Excision of an Unstable Pathogenicity Island in
Salmonella enterica Serovar Enteritidis Is Induced during Infection of Phagocytic Cells

Tania S. Quiroz1, Pamela A. Nieto1, Hugo E. Tobar1, Francisco J. Salazar-Echegarai1, Rodrigo J. Lizana1, Carolina P. Quezada2, Carlos A. Santiviago2, Daniela V. Araya1, Claudia A. Riedel2, Alexis M. Kalergis1,4, Susan M. Bueno1*

1 Departamento de Genética Molecular y Microbiología, Facultad de Ciencias Biológicas, Millennium Institute on Immunology and Immunotherapy, Pontificia Universidad Católica de Chile, Santiago, Chile, 2 Departamento de Bioquímica y Biología Molecular, Facultad de Ciencias Químicas y Farmacéuticas, Universidad de Chile, Santiago, Chile, 3 Facultad de Ciencias Biológicas y Facultad de Medicina, Millennium Institute on Immunology and Immunotherapy, Universidad Andrés Bello, Santiago, Chile, 4 Departamento de Reumatología, Facultad de Medicina, Pontificia Universidad Católica de Chile, Santiago, Chile

Abstract
The availability of the complete genome sequence of several Salmonella enterica serovars has revealed the presence of unstable genetic elements in these bacteria, such as pathogenicity islands and prophages. This is the case of Salmonella enterica serovar Enteritidis (S. Enteritidis), a bacterium that causes gastroenteritis in humans and systemic infection in mice. The whole genome sequence analysis for S. Enteritidis unveiled the presence of several genetic regions that are absent in other Salmonella serovars. These regions have been denominated “regions of difference” (ROD). In this study we show that ROD21, one of such regions, behaves as an unstable pathogenicity island. We observed that ROD21 undergoes spontaneous excision by two independent recombination events, either under laboratory growth conditions or during infection of murine cells. Importantly, we also found that one type of excision occurred at higher rates when S. Enteritidis was residing inside murine phagocytic cells. These data suggest that ROD21 is an unstable pathogenicity island, whose frequency of excision depends on the environmental conditions found inside phagocytic cells.

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* E-mail: sbueno@bio.puc.cl

Introduction
S. Enteritidis is a Gram negative bacterium and the etiological agent of several foodborne diseases in humans [1]. This bacterium belongs to the subspecies I of the species Salmonella enterica, whose members cause systemic diseases in warm-blooded animals [2,3]. The ability of S. Enteritidis to cause a systemic disease in the host is due to its capacity to survive and replicate inside eukaryotic cells, especially within epithelial and phagocytic cells [4]. This feature of S. Enteritidis promotes the establishment of systemic disease in mammals and birds after ingestion of contaminated food or water [5].

As for many other Enterobacteria, the complete genome of S. Enteritidis has been sequenced and analyzed [2]. That information has allowed the identification of several genetic regions absent in the genome of other Salmonella serovars, such as Typhimurium. These distinctive gene clusters, denominated “Regions of difference” (ROD), could have been acquired by means of lateral gene transfer [2]. One of such regions is ROD21, a pathogenicity island found only in the chromosome of S. Enteritidis, S. Gallinarum and S. Dublin, but absent in other Salmonella serovars whose whole genome has been sequenced [2,6]. Similar to other pathogenicity islands described in Enterobacteria, ROD21 is located next to a gene coding for a tRNA. Previous reports have shown that genomic islands of Enterobacteria located near tRNA genes are unstable, because they excise from the bacterial chromosome [7–9]. For instance, it has been described that pathogenicity islands SHI-1 and SRL of Shigella flexneri excise from the bacterial chromosome in laboratory growth conditions [10], as well as the high pathogenicity island of Yersinia pseudotuberculosis [11,12]. In Salmonella, previous reports have shown that SPI-7 of serovar Typhi, a 133 kb genomic island adjacent to a pheU tRNA gene, excises from the chromosome and gets lost at low rate in laboratory growth conditions [13]. In addition, it has been recently described that the prophage-like element pSE14 of S. Enteritidis (another S. Enteritidis ROD) also excises spontaneously from the chromosome under standard culture conditions [14].

