset of the defence systems in the original T4 E. coli isolates used.

We then sought to classify whether each system functioned via Abi or provided direct immunity such that infected cells could survive an infection. Because Abi systems require killing of infected cells, these systems typically only provide defence at a low multiplicity of infection (MOI) in which bacteria outnumber phages, whereas direct immunity provides defence to the infected cell and thus allows comparable, although not identical, rates of growth at MOIs above and below 1.0. We thus tested the growth of strains harbouring each defence system infected at an MOI of 0.05 and 5. Of the 21 systems tested, 9 provided direct inhibition of phage infection, producing comparable protection at both MOIs; the other 12 probably use an Abi mechanism of protection with stronger protection at the lower MOI (Fig. 3b and Extended Data 3a). These results validate the ability of our screening strategy to detect Abi defences and underscore the notion that Abi systems contribute considerably to E. coli immunity.

One advantage to our screen is that we have the strains from which these defence systems originated, in contrast to computational screens that have identified many defence systems from species that have sequenced genomes but are not readily available.
We were thus able to delete candidate systems in their originating strains and test whether, in their native context, they protect against phage. Specifically, we tested the native role of PD-T4-2 and PD-T4-9, which originate from ECOR65 and ECOR22, respectively, strains to which T4 can adsorb but not infect. Deleting each system dramatically increased the plaquing efficiency of T4, demonstrating that these systems provide defence in both their original, native context and when introduced into E. coli K12 (Fig. 3c). We also asked whether our defence systems work in E. coli strains other than MG1655 by testing four candidate defence systems in ECOR13 and E. coli C, which are natively susceptible to the three phases. All four systems provided protection, indicating that the function of the systems identified is not strictly dependent on strain background (Extended Data Fig. 3b).

In total, 26 of 32 proteins in the 21 systems identified here were annotated in GenBank as either ‘hypothetical protein’ or as containing domains of unknown function and had no primary sequence homology to any characterized anti-phage defence system. To more sensitively characterize each protein, we used HHpred to detect even remote similarity to PFAM domains\(^9\). This did not reveal any homology to a known system, and in the majority of cases, most of each protein remained uncharacterized. We were, however, able to detect potential similarity to some motifs or domains characteristic of defence systems, for example, nucleic-acid binding or cleavage domains (Fig. 3a and Supplementary Table 3). In most cases, this was limited to small regions hinting at enzymatic function, but not a mechanism of activation or specific targets.

Remote homology detection revealed several intriguing features uncharacteristic of known defence systems. These included (1) similarity to a ribosome-dependent ribonuclease (ReIle) in conjunction with a phage-sheath-like domain, (2) a three-gene operon encoding an exotoxin A-like domain known to participate in bacterial virulence although not phage defence, an unknown protein and a SecB-like chaperone, (3) a putative membrane-anchored protein with a central coiled-coil domain (DUF4041) and a C-terminal DNA binding/cleavage domain, (4) a beta propeller fused to a DNA binding/cleavage domain, (5) a P4 phage beta-like protein, (6) a lipoprotein, (7) a zinc-finger-like domain fused with a C-terminal domain of unknown function that belongs to an extended family of ribonucleases, (8) a CoiA domain and (9) DUF6575.

Several defence systems showed similarity to domains that are less frequently associated with defence systems compared with domains such as nucleases and helicases. These less frequent domains included a peptidase, a eukaryotic-like Ser/Thr kinase\(^9\), a NAD\(^+\)-binding Sir2 homologue\(^9\), a GIY-YIG nuclease\(^9\) and a putative DNase\(^9\). Four of the systems identified contained a component with similarity to a toxin-antitoxin (TA) system, but either none of these systems were found in existing TA databases or they encoded additional uncharacterized components aside from simply toxin and antitoxin. Collectively, our results reveal a striking diversity of proteins involved in bacterial defence and highlight the vast, unexplored landscape of antiviral immunity in bacteria.

In total, 11 defence systems had a component suggestive of DNA binding or cleavage activity (often including distant similarity to the restriction endonuclease-like fold PD(D/E)XK), although this similarity was, as noted above, typically restricted to a small region or motif. These putative DNases are unlikely to be part of restriction-modification systems as only one included a predicted methylase. Notably, 7 of 11 of these provided direct immunity (not Abi), suggesting non-self, nucleic-acid-targeting activity in potentially novel ways. Remote similarity to HEPN motifs or domains were found in 6 of the 21 systems. These domains are also present in the ribonucleases associated with CRISPR-Cas and toxin-antitoxin systems, supporting the notion that they are common, versatile components of defence systems\(^9\). In four cases, we mutagenized key conserved residues in the predicted domains and found that they were essential to phage protection, suggesting that the remote domain predictions feature in the function of these systems (Fig. 3d).

To assess the conservation of the 21 systems identified, we investigated the phylogenetic distribution of each (Fig. 3e). Homologues of each system (that is, those that encode all components) were found in other gamma-proteobacteria, with 16 and 18 having homologues in alpha- and beta-proteobacteria, respectively. More than half were also abundant in Firmicutes, Actinobacteria, Bacteroidetes and Spirochaetia, suggesting that many of the systems represent new, widespread classes of anti-phage defence systems. Seven of the systems we identified were restricted to proteobacteria, and three were exclusively found in \(\gamma\)-proteobacteria. Thus, the phage immune landscape of E. coli is composed of both widespread and clade-specific systems.

