Phage-inducible chromosomal islands are ubiquitous within the bacterial universe

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Abstract
Phage-inducible chromosomal islands (PICIs) are a recently discovered family of pathogenicity islands that contribute substantively to horizontal gene transfer, host adaptation and virulence in Gram-positive cocci. Here we report that similar elements also occur widely in Gram-negative bacteria. As with the PICIs from Gram-positive cocci, their uniqueness is defined by a constellation of features: unique and specific attachment sites, exclusive PICI genes, a phage-dependent mechanism of induction, conserved replication origin organization, convergent mechanisms of phage interference, and specific packaging of PICI DNA into phage-like infectious particles, resulting in very high transfer frequencies. We suggest that the PICIs represent two or more distinct lineages, have spread widely throughout the bacterial world, and have diverged much more slowly than their host organisms or their prophage cousins. Overall, these findings represent the discovery of a universal class of mobile genetic elements.

Introduction
The Staphylococcus aureus pathogenicity islands (SaPIs) are a novel class of phage satellites that are intimately related to certain temperate (helper) phages, of whose life cycles they parasitize. Following infection by a helper phage or SOS induction of a helper prophage, the SaPI genome excises from the bacterial chromosome, using SaPI-encoded integrases (int) and excision functions (xis) [1, 2], replicates extensively using its own replicon [3], and is efficiently packaged into infectious particles composed of phage virion proteins [4–6]. Some SaPIs encode capsid morphogenesis functions that remodel the phage capsid to only fit their smaller genomes [7, 8]; others simply use full-size phage capsids (for a recent review, see [9]). The hallmark of this parasitism is a key SaPI regulatory gene that diverts the phage reproduction cycle to its own end. This gene encodes a master repressor (Stl) that governs expression of the SaPI genome [10]. Unlike the classical phage repressor, the SaPI Stl repressor is not cleaved following activation of the SOS response; rather the repression is lifted by the formation of a complex between the repressor and a specific helper phage protein [11–13]. This serves to couple the SaPI life cycle with that of the helper phage, ensuring that the SaPI is not activated unless the reproductive cycle of a helper phage is in progress. Once induced, different SaPIs use different strategies to initiate specific packaging from the cognate SaPI genome [7, 14], ensuring their high intra- and inter-generic transfer [15–17]. Moreover, SaPIs have a huge impact on the biology of their helper phages by interfering with phage reproduction using different and complementary strategies [18–20]; they are also key elements driving phage evolution [21].

Not surprisingly, SaPI-like elements are not unique to staphylococci, and we have recently demonstrated that they are widespread in Gram-positive (GP) cocci [22]. This new family of mobile genetic elements (MGEs), that we have
called generically phage-inducible chromosomal islands (PICIs), has very well conserved features [9, 23]. All PICIs have a conserved gene organization (Fig. 1), and encode a pair of divergent regulatory genes, including the PICI master repressor Rpr (called Stl in the SaPIs). Left of rpr, and transcribed in the same direction, PICIs encode a small set of genes including an integrase (int) gene. Right of rpr, and transcribed in the opposite direction, the PICIs encode an excision function (xis), and a replication module consisting of a primase homolog (pri) and a replication initiator (rep), which are sometimes fused, followed by a replication origin (ori). Next to these genes, and also transcribed in the same direction, PICIs encode genes involved in phage interference, and sometimes, a terminase small subunit homolog (terS) which is responsible for the high efficiency of the SaPI packaging [7]. In the SaPIs, accessory genes (usually involved in virulence; [9]) can be found either at the 3′ end of the elements or between the int and stl genes (Fig. 1).

In addition to this well-conserved gene organization, the GP PICIs have other common features that distinguish them from their cousin prophages: (i) unique attachment (att) sites that are never occupied by prophages; (ii) the absence of phage structural and lytic genes; (iii) size typically around 15 kb; (iv) sophisticated strategies to interfere with helper phage reproduction and (v) genes unique to PICIs. Orthology analyses show that PICI genes belong to large sets of orthologs, within which the first 8–10 are almost always PICI genes, most of which never appear in other genetic elements [22, 24].

The preliminary analysis of the PICIs present in the GP cocci suggested that these elements could have evolved de novo in the different genera [22], suggesting their existence is strongly selected in nature. Therefore, and as happens with the classical MGEs, we hypothesized that these elements are likely not just confined to the GP cocci but widespread in nature. We now report the identification and characterization of a large number of PICI elements in various Gram-negative (GN) bacteria. Because they have certain key features in common and because they are so widespread, we suggest that their lifestyle may have strong selective value and that they represent a novel agency of horizontal dissemination of virulence and other important mobile genes among bacteria.

**Results**

**Identification of PICI candidates**

Since PICIs are a successful biological strategy in GP cocci, we hypothesized that similar elements will be widespread in GN bacteria. Using the aforementioned criteria defining the PICI elements in the cocci (see material and methods for more details), we have assembled a representative collection of putative PICIs in the GN bacteria by genome searching; their genomes and characteristics are depicted in Fig. 1 and S1, and in Table S1. Following the nomenclature proposed for the elements present in the GP bacteria, the individual PICIs are designated with reference to their species – thus, EeCl or PmCl would be used for PICIs of *Escherichia coli* or *Pasteurella multocida*, respectively, where “n” would be used for the specific PICI-containing strain.