The few reports that have identified specific conditions that promote the excision and transfer of genomic islands have focused on bacteria infecting plants. For instance, the Pseudomonas syringae pv. Phaseolicola genomic island 1 (PPHGI-1) excises from the chromosome and is transferred to recipient strains at high rates...
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During the infection of host plants [15]. In addition, the frequency of transfer to different strains of the same species is enhanced in *vitro* when plant apoplastic fluids are added to bacteria undergoing transformation, suggesting that excision and transfer of this genomic island can be influenced by components derived from the host [15]. Also, it has been described that the density of the bacterial population is another factor that influences the excision rate of genomic islands, as it was observed for the symbiotic bacterium *Mesorhizobium loti* [16]. Furthermore, there is one single study in bacteria causing disease in animals describing that the excision of the *Vibrio Pathogenicity Island 2* from *Vibrio cholerae* can be induced at low temperatures and after UV irradiation [17]. However, the conditions or signals responsible of promoting or preventing the excision of unstable genomic islands in *Salmonella* remain largely unknown. Furthermore, whether the excision of these genetic elements contributes to pathogenicity is an important question that requires to be addressed.

Here we show that ROD21 is an unstable *Salmonella* pathogenicity island that can excise from the bacterial chromosome due to at least two different and independent recombination events. However, only site-specific recombination could lead to ROD21 loss, suggesting that this pathogenicity island may be kept as an episomal element inside the bacterium. Of major importance was the observation that the excision rate of ROD21 increases when *S. Enteritidis* resides inside phagocytic cells, such as dendritic cells and macrophages. These results suggest that the excision frequency of ROD21 can be enhanced by specific environmental conditions taking place inside phagocytic cells during the oxidative stress response against intracellular bacteria.

**Results**

Characterization of ROD21 in the genome of *S. Enteritidis*

ROD21 is a 26,687 bp DNA fragment located between coordinates 2,061,170 and 2,087,657 in the *S. Enteritidis* PT4 NCTC13349 genome. ROD21 is found in a region of the *S. Enteritidis* chromosome that is common to the genome of *S. Typhimurium* strains LT2 and 14028 (Fig. 1A). ROD21 is found next to an asparagine tRNA gene (*asnT-2* or *attL*) and at its right end is delimited by 24 base pairs (bp), 22 of which are identical to the 3′ end of *asnT* (Fig. 1B). This element was denominated direct repeated sequence (DRS or *attP*). Furthermore, near ROD21 there are two asparagine tRNA genes located in the same orientation as *asnT-2* (*asnT-1* and *asnT-3* in this study). *asnT-1* is located 971 bp upstream of *asnT-2*, while *asnT-3* is located 11,487 bp downstream of the DRS. A fourth asparagine tRNA gene (*asnW*) is located 9,943 pb downstream of the DRS, but in the opposite direction when compared to the other *asnT* genes.

ROD21 harbors 29 coding sequences that are similar to genes previously described in other bacteria, or that have functions already assigned in *Salmonella* (Table 1). Interestingly, some regions of this pathogenicity island have a genetic organization similar to a genomic island found in Uropathogenic *Escherichia coli* strain CFT073 [18] (Table 1). One of the genes in ROD21 that has an assigned function is *SEN1973* (or *tlpA*), which encodes a virulence protein that has a Toll/Interleukin-1 receptor (TIR) domain [19]. A previous report has described that this protein interferes with the signaling derived from Toll-like receptor engagement and NF-κB activity in epithelial cells. Furthermore, mutant strains of *S. Enteritidis* lacking this gene cause a less severe disease in mice [19]. ROD21 also contains the gene *SEN1993*, encoding a protein homologous to HnsT from uropathogenic *E. coli*, which releases the suppression of virulence genes in these pathogenic bacteria [20]. Another gene found in this pathogenicity island is *SEN1978* that encodes a putative type IV pilin protein. In addition, ROD21 harbors genes coding for proteins belonging to conjugation systems: TraD (*SEN1979*) and MobA/MobL (*SEN1980*) (Fig. 1 and Table 1). Another CDS with assigned function is *SEN1970*, which encodes a putative integrase sharing 74% similarity to the integrase in prophage P4. Based on the presence of *att* sites and the integrase gene found in ROD21, we hypothesized that this pathogenicity island would excise from the *S. Enteritidis* chromosome.