Mobile elements dominate the defence system landscape in E. coli. Previous searches for proteins enriched in defence islands identified and validated 38 novel defence systems\(^12,13\). None of the 21 systems identified here are homologues of those systems, and only one component of (PD-T7-2) resembles (32% identity) a protein of a previously validated multi-component system. PD-T4-8 has a DUF4263 domain in common with the Shedu defence system but is not homologous to the validated Bacillus cereus system. The computational approach in ref. \(^13\) also identified 7,472 protein families enriched in defence islands that have yet to be validated. Only 14 of 32 proteins identified here have homology to those, and often with <35% identity over limited regions of the proteins (Supplementary Table 4). These observations suggest that our experimental-based selection may uncover different types of defence systems than can be found computationally by searching for enrichment in defence islands.

To further probe this idea, we analysed the native genomic context of the 21 systems identified here. We found that 12 of the 21 systems were located in intact prophages (Extended Data Fig. 4). Seven of these systems were found within P2-like prophages, with four located in the same position of the P2 genome, directly between the genes encoding the P2 replication endonuclease and...
**a**

| T2 | T4 | T3 | T7 |
|----|----|----|----|
| SEC | uni03D5 | 17 | SEC | uni03D5 | 18 | SEC | uni03D5 | 27 |
| vir | 0 | –1 | –2 | –3 | –4 | –5 |

| PD-T4-1 | PD-T4-2 | PD-T4-3 | PD-T4-4 | PD-T4-5 | PD-T4-6 | PD-T4-7 | PD-T4-8 | PD-T4-9 | PD-T4-10 |
|---------|---------|---------|---------|---------|---------|---------|---------|---------|---------|
| S       | S       | S       | S       | S       | S       | S       | S       | S       | S       |

**b**

- **Abi**: Abortive infection
- **No defence**: Empty vector

**c**

- **ECOR22**
- **ΔPD-T4-9**
- **ECOR65**
- **ΔPD-T4-2**

**d**

- **EV**
- **PD-T4-2**
- **H127A**
- **PD-T4-3**
- **Y19A**
- **PD-T4-4**
- **K48A**
- **PD-T4-6**
- **K101A**

**e**

- ** pll-NTD**
- **DUF262 nuclease**
- **DUF4041 TM**
- **GIY-YIG nuclease**
- **P(D/E)XK nuclease**
- **Lipoprotein**
- **RelE**
- **Ser/Thr kinase**
- **HEPN**
- **PD(D/E)XK nuclease**
- **Phage sheath**
- **ADP-ribosyltransferase**
- **SecB**
- **Mrr**
- **DUF4145 propeller**
- **PD(D/E)XK nuclease**
- **Mrr P-loop NTPase**
- **CoiA**
- **ParBc nuclease**
- **HEPN/DUF4145 Methyltransferase**
- **Zn finger**
- **ADP-ribosyltransferase SecB**
- **H127A**
- **Y19A**
- **K48A**

**f**

- **Alpha-proteobacteria**
- **Beta-proteobacteria**
- **Delta-proteobacteria**
- **Nitrospirae**
- **Acidobacteria**
- **Aquificae**
- **Placmomyotes**
- **Verrucomicrobia**
- **Chlamydiae**
- **Bacteroidetes/chlorobi**
- **Spongochaeta**
- **Chloroflexi**
- **Firmicutes**
- **Actinobacteria**
- **Tenericutes**
- **Deinococcus-Thermus**
- **Thermotogae**
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portal protein (Fig. 4a). This location has been previously found to harbour the defence systems tin and old, the former being also identified in our screen against \( \lambda \)\textsuperscript{old}\textsuperscript{24} (Supplementary Table 1). More recently, this location was found to encode a wide array of previously uncharacterized defence systems\textsuperscript{25}. We also observed a second defence-enriched locus in P2-like phages, which contained three of the systems discovered here, and the previously identified defence gene \( \text{fun} \) in the P2 reference genome\textsuperscript{24} (Fig. 4a). This hotspot was not noted in the recent analysis of P2 defences. Notably, 47 of our 71 \( E. \ coli \) strains together encode 63 P2 portal proteins, with 110 unique proteins found in the adjacent hotspot. Only 13 of these 63 hotspots contain previously known defence systems, and most genes are annotated as hypothetical. These findings suggest that not only do P2 prophages encode a rich diversity of anti-phage proteins\textsuperscript{26} but that P2 defence hotspots make up a substantial fraction of the immune landscape in \( E. \ coli \) (Fig. 4b).