**Features of PICI candidates**

The putative PICIs described here have a number of features in common. (i) Occurrence: they are very common among GN bacteria, especially in members of the *Enterobacteriaceae* and *Pasteurellaceae* (Gammaproteobacteria) (Table S1). (ii) Exclusivity: the KEGG orthology analyses confirmed that the GN PICI-encoded genes are PICI specific, not being present in other families of MGEs (Tables S2–S4). These analyses were performed using three unrelated PICI elements, two from *E. coli*, EcCICFT073 and EcCl042, and one from *P. multocida*, PmCIATCC43137, as representatives of the GN PICIs. (iii) Integration: in each analyzed species, the PICI attC sites are never used by temperate prophages (Fig. S2, Table S1). (iv) *Transcriptional organization*: contrary to the PICIs from the GP bacteria, the PICIs in the GN species are transcribed unidirectionally and rightward, unlike the vast majority of temperate phages but similar to a few (for example, phages Mu or Sp18, in *E. coli* O157 Sakai [25]). This organizational differentiation suggests that the elements from GN and GP bacteria comprise at least two different lineages. (v) Replication origins: The region immediately 3′ to the replication initiation gene in the GP PICIs represents the replication origin [3]. Examination of this region for several of the PICIs from the GN organisms reveals an organization and functionality identical to that of the GP PICIs (Fig. 2a). (vi) Capsid morphogenesis: some of the elements encode homologs of the PICI capsid morphogenesis genes (Fig. 1). (vii) Interference: The two PICIs analyzed in detail show interference mechanisms similar to those previously identified in the SaPIs and in other PICI elements from Gram-positive bacteria, representing a fascinating example of convergent evolution (see below). (viii) Accessory genes: Many of the putative PICIs carry identifiable genes that do not appear to be involved in the PICI lifecycle. These accessory genes are carried exclusively by the PICIs. As noted, some are known to be involved in virulence.

**Induction and transfer**

GP PICI DNA is packaged in particles composed of phage virion proteins [5, 6]. While some GP PICIs express a small terminase subunit (TerS) homolog [7], others are highly
**Staphylococcus aureus**

**Pasteurella multocida**

**Enterococcus faecalis**

**Escherichia coli**

![Genome maps for PICIs](image)

**Fig. 1** Genome maps for PICIs. Genomes are aligned according to the prophage convention, with the integrase gene (int) at the left end. Genes are colored according to their sequence and function: int is yellow; transcription regulator (stl or rpr (GP), and alpA or merR (GN)) is dark blue; replication genes are purple; encapsidation genes are green, with the terminase small subunit gene (terS) in light green; superantigen and other virulence genes are pink; genes encoding putative phage resistance proteins are black; other accessory genes are red; genes encoding hypothetical proteins are white.

**Flanking genes**  
**Attachment sites (attL/attR)**  
**Replication origin (ori)**  
**Integration (int)**  
**Regulatory (alpA/stl/rpr)**

**Replication (pri)**  
**Virulence (rdgR/terS)**  
**Phage interference (abiF)**  
**Accessory**  
**Hypothetical**  

**cos site**

**Staphylococcus aureus**

**Pasteurella multocida**

**Enterococcus faecalis**

**Escherichia coli**
there are always two sets of iterons represented by arrows, and their sequences are shown at left. Note that there are always two sets of iterons flanking an AT-rich region, which could be the melting site. Some GP PICIs direct the formation of capsids with putative PICI-like elements in different phages. Since our genomic scrutiny had revealed the presence of PICIs. The iterons are represented by arrows, and their sequences are shown at left. Note that there are always two sets of iterons flanking an AT-rich region, which could be the melting site. Some, but not all, of the newly identified PICIs also contain a tetS homolog (shown in light green); we have to date identified both the helper phages and the interfering mechanisms for two of these: EcCICFT073 and PmCII72.

ECICFT073

Since our genomic scrutiny had revealed the presence of putative PICI-like elements in different E. coli strains, we wanted to characterize some of them in more detail. For that, we inserted a cat marker into the elements present in strains IHE3034, 18098 and O42, and a tetA marker into the element present in the CFT073 strain. The localization of the antibiotic resistant markers in the PICI elements is shown in Fig. S3. While the two first islands encode TerS, the last two do not. The strains carrying the putative GN PICIs were treated with mitomycin C (MC) and the transfer of these elements after induction of the resident prophages was tested. As shown in Table 1, the ECICFT073 element, but not the other three, was transferred at high frequency to the non-lysogenic DH5α (recA mutant), C600, 594 and WG5 strains. Moreover, MC induction of a CFT073 culture resulted in PICI DNA amplification, although no “PICI monomer” band appeared (Fig. 3). There was, however, a band containing covalently closed circular (CCC) DNA, indicative of PICI excision. On the basis of the following tests, we suggest that these results are indicative of helper phage induction of PICI excision, replication and packaging: we inserted a tetA marker into the chromosomal lac gene of the ECICFT073 host strain, the uropathogenic CFT073 strain, and tested the transfer of this marker by the resident prophages. Contrary to the transfer observed for the ECICFT073 tetA element, we did not detect transfer of the chromosomal tetA to the recipient strains, suggesting that none of the resident prophages is a generalized transducing phage. To further confirm that the transfer of ECICFT073 tetA was PICI-specific rather than generalized transduction, we mutationally inactivated the ECICFT073 tetA int gene and found that this abolished detectable excision and transfer (Fig. 3; Table 1). We also mutated the pri gene and found that this greatly reduced but did not completely eliminate transfer (Fig. 3; Table 1).

We next tested for MC induction and PICI transfer with the non-lysogenic C600 derivative carrying the ECICFT073 tetA element, and failed to detect amplification, lysis, or transfer, confirming that the ECICFT073 transfer depends on a helper phage. Finally, we identified the ECICFT073 helper phage. The CFT073 genome contains at least 5 prophage-like elements [28], of which #3 is actually ECICFT073 and is listed as O-island 51. We deleted each of the other 4 and found that deletion of #4 eliminated detectable ECICFT073 tetA induction and transfer (Fig. 3; Table 1), whereas none of the other prophage deletions had any effect.