At least two different recombination events promote ROD21 excision from the bacterial chromosome

Next, we tested whether ROD21 is able to undergo spontaneous excision from the bacterial chromosome. Because ROD21 is flanked by three copies of *asnT* and delimited by the DRS, at least two types of recombination events may take place: recombination between *asnT-2* and the DRS (excision type 1) and recombination between *asnT-2* and *asnT-3* (excision type 2). Either one of these recombination events might result in the complete excision of this new pathogenicity island (Fig. 2).

To evaluate whether these potential recombinations can occur, we performed a PCR reaction using primers that hybridize upstream and downstream of *asnT* genes and the DRS. As shown in Fig. 2B and 2C, these hypothetical excision events yield two different *attB* and *attP* sequences, which could be detected by PCR using several different primer combinations. To detect these excisions, the genomic DNA of four *S. Enteritidis* strains was obtained as described in materials and methods and tested by PCR. Conventional PCR amplifications failed to produce measurable amounts of the expected PCR products for each of the *attB* and *attP* sequences in each of the *S. Enteritidis* strains evaluated (data not shown). To increase the sensitivity of detection, nested PCRs were performed as described in material and methods and the expected sized amplicons were obtained: 591 and 657 bp for the chromosomal *attB-1* and *attB-2* respectively (Fig. 3A) and 958 and 1058 bp for the episomal *attP-1* and *attP-2* respectively (Fig. 3B). To corroborate the specificity of these PCR products, the DNA fragments obtained from *S. Enteritidis* PT1 were sequenced. As shown in Fig. 3, each of the obtained PCR products matched the expected *attB* (Fig. 3A) and *attP* (Fig. 3B) sequences. These data suggest that both recombination events occurred at low frequency when bacteria grew to stationary phase in LB medium.

Excision type 2 results in the loss of ROD21 from the chromosome of *S. Enteritidis*

Then, we evaluated whether the excision of ROD21 results in the loss of this pathogenicity island from the bacterial genome. To evaluate this possibility, we inserted the genes *tetA* and *tetR* downstream the gene *SEN1973* in the strain of *S. Enteritidis* Phagetotype 1 (PT1), to generate the ROD21::*tetRA* strain (Fig. 4A). The genes *tetA* and *tetR* confer resistance to tetracycline, but also prevent the growth of tetracycline-resistant bacteria in a medium containing zinc chloride and fusaric acid (BM medium), as described previously [21]. Therefore, only those *Salmonella* strains sensitive to tetracycline will grow in BM medium [21]. Nested PCR reactions showed that excision type 1 and type 2 (Fig. 2A) occurred in the modified ROD21::*tetRA* strain, as efficiently as observed for the wild type (WT) strain (data not shown). To evaluate if these excisions caused ROD21 loss, the ROD21::*tetRA* strain was grown in LB medium and then seeded on solid BM medium, as described in materials and methods. Bacteria were incubated 36 h at 37°C to select for tetracycline sensitive bacteria.
A total of 35 tetracycline sensitive colonies out of 137 seeded plates were tested by PCR to evaluate ROD21 loss. We observed that only 6 out of the 35 colonies isolated had lost ROD21 from the chromosome, indicating that the frequency of ROD21 loss is \(4.38 \times 10^{-2}\). Further, PCR reactions showed that only the attB sequence generated by excision type 2 could be detected in the genome of all isolated S. Enteritidis strains (Fig. 4B), suggesting that excision type 2 was responsible of ROD21 loss in all isolated CFUs. With regard to the other 29 colonies that lost tetracycline resistance without excising ROD21, it is possible that they might have undergone other type of mutations resulting in tetracycline sensitivity, such as mutations in tetA or tetR genes [22,23] or changes in cell membrane permeability. However, further research would be required to clarify this issue.

