Conjugal Transfer of the Pathogenicity Island ROD21 in *Salmonella enterica* serovar Enteritidis Depends on Environmental Conditions

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Abstract

Unstable pathogenicity islands are chromosomal elements that can be transferred from one bacterium to another. *Salmonella enterica* serovar Enteritidis (S. Enteritidis) is a pathogenic bacterium containing such unstable pathogenicity islands. One of them, denominated ROD21, is 26.5 kb in size and capable of excising from the chromosome in certain culture conditions, as well as during bacterial infection of phagocytic cells. In this study we have evaluated whether ROD21 can be effectively transferred from one bacterium to another. We generated a donor and several recipient strains of S. Enteritidis to carry out transfer assays in liquid LB medium. These assays showed that ROD21 is effectively transferred from donor to recipient strains of S. Enteritidis and S. Typhimurium. When *Escherichia coli* was used as the recipient strain, ROD21 transfer failed to be observed. Subsequently, we showed that a conjugative process was required for the transfer of the island and that changes in temperature and pH increased the transfer frequency between *Salmonella* strains. Our data indicate that ROD21 is an unstable pathogenicity island that can be transferred by conjugation in a species-specific manner between *Salmonellae*. Further, ROD21 transfer frequency increases in response to environmental changes, such as pH and temperature.

Introduction

Bacterial genomes evolve by various processes, such as mutations, chromosomal rearrangements, deletions of one or more genes or acquisition of foreign genetic material by horizontal gene transfer (HGT) [1]. The loss or gain of genetic material generates genomic changes that can rapidly and radically alter the lifestyle of the bacterium [2,3]. These appear to be key mechanisms used by bacteria to genetically adapt to new environments, which can have a strong impact for the emergence of new pathogenic versions of a particular bacterial strain. The processes of horizontal gene flow include mobile genetic elements, such as conjugative plasmids, transposons, insertion sequences, prophages, integrons and genomic islands. *Salmonella enterica* is one of the most interesting examples of HGT; due to the large number of plasmids, genomic islands and prophages that these pathogenic bacteria contain [4]. *Salmonella enterica* is a facultative intracellular enteric pathogen that affects a wide host range and has more than 2,500 serovars, of which the serovars Typhimurium and Enteritidis are of great impact in health in humans, as well as in animals of commercial importance.

The availability of the complete genome sequence of several *Salmonella enterica* serovars has allowed comparative genomic analyses that led to the identification of regions acquired through HGT. One of such HGT-acquired regions is ROD21 (Region of Difference 21, [5]), which is a 26.5 kb genomic island, located between the 2,061,170 and 2,087,657 coordinates of the *S. enterica* genome (NCTC13549). ROD21 can also be found in the genomes of *S. Gallinarum*, *S. Dublin* and *S. Typhi* [5]. The ROD21 region shows features that are characteristic of a pathogenicity island, such as insertion near to three copies of the gene coding for asparagine tRNA (*asnT*); a gene coding for an integrase at the 5′ end, similar to those present in phage P4; the region is flanked by direct repeat sequences (DRS) of 24 bp, which are identical to the last 22 bp of the gene coding for *asnT*; and a G+C percentage equal to 37.5% that is significantly different to the rest of the *Salmonella* genome (52.2%) [5,6]. Further, ROD21 is considered a pathogenicity island because it contains the gene *tlpA*, which encodes a protein involved in the virulence of the bacterium through modulation of the host defense mechanisms that determine the regulation of NF-kB and caspase-1 activation [7]. Furthermore, recent studies have suggested that other genes encoded in this pathogenicity island are required for *S.* Enteritidis to cause disease in mice [6,8]. Previous work from our laboratory have shown that ROD21 is an unstable pathogenicity island that excises from the chromosome by site-specific recombination, a process catalyzed by the P1 integrase and a putative excisionase (SEN1998), also encoded in this region [6,9]. These enzymes are
thought to recognize DRSs that flank the pathogenicity island [10] and produce a circular intermediate, which can either remain as an episomal element inside the bacterium or be degraded and disappear. Despite of this fact, previous studies have described that most clinical strains of S. Enteritidis isolated in various countries carry this pathogenicity island [6,11], suggesting that it is possible that ROD21 can be transferred between S. Enteritidis strains. Consistent with this notion, bioinformatic analyses have identified ORFs in this pathogenicity island that show sequence identity with proteins involved in conjugative mobilization of DNA, such as genes found in several conjugative plasmids, such as MoLA and TraD [6].

Based on the evidence described above, here we have determined whether this unstable pathogenicity island is transferable to recipient bacteria in vitro. Further, we have measured the transfer frequency and identified the mechanism responsible for this process. We observed that ROD21 could be transferred by conjugation between S. Enteritidis and between S. Enteritidis and S. Typhimurium. In addition, the frequency of ROD21 transfer can be modulated by changes in pH and temperature. Our data suggest that ROD21 can be transferred by conjugation between different Salmonella strains in a species-specific manner, a process that could be modulated by the environmental conditions confronted by bacteria.

Materials and Methods

Ethics Statement

All experimental procedures performed in this study were revised and approved by the Bioethics and Biosafety Committee of the School of Biological Sciences, Pontificia Universidad Católica de Chile. All animal work was performed according to the Guide for Care and Use of Laboratory Animals (National Institute of Health, USA) and Institutional guidelines. All experiments using animals were overseen by a Veterinarian at all times.

Strains and plasmids

Strains used in this study are listed in Table 1. Salmonella enterica serovar Enteritidis PT1 (S. Enteritidis) was provided by Mrs. Alda Fernández from Public Health Institute of Chile, Santiago, Chile; S. Enteritidis PT4 was provided by Dr. Carlos Santiviago from University of Chile, Santiago, Chile; Salmonella enterica serovar Typhimurium (S. Typhimurium) 14028s was provided by Dr. Guido Mora from University Andrés Bello, Santiago, Chile; and S. Typhimurium pSLT+ was provided by Dr. Josep Casadesús from Universidad de Sevilla, Sevilla, Spain. E. coli strains H10407 and E. coli EDL933, were both provided by Dr. Roberto Vidal from University of Chile and E. coli strain DH5α (naldixic acid resistant bacteria, NalR) was purchased from Invitrogen. These strains were stored at -80°C in LB medium supplemented with 20% glycerol and grown in liquid LB medium at 37°C with aeration provided by a shaker. Plasmids pKD3, pKD4 and pKD46 were obtained from the E. coli Genetic Resources at Yale CGSC, The Coli Genetic Stock Center. Plasmid pCLF1 was provided by Dr. Carlos Santiviago from the University of Chile, Santiago, Chile.