Five defence systems were found associated with other types of prophages or their remnants, including P4 satellite prophages or related integrases, Mu-like phages and lambdoid phages (Extended Data Fig. 4). We also observed a defence-enriched locus within an integrative and conjugative-like element (ICE), containing two systems we identified—PD-T4-4 and PD-T7-2 (Fig. 4a). This element is not widely distributed in our \( E. \ coli \) strain collection, but similar elements in other genomes encode known defence systems at this locus, as well as hypothetical proteins that may also function in phage defence (Fig. 4a). PD-T7-4 and its homologues often overlap an integrase gene, while PD-T4-5 was identified on a plasmid. The remaining 4 systems did not appear to reside within active MGEs, but each had a nearby integrase gene suggesting that they may be part of decaying MGEs (Extended Data Fig. 4).

Our study supports previous findings that in addition to defence islands, prophages and other MGEs are a rich reservoir of defence systems\textsuperscript{26}. However, these categories are not mutually exclusive, as some defence islands may be carried on or derived from MGEs\textsuperscript{27}. To more systematically document the different genomic contexts for the systems identified here, we collected, for each of the 21 systems, all homologues in a set of 544,603 publicly available bacterial genomes. We classified the genes within 10 kb upstream and downstream of each homologue as either defence-related, prophage-related or neither (Methods). We then tabulated the number of defence- and prophage-related genes flanking each homologue (Fig. 4c,d and Extended Data Fig. 5).

We detected two distinct patterns. For systems that we found outside of prophages in our strains, the homologues were also typically not prophage-associated (Fig. 4c and Extended Data Fig. 5) and were often near several other genes encoding defence-associated domains. Thus, these systems do appear in defence islands, even though they were not previously detected as enriched in them. For the systems we identified within \( E. \ coli \) prophages, some of their homologues were also found in prophages, as evidenced by dozens of flanking prophage-related genes (Fig. 4d and Extended Data Fig. 5). These prophage-associated homologues were typically near 1–2 defence-associated genes but rarely more than 2, suggesting that some systems reside in small defence islands or clusters within a prophage, as with the P2 hotspots (Fig. 4d)\textsuperscript{28}.

Notably, there are homologues of each system we identified that can be found in defence islands (Fig. 4e), some more rarely than others, indicating that they do not require a prophage context to function. In aggregate, we observed an inverse correlation between the number of neighboring defence-associated genes for homologues of systems commonly found in prophages versus those that are not (Fig. 4f,g). This highlights the constraint on how many defence systems can be carried by prophages, or within a given hotspot, due to size limitations in DNA that can be packaged. We suspect that this is one reason why many of these genes are enriched in prophages as compared with defence islands.

Finally, we compared defence island and prophage enrichment (Methods) between systems discovered here and those previously predicted computationally and validated experimentally\textsuperscript{25,26}. We found that our experimentally selected systems were on average less frequently associated with known defence genes but more frequently associated with prophage genes (Extended Data Fig. 6). These analyses suggest that defence-island enrichment methods may be less sensitive in identifying defence systems frequently found in prophage.

**Previously uncharacterized toxin-antitoxin-like systems.**

Toxin-antitoxin (TA) systems are typically composed of a protein toxin that can arrest cell growth but is normally neutralized by a cognate, co-expressed antitoxin. TA systems are extremely prevalent in bacterial genomes and MGEs, but their functions remain poorly understood\textsuperscript{26,27}. A handful of TA systems have been found to provide anti-phage defence through an Abi mechanism\textsuperscript{26,27,28}. Our selection yielded four different systems that were recognizable as TA-like in nature. These encoded gene products with sequence similarity to toxic proteins, mostly featured multiple components, and provided Abi defence (PD-T4-5, PD-T4-7, PD-T4-9 and PD-\( \lambda \)-2). A fifth (PD-T4-10) facilitated Abi defence and had two overlapping ORFs, reminiscent of many TA systems. As noted above, none of these systems were previously annotated as TA systems, so we sought to validate the three featuring multiple components: PD-T4-10, PD-\( \lambda \)-2 and PD-T4-9 (Fig. 5a).

For PD-T4-10, neither of the two proteins had a predicted function. We expressed each component from an inducible promoter and found that the second component, PD-T4-10B, was toxic. This toxicity was fully neutralized when PD-T4-10A was co-expressed (Fig. 5b). PD-T4-10A could not be deleted, consistent with it being an antitoxin, whereas deletion of the toxin PD-T4-10B abolished resistance to T4 infection (Fig. 5c). Thus, this system comprises a novel, bona fide TA system that provides strong protection against T-even phages.

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**Fig 4 | Prophages and MGEs are major sources of defence systems.** a. Hotspots of previously uncharacterized defence systems. Left: the native genome context of seven defence systems identified here, showing the boundaries of P2-like prophages in the genome from which they originated. Genes are colour-coded as indicated at the bottom. Right: two defence systems were identified in the accessory region of an ICE-like element within the indicated genome. Homologous elements from other bacterial genomes contain known and putative defence systems in the same location. b. All identified instances of P2 defence hotspot #1 in our 73-strain \( E. \ coli \) collection. White genes are flanking conserved P2 genes. Each colour of gene within the hotspot represents a protein cluster (30% identity). All grey genes belong to a lone cluster. Double slashes denote the end of a contig. c. Number of defence and prophage-associated genes \( \pm 10 \text{kb} \) from system homologues. The scatterplots indicate, for each homologous system, the numbers of prophage and defence-associated genes within \( \pm 10 \text{kb} \). Examples in c represent systems that were found outside of prophages in our genome collection. For all 21 systems, see Extended Data Fig. 5. d. Same as c but for systems we found in prophages. e. Examples of the \( \pm 10 \text{kb} \) context for PD-T7-1 homologues in a defence island or prophage. f. Distribution of the number of nearby defence-domain-containing genes in homologues of systems commonly found in prophages (>10% homologue-containing regions with 8+ prophage genes in proximity) or not; \( n = 12 \) and 9 systems, respectively. Boxes indicate bounds of the distribution as median \( \pm \) quartiles, and limits exclude outliers. g. Linear least squares regression for total nearby prophage gene and defence-domain-containing genes for each system. Pearson \( r = -0.442, P = 0.045 \); error indicates 95% CI.
Figure 1: Integrated and conjugative-like elements