**Functionality of the ECICFT073 genes**

Having established that the ECICFT073 island can be phage-induced, we looked in more detail at the ECICFT073-encoded genes to confirm that these ORFs encode proteins functional in the various aspects of the PICI life cycle as previously defined for the SaPIs. There are 5 key features of the PICI life cycle: (i) excision, which may be spontaneous, (ii) replication, (iii) specific packaging, (iv) interference, and (v) induction. Of these, induction depends on the helper phage and in the GP PICIs controls all of the aforementioned features.

**Integration**

To test for int (excision/circularization/integration) function, we initially performed PCR analysis with inward- and outward-directed primers as shown in Supplementary Fig. 4. Amplicons were obtained indicative of spontaneous excision and circularization, suggesting int functionality. To clearly prove this, we cloned the int-attP$_{int}$ segment into the

![Fig. 2 Characterization of GN PICI replication origins.](image-url)

**Fig. 2** Characterization of GN PICI replication origins. a Comparative map of the replication origins of several E. coli PICIs. The iterons are represented by arrows, and their sequences are shown at left. Note that there are always two sets of iterons flanking an AT-rich region, which could be the melting site. Some GP PICIs direct the formation of capsids with putative PICI-like elements in different phages. Since our genomic scrutiny had revealed the presence of PICIs. The iterons are represented by arrows, and their sequences are shown at left. Note that there are always two sets of iterons flanking an AT-rich region, which could be the melting site. Some, but not all, of the newly identified PICIs also contain a tetS homolog (shown in light green); we have to date identified both the helper phages and the interfering mechanisms for two of these: ECICFT073 and PmCII72.

### Table 1

| PICI          | Iteron sequence | Iteron organization |
|--------------|-----------------|---------------------|
| ECICFT073    | ctaccc          |                     |
| ECIO42       | aagtg            |                     |
| ECIE1D1a     | gcgggtt         |                     |
| ECIRM13514   | gttcac           |                     |

packed because they carry the phage pac or cos sequence [14, 22]. Some GP PICIs direct the formation of capsids 1/3 the size of the helper phage capsid, using different strategies [7, 8, 22, 26, 27]. Some, but not all, of the newly identified PICIs also contain a tetS homolog (shown in light green); we have to date identified both the helper phages and the interfering mechanisms for two of these: ECICFT073 and PmCII72.

### ECICFT073

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![image-url]
thermosensitive pMAK700 plasmid and tested it for integration in DH5α (recA mutant) by overnight growth followed by plating on selective agar at the restrictive temperature. As controls, we generated pMAK700 derivative plasmid mutants in the int gene or in the attP site. We obtained colonies only from plasmids carrying an intact int-attP complex, confirming int functionality.

Table 1 Effects of PICI and phage mutations on PICI transfer

| Donor strain            | C600   | WG5    | DH5α   | 594    |
|-------------------------|--------|--------|--------|--------|
| CFT073 c1501::tetA      | 8 × 10⁵ | 3.4 × 10⁵ | 4.07 × 10⁵ | 4.2 × 10⁵ |
| CFT073 lacZ::tetA       | < 1    | < 1    | < 1    | < 1    |
| CFT073 c1499::cat       | 1 × 10⁵ | 3.6 × 10⁵ | 2 × 10⁵  | 2.1 × 10⁵ |
| IHE3034 RS04635-RS04640::cat | < 1 | 40     | < 1    | < 1    |
| O42 1749::cat           | < 1    | < 1    | < 1    | < 1    |
| 18098 ORF17-ORF18::cat  | 80     | 110    | < 1    | 10     |
| PICICFT073 mutants      |        |        |        |        |
| CFT073 c1501::tetA Δint | < 1    | < 1    | < 1    | < 1    |
| CFT073 c1501::tetA ΔalpA| < 1    | < 1    | < 1    | < 1    |
| CFT073 c1501::tetA Δpri-rep| 50 | 87    | 45     | 37     |
| CFT073 c1501::tetA Δc1499| 3.8 × 10⁵ | 1.64 × 10⁵ | 2.37 × 10⁵ | 2.5 × 10⁵ |
| CFT073 c1501::tetA Δc1500| 8.7 × 10⁵ | 2.17 × 10⁵ | 5.37 × 10⁵ | 6.87 × 10⁵ |
| Other helper phages     |        |        |        |        |
| C600 λ EcCICFT073 c1499::cat | 2.2 × 10³ | 1.3 × 10³ | 1.1 × 10³ | 1 × 10³ |
| 594 80 EcCICFT073 c1499::cat | 1.33 × 10⁶ | 2.68 × 10⁶ | 8 × 10⁵  | 1.02 × 10⁶ |
| 594 80 ΔterS EcCICFT073 c1499::cat | < 1 | < 1    | < 1    | < 1    |
| CFT073 Prophage mutants |        |        |        |        |
| CFT073 c1501::tetA Δø1  | 1.01 × 10⁴ |        |        |        |
| CFT073 c1501::tetA Δø2  | 5.19 × 10³ |        |        |        |
| CFT073 c1501::tetA Δø4  | < 1    |        |        |        |
| CFT073 c1501::tetA Δø5  | 3.90 × 10³ |        |        |        |
| CFT073 c1499::cat Δø4   | < 1    |        |        |        |

aTransductants/ml of lysate, using E. coli C600, WG5, 594 or DH5α (recA mutant) as recipient strains. The means of results from three independent experiments are presented. Variation was within ± 5% in all cases.
bPhages deleted in strain E. coli CFT073