Donor strain (S. Enteritidis ROD21::aph) and recipient strains of ROD21 (S. Enteritidis AROD21 putAP::cat, S. Typhimurium putAP::cat, S. Typhimurium pSLT+ phoN::dhfr, H10407 putAP::cat, EDL933 putAP::cat) were generated by Lambda Red-mediated recombination, as described by Datsenko and Wanner [12]. Briefly, a PCR product encoding cat, aph or dhfr gene was generated by PCR amplification from pKD3, pKD4 or pCLF1 plasmid, respectively. The primers used in this study are listed in Table 2. To insert the aph gene into ROD21, we used primer ROD21::kan (H1+P1) and primer ROD21::kan (H2+P2) to generate a PCR product that was inserted between genes SEN1975 and SEN1976. We also designed a set of primers to insert the cat gene into the chromosome of S. Enteritidis PT1 strain that spontaneously lost ROD21, as described before [6]. These primers were putAP::Cm (H1+P1) and primer putAP::Cm (H2+P2) (Table 2). To insert the cat gene into S. Typhimurium 14028s chromosome we designed primers STM14028 putAP::Cm (H1+P1) and STM14028 putAP::Cm (H2+P2) (Table 2). To insert dhfr gene upstream to phoN gene into S. Typhimurium pSLT+, we designed primers STM_phoN_tmmp (H1+P1) and STM_phoN_tmmp (H2+P2) (Table 2). We also designed a set of primers to insert cat gene into E. coli H10407 and E. coli EDL933 strains. These were EHEC-ETEC putAP::Cm (H1+P1) and EHEC-ETEC putAP::Cm (H2+P2) (Table 2). The first 40 bp of all these primers aligned with the chromosome of the manipulated strains and the last 20 bp of these primers aligned with pKD3, pKD4 or pCLF1. Competent strains harboring the thermosensitive plasmid pKD46 were prepared as previously described [12] and PCR products containing cat, aph or dhfr genes were electrotropinized into these competent cells. After electrotropinization, bacteria were incubated for 1 h at 37°C with aeration in 1 mL of LB medium and then seeded in solid LB medium supplemented with 50 μg/mL kanamycin, 10 μg/mL chloramphenicol or 20 μg/mL trimethoprim. To verify the correct insertion of aph gene in the donor strain, a PCR amplification was performed with genomic DNA of the mutant strains and primers SEN1975 Fw and SEN1970. The primers used to generate the PCR product from pKD4 were SEN1980::Kan (H1+P1) with SEN1980::Kan (H2+P2). Capital letters account for sequence that hybridizes with the S. Enteritidis chromosome and lowercase letters account for sequence that hybridizes with pKD4 plasmid. To generate strain S. Enteritidis ROD21::aph/pBAD-SEN1998, donor strain S. Enteritidis ROD21::aph was transformed by electroporation with either plasmid pBAD-empty or pBAD-SEN1998, as described previously [9] and seeded in solid LB medium supplemented with 50 μg/mL kanamycin and 100 μg/mL ampicillin.

Molecular biology techniques

Bacterial genomic DNA was prepared using the phenol-chloroform method described by Sambrook et al [13]. Using genomic DNA from bacterial strains and primer pairs described in Table 2, PCR amplifications were performed in a MaxiGene Gradient Thermocycler (Axygen), using approximately 1 ng/mL of DNA, 1 nmol/mL of each primer, 0.2 mM deoxynucleoside triphosphates, 1.5 mM Magnesium Chloride and 50 U/mL of Taq DNA polymerase (Invitrogen), using standard PCR amplification cycles. PCR products were resolved by electrophoresis in 1% agarose gels containing 0.5 μg/mL ethidium bromide and visualized under a UV light transilluminator (UVP, Inc).

Broth mating assays

To assess the potential for horizontal transfer of ROD21 in LB broth (1% tryptone, 0.5% yeast extract, 1% NaCl), mates were conducted as described previously by Coburn et al [14]. In these
Table 1. Strains and plasmids used in this study.

| Strains                                         | Relevant genotype                                                                 | Reference |
|------------------------------------------------|------------------------------------------------------------------------------------|-----------|
| Salmonella enterica                             |                                                                                    |           |
| S. Enteritidis PT4                              | Reference strain NCTC 13349                                                       | [5]       |
| S. Enteritidis PT1                              | Clinical isolate, wild type strain                                                | [6]       |
| S. Enteritidis PT1 ROD21::aph                   | S. Enteritidis with aph gene inserted between 1975–1976 ORFs                     | This work |
| S. Enteritidis PT1 ROD2 pBAD-SEN1998            | S. Enteritidis strain that carries pBAD-SEN1998 plasmid                           | [9]       |
| S. Enteritidis PT1 ROD21::aph pBAD-SEN1998      | S. Enteritidis strain with aph gene inserted between 1975–1976 ORFs and carries pBAD-SEN1998 plasmid | This work |
| S. Enteritidis PT1 ROD21::aphΔ aph trg::mudQ    | S. Enteritidis strain lacking ROD21 and the cat gene inserted between putA and putP genes | This work |
| S. Typhimurium                                  | Reference strain ATCC 14028s, wild type                                            | [38]      |
| S. Typhimurium putAP::cat                       | S. Typhimurium strain with the cat gene inserted between putA and putP genes      |           |
| This work                                       |                                                                                    |           |
| S. Typhimurium pSLT::aph                        | S. Typhimurium strain that lacks the virulence plasmid                            | [16]      |
| S. Typhimurium pSLT::phoN::dhfr                 | S. Typhimurium strain that lacks the virulence plasmid and is resistant to trimethoprim | This work |
| Escherichia coli                                |                                                                                    |           |
| H10407                                          | Clinical isolate enterotoxigenic pathotype, wild type strain                      | [39]      |
| H10407 putAP::cat                               | E. coli strain with cat gene inserted between putA and putP genes                 | This work |
| EDL933                                          | Clinical isolate enterohaemorragic pathotype, wild type strain                    | [40]      |
| EDL933 putAP::cat                               | E. coli strain with cat gene inserted between putA and putP genes                 | This work |
| DH5α™                                           | E. coli strain (F−, Δ80lacZAM15, ΔlacZYA-argF) u169 recA1 endA1 hsdR17 (K−, mK+) phoA supE44 λ− thi-1 gyrA96 relA1) Invitrogen |
| Plasmids                                        |                                                                                    |           |
| pKD3                                           | Plasmid that contains cat gene                                                    | [12]      |
| pKD4                                           | Plasmid that contains aph gene                                                    | [12]      |
| pKD46                                          | Plasmid that contains Lambda red recombinase genes from phage Lambda              | [12]      |
| pCLF1                                          | Plasmid that contains dhfr gene                                                   | GenBank: HM047090.1 |

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experiments, strain S. Enteritidis ROD21::aph (donor) and S. Enteritidis ΔROD21 putAP::cat (recipient) were grown to stationary phase in LB broth, washed with phosphate-buffered saline (PBS) pH 7.4 before mixing 0.15 mL of S. Enteritidis ROD21::aph and 0.45 mL of S. Enteritidis ΔROD21 putAP::cat with 4.5 mL of pre-warmed LB broth. After incubation at 37°C for 4 h, aliquots were removed for quantitative determination of donors, recipients and transconjugants by seeding on solid LB medium supplemented with the appropriate antibiotics. Donor S. Enteritidis ROD21::aph was selected on solid LB medium supplemented with kanamycin, recipient S. Enteritidis ΔROD21 putAP::cat was selected on solid LB medium supplemented with chloramphenicol and transconjugants were selected on solid LB medium supplemented with kanamycin and chloramphenicol, or kanamycin and trimethoprim. Transfer frequency measurements were calculated by dividing the number of transconjugants by the viable counts of donor cells in the mating mixture, determined at the end of the 4 h period. Using the same approach, we performed broth mating assays to determine whether ROD21 was able to transfer to the other recipient strains as mentioned above: S. Typhimurium 14028s putAP::cat, S. Typhimurium pKD4, H10407 putAP::cat and EDL933 putAP::cat and E. coli DH5α NalR.