(a) Schematic representation of known systems, screen hits, and candidate uncharacterized systems. Examples include
- **Known systems**: PD-T7-2, PD-T4-5, PD-T4-9, PD-T7-2
- **Screen hits**: PD-T4-5, PD-T7-2, PD-T7-2
- **Candidate uncharacterized systems**: PD-T7-2, PD-T7-2

(b) P2 portal protein and P2 defence hotspot
- **Known systems**: Sir2, HerA
- **Screen hits**: Sir2, HerA

(c) Distribution of defence-associated domains
- **PD-T7-3** homologues
- **PD-T7-4** homologues
- **PD-T7-5** homologues

(d) Distribution of prophage genes
- **PD-T7-6** homologues
- **PD-T7-7** homologues
- **PD-T7-8** homologues

(e) Representative biological systems from different bacterial species
- **Salmonella enterica**
- **Pseudomonas aeruginosa**
- **Klebsiella pneumoniae**
- **Stenotrophomonas ureilytica**

(f) Systems common vs. uncommon in prophage
- **Systems common in prophage**: 12 counts × 1,000
- **Systems uncommon in prophage**: 0.6 counts

(g) Mean fraction nearby prophage
- **Mean fraction nearby prophage**: 0.20

Additional notes:
- **2.5 kb marker**: Indicates scale for all genomic regions.
- **Integration**: P2 portal protein, P2 defence hotspot.
- **Replication**: SIR2, RM-like ATPase, Fun, Lysis, Integration Transfer, Replication, Lysogeny, Old Tin.
- **Type-I RM**, **Type-IV RM**, **QatA**, **CBASS**, **TRIP13**.
- **Rhizobium leguminosarum**.
- **Salmonella enterica**.

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PD-λ-2 features three components. The first is similar to HigB toxins that inhibit translation, the second has a HigA antitoxin-like domain (Xre-family helix-turn-helix) fused to a C-terminal peptidase domain and the third component is related to a P4 phage antitoxin. An Xre-peptidase fusion, co-expressed with an upstream toxin-like gene, has been observed in sequence-based TA searches but the function of these loci is unknown. Unexpectedly, we found that overexpressing the second component, PD-λ-2B, alone was toxic, with toxicity rescued by co-expression with PD-λ-2A (Fig. 5d). We also confirmed that PD-λ-2C, although not required to neutralize the toxin, is required for defence against λ vir (Fig. 5e).

PD-T4-9 also contains a third component, a SecB-like chaperone, suggesting that it is related to an enigmatic class of TA systems called toxin-antitoxin-chaperone (TAC) systems, so we renamed this system CmdTAC for chaperone-mediated defence. The antitoxins of TAC systems are typically homologous to canonical antitoxins but feature an unstructured extension of their C termini called the chaperone addiction domain (ChAD). In the absence of the cognate SecB-like chaperone, the ChAD renders the antitoxin prone to aggregation and proteolytic degradation, thereby freeing the toxin. Inducing the expression of just CmdT, a previously uncharacterized toxin that has an ADP-ribosyltransferase-like domain, was not toxic, but co-expressing CmdT with the presumed antitoxin, CmdA, was toxic to cells. Co-expression of CmdT with both the putative antitoxin and chaperone components largely restored viability (Fig. 5f). Together, these results suggest that CmdTAC is a novel TAC system that protects against phage.

To further characterize the role of the chaperone, CmdC, we overproduced it during T4 infection of CmdTAC+ cells. Interestingly, an oversupply of CmdC abolished phage protection by CmdTAC, suggesting that destruction of CmdC, or sequestration of CmdC from the complex may allow this TA system to activate in response to phage infection (Fig. 5g). The chaperone normally promotes neutralization of CmdT by CmdA but, following infection, could be

Fig. 5 | Previously uncharacterized toxin-antitoxin-derived defence systems. a, Schematics of PD-T4-9, PD-T4-10 and PD-λ-2 defence system operons and their domain predictions. Representative homologues of the systems are shown in their genomic contexts and indicate conservation and order of the system components. Blue, putative antitoxin; red, putative toxin; green, accessory factor. b, d, f, i, Each component or pair of components indicated was expressed (+) or not (−) from an inducible promoter and assayed for viable colony-forming units in tenfold serial dilutions. c, e, Plaquing assays for the phage indicated on cells harbouring an empty vector or a vector containing a given defence system with all components (WT) or lacking the component indicated. g, j, Plaquing of phages on TAC-containing strains expressing a second copy of the chaperone component to varying levels during infection. h, Schematic and T2 plaquing assay of the mqsRAC TAC system.
depleted or sequestered by a phage product, leading to liberation of the toxin and abortive infection. Providing additional CmdC prevents the loss or complete sequestration of chaperone, thereby preventing release of CmdT from CmdA. Additional work is needed to further dissect the mechanistic nature of chaperone-phage interactions.