Fig. 3 Phage induction of EcCICFT073. CFT073 strains were SOS induced with MC (2 μg/ml). Samples were removed at the indicated time points and used to prepare minilysates, which were resolved on an 0.7% agarose gel (upper panel), and Southern blotted (lower panel) with an EcCICFT073 probe. M: Southern blot molecular marker (DNA molecular weight marker VII; Roche)

Replication

The previous results suggested that the GN PICIs encode a functional replicon. We have shown previously that the Pri-Rep-ori segment of SaPlbov1, SaPIII or EcIV583 can drive autonomous replication of a plasmid in different GP bacteria [3, 10, 22]. Since the putative PICIs seem to have the same replicon organization (Fig. 2a), we cloned the replication region of EcCICFT073, including the rep gene and the putative ori site, into the thermo-sensitive plasmid pKO3-blue [29], with rep under the control of an arabinose-inducible promoter. We also constructed plasmids carrying mutations in the rep gene or in the ori site. These plasmids were transformed into E. coli DH5α, and 5 × 10⁵ cells were plated on selective agar and incubated either at the permissive (30 °C) or the restrictive temperature (44 °C). Strains carrying the putative PICI rep-ori replicon generated colonies in E. coli DH5α (Fig. 2b), while the strains carrying the Δrep mutant plasmid did not. Note that at 44 °C, the number of colonies obtained was reduced compared to 30 °C, suggesting defects on plasmid segregation. Since the deletion of the ori significantly reduced the number of colonies obtained compared to the
that in the EcCICFT073 resistant marker did not change the size of the island. Note additional version of the PICI in which the antibiotic of the island (Fig. S3). In the new variant, the size increased by the insertion of the cassette in the middle of plasmid, these results confirm that the GN PICIs encode a functional replication module.

Packaging

As with the phages, there are two strategies by which the SaPIs can be packaged. Some SaPIs that use the headful mechanism encode a homolog of the phage terminase small subunit (TerS) that directs the preferential packaging of the SaPIs [7]. The TerS has the same function as the phage enzyme (TerSp), recognizing a specific (pac) site on its cognate genome to initiate packaging. Other SaPIs carry the helper phage cos sequence in the SaPI genome, which ensures SaPI packaging depending on the helper cos phage machinery [14]. The λ cos sequence contains three regions required to interact with the packaging machinery: cosQ, cosN, and cosB. cosQ is required for the termination of chromosome packaging. cosN is the site at which terminase cuts the DNA. Initiation of DNA packaging requires cosN and the adjacent cosB site. cosB consists of three binding sites, R3, R2, and R1, for λ TerS [30] (Fig. S5). EcCICFT073 does not encode a TerS homolog, suggesting that in order to be efficiently packaged EcCICFT073 probably uses the cos packaging strategy. As shown in Fig S5, EcCICFT073 carries two cosN sequences that are identical to those present in its inducing phage 4 (Fig. S5). Remarkably, these EcCICFT073 cosN sequences resemble that present in the archetypical E. coli λ and φ80 phages, used during decades as models in phage studies. However, we were unable to identify the phage cosB region in the EcCICFT073 element (Fig. S5), questioning if the island carries a bona fide cos sequence. To test the EcCICFT073 and phage cos sites for function, we introduced independently the λ, φ80, phage 4, and each of the two EcCICFT073 cos sites (containing the putative cosQ, cosN and cosB) into plasmid pET28a, which was not transferrable by λ, and found that the cloned cos sites enabled transfer of the plasmids by phages λ and φ80 (Fig. S5). Consistent with the presence of a completely different cosB sequence, transfer of the plasmids carrying the EcCICFT073 cos sites was reduced compared to that observed with the plasmids carrying the cognate phage cos sequences (Fig. S5).

The results described above raised the exciting possibility that phages λ and φ80 could work as helpers for EcCICFT073. To test that, and assuming EcCICFT073 is transferred using a cos mechanism, we generated one additional version of the PICI in which the antibiotic resistant marker did not change the size of the island. Note that in the EcCICFT073 tetA island the size of the element was increased by the insertion of the cassette in the middle of the island (Fig. S3). In the new variant, the cat marker replaced gene EcCICFT073 c1499 (Fig. S3), for which we have not been able to find a phenotype. The new island, generated in strain CFT073, was also introduced in the CFT073 mutant in the helper phage 4 and in the λ and φ80 lysogens. The different strains were SOS (MC) induced, and the transfer of the EcCICFT073 cat island analyzed. The analysis of the lysates obtained from the different CFT073 derivative strains revealed that transfer of EcCICFT073 cat by helper phage 4 was higher than that previously observed for the EcCICFT073 tetA island (Table 1), supporting the idea that EcCICFT073 uses a cos mechanism for packaging. Remarkably, phages λ and 80 transferred the EcCICFT073 cat at frequencies, for phage 80, significantly higher than those observed with the helper phage 4 (Table 1).

We next generated a set of mutant strains in which either one of each or both cos sites present in EcCICFT073 were deleted. Deletion of the EcCICFT073 cos1 site significantly reduced transfer of the element by phages λ and φ80, while deletion of the EcCICFT073 cos2 site slightly increased phage φ80-mediated transfer of the island but significantly increased transfer mediated by phage λ; deletion of both cos sites eliminated phage-mediated transfer of the island, confirming the identity of these sequences as cos sites and that both cos sites are functional (Fig. 4). This situation mirrors SaPIbov4 [31], which also has two putative cos site in its sequence.

Finally, we generated a phage 80 terS mutant, and tested for both phage and EcCICFT073 transfer (Table 1). Deletion of the phage-encoded terS gene abolished packaging and transfer of both elements, confirming that EcCICFT073 hijacks the phage proteins for transfer.
Interference

EcCICFT073 severely interferes with phage λ reproduction (Fig. 5). A conserved mechanism of phage interference in many PICIs is the production of PICI-sized particles that are too small to package an intact phage genome [27, 32]. To test this possibility, we MC-induced the lysogenic strain carrying phage λ and EcCICFT073, the virions were pelleted and the extracted DNA was analyzed. All of the packaged DNA was phage-sized, ruling out the possibility that the observed interference was generated by capsid size redirection.