To rule out transfer of ROD21 by naked DNA uptake due to lysis of donor cells, we purified DNA from the donor strain S. Enteritidis ROD21::aph using the phenol-chloroform methodology. Then, 10 ng, 100 ng and 1,000 ng were resuspended in LB medium and mixed with 450 μL of stationary phase, PBS-washed, Salmonella recipient strains. The mixture was inoculated in 4.5 mL pre-warmed LB medium in the same conditions described above. To test whether ROD21 was transferred by phage transduction, a liquid culture of the donor strain S. Enteritidis ROD21::aph was filtered using a 0.2 μm pore size polycarbonate filter (Orange Scientific Braine-l’Alleud, Belgium) and the supernatants containing phages were treated with 10 units of RQ1 DNase (Promega, Madison, WI). For these experiments, 150 μL of the treated supernatants were mixed with 450 μL of stationary phase, PBS-washed, Salmonella or E. coli recipient strains. Then, the mixture was inoculated in 4.5 mL pre-warmed LB medium under the same conditions described above.

Filter mating assays

Filter mating assays were carried out employing a donor to recipient ratio equal to 1:10. Stationary phase cultures of the donor (S. Enteritidis ROD21::aph, S. Typhimurium ROD21::aph tyg::mudQ pSLT+) or S. Typhimurium ROD21::aph tyg::mudQ pSLT−) and recipient (S. Enteritidis ΔROD21 putAP::cat, S. Typhimurium 14028s putAP::cat, S. Typhimurium pKD4, DH5α NalR), S. Typhimurium pKD4, DH5α NalR, S. Typhimurium pKD4, DH5α NalR, E. coli H10407 putAP::cat, E. coli EDL933 putAP::cat or E. coli DH5α NalR bacteria were mixed in LB medium (0.5 mL of the donor and 4.5 mL of the recipient) and then collected on a 0.2 μm pore size mixed cellulose/ester membrane filter (Advantec). The filter was
placed with bacteria side down on a LB agar plate and incubated at 37°C for 4 h. Bacterial cells were harvested from the filter and resuspended to a final volume of 1 mL in LB and seeded on LB agar plates containing the respective antibiotics for selection. Transconjugants were selected with kanamycin and chloramphenicol or kanamycin and nalidixic acid. Transfer frequency measurements were obtained by dividing the number of transconjugants by the viable counts of donor cells in the mating mixture, determined at the end of the mating assay.

Oxidative stress, temperature and pH treatments

To evaluate whether environmental conditions, such as oxidative stress, temperature and pH could influence the transfer of ROD21, we measured the transfer frequency in bacteria grown in the presence of hydrogen peroxide (H₂O₂), in bacteria grown at 23°C, 37°C or 43°C and in bacteria grown at pH 7 or pH 5. All treatments were performed in separate experiments. As mentioned above, donor and recipient bacteria were grown overnight at 37°C to stationary phase and aliquots were co-incubated for 4 h in LB medium. For oxidative stress assays, co-cultures were incubated at

| Primers used to verify the genotype | N° | Primer | Sequence (5’→3’) |
|-----------------------------------|----|--------|------------------|
| 1 | SEN 1988 Fw | GACGGAATCTTTTTGCCCTG |
| 2 | SEN 1970 (integrase) Rev | CGGCAATCTTTCCTGACCAT |
| 3 | SEN 1999 Fw | CAGCAAGACCCCTGCAAGAGT |
| 4 | SEN 2001 Rev | AGTGGGCCATTTGGGATCAG |
| 5 | SEN yeeO Fw | TCATAATACACGCACTGG |
| 6 | SEN 2013 Rev | CGTCCACGACCTCATTAAAC |
| 7 | putAP Fw | ATGTGACCTGCGTTGGCAGC |
| 8 | putAP Rev | GCCCCCTTGAGCATCTGCGA |
| 9 | SEN1975 Fw | TTCTGATGACGACGTAAAGAGG |
| 10 | SEN1976 Rev | TGGTGTTGAAAGAAGATGATCCTT |
| 11 | LepA Gifsy-1 Fw | GCCCAGGATTTATCGGAACAGA |
| 12 | GobB Gifsy-1 Rev | GGTGATGTCGCCCTGTACT |

Table 2. Primers and probes used in this study.

| Primers used to insert resistance genes | Primer | Sequence (5’→3’) |
|----------------------------------------|--------|------------------|
| SEN 1970 (H1+P1)                      | ATGTGCGTAAAGCTGTCATGCTTGATTTGGTACCGACCGGtcgtagcgggagtccc |
| putAP:cm                              | AAATGTTAAAAAGAGAAGGGGTCGTAGCCGGGAAAAGtgtagcgggagtccc |
| SEN 1980:kan (H1+P1)                  | AAAACTACGCGACCTTGGTTATGTTGAGTTTACTTTGAgatcgggagtccc |
| SEN 1980:kan (H2+P2)                  | CAGGTATGACGCTTAAATTTGATTTCTACATTGatcgggagtccc |
| EHEC-ETEC putAP::Cm                   | ATTCACAATGATTTAAACACACACCATATTATTTAatcgggagtccc |
| ROD21::kan (H1+P1)                    | GGAATCTCCATTGGAGAGAAATACATATACCTGGTGGAAAAAGtgtagcgggagtccc |
| ROD21::kan (H2+P2)                    | ATGTGTTGTTATAACATTATAAAAAATTTTACATTTTAAAGtgtagcgggagtccc |
| STM 14028 putAP::Cm                   | AAAAGTGCTCAGCTGAAATTGAGAGTGAACCTGAACCTATTTTAAAGtgtagcgggagtccc |
| STM 14028 putAP::Cm                   | CAGTGGCAAGGGTGTTATTGGCTACCGGCGACGTACCCGtcgtagcgggagtccc |
| STM_phoN_tmtp (H1+P1)                 | CAGACCGCAGTTTACCCGTGCTACCGCCGACCGACGTATGcgaatcttccttcagcgg |
| STM_phoN_tmtp (H2+P2)                 | GGGCGCCGGGGCTGGAATGCCTTCGTTTTCCCGGATTTGGGCAGGxcgtagcgggagtccc |

| Primers and probes used to detect attB1 | Primer | Sequence |
|----------------------------------------|--------|----------|
| attB1-nested-RT-Fw | GTTACTATGCCGGCGCGCTTCAAC |
| attB1-nested-RT-Rev | CCGATTAGCCCCAAAAACTATAG |
| rpoD-RT-Fw | GTGAGCCGGAGAGGCGAAAA |
| rpoD-RT-Rev | CAGAACCGGAGTGTGCAGG |
| attB1 probe | TTCGAGTACCAGTCAAGGA |
| rpoD probe | CGACATGCCTACAC |

The location of primers pairs from 1 to 6 is shown in Figures 1 and 2. Lowercase indicate the region that hybridizes to the 5’ or 3’ end of the pKD3 or pKD4 plasmids.