We hypothesized that TAC systems may be a broad class of phage defence systems. To test whether other TAC systems can protect *E. coli* MG1655 against phage infection, we cloned and tested an MsqRAC system from *E. coli* C496_10. Although completely unrelated in toxin and antitoxin sequence to CmdTAC, MsqRAC is a canonical TAC system, has been characterized in Mycobacteria and includes a SecB-like chaperone homologous to CmdC. This system conferred robust protection against T2 (Fig. 5h) but not T4 as with *cmdTAC*. Toxicity of MsqR could only be rescued by expressing MsqA and MsqC (Fig. 5i), and inducing additional MsqC in MsqRAC⁺ cells inhibited phage protection (Fig. 5j). Thus, our work indicates that TAC systems may be a widespread and diverse new class of phage defence system.

**Discussion**

Our work indicates that a large reservoir of diverse, previously unknown phage defence genes is distributed across the *E. coli* pan-genome. Like many bacteria, there is tremendous variability in the ‘accessory’ genomes of different strains of *E. coli*. Many of these accessory genes are probably associated with phage defence, as recently suggested for marine *Vibrio* species. Although efforts to find new defence systems on the basis of proximity to known systems have proven fruitful, our work reveals that many phage protective systems remain unidentified.

Our functional screening, which is agnostic to the genomic context of defence systems, indicates that many systems are not commonly or detectably enriched among known defence islands. Indeed, none of the 21 systems we identified were previously reported to provide phage defence in previous discovery efforts based on defence island enrichment. Moreover, only three of the systems we identified were within 10 kb of other known defence systems in our *E. coli* genomes. However, notably, 15 of the 21 systems we identified were present in apparently active or recently active MGEs (prophages, ICEs or plasmids) (Extended Data Fig. 4), with the other 6 located in regions suggestive of MGEs in the late stages of decay. Homologues of the systems discovered here are sometimes present in defence islands (Figs. 4d,e and 5a), but these associations are often relatively rare.

It is well established that MGEs contribute to antiviral defence in bacteria. By providing a glimpse into the relative contributions of MGEs and defence islands to immune system context, our results support the notion that active prophages and other MGEs are likely the primary reservoirs of anti-phage defence systems in *E. coli*. This idea is consistent with studies that have identified diverse anti-phage systems in *P. aeruginosa* and *P. stutzeri* and another recent study revealing how the transfer of defence-containing ICEs drove the emergence of phage resistance in clinical isolates of *V. cholerae*. The defence systems found in functional MGEs probably help these elements to protect themselves and host resources by preventing infection by other phages. However, given genome size and packaging constraints, there is a limit to the number of defence systems carried by a given prophage. Such constraints may help explain why increased prophage association was correlated with lower defence gene association (Fig. 4g). Future investigation should work towards further uncovering the vast reservoir of anti-phage elements carried on MGEs.

Of the 32 proteins in 21 systems found here, 13 feature domains never before reported to function in phage defence. These include a protein with distant similarity to ReE and a region of a phage tail sheath, an Exotoxin A-like domain previously only shown to function in bacterial virulence, a SecB-like chaperone, DUF6575, DUF4041, a CoIA competence-related domain, a β-propeller fold, an ImmA/IrrE peptidase, a HigB toxin and three proteins with no ascertainable similarity to deposited systems. In addition, ten proteins contained large regions with no predicted domains (Fig. 3a).

Although some regions of the proteins identified have distant similarity to known nuclease motifs, these domains are found here in new or unusual contexts and associations, which raises fascinating questions for future investigation. For example, a 7-bladed beta propeller with a separate N-terminal Mrr-family nuclease domain. The β-propeller fold consists of separate modules that adopt a disc-like, circularly arranged structure with a central channel that can accommodate many substrates including protein and DNA. Determining how this domain aids in activation or target specificity of this system opens many avenues of future discovery. We also uncovered five single-protein systems that exclusively contain a putative DNase domain. Although these domains are found in defence systems such as RM and CRISPR, how orphan DNase-like proteins sense and respond to phage infection, especially to provide direct defence, is unclear. As all but one is encoded without a DNA methylase, they probably do not distinguish phage and host DNA in the same way as RM systems. Additionally, these DNAse-containing systems protected against T4 and T7, which are intrinsically resistant to most RM systems. Type-IV RM systems are single-component defences known to target the modified cytosine of T-even phages, but none of the five proteins discussed here resemble these systems. How those that are specific to T7 might target T7 DNA, which is not modified, is unclear. The existence of unconventional, phage-targeting, nucleic-acid degradation systems underscores a knowledge gap in the molecular mechanisms of viral resistance and self/non-self recognition.