Next we searched for putative genes in the EcCICFT073 element that could be involved in this interference. EcCICFT073 encodes a putative capsid protein (ORF c1499; AAN79968) and a head decoration protein (ORF c1500; AAN79969.1). Since in SaPIbov5 the capsid protein homolog (Ccm) blocks phage reproduction by generating SaPI-sized capsids [8], we analyzed whether these proteins could be involved in the observed interference using an alternative mechanism. To do that, we deleted genes c1499 or c1500 in the EcCICFT073 tetA element present in the non-lysogenic strain C600, and tested the ability of λ to generate plaques in these strains. None of the mutants allowed normal phage reproduction (Fig. 5), suggesting that other EcCICFT073 genes are responsible for the observed interference. Moreover, deletion of the c1499 or c1500 genes in the CFT073 strain did not affect the transfer of the island by the resident prophages (Table 1).

Induction

The main difference between the GP and GN PICIs lies in the regulatory region (Fig. 1 and S1). While the GP elements encode a master repressor (called Stl in the SaPIs or generically Rpr in the other GP elements) [10, 22], the existence of a global repressor in the GN elements is unclear. One attractive candidate is AlpA, homologs of this protein being encoded by all the *E. coli* PICIs (Fig. 1 and S1). AlpA is predicted to act as a DNA-binding transcriptional regulator, and previous work suggested that it controls the expression of *int* in a P4-like phage [33, 34]. We hypothesized that AlpA could act as the main repressor of the PICIs, blocking expression of the PICI genes. To test this, we initially generated different transcriptional fusions in which the *lacZ* reporter was fused either to EcCICFT073 ORF3 or was located after the stop codon of *alpA* (see scheme in Fig. 6a). Expression of the reporter was measured in the presence or absence of *alpA*. Contrary to our hypothesis, AlpA is not a repressor but seems to be an activator of the EcCICFT073 genes. To confirm this, and also to know if AlpA controls *int* expression, we fused the promoter regions of *alpA* (plasmid P*alpA*) or *int* (plasmid P*int*) to *lacZ*, using plasmid pRW224 (see scheme in Fig. 6c). Simultaneously, the *alpA* gene was cloned in plasmid pBAD18 under the control of the P*BAD* promoter, inducible by arabinose, generating plasmid pBAD-alpA. Plasmids P*alpA* or P*int* were introduced into the strains carrying plasmids pBAD-alpA or pBAD18, and the expression of the different promoter regions monitored in presence or absence of arabinose. As expected, AlpA activates its own expression. Contrary to previous results [33, 34], AlpA does not seem to control *int* expression in EcCICFT073.

Our results suggested that induction of the GN PICIs requires the expression of the AlpA activator. In support of this, inactivation of *alpA* significantly diminished induction and transfer of the EcCICFT073 island present in the CFT073 strain (Fig. 3, Table 1). Moreover, over-expression of AlpA in the non-lysogenic *E. coli* C600 carrying the EcCICFT073 island resulted in induction and uncontrolled replication of the EcCICFT073 element (Fig. 6e). In support of the previous results, this uncontrolled replication requires the expression of the EcCICFT073 encoded *pri* and *int*.
genes (Fig. 6e). Note that in this case, and since the strain is not lysogenic, just the closed circular (CCC) and the concatemeric (bulk) forms are observed because the element is not packaged. As occurred with the SaPIs [10], the uncontrolled replication of the EcCICFT073 affected both the growth curve and the viability of the E. coli cells (Fig. S6).

Finally, we introduced the pJP1290 plasmid, which measures expression of alpA, in strains C600 and 594 (non-lysogenic) and the C600 and 594 derivatives lysogenic for phages λ and φ80, respectively. Since AlpA is absolutely required for the transfer of the island, we hypothesized that the helper phages would induce expression of alpA. To test this, the different strains were MC-induced and the
expression of the blaZ reporter monitored. As expected, alpA expression does not depend on the SOS response but on the presence of a helper phage. Thus, alpA expression was increased in the induced lysogenic strains, compared to the induced non-lysogenic C600 and 594 strains (Fig. 6f). Note that alpA expression was also increased in the lysogenic strain carrying the phage φ80 mutant in the terS gene. Since this phage mutant can’t mobilize the island, this strain was used to propose that the EcCICFT073 element hijacks the phage machinery for packaging. However, another possibility could be that terS was the EcCICFT073 inducer. The fact that this mutant induces alpA expression rule out this possibility. The identification of the phage-encoded EcCICF073 inducer is now under study.

**PmCl172**

In a complementary study aimed at characterizing prophages present in the animal pathogen *P. multocida*, we MC-induced different lysogenic strains, the particles were pelleted and the extracted DNA was separated. As shown in Fig. 7, two bands, one of phage monomer size and the other of PICI monomer size, appeared in the analysis of strains Pm86 and Pm172. This is the classical pattern observed when a strain containing a PICI-helper phage pair is MC-induced, suggesting that *P. multocida* contains bona fide PICI elements, which are packaged in small capsids as are many of the analyzed PICIs [8, 27, 32, 35]. To confirm that this was the case, we initially sequenced both the *P. multocida* strains as well as the DNA obtained from the induced lysates. As expected, we identified a PICI element in both strains, which incidentally were identical (Fig. 1; Table S1). While two phages were present in strain Pm86 (Fig. S7), strain Pm172 only contained one (Fig. S7). The latter phage was also present in strain Pm86, suggesting this is the helper phage for the PmCl172 and PmCl186 PICIs. This phage belongs to the Mu family of phages, demonstrating for the first time a Mu-like phage, belonging to the *Myoviridae* family, as helper phage for a PICI element.