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Figure 1. Schematic representation of the modified genetic regions of the strains used in the _S. Enteritidis-to-S. Enteritidis_ mating assays. (A) Donor strain, showing excision type 1 and 2; (B) recipient strain and (C) transconjugant strain. Numbers above of each scheme represent the coordinates in the chromosome of _S. Enteritidis_. Asparagine tRNA genes are designated as _asnT-1_, -2 and -3. DRS stands for Direct Repeated Sequence. Black arrows represent the inserted antibiotic gene (_aph_ or _cat_). Thin arrows represent primers used for PCR verification of the strains: (1)
left-end of ROD21 (1,011 bp), (2) DRS (914 bp), (3) right-end (903 bp), (4) insertion of cat gene (1,030 bp without cat insert, 1,943 bp with cat insert), and (5) insertion of aph gene (2,469 bp). Below each scheme, agarose gels show PCR products obtained for the recipient and transconjugant S. Enteritidis strains.

37°C in LB medium supplemented with 0.25 mM H2O2 (Merck). For temperature treatments, co-cultures were incubated in LB medium at 23°C, 37°C or 43°C. Finally, for pH treatments, donor and recipient bacteria were grown at 37°C in LB medium adjusted to pH 5 with HCl (Merck). The frequency of ROD21 transfer was determined as mentioned above.

To detect ROD21 excision in the conditions described above, 1 mL of each bacterial culture was used to prepare genomic DNA and detection of ROD21 excision was performed with TaqMan MGB probes and TaqMan Universal PCR Master Mix (Applied BioSystems), as recommended by the manufacturer. Primers and probes used to detect ROD21 excision (attB-1) were: attB1-nested-RT-Fw, attB1-nested-RT-Rev and attB1 probe (Table 2). Primers and the probe used to detect tpoD were: tpoD-RT-Fw, tpoD-RT-Rev and tpoD probe (Table 2). The reaction mixture contained 1 μL of genomic DNA as a template, 0.3 pmol/μL of each primer and 0.3 pmol/μL of TaqMan MGB probes. Standard curves for attB-1 and tpoD were generated using serial dilutions of a plasmid containing the corresponding PCR fragment for attB-1 and tpoD. Thermal cycling conditions were: 1 cycle of 5 min at 50°C and 10 min at 95°C and 40 cycles of 15 sec at 95°C, 15 sec at 58°C and 1 min at 60°C. The copy number of attB-1 and tpoD was calculated as follows: the number of attB-1 and tpoD copies was determined by the ratio between the amount of DNA in a sample and the weight of one molecule of the plasmid with the insert. To graph the standard curve, the value of the threshold cycle (Ct) was confronted with the log10 of the initial copy number of each sample, generating a linear relationship to calculate the number of copies in the sample, which has a specific Ct. The values were expressed as the ratio of attB-1 copy number/tpoD copy number and for each condition evaluated, the fold increase over the basal condition (pH 7, 37°C and absence of hydrogen peroxide) was plotted.

Survival assays

Groups of 4 male C57BL/6 mice (5–6 week age) were used to evaluate the virulence of WT S. Typhimurium and the strain S. Typhimurium-ROD21. To perform infection assays, bacteria were grown in LB medium at 37°C until an OD600 equal to 0.6 was reached. The volume of bacterial culture containing 1 x 10^7 CFUs was centrifuged in a refrigerated microcentrifuge (CT15RE Hitachi) at 10,000 x g for 5 minutes. Bacterial pellets were thoroughly resuspended in 200 μL of PBS and used to infect mice intragastrically. Serial dilutions of the bacterial inoculum were seeded in LB plates to corroborate infective doses. After infection, survival rate was recorded every 12 h. Mice that lost more than 25% of their initial body weight after infection were euthanized.

Statistical analyses

For all experiments, 3 independent assays were performed (n = 3). Data and statistical analysis were performed using Prism 4 software (Graph Pad Software, Inc.). For statistical analyses unpaired Student’s t test, Wilcoxon signed rank test and Mann-Whitney test were used when the data were not in a Gaussian distribution, or two ways ANOVA, when the distribution of data was Gaussian. The survival analyses were performed with Kaplan-Meier with Mantel-Cox post-test. In any case, p values below 0.05 were considered statistically significant.

Results

ROD21 is transferred between Salmonella strains

As we have shown previously, ROD21 can excise from the chromosome of S. Enteritidis in at least 2 different ways (excision type 1 and excision type 2) [6]. When excision type 1 takes place, genes from SEN1970 to SEN1999 excise from the chromosome and form an episomal element. On the other hand, excision type 2 promotes the excision of an episomal element including genes SEN1970 to SEN2011 (Figure 1A). To evaluate which episomal element can be transferred from S. Enteritidis to a ROD21-deficient S. Enteritidis strain, we generated donor and recipient strains. As shown in Figure 1A, to generate the donor strain, a kanamycin resistance gene (aph) was inserted between genes SEN1975 and SEN1976 in a strain of S. Enteritidis that has ROD21 inserted on the chromosome. (S. Enteritidis ROD21::aph). Similarly, to generate a recipient strain, a chloramphenicol resistance gene (cat) was inserted between genes SEN0986A (putA) and SEN0987 (putP) in a S. Enteritidis strain that had lost ROD21 due to excision type 2 (S. Enteritidis ∆ROD21 putAP::cat) (Figure 1B). To evaluate whether ROD21 was transferred from the donor to the recipient strain, we co-cultured both strains as described in Materials and Methods and selected transconjugants S. Enteritidis. In these assays, we found transconjugant bacteria, suggesting that ROD21 was effectively transferred from the donor to the recipient strain. To corroborate that these transconjugant strains of S. Enteritidis have incorporated ROD21 in their chromosomes, we detected the insertion of this pathogenicity island by PCR at attB site in the chromosome of the recipient strain. As shown in Figure 1C, we observed that the recipient strains had incorporated ROD21 at the attB site. Interestingly, all transconjugant bacteria evaluated in these assays acquired genes SEN1970 to SEN2011, suggesting that the episomal element was generated due to excision type 2, transferred from the donor strain and successfully integrated in the chromosome of recipient bacteria.