S. Enteritidis strains that have lost ROD21 show reduced virulence

To evaluate the impact of ROD21 loss in the virulence of S. Enteritidis, groups of C57BL/6 mice were orally infected with either WT S. Enteritidis or one of the isolated strains of S. Enteritidis lacking ROD21 (ΔROD21). After infection, the survival rate of mice was evaluated on a daily basis. As shown in Fig. 4C, mice infected with the ΔROD21 strain survived longer than did mice infected with the WT S. Enteritidis strain. However, 15 days after the infection, over 80% of the mice in both groups had died due to Salmonella infection. These data suggest that the loss of ROD21 causes an apparent mild reduction of the S. Enteritidis virulence in mice. To further study the attenuation of the ΔROD21 strains, we performed in vivo competition assays consisting of either oral or intravenous infection of mice with a mixture of WT and ΔROD21 strains (at a ratio equal to 1). After 72 h of infection, colonizing bacteria were recovered from spleens and livers of infected mice to evaluate the ratio of WT/ΔROD21 in these organs. As shown in Fig. 4D and 4E, the WT strain was recovered in larger proportions in spleens and livers of infected mice 72 h after infection, as compared to the ΔROD21 strain. Because similar data were obtained independently of the route of infection (orally or intravenously), it is likely that the genes encoded by ROD21 might be required for the systemic phase of Salmonella infection.

ROD21 excision is induced during infection of phagocytic cell

To determine whether the rate of ROD21 excision changes during infection, quantitative real time PCR assays were performed to measure the number of bacteria that underwent excision type 1 under different growth conditions. Because an important step in S. Enteritidis infective cycle is the invasion and survival in phagocytic cells, ROD21 excision frequency was determined for S. Enteritidis while infecting phagocytic cells, such as macrophages and dendritic cells (DCs). Genomic DNA was obtained from S. Enteritidis PT1 strain recovered at different steps during a gentamicin protection assay, as described in materials and methods. Then, the copy number of the attB sequence for each sample was quantified by using quantitative real time PCR (qPCR). In these assays, the copy number of attB sequences represents the amount of bacterial chromosomes that underwent ROD21 excision. qPCR data were normalized based on the total amount of DNA for each sample. Further, the copy number of the

Figure 1. Schematic representation of ROD21 in the chromosome of S. Enteritidis. (A) Representation of the genetic location of the genes coding for the asparagine tRNA (asnT-1, -2 and -3) in the chromosome of S. Typhimurium and S. Enteritidis and the exact location of ROD21 in the chromosome of S. Enteritidis. Black and dark gray arrows represent those genes shared between both serovars and light gray arrows represent genes found only in ROD21 of S. Enteritidis. Numbers next to each scheme are coordinates in the chromosome of S. Typhimurium and S. Enteritidis. DRS stand for Direct Repeated Sequence (attR). (B) The alignments of DRS and attB-1, attB-2 (attI) and attB-3 show that the DRS is identical to the last 22 bp of the attI gene.