Next, we confirmed that the PmCl172- and Pm86-Mu phage are a bona fide PICI-helper phage interaction by performing a Southern blot analysis of the packaged DNA observed in Fig. 7, using PmCl172- and phage Mu-specific probes. Southern blot analyses revealed that the Mu phage probe hybridized both with the PICI-sized (small) and the phage-sized (large) DNA bands (Fig. 7), suggesting that the phage DNA can be packaged in the PmCl-sized particles. Packaging of a significant proportion of phage DNA in the small particles generates defective phages, confirming that PmCl172 and PmCl186 PICIs interfere with phage reproduction using a very well conserved strategy, production of PICI-sized capsids, which is a key feature of the previously analyzed GP PICI elements [8, 27, 32, 35].

Finally, to confirm the existence of the PICI-sized capsids, we subjected the virions to electron microscopy (EM). The phage-sized particles had the characteristic size and shape of this class of bacteriophages: hexagonal capsids of approximately 59 to 63 nm length and 51 to 56 nm width and long contractile tails, which are approximately 150 nm long and 15 to 16 nm wide (Fig. 7c). As expected, the majority of the particles had small, hexagonal heads, about 38–39 nm length and 37 nm wide, attached to approximately 150 nm long and 15 nm wide tail (Fig. 7c). This result confirms that PmCl172 causes the formation of small capsids, consistent with its smaller genome size. As occurs with SaPIbov5, which encodes a capsid protein responsible of the production of the SaPI-sized small capsids [8], PmCl172 also encodes a major capsid protein homolog (Fig. 1), which could be involved in this process. Unfortunately, the absence of tools to manipulate *P. multocida* has impaired us deciphering in detail the biology of the PmCl172 element.

**Discussion**

In this report we demonstrate that PICIs, of which the staphylococcal SaPIs are a subset, are not just confined to GP
bacteria but are widespread among other bacteria. Although all the GN PICIs identified in this study are in Gamma-proteobacteria, and more specifically in members of the Enterobacteriaceae and Pasteurellaceae, the fact that similar elements can be found both in GP and GN bacteria confirms the universality of this novel family of MGEs. PICIs are phage satellites with a unique genomic organization which is easy to see in KEGG genome maps and sets them apart from other mobile genetic elements. They interact with certain “helper” bacteriophages in a unique and highly characteristic manner: in the GP cocci, the PICI encode an SOS-insensitive master repressor, Stl, that maintains it in a

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**Fig. 7** Induction of PmCI172 and PmCI86. **a** DNA extracted from a phage lysates of MC-treated cultures of *P. multocida* strains 172 or 86. **b** Southern blot of the DNA shown in panel (a), using a phage- or PmCI-specific probe. **c** Electron microscopy analysis of the MC-induced *P. multocida* 172 lysate. Several different fields are shown, containing normal Mu phage particles and PmCI172 particles (arrows), which have smaller heads. Scale bars are 50 nm.

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**a**

*P. multocida* strains

|   | Pm86 | Pm172 |
|---|------|-------|
| Agarose gel | ![Agarose gel](image1) |

12Kb

**b**

*P. multocida* 172

|   | Mu phage | PmCI172 probe |
|---|----------|---------------|
| Southern blot | ![Southern blot](image2) |

**P. multocida* 86

|   | Mu phage | PmCI172 probe |
|---|----------|---------------|
| Southern blot | ![Southern blot](image3) |

**c**

Phage

![Phage analysis](image4)

PmCI172 sized particles

![PmCI172 sized particles](image5)
quiescent state and is counteracted by a specific phage protein, initiating the PICI life cycle [13, 22]. In the GN bacteria, it seems the PICIs use a completely different strategy coupling the phage and the PICI cycles. In this case, the GN PICIs encode AlpA, an activator of the PICI cycle whose expression depends on the helper phage. Why GP and GN bacteria use different strategies to be induced is a mystery. In both cases, however, PICI induction is followed by PICI replication, encapsidation of PICI DNA in infectious particles composed of phage proteins and finally high transfer of these elements.

Like their GP counterparts, GN PICIs also severely interfere with helper phage reproduction, this probably being one of the *raisons d’être* of these elements. The PICI genomes are usually about 1/3 the size of their helper phage genomes and among the GP PICIs, many encode proteins that redirect the helper phage to form capsids 1/3 the size of the phage capsid [7, 8, 27]. This strategy is also used by the GN PICIs. Although we have not identified the genes responsible for this process, none of the proteins encoded by the PmCI172 element resemble previously characterized proteins, suggesting that capsid size redirection is a convergent evolutionary process characteristic of all the PICI elements, which in turn defines the size of these elements.

As occurs with GP PICIs, GN PICIs have an important role in virulence and several of the newly identified PICIs present in GN bacteria contain recognizable accessory genes. These are listed in Table S1. The identities of many of these genes are based on annotations; just a few of them have been tested experimentally. Several elements, including islands harbored by variants of the notorious *E. coli* O157:H7 encode RgdR, which regulates expression of the type III secretion (T3S) system, encoded by the locus of enterocyte effacement (LEE) of the *E. coli* pathogenicity island, PAI-I. Deletion of this island in *E. coli* O157 strain TUV93-0 resulted in a reduction in LEE expression and T3S [36]. This deletion also reduced the capacity of the bacteria to attach to epithelial cells and reduced the shedding of *E. coli* O157 by sheep. RgdR also controls the expression of other genes, including those involved in motility, suggesting a global role in *E. coli* gene expression [36]. It is especially relevant for this report that EcCICFT073, and the related EcCI042, two of the *rgdR*-containing elements in *E. coli*, correspond to O-island 51, a 15 kb O157:H7 island annotated, as usual, as a defective prophage [36]. In addition to RgdR, which is encoded by all the analyzed *E. coli* PICIs, some *E. coli* PICIs encode allelic variants of PerC. Remarkably, some but not all the PICI-coded PerC homologs also activate expression of the LEE pathogenicity island [37].