ROD21 is transferred from S. Enteritidis to S. Typhimurium

Next, we evaluated whether ROD21 can be transferred from S. Enteritidis to S. Typhimurium. Recipient strains of S. Typhimurium were generated by insertion of the cat gene between the genes STM14_1278 (putA) and STM14_1281 (putP), respectively (Figure 2A). Recipient and donor strains were mixed and incubated in LB medium, as described above. In these assays, we obtained recipient S. Typhimurium strains that acquired ROD21 from donor S. Enteritidis. To corroborate that these transconjugant S. Typhimurium strains effectively acquired ROD21, the borders of the island and several regions inside ROD21 were amplified by PCR (Figure 2A). To demonstrate that the double resistant S. Typhimurium strains had acquired ROD21 and to rule out the transfer of the chloramphenicol resistance gene from S. Typhimurium to S. Enteritidis, we successfully amplified segments of DNA by PCR that were present only in S. Typhimurium but absent in S. Enteritidis (genes in the prophage Gifsy-1). These PCR reactions were performed in all the transconjugant colonies obtained in each assay and in all of them the prophage Gifsy-1 was detected (data not shown). These data
support the notion that ROD21 can be transferred from S. Enteritidis to S. Typhimurium.

To determine which episomal ROD21 element was transferred and integrated in the chromosome of S. Typhimurium (episomal ROD21 produced either by excision type 1 or type 2), we amplified a portion of the DNA included between \textit{attR} and the \textit{asnT-2} of ROD21, which is present in the episomal element generated by excision type 2 (Figure 2B) and shared by S. Enteritidis and S. Typhimurium. In that region there is a fragment of 50 bp that is present only in S. Enteritidis but absent in S. Typhimurium and therefore allowed us to discriminate whether or not the DNA fragment between DRS and \textit{asnT-2} present in the genome of the transconjugant S. Typhimurium came from S. Enteritidis. Thus, we sequenced the PCR product generated to detect the 50 bp that are present in S. Enteritidis but absent in S. Typhimurium. All the S. Typhimurium transconjugants obtained in these assays were sequenced and none of them showed the sequence specific for S. Enteritidis (Figure 2B). These data suggest that the acquired episomal element from S. Enteritidis was the one generated due to excision type 1 and that the integration site of ROD21 in S. Typhimurium was \textit{asnT-2}.

To determine if the virulence of S. Typhimurium could be affected due to the acquisition of ROD21, we infected C57BL/6 mice intragastrically with both S. Typhimurium and S. Typhimurium-ROD21 and measured survival. As shown in Figure 2C, we observed that mice infected with S. Typhimurium-ROD21 showed reduced survival as compared to mice infected with WT S. Typhimurium. These results support the notion that acquisition of ROD21 by S. Typhimurium increase bacterial virulence, indicating that the genes contained in this pathogenicity island can work as virulence factors.

**Excision of ROD21 is required for the transfer to recipient strains**

To determine if the transfer of ROD21 requires the excision of this pathogenicity island and to rule out chromosomal transfer, we used a S. Enteritidis strain unable to excise ROD21 that we have described in a previous study [9]. This bacterial strain lacks the ORF SEN1970, which encodes for an integrase belonging to the family of P4 coliphages. Further, this strain lacks the DRS/\textit{attL} sequence required for excision. When transfer assays were performed using this mutant strain, no transconjugant bacteria were obtained (Figure 3A). In addition, we tested ROD21 transfer using a S. Enteritidis strain that excises ROD21 with higher frequency due to over expression of SEN1998, an ORF that encodes for a putative excisionase. We have previously shown that a strain of S. Enteritidis harboring the plasmid pBAD-SEN1998 showed enhanced ROD21 excision due to arabinose-induced.

**Figure 2. Transfer of ROD21 to S. Typhimurium.** (A) Scheme of the genome of the S. Typhimurium recipient strain with cat gene inserted in the putAP region (black arrow) and double-resistant strain with \textit{aph} gene inserted in ROD21 and \textit{cat} gene inserted in the putAP region (black arrows). Numbers above each scheme represent the coordinates in the chromosome of S. Typhimurium. Asparagine tRNA genes are designated as \textit{asnT-1}, \textit{asnT-2} and -3. DRS stands for Direct Repeated Sequence. Thin arrows represent primers used for PCR verification of the strains: (1) left-end of ROD21 (1,011 bp), (2) DRS (914 bp), (3) Tight-end of ROD21 (903 bp), (4) insertion of cat gene (1,838 bp), (5) insertion of \textit{aph} gene (2,469 bp), and (6) left-end of Gifsy-1 (1,110 bp). Below the scheme, agarose gels show PCR products obtained for the recipient and transconjugant S. Typhimurium strains. (B) Schematic representation showing the location of a 50 bp region present in S. Enteritidis, downstream DRS, but absent in S. Typhimurium. The figure below shows the alignments of the theoretical sequence of the above mentioned region for S. Enteritidis PT4 and S. Enteritidis 14028 and the experimental sequences obtained for S. Enteritidis PT1, S. Typhimurium 14028 and 7 transconjugants of S. Typhimurium-ROD21 obtained in a representative transfer assay. (C) Survival of C57BL/6 mice intragastrically infected with \(1 \times 10^6\) CFU of S. Typhimurium wild type and S. Typhimurium-ROD21 strain. Data shown are the averages of three independent experiments, which included 4 mice per group. ***, P<0.05 Kaplan-Meier and Mantel-Cox post test.

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**Figure 3. ROD21 transfer requires excision.** Transfer frequency of ROD21 from donor S. Enteritidis, from S. Enteritidis \textit{ΔDRSSEN1970}, from S. Enteritidis ASEN1980 and from S. Enteritidis/pBAD-SEN1998 supplemented with arabinose to (A) recipient S. Enteritidis and (B) recipient S. Typhimurium. Graph shows the frequency of ROD21 transfer, expressed as the ratio between transconjugant bacteria and the total amount of donors calculated in each assay. Results are the average of four independent experiments (Y axis are expressed as reverse of log values). ***, P<0.001, Student’s t test. ND = Non-detected.

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SEN1998 overexpression (Figure S1) [9]. As shown in Figure 3, the transfer rate of ROD21 from S. Enteritidis/pBAD-SEN1998 to recipient strains increased by almost 13 times (Figure 3A–B), suggesting that ROD21 excision is required for its transfer.

ROD21 is transferred from donor to recipient strains by conjugation

After showing that ROD21 could be transferred from S. Enteritidis to S. Enteritidis and S. Typhimurium, the HGT mechanism responsible for mobilization of this pathogenicity island was explored. Recipient S. Enteritidis was incubated with 10 ng, 100 ng or 1,000 ng of purified DNA from donor strain and with the filtered supernatant of the donor strain, as described in Materials and Methods. In both cases we failed to obtain bacteria resistant to both antibiotics (data not shown), ruling out that ROD21 could be transferred by natural transformation or by phage transduction. However, when we performed filter mating assays, as described in Materials and Methods, we found a higher frequency of transfer from S. Enteritidis to both S. Enteritidis and S. Typhimurium (Figure 3A–B). These data suggest that ROD21 transfer takes place by means of bacterial conjugation.