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| ORF   | Size | Location       | Hypothetical role                                                                 | serU island of the UPEC CFT073 | % identity |
|-------|------|----------------|-----------------------------------------------------------------------------------|---------------------------------|------------|
| SEN1970 | 1275 | 2061363–2062637 | Integrase. P4-like integrase.                                                      | c2392                           | 25         |
| SEN1971 | 252  | 2062709–2062960 | Exported protein (S. Dublin) Identities: 100%                                    | -                               | -          |
| SEN1972 | 838  | 2063439–2064276 | Pseudogen (putative membrane protein, S. Gallinarum. Identities: 99%)            | -                               | -          |
| SEN1974 | 609  | 2064296–2064906 | Hypothetical protein (SeD_A2308. S Dublin) Identities:99%                        | -                               | -          |
| SEN1975 | 882  | 2065272–2066153 | TipA. Cytoplasmic protein with TIR domain (Salmonella sp.) Identities: 100%       | c2398                           | 45         |
| SEN1976 | 2143 | 2066381–2068523 | Pseudogen (putative type IV prepilin protein, S. Gallinarum. Identities: 99%)   | c2394                           | 76         |
| SEN1978 | 558  | 2068581–2069138 | Type IV Pilin (S. Enteritidis). Identities: 100%. (N-terminal PilS domain).      | c2395                           | 93         |
| SEN1979 | 306  | 2069546–2069851 | Conserved hypothetical protein (S. Enteritidis and S. Dublin). Identities: 100%. (Conjugal transfer protein TraD domain) | c2396                           | 87         |
| SEN1980 | 1521 | 2070488–2072008 | MobA/MobL family protein (S. Dublin) and possible Conjugal transfer protein (S. Enteritidis). Identities: 100%. (MobA/MobL family). | c2397                           | 84         |
| SEN1981 | 915  | 2072032–2072946 | Conserved hypothetical protein (S. Enteritidis). Identities: 99%                  | -                               | -          |
| SEN1981A | 264  | 2073108–2073371 | Membrane protein                                                                  |                                 |            |
| SEN1982 | 543  | 2073382–2073924 | Lipoprotein (S. Enteritidis). Identities: 100%.                                   | c2401                           | 96         |
| SEN1983 | 488  | 2074332–2074819 | Pseudogen (exported protein. S. Gallinarum. Identities: 99%)                     | -                               | -          |
| SEN1984 | 378  | 2074852–2075229 | Exported protein (S. Enteritidis). Identities: 100%                               | -                               | -          |
| SEN1985 | 1611 | 2075307–2076917 | Hypothetical protein (S. Gallinarum and S. Enteritidis). Identities: 100% (S-adenosylmethionine-dependent methyltransferases (SAM or AdoMet-MTase) domain) | -                               | -          |
| SEN1986 | 963  | 2076969–2077931 | Hypothetical protein (S. Dublin and S. Enteritidis). Identities: 100%             | c2406                           | 86         |
| SEN1987 | 423  | 2077987–2078409 | Hypothetical protein (S. Enteritidis). Identities: 100%                          | -                               | -          |
| SEN1988 | 270  | 2078458–2078727 | Hypothetical protein (S. Gallinarum and S. Enteritidis). Identities: 100%        | -                               | -          |
| SEN1989 | 300  | 2079026–2079325 | Hypothetical protein (S. Enteritidis). Identities: 100%                          | -                               | -          |
| SEN1990 | 735  | 2080087–2080821 | DNA-binding protein (S.Dublin and S. Enteritidis). Identities: 100% (Domain: helix_turn_helix multiple antibiotic resistance protein) | -                               | -          |
| SEN1991 | 792  | 2080852–2081643 | Hypothetical protein (S.Gallinarum and S. Enteritidis). Identities: 100%         | -                               | -          |
| SEN1992 | 480  | 2081729–2082208 | Hypothetical protein (S. Gallinarum). Identities: 98%.                          | c2410                           | 92         |
| SEN1993 | 405  | 2082369–2082773 | DNA-binding protein (histone-like protein hlp-II) (S. Gallinarum and S. Enteritidis). Identities: 100% (Domain: global DNA-binding transcriptional dual regulator H-N5; Provisional) | c2411                           | 91         |
| SEN1994 | 567  | 2083189–2083755 | Membrane protein (S. Gallinarum and S. Enteritidis). Identities: 100%             | -                               | -          |
| SEN1995 | 1272 | 2083802–2085073 | Conserved Hypothetical protein (S. Dublin and S. Enteritidis). Identities: 100% | -                               | -          |
| SEN1996 | 297  | 2085319–2085615 | Hypothetical protein (S. Gallinarum and S. Enteritidis). Identities: 100%        | c2414                           | 95         |
| SEN1997 | 303  | 2085660–2085962 | Hypothetical protein (S. Gallinarum and S. Enteritidis). Identities: 100%        | c2415                           | 95         |
| SEN1998 | 219  | 2086032–2086250 | Phage regulatory protein (Salmonella sp.). Identities: 100%. (Domain: Prophage CP4-5 regulatory protein (AlpA)/Predicted transcriptional regulator (Transcription)) | -                               | -          |
Excision type 1 of ROD21 was determined for extracellular bacteria, which were recovered from the supernatants of phagocytic cells 2 h after infection. Further, ROD21 excision was also determined for intracellular bacteria recovered from phagocytic cells at different times post-infection (2, 18 and 24 h). Then, the ratios of the ROD21