We suggest that all of the PICIs in the GN species are co-ancestral and represent a separate but very similar lineage, since their transcriptional organization is mirrored in rare phages from GN organisms. At the same time, the unique organization of the replication origin is the same for both the GP and GN lineages, which is consistent with co-ancestry. Although this organization was not a criterion for the identification of the PICIs, it is shared by all those identified in this study and by no other known bacterial replicon. Coancestry could mean that the divergent evolution of PICIs and host chromosomes has simply been parallel, or that the PICIs have been horizontally transferred and have retained greater similarities than their host bacteria. Current studies in our labs are focused on deciphering these interesting evolutionary questions. Our overall view, therefore, is that the PICIs represent two or more families of unique mobile elements whose biological features have been strongly selected during long-term evolution, so that they represent coherent clades that have diverged from and evolved in parallel with temperate phages (or proto-phages) and have spread widely among diverse bacterial genera. If this spread has been via helper phage induction and packaging, then it is likely that spread would have occurred between species that share phage adsorption receptors, leading to a chain of island-hopping (or rather host-hopping) events that could, in principle, be tracked by comparing PICI gene similarities as a function of the relatedness of host bacterial phage receptors.

The P2-P4 system in *E. coli* represents a theoretical precedent for phage satellites, in which P4 utilizes P2 virion proteins to form small capsids into which its small genome is packaged [38, 39]. PICIs and P4 share some features in common: (i) phage-mediated activation of their life cycles, which in the case of P4 depends on the P2 Cox protein [40], while for the GN PICIs the activator remains to be identified; (ii) autonomous replication, which in the case of P4 depends on the α protein [41] and its cognate ori site (named ccr [42, 43]); (iii) carriage of the helper phage cos sequence in their genomes, which allows these elements to parasite the packaging machinery of their cognate helper phages; (iv) interference with phage reproduction, a process that in P4 depends on Sid, a protein that redirects capsid formation generating P4-sized particles [44].

However, and in spite of these similarities, P4 and the PICIs present in *E. coli* show important differences supporting the idea that they have evolved independently: (i) P4 and PICIs are fundamentally different both in genomic structure (see Fig. S8) and in gene content. As occurred with the PICIs, the KEGG analysis revealed that the P4 genes are exclusive of this family of elements (Table S5). Importantly, only three PICI-coded proteins (AlpA, Pri-Rep and Int) showed distant homology with the P4-coded proteins, corresponding to proteins that have similar roles in both elements. We have identified 4 allelic variants of AlpA in the *E. coli* PICIs (Fig. S9), all of them being remotely related to the AlpA-homolog (named Vis) encoded by P4.
(Fig. S9). In fact, these proteins seem to have a different role in each of the elements: while Vis is required for P4 excision [45], AlpA controls induction of the EcCICFT073 cycle. In the case of the Pri-Rep proteins, a similar scenario is observed: 4 allelic variants are found in the PICI elements from *E. coli* (Fig. S10), which are different in sequence from the α protein of P4 (Fig. S10). The situation with the Int proteins is even more interesting. We have identified in *E. coli* 36 attC sites for phages, 8 for PICIs and 4 for P4 elements (Fig. S2). All the attC sites are exclusive to one type of element except one, which is shared by PICIs and P4. Importantly, and with the exception of the int gene, the P4 and PICI elements that integrate in the same attC conserved their own features and are completely different both in sequence and structure (Fig. S11). (ii) PICIs do not have a plasmid state, nor encode genes to control plasmid copy number or plasmid segregation. (iii) PICIs play an important role in virulence, with many *E. coli* elements encoding two activators of the LEE pathogenicity island. (iv) P2 and P4 mutually cross induce one another through back-and-forth transcriptional activation. Indeed, P4 forms plaques on a P2 lysogen, but these are P2 plaques resulting from P4 induction of P2, whereas we have not been able to see phage induction mediated by an incoming GN PICI. (v) In *E. coli*, P4 and PICI elements seem to parasitize two completely different families of helper phages: *Myoviridae* and *Siphoviridae*, respectively. Taken together, all these similarities and divergences highlight the idea that although these elements have evolved independently, their parallel modus vivendi has been strongly selected in nature.

A similar scenario occurs with a family of pathogenicity islands recently identified in *Vibrio cholerae*. These elements, denominated PLEs for phage-inducible chromosomal island-like elements, are widespread genomic islands that share some features with the classical PICIs, including mobilization by helper phages and severe interference with phage reproduction [46, 47]. In contrast to the GN PICIs, PLEs are larger in size (18–19 kb) and have both different genetic organization and gene content. Thus, although they encode an integrase, they do not encode identifiable regulatory, replication or packaging modules, even though infection by a helper phage leads to PLE excision, replication and packaging [46]. Whether these functions depend on the helper phage machinery, or by contrast are provided by still-unidentified PLE genes remains to be determined. Further studies are required to understand the origins of, and relationships among, these subcellular elements and their cognate phages.