To evaluate if ROD21 encodes its own conjugation system, we generated a S. Enteritidis strain lacking ORF SEN1980, which encodes a protein similar to mokA from the E. coli plasmid RSF1010 and which has been involved in mobilization of genetic elements. We observed that deletion of SEN1980 did not alter the transfer frequency of ROD21 (Figure 5A). This result suggests that ROD21 transfer relies on a conjugation system encoded by genes outside this pathogenicity island. Supporting this hypothesis is the fact that S. Enteritidis harbors a virulence plasmid similar to the virulence plasmid pSLT of S. Typhimurium, which contains ORFs homologous to tra genes with an organization similar to the F plasmid [15]. Given that strains of S. Enteritidis lacking the virulence plasmid are not available, we tested whether absence of the virulence plasmid could affect the transfer of ROD21 in a S. Typhimurium strain lacking this plasmid [16,17]. First, we tested S. Typhimurium ROD21:aph trg::mudQ pSLT+ as a donor strain and S. Typhimurium phoN::dhfr pSLT+ as recipient strain, and we effectively observed transfer of ROD21 between these bacteria, at a frequency equal to 1.1 × 10^-6. Then, we used the strain S. Typhimurium ROD21::aph trg::mudQ pSLT+ as a donor strain and S. Typhimurium phoN::dhfr pSLT- as a recipient strain. When coculture assays in LB medium were performed, no transconjugant bacteria were obtained, suggesting that ROD21 transfer requires conjugative machinery, which is possibly encoded by the virulence plasmid.

Changes in temperature and pH affect the frequency of ROD21 transfer

Next, we evaluated what environmental factors could influence the transfer frequency of ROD21. Previously we had reported that ROD21 excision increased when bacteria were exposed to hydrogen peroxide and when bacteria resided inside phagocytic cells [6]. Therefore, we performed transfer assays in the presence of H2O2. As shown in Figure 4A, no significant changes in the transfer frequency of ROD21 were observed neither to S. Enteritidis nor to S. Typhimurium, suggesting that in the conditions used in this study oxidative stress failed to alter the transfer frequency of this pathogenicity island. Next, we evaluated whether changes in temperature or pH could affect the transfer of ROD21. Transfer assays were performed incubating the recipient and donor strains at either pH 7 or 5 and observed a significant increase of the transfer frequency from S. Enteritidis to both S. Enteritidis (Figure 4B, left panel) and S. Typhimurium (Figure 4B, right panel). Further, the frequency of ROD21 transfer was measured at 37°C, 23°C and 42°C, and observed that the transfer of this pathogenicity island from S. Enteritidis to S. Enteritidis (Figure 4C, left panel) or to S. Typhimurium (Figure 4C, right panel) increased when bacteria were incubated at 23°C and 43°C, as compared to 37°C.

To evaluate whether the increased transfer of ROD21 correlated with an enhanced ROD21 excision, we used qPCR to quantify the amount of attl sequence in bacteria grown in the presence of H2O2, as well as in bacteria grown at different temperatures and at different pH values. As shown in Figure 5, the presence of H2O2 (A) produced a non-significant increase of ROD21 excision. However, when bacteria were exposed to 23°C, 43°C (B) or pH 5 (C) the level of ROD21 excision showed a significant increase. As expected, higher excision was detected when we observed increased transfer. These results suggest that changes in environmental conditions could increase the excision and transfer frequency of ROD21.

ROD21 is not transferred to E. coli

Because the organization and sequence of the genes coding for asparagine tRNA in the genome of E. coli is similar to S. Enteritidis (Figure S2), we thought that ROD21 could be transferred to E. coli and integrated in its chromosome. To evaluate this possibility, we performed transfer assays using S. Enteritidis as a donor and E. coli as a recipient strain (Figure S2). However, no E. coli strains that were resistant to both antibiotics were obtained after co-cultures between donor S. Enteritidis and each of the recipient strains of E. coli. Moreover, the nalidixic acid resistant E. coli DH5α was used as an avirulent and permissive strain as a recipient, but failed to produce transconjugants. We also tested ROD21 transfer from S. Enteritidis to each of the donor E. coli strains at different pH and temperatures and did not detect double-resistant strains (data not shown). These data suggest that E. coli is not permissive for ROD21 transfer from S. Enteritidis.

Discussion

HGT has been highlighted as one of the main mechanisms responsible for bacterial microevolution defined as a rapid adaptation to environmental changes [18]. Along these lines, Salmonella enterica is an important example because recent genome sequencing studies have suggested that about 10% of their genetic material has been acquired by HGT [4,19]. These HGT elements have caused great impact on the variability of the Salmonella genus, in which more than 2,500 serovars have been identified.

The aim of this study was to evaluate the transfer of the pathogenicity island ROD21, present in the chromosome of S. Enteritidis, to other strains of Salmonella. ROD21 was described as a region present in Salmonella serovars Gallinarum, Dublin, Typhi and Enteritidis [5]. These data suggested that these serovars had acquired ROD21 by HGT. We have reported that ROD21 is an
unstable element, due to its ability to undergo spontaneous excision from the bacterial chromosome during growth in liquid culture medium and during infection of murine immune cells [6]. It has been reported that pathogenicity islands, that are capable of spontaneous excision from the genome of Gram positive and Gram negative bacteria, have the ability to be transferred to other bacteria [14,20,21].

By using transfer assays in liquid LB medium, we obtained recipient strains of *S.* Enteritidis that have acquired the pathogenicity island, which integrated to the chromosome of these bacteria. *S.* Typhimurium was also found to acquire and integrate the pathogenicity island into its genome. Furthermore, our results suggest that transfer of ROD21 requires excision of the pathogenicity island from the chromosome. Consistently, conditions that increase ROD21 excision also cause an increase of the transfer frequency of this genetic element. These results support the notion that ROD21 is an unstable and mobile genetic element that can be transferred between *Salmonella* strains.

According to our previous studies, it is likely that ROD21 excises from the *S.* Enteritidis chromosome by at least two different recombination events [6], leading to the generation of two episomal elements. Here, the strain of *S.* Enteritidis used as a recipient strain spontaneously lost the island due to excision type 2 (Figure 1). We found that transconjugant *S.* Enteritidis recovered the entire region (comprising genes from SEN1970 to SEN2011). Previously, we reported that ROD21 loss was detectable only in bacteria that underwent excision type 2 [6], suggesting that the episomal element produced by this event of recombination was transferred and integrated with higher frequency than episomal elements generated by recombination between the different *asnT* genes or between *asnT* genes and the DRS. While these different recombination events could generate at least 5 different episomal elements, we found that the episomal element generated by excision type 2 was able to be integrated in the chromosome of *S.* Enteritidis and detected using our experimental approach.

In the case of *S.* Typhimurium, this serovar could act as recipient for ROD21 because the organization of the *asnT* genes was similar to *S.* Enteritidis, but only lacks this genetic island (genes SEN1970 to SEN1999) [6]. Analysis of double resistant transconjugant bacteria showed that they had acquired the complete pathogenicity island. Moreover, sequence assays suggest that transconjugant *S.* Typhimurium incorporated ROD21 by site-specific recombination of the episomal element formed by excision type 1.