We also identified and validated three TA systems as phage defence elements including *cmdTAC*, prompting the discovery that unrelated TAC systems such as *msqRAC* also function in antiviral immunity. MsqRA is a well-studied TA system but has no documented role in phage defence; our results suggest that bacteria have co-opted this TA system by addition of a chaperone-dependent antitoxin to activate the toxin in response to infection. Our findings support the notion that TA systems play a central role in phage defence.

Finally, we note that four systems identified here probably belong to classes of previously described systems but have diverged quite significantly such that they share little significant sequence homology, reinforcing the extreme divergence and adaptations that typify many immune systems. These divergent systems include PD-T7-2 in which the second protein is similar to HerA of Sir2/HerA, PD-T4-8 whose central domain bears distant resemblance to Shedu, PD-λ-5 which appears to be a highly compacted phage version of an RM+Abi system and PD-T4-5—a plasmid-borne gene that is a distant relative of Abi/F/D.

Our screening methodology enables the experimental discovery of anti-phage defence genes and has several powerful features. First, we can return to the strain of origin for a given system and demonstrate that it provides defence in its native context. Second, there is a built-in pairing of defence systems to the phage they defend against, whereas with computational studies, the phage(s) a given system defends against must be subsequently identified. Third, in the pipeline described here, the source DNA comes from other *E. coli* strains, which probably minimizes false negatives that can arise from producing candidates in a heterologous host. Finally, our experimental approach is not limited to genes that are detectably enriched in defence islands. As noted, only three of the systems we identified were natively associated with obvious defence islands, and some also do not appear to have many close homologues in defence islands. Some defence systems may not associate with defence islands, while some may have arisen too recently or not be
widespread enough to detect an association. Indeed, some of the systems identified here show a relatively limited phylogenetic distribution. However, the phage defence capabilities of bacteria probably include both broadly conserved and clade-specific systems adapted to the unique biology of a given organism and its phages.

A limitation of our method is that the initial selection does not distinguish between clones that lead to cell surface modifications that block adsorption and those providing intracellular defence systems; these must be distinguished in downstream steps. Further, at present, our method probably does not reach saturation as some systems are strongly overrepresented among the positive clones, which probably hinders the identification of others. The reason for this uneven distribution is unclear, but efforts are underway to increase throughput and sensitivity of the method.

The methodology developed here can be powerfully extended in several ways. First, genomic DNA from other sources, including metagenomic DNA, could be used as input material. From just 71 strains, we identified 21 new defence systems, suggesting that fertile ground remains to be discovered both in and beyond E. coli. Second, the panel of phages tested here was limited to three and could easily be expanded, particularly given the enormous diversity of phages. Finally, with only small modifications, any transformable bacterium could be used as the host strain. Further identification and characterization of bacterial immune systems promises to shed new light on the ancient arms race between bacteria and their viral predators, and may also have practical applications, providing the foundation for precise molecular tools and helping to inform future efforts to develop phage as therapeutics.

**Methods**

**Bacteria and phage growth and culture conditions.** Cultures were routinely grown at 37 °C in LB medium unless otherwise stated. Phage stocks were propagated on MG1655, filtered through a 0.2 µm filter and stored at 4 °C. Select ECOR strains were obtained from the Thomas S. Whittam STEC Center at Michigan State University, and UMB isolates were obtained from Alan J. Wolfe at Loyola University Chicago (Supplementary Table 5). Other strains, plasmids and primers + synthetic gene fragments are listed in Supplementary Tables 6–8, and primers can be obtained from TIB Molbiol (Invitrogen). Synthesis of gene fragments was performed by IDT. Synthetic gene fragments were linearized by restriction digestion with the restriction enzyme NdeI, EcoRI or SfiI for Oxford Nanopore sequencing library construction. Long-read sequences were chromatographed and quality checked (Agilent Technology) to confirm integrity, and high-quality samples were indexed by native barcoding (ONT kits EXP-NB104114) with fluorescent reagents from New England Biolabs. Libraries were prepared using the LSK-109 chemistry, and samples were run on either a MinION (R9) or PromethION (R9.4) flowcell. Basecalling was done using built-in ONT tools. Processed reads were aligned to public reference genomes of the source organisms, or the reference genomes contained in the fosmid inserts, using the Minimap2 plugin within the Geneious Prime 2020.2.4 software suite. Fosmids that could not be mapped to their genomes due to contig gaps were also sequenced identically, and de novo assembly was conducted using Flye, also in Geneious. Candidate defense systems were predicted to be any gene or operon residing under the coverage maxima.