Although cos sites were long ago cloned into *E. coli* plasmids, generating cosmids, which are widely used as cloning vectors owing to their efficient transfer by phage λ, our results represent the first demonstration of naturally-occurring phage λ-mediated transfer. Since SaP1bov5 is also transferred by helper cos phages, our results with the PICIs represent a paradigm shift involving cos phages in gene transfer. The PICIs are a special case because they undergo phage-like replication, generating concatemeric post-replicative DNA, which would have cos sites spaced one genome apart. However, EcCICFT073 lacks the capsid morphogenesis genes carried by most SaPIs and is therefore packaged in full-sized phage capsids, which are designed to accommodate the 45 kb phage genomes. This would mean that the phage terminase skips two successive cos sites during packaging; the mechanism by which this occurs remains to be determined, although it is likely that the expansion generated once the capsid is fulfilled controls this phenomenon.

Bacteria are successful as commensal organisms or pathogens in part because they adapt rapidly to selective pressures. MGEs play a central role in this adaptation process and are a means to transfer genetic information among and within bacterial species. Here, and as is the case with other classical MGEs like plasmids, transposon or phages, we demonstrate that the PICIs are widespread in nature. These findings represent the discovery of a universal agency of horizontal dissemination of important accessory genes in the bacterial universe.

**Materials and methods**

**Identification of PICI candidates**

The following criteria were initially used to identify putative PICIs in the GN genomes: (i) well-conserved gene organization, including an integrase, a replicating module and a master transcriptional regulator; (ii) unique attachment (att) sites that are never occupied by prophages; (iii) the absence of phage structural and lytic genes; and (iv) size typically around 12–15 kb. After this initial screening, the analysis of orthologies points to elements that might correspond to PICIs. Examination of the corresponding KEGG genome maps ([http://www.genome.jp/kegg](http://www.genome.jp/kegg); release May 1 2016) was used to confirm the identifications. In these analyses we examined the *E. coli* and the *P. multocida* genomes that have been coded for KEGG because the KEGG genome maps enable PICI-like elements to be readily identified.

**Bacterial strains and growth conditions**

Bacterial strains used in these studies are listed in Supplementary Table S7. *E. coli* strains were grown at 37 °C or 30 °C overnight on Luria-Bertani agar and on Luria-Bertani broth with shaking (180 r.p.m.). *P. multocida* strains were grown at 37 °C overnight on Brain-Heart-
Infusion agar and on Brain-Heart-Infusion broth with shaking (250 r.p.m.). Ampicillin (100 µg ml⁻¹), Kanamycin (30 µg ml⁻¹), Chloramphenicol (20 µg ml⁻¹) or Tetracycline (20 µg ml⁻¹), from Sigma, were added when appropriate. Phage and PICI analyses were performed, as described previously [11–13].

DNA methods

Gene insertion was performed as described [48]. The tetracycline resistance marker (*tetA*) or chloramphenicol resistance marker (*cat*) was amplified by PCR with primers listed in Supplementary Table S6. The resulting PCR products were used to transform the recipient strain harboring plasmid pKD46 [48], which expresses Lambda Red recombinase. PCR was performed to verify the insertion of markers.

Plasmid construction

Plasmids used in this study were constructed by cloning PCR products amplified with oligonucleotide primers purchased from Invitrogen. Clones were sequenced by Eurofins. Plasmids and primers are listed in Supplementary Table S8 and Table S6.

Southern blot

Following plasmid (arabinose) or phage (MC) induction, samples were taken and pelleted at the indicated times. Samples were re-suspended in 50 µl lysis buffer (47.5 µl TES-Sucrose and 2.5 µl lysozyme [10 µg/ml]) and incubated at 37 °C for 1 h. Subsequently, 55 µl of SDS 2% proteinase K buffer (47.25 µl H2O, 5.25 µl SDS 20%, 2.5 µl proteinase K [20 mg/ml]) was added and incubated at 55 °C for 30 min. Samples were vortexed with 10 µl of 10× loading dye. Cycles of incubation in dry ice with ethanol and at 65 °C were performed. Chromosomal DNA was separated by agarose gel electrophoresis. Samples were run on 0.7% agarose gel at 30 V overnight. Nylon membranes (Hybond-N 0.45 mm pore size filters; Amersham Life Science) were used for the transfer of DNA. DNA was detected using a DIG-labeled probe and anti-DIG antibody.

β-galactosidase assays

Strains were grown at 37 °C in LB medium containing the appropriate antibiotics. Following plasmid (arabinose) or phage induction (MC), samples were taken and pelleted at the indicated times. The Miller method [49] was used to measure β-galactosidase levels. The average of at least three independent experiments is shown in Miller units.

Electron microscopy (P. multocida)

Ten milliliters of filtered *P. multocida* Pm172 phage lysate were centrifuged at 40,000 × g for 90 min at 4 °C. The supernatant was carefully removed and the pellet was re-suspended gently with 0.5 ml of 0.1 M ammonium acetate (pH 7.3). The suspension was stored overnight at 4 °C for negative staining and visualization by TEM. Three hundred-mesh carbon-coated nickel grids were dropped onto 50–100 µl of phage suspension. The grids were allowed to adsorb phage suspension for 2 min. The grids were washed 3 times with dH2O for 10°, excess fluid was removed using Whatman filter paper and the grids were negatively stained with 2% ammonium molybdate for 30°. Excess staining solution was removed with Whatman filter paper. The grids were allowed to dry at room temperature for ~15–20 min. The samples were examined by TEM (FEI Tecnai TF20) at 200 kV using Gatan Microscopy Suite Software.

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Author contributions JRP and RD designed the study; AF-S, RM-R, RFA, and JC performed the experiments; JRP and RD wrote the manuscript. All the authors approved the final manuscript.

Compliance with ethical standards

Conflict of interest The authors declare that they have no conflict of interest.

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Phage-inducible chromosomal islands are ubiquitous within the bacterial universe

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