When experiments were performed using *E. coli* as recipient bacteria, no transconjugant bacteria were obtained. It was previously described that one of the greatest barriers to the transfer of genetic material is the presence of restriction systems, especially for the acquisition of DNA from a different species [22,23]. However, we did not obtain transconjugant bacteria when we used the *E. coli* strain DH5α, which has a mutation in the Endonuclease I gene (*endA*). It has also been proposed that the main obstacle to the acquisition of genetic material between *E. coli* and *S.* Enteritidis remains the restriction systems of the recipient strain, which present a major limitation for the transfer of genetic material.
and Salmonella is the level of sequence identity between the donor and recipient bacteria, where the SOS system would play an important role [24,25,26]. These explanations could account for the lack of ROD21 integration in the genome of recipient E. coli strains.

As mentioned above, transfer of genetic material may occur by transformation, transduction or conjugation. The presence in ROD21 of CDS with homology to proteins involved in bacterial conjugation, such as MobA, TraD, PisS and Piv, suggests conjugation as the mechanism responsible for the transfer. However, it is known that complete operons encoding the conjugation machinery are required to mediate the transfer of unstable episomal elements [27]. Consistent with this notion, deletion of the mobA homolog SEN1980 did not alter the frequency of ROD21 transfer in our assays. By analyzing the mechanism by which ROD21 is transferred, we ruled out natural transformation and transduction. Results of filter transfer assays, which increase the frequency of transfer to S. Enteritidis and S. Typhimurium, in two and one order of magnitude respectively, suggest that the event of mobilization occurs via conjugation. This increase is possibly due to a higher probability of intimate bacterial contact.

To date, many mobile genetic elements have been classified as Integrative and Conjugative Elements (ICE) because they possess the complete machinery to excise and transfer by themselves to other bacteria [28,29,30]. When we used a strain of S. Typhimurium lacking the virulence plasmid pSLT, which has been described as similar to F plasmid [31], transfer of ROD21 did not occur. This result supports the notion that ROD21 requires conjugative machinery, likely provided in trans, to transfer to other bacteria. Some genetic elements capable of being transferred have been classified as Integrative and Mobile Elements (IME), despite lacking the genes encoding for the transfer machinery in their sequence. Such is the case of Salmonella genomic island 1 [32], thus it would seem appropriate to classify ROD21 as an IME.

As we have previously shown, environmental conditions can affect ROD21 excision [6], suggesting that they might modulate the transfer of this pathogenicity island. When analyzing the effect of the temperature, we observed an increase in the efficiency of ROD21 transfer from S. Enteritidis to S. Typhimurium both at 23°C and 43°C. Consistent with our data, transfer of plasmid R27 from E. coli increases at low temperatures (25°C to 30°C) due to an H-NS-mediated overexpression of genes involved in mobility [33]. CDS SEN1993 encoded in ROD21 shows significant sequence homology with the H-NS protein, which could explain the increased transfer frequency in response to temperature changes. Although it has been described that high temperatures decrease the transfer of genetic elements, the natural reservoir of S. Enteritidis is poultry, whose body temperature is approximately 45°C [34]. Thus, it is possible that transfer of ROD21 in poultry is a constant and dynamic phenomenon, possibly to ensure the presence of the pathogenicity island in the chromosome of S. Enteritidis.

It is known that pH can affect the transfer of genetic elements from one bacterium to another [35,36,37]. Salmonella passes through the digestive tract encountering various pH values, which could be an important aspect for the infectious cycle. Furthermore, pH modulation of ROD21 transfer could play a role while located in phagosomes inside phagocytic cells, which usually show low pH. Consistent with this notion is the observation that the transfer of ROD21 to S. Enteritidis and S. Typhimurium peaked at pH 5. It is likely that this process is modulated by the differential expression of genes involved in genetic transfer at various pH values. In agreement with these observations, we detected higher levels of ROD21 excision in the conditions where we observed higher transfer.

In conclusion, our data suggest that ROD21 is an unstable pathogenicity island that can be transferred to strains of S. Enteritidis and S. Typhimurium by conjugation. Such a transfer is a species-specific event that is affected by environmental conditions influencing both the natural life cycle of bacteria, as well as their infective process, ensuring the maintenance of this pathogenicity island by the bacterial population.

Supporting Information

Figure S1 Excision of ROD21 in donor strain overexpressing ORF SEN1998. (A) Expression of excisionase (SEN1998) in wild type strain and donor strain transformed with the plasmids pBAD-empty or pBAD-SEN1998, when arabinose (1 mM) is added. The results are expressed as the ratio of SEN1998/podD expression (relative quantification or RQ). **, p<0.005 one-way ANOVA and Tukey post-test. (B) Quantification of excision of ROD21 in S. Enteritidis wild type, S. Enteritidis transformed with the plasmid pBAD-TOPO or pBAD-SEN1998, with or without arabinose (1 mM). The results are expressed as the ratio of attB1/podD (relative quantification or RQ). ***, p<0.001 one-way ANOVA and Tukey post-test.

(TIF)

Figure S2 Generation of recipient E. coli strains. (A) Scheme of the region of possible integration of ROD21 in E. coli strains H10407 and EDL933. Numbers on top of each scheme represent the coordinates in the chromosome of both strains. DRS stands for Direct Repeated Sequence (attR). Thin arrows represent primers used for PCR verification of the strains: (1) left-end of ROD21 (1,011 bp), (2) DRS (914 bp), (3) right-end of ROD21 (903 bp), (4) insertion of cat gene (1,958 bp) and (5) insertion of aph gene (2,469 bp). (B) Comparison of the sequences of asparagine tRNA genes of S. Enteritidis and E. coli strains used in this study.

(TIF)

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Author Contributions

Conceived and designed the experiments: SMB FJS. Performed the experiments: FJS HET PAN. Analyzed the data: SMB FJS. Contributed reagents/materials/analysis tools: CAR. Wrote the paper: SMB FJS CAR.

References

1. Juhas M, Van Der Meer JR, Gaillard M, Harding RM, Hood DW, et al. (2009) Genomic islands: tools of bacterial horizontal gene transfer and evolution. FEMS Microbiology Reviews 33: 376–393.

2. Broudy TB, Fischetti VA (2003) In vivo lysogenic conversion of Tox(−) Streptococcus pyogenes to Tox(+) with Lysozyme Streptococci or free phage. Infect Immun 71: 3782–3786.
3. Yasmin A, Keme JC, Shankar J, Darby AC, Hall N, et al. (2010) Comparative genomics and transcription potential of Enteroctococcus faecalis temperate bacterio- phages. J Bacteriol 192: 1122–1130.

4. Porvollik S, McClelland M (2003) Lateral gene transfer in Salmonella. Microbes Infect 5: 977–989.

5. Thomson NR, Clayton DJ, Windhorst D, Vernikos G, Davidson S, et al. (2008) Comparative genome analysis of Salmonella Enteritidis PF4 and Salmonella Gallinarum 287/91 provides insights into evolutionary and host adaptation pathways. Genome Res 18: 1624–1637.

6. Quirouz TS, Nieto PA, Tobar HE, Salazar-Echegaray IF, Lizana RJ, et al. (2011) Excision of an unstable pathogenicity island in Salmonella enterica serovar Enteritidis is induced during infection of phagoctytic cells. PLoS One 6: e25031.