**Library construction.** Genomic DNA was collected from pooled overnight cultures of each E. coli isolate using the PureLink genomic DNA mini kit (Invitrogen). From each isolate, a fosmid library was constructed by ligation reactions using the CopyControl fosmid library production kit (Lucigen) according to the manufacturer’s protocol. Plasmid sub-libraries were constructed first by extracting fosmid DNA from select positive clones using the ZR plasmid miniprep kit (Zymo Research). Equimolar pooled fosmids were sheared to an average of 8 kb using g-TUBEs (Covaris). Sheared DNA was end-repaired and 5’-phosphorylated using the ZR plasmid miniprep kit (Zymo Research). The plasmid vector was prepared by PCR, and blunt-ended fragments were ligated to the plasmid using T4 DNA Ligase (NEB) for 2 h at room temperature. The ligation reaction was electroporated into MegaX DH10B T1R Electrocomp cells (Thermo Fisher) and selected on LB with 25 µg ml−1 chloramphenicol and grown to stationary phase at 37 °C (approximately 4 h, optical density (OD)600=1.0). The ligation reaction was electroporated into the End-It DNA end-repair kit (Lucigen) and then purified using the DNA Clean and Concentrator kit (Zymo Research). The plasmid vector was prepared by PCR, and blunt-ended fragments were ligated to the plasmid using T4 DNA Ligase (NEB) for 2 h at room temperature. The ligation reaction was electroporated into MegaX DH10B T1R Electrocomp cells (Thermo Fisher) and selected on LB with 25 µg ml−1 kanamycin. 

**Defence system selection.** We used a variant of the previously described ‘tab’ selection procedure to select for fosmids that provide resistance to phage infection. A heavy inoculum (>30 µl) of a high-titre library freezer stock or empty vector freezer stock was inoculated into 5 ml LB containing 20 µg ml−1 chloramphenicol and grown to stationary phase at 37 °C (approximately 4 h, optical density (OD)600=2–3). Cultures were adjusted to OD600=1.0, and 0.1 ml (~8×107 cells) was pipetted onto one side of 3–6 empty 15 cm Petri dishes. A light inoculum (~50 µl) was pipetted onto the other side of the Petri dishes. Plates were incubated at 37 °C overnight, except for one T4 screen which was conducted at 30 °C. The method is adapted from ref. 42. Overnight bacterial cultures were diluted 1:100 and grown to OD600=0.5. Cultures were incubated at 37 °C for 1 h, then incubated at 25 °C for 15 min or 25 °C for 25 °C (T7) for 15, 25 or 15 min for T4, λ, and T7, respectively. Samples (500 µl) were then added to a tube of ice-cold chloroform and vortexed, and unadsorbed phage were enumerated by the top agar overlay method using a susceptible
indicator strain. Percent adsorption was determined relative to a simultaneous mock control experiment that contained growth medium but no host cells.

**Ablative infection assays.** Overnight cultures were normalized to OD₆₀₀ = 1.0 and then diluted 100-fold. Diluted cultures (150 µl) were dispensed in a flat-bottomed 96-well plate. Phage dilutions (10 µl) were added to each well such that the MOI varied from 50 to 0.005. Wells were then overlayed with 20 µl mineral oil, and plates were covered with a breathable membrane. Plates were incubated at 37°C in a Biotek Synergy H1 microplate reader. OD₆₀₀ was measured every 15 min. Three technical replicates were conducted for each strain.

**Toxicity assays.** Strains containing plasmids with inductive promoters were grown overnight at 37°C in LB under repressing conditions (LB or LB + 0.2% glucose). Cultures were washed in LB, and tenfold serial dilutions were spotted on LB agar with or without inducer (0.2% arabinose, 200 µg·ml⁻¹ vanillate, or 100 µg·ml⁻¹ anhydrotetracycline). Plates were incubated overnight at 37°C.

**Bioinformatic analyses.** Sanger sequences of fosmid ends were mapped to their strains of origin using BLASTn⁴³, followed by manual inspection. General remote homology was determined using the function cluster within MMseqs2⁵⁷ with the following parameters: –evalue 0.001 -qcov_hsp_perc 80 of the predicted domain were taken from the top hit, while the label was chosen according to the 80% identity cut-off. The local genomic context of a defence system homologue was defined as all adjacent genes that were part of a single multi-component defence system. For the defence-related domain search, sequences were searched against defence-related pfam and COG domains identified and used in refs. 11–13. For scatterplots and marginal histograms in Fig. 4 and Extended Data Fig. 5, domains available as of May 2021 when the pVOG database was downloaded. For the defence-related domain search, sequences were searched against defence-related pfam and COG domains identified and used in refs. 11–13.

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

**Data availability**

All data generated or analysed during this study are included in the published article (and its Supplementary Information).

**Code availability**

Code used in this study is available at https://github.com/chrisdoering197/phagedefense.

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Author contributions

M.T.L. and C.N.V. designed the experiments and wrote the manuscript. C.N.V designed and implemented the screen. C.R.D. wrote code to collect homologue data. C.N.V and C.R.D. analysed data. C.N.V. and M.L.L. contributed to routine and implemented the screen. C.N.V., C.R.D., G.I.C.T. and M.L.L. contributed to routine experiments and cloning.

Competing interests

The authors declare no competing interests.

Additional information

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Extended Data Fig. 1 | Diversity of the *E. coli* isolate pangenome used in this study. (a) (Left) Phylogenetic tree of *E. coli* strain collection used to construct the genomic library screened. *E. coli* K-12 (MG1655) and B (REL606) are also included. (Right) Bars indicate presence/absence (red/white) of individual gene clusters (95% identity threshold). (b) Plot of the number of gene clusters versus the number of strains they are found in, for example ~8,000 clusters are each found in only one genome. These sparsely conserved clusters represent the accessory genome, whereas ~3,000 clusters are found in all 73 genomes and represent the *E. coli* core genome.
**Extended Data Fig. 2 | Adsorption of bacteriophage on various strains.** (a) Mean adsorption of T4 on control (EV) and LPS-containing fosmid strains (T4F17). Error bars represent standard deviation of three biological replicates. (b) Adsorption of T4, λ, or T7 on strains expressing defense systems against their respective phages.
Extended Data Fig. 3 | Mechanisms of defense and defense system function in other strains. (a) Growth of control (EV) or defense system-expressing strains with phage MOIs of 0, 0.05 or 5. Phages used in each experiment are shown on the right. Asterisks indicate systems not showing direct immunity and likely representing abortive infection mechanisms. Lines represent the mean of three technical replicates with shaded regions indicating standard deviation. (b) EOP measurements for T4 and \( \lambda \) on \( E. \ coli \) strains MG1655, ECOR13, or C expressing various defense systems indicated.
Extended Data Fig. 4 | Native genomic neighborhoods of newly identified defense systems. Genome maps of native locations of the defense systems, showing the flanking 10 kb regions, unless interrupted by the end of a contig. Prophage and defense-domain containing genes were called as described in the Methods.
Extended Data Fig. 5 | Genome context of defense system homologs. Data as in Fig. 4c, extended to all systems discovered here and sorted by the MGE context in which they were found.
Extended Data Fig. 6 | Comparison of defense and prophage enrichments between experimentally and computationally discovered systems. (a) Overview of method. Genes +/- 10 kb of homologs of defense systems were predicted as defense- or prophage-associated. To minimize the effects of sequence/genome redundancy, proteins were clustered to 95% identity. % scores were calculated as the number of prophage or defense-associated clusters over total clusters. (b) Boxplots of % prophage- and defense-associated genes near the experimentally discovered systems (this study) or computationally predicted and validated systems. P values indicate significance from two-sided Mann-Whitney U test. Boxes indicate bounds of the distribution as median +/- quartiles, and limits exclude outliers.
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☑☐ A statement on whether measurements were taken from distinct samples or whether the same sample was measured repeatedly

☑☐ The statistical test(s) used AND whether they are one- or two-sided

☐☐ Only common tests should be described solely by name; describe more complex techniques in the Methods section.

☑☐ A description of all covariates tested

☑☐ A description of any assumptions or corrections, such as tests of normality and adjustment for multiple comparisons

☐☐ 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)

☐☐ For null hypothesis testing, the test statistic (e.g. F, t, r) with confidence intervals, effect sizes, degrees of freedom and P value noted

Give P values as exact values whenever suitable.

☐☐ For Bayesian analysis, information on the choice of priors and Markov chain Monte Carlo settings

☐☐ For hierarchical and complex designs, identification of the appropriate level for tests and full reporting of outcomes

☐☐ 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
BLAST v. 2.11.0+ for homolog searches. HMMER3 v. 3.1b2 for domain searches. Taxonkit v. 0.8.0 to gather taxonomy information. MMseqs2 v. 13-45111 for protein sequence clustering. Custom code was generated, using these tools, to detect homologous proteins and their genome context from NCBI genomes. This code is publicly available at https://github.com/chrisdoering8197/phagedefense

Data analysis
HHsuite v. 3.1.0 for remote homology detection (includes HHblits and HHSearch). Prokka v. 1.14.6 for genome annotation. Roary v. 3.12.0 for pangenome analysis. FastTree v. 2.1.10 for phylogenetic tree construction. DefenseFinder v1.0 for prediction of known defense systems.

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

All data generated or analyzed during this study are included in the published article (and its supplementary information files).
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

Life sciences study design

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

| Sample size | No sample size calculation was performed. Given large effect sizes, we chose to replicate experiments in triplicate as is standard to indicate reproducibility. |
|-------------|------------------------------------------------------------------------------------------------------------------------------------------|
| Data exclusions | Only representative sub-library peaks were included in Fig. 2C because the final identification of genes and their functions is confirmed in the subsequent figures. |
| Replication | All experimental findings were repeated at least twice. All reported results were successfully reproduced. |
| Randomization | No experimental groups or control groups were subjectively chosen and there are no covariates to control for as experiments were done in isogenic strains. No experiments required randomization: |
| Blinding | Blinding was not relevant because all data were obtained objectively and had strong effect sizes and raw data is reported in the manuscript. All comparative analyses were performed in isogenic strain backgrounds. |

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 | Methods |
|---------------------------------|---------|
| n/a | n/a |
| ☒ Antibodies | ☒ ChIP-seq |
| ☒ Eukaryotic cell lines | ☒ Flow cytometry |
| ☒ Palaeontology and archaeology | ☒ MRI-based neuroimaging |
| ☒ Animals and other organisms | |
| ☒ Human research participants | |
| ☒ Clinical data | |
| ☒ Dual use research of concern | |