7. Newmann RM, Salunkhe P, Godzik A, Reed JC (2006) Identification and characterization of a Novel Bacterial Virulence Factor That Shares Homology with Mammalian Toll/Interleukin-1 Receptor Family Proteins. Infect Immun 74: 594–601.

8. Silva CA, Blouedel CJ, Quezada CP, Porvollik S, Andrews-Polymenis HL, et al. (2012) Infection of mice by Salmonella enterica serovar Enteritidis involves additional genes that are absent in the genome of serovar Typhimurium. Infect Immun 80: 839–849.

9. Tobar HE, Salazar-Echegaray IF, Nieto PA, Palavecino CE, Sebastian VP, et al. (2013) Chromosomal excision of a new pathogenicity island modulates Salmonella virulence in vivo. Curr Gene Ther 13: 240–249.

10. Turner SA, Luck SN, Sakellariou H, Rajakumar K, Adler B (2004) Role of attP in integrase-mediated integration of the Shigella resistance locus pathogenicity island of Shigella flexneri. Antimicrob Agents Chemother 48: 1028–1031.

11. Betancor L, Yim L, Fookes M, Martinez A, Thomson NR, et al. (2009) Genomic and phenotypic variation in epidemic-spanning Salmonella enterica serovar Enteritidis isolates. BMC Microbiol 9: 237.

12. Datsonko KA, Wanner BL (1999) One-step inactivation of chromosomal genes. Curr. Gene Ther 9: 15–23.

13. Coburn PS, Baghdayan AS, Dolan GT, Shankar N (2007) Horizontal transfer of Enterococcus faecalis mobile genetic elements enabling dynamic lateral gene flow. Nat Rev Microbiol 5: 552–563.

14. Garcia-Quintanilla M, Casadesus J (2011) Virulence plasmid interchange in Enterococcus faecalis strains. Microbes Infect 13: 715–724.

15. rotate R, Casadesus J (1999) The virulence plasmid of Salmonella. Int Microbiol 2: 177–184.

16. Garcia-Quintanilla M, Casadesus J (2011) Virulence plasmid interchange among strains of Salmonella enterica serovar Typhimurium. Plasmid 65: 169–175.

17. Tinge SA, Curtis R 3rd (1990) Conservation of Salmonella typhimurium virulence plasmid maintenance regions among Salmonella serovars as a basis for plasmid curing. Infection and immunity 58: 3084–3092.

18. Medini D, Donati C, Tettelin H, Masignani V, Rappuoli R (2005) The microbial pan-genome. Curr Opin Genet Dev 15: 589–594.

19. Edwards RA, Olsen GJ, Maloy SR (2002) Comparative genomics of closely related salmonellae. Trends Microbiol 10: 94–99.

20. Ramay JP, Sullivan JT, Jamhari N, Ortori CA, Herb S, et al. (2009) A LuxR-family regulatory system controls excision and transfer of the M. chelonae lon strain. J Bacteriol 187: 3950–3959.

21. Schubert S, Darlu P, Clermont O, Wieser A, Magistret G, et al. (2009) Role of intraspecies recombination in the spread of pathogenicity islands within the Salmonella enterica species. PLoS Pathog 5: e1000257.

22. Corvaglia AR, Francois P, Hernandez D, Perron K, Linder P, et al. (2010) A type III-like restriction endonuclease function as a major barrier to horizontal gene transfer in clinical Staphylococcus aureus strains. Proc Natl Acad Sci U S A 107: 11954–11958.

23. Thomas CM, Nielsen KM (2005) Mechanisms of, and barriers to, horizontal gene transfer between bacteria. Nat Rev Microbiol 3: 711–721.

24. Matie I, Taddei F, Radman M (2008) No genetic barriers between Salmonella enterica serovar Typhimurium and Escherichia coli in SOS-induced mismatch repair-deficient cells. J Bacteriol 190: 5922–5924.

25. Rassier D, Delft C, Radman M (1991) Interspecific recombination between Escherichia coli and Salmonella typhimurium occurs by the RecABC pathway. Biochimie 73: 371–374.

26. Rassier D, Thaler DS, Radman M (1989) The barrier to recombination between Escherichia coli and Salmonella typhimurium is disrupted in mismatch-repair mutants. Nature 342: 596–601.

27. Wolfraテスト RA, Waldor MK (2010) Integrative and conjugative elements: mosaic mobile genetic elements enabling dynamic lateral gene flow. Nat Rev Microbiol 8: 552–563.

28. Butlermann P, Covarrubias PC, Lexican G, Katz A, Tapia P, et al. (2012) ICE Af 1, an actively excising genetic element from the bioninonizing bacterium Acidithiobacillus ferrooxidans. J Mol Microbiol Biotechnol 22: 399–407.

29. Moreno Switt AI, de Bakker HC, Cummings CA, Rodriguez-Rivera LD, Govoni G, et al. (2012) Identification and characterization of novel Salmonella mobile elements involved in the dissemination of genes linked to virulence and transmission. PLoS One 7: e41247.

30. Sencillo V, Czeczowski K, Pradovnd N, Minoia M, Miyazaki R, et al. (2009) Intracellular excision and reintegration dynamics of the ICEc1 genomic island of Pseudomonas knackmussi. Infect Immun 77: 1293–1306.

31. Garcia-Darriana MP, Francia MV, de la Cruz F (2009) The diversity of conjugative relaxases and its application in plasmid classification. FEMS Microbiol Rev 33: 657–687.

32. Doublier B, Boid D, Mulvey MR, Cloeckaert A (2005) The Salmonella genomic island 1 is an integrative mobilizable element. Mol Microbiol 55: 1911–1924.

33. Forns N, Banos RC, Balsalobre C, Juarez A, Madrid C (2005) Temperature-dependent conjugative transfer of R27: role of chromosome- and plasmid-encoded Hha and H-NS proteins. J Bacteriol 187: 3950–3959.

34. Giloh M, Shander D, Yahav S (2012) Skin surface temperature of broiler chickens is correlated to body core temperature and is indicative of their thermoregulatory status. Poult Sci 91: 173–188.

35. Davidson J (1999) Genetic exchange between bacteria in the environment. PLoS 42: 73–91.

36. Lorenz MG, Wackernagel W (1994) Bacterial gene transfer by natural genetic transformation in the environment. Microbew Rev 58: 563–602.

37. Williams HG, Day MJ, Fry JC, Stewart GJ (1996) Natural transformation in river epithile. Appl Environ Microbiol 62: 2994–2998.

38. Jarvik T, Smolle C, Grosman EA, Ochman H (2010) Short-term signatures of evolutionary change in the Salmonella enterica serovar Typhimurium 14028 genome. J Bacteriol 192: 560–567.

39. Crossman LC, Chaudhuri RR, Beatson SA, Wells TJ, Desvaux M, et al. (2010) A commensal gone bad: complete genome sequence of the prototypical enterotoxigenic Escherichia coli strain H10407. J Bacteriol 192: 3822–3831.

40. Perna NT, Plunkett G 3rd, Burland V, Ma B, Glaser JD, et al. (2001) Genome sequence of enterohaemorrhagic Escherichia coli O157:H7. Nature 409: 529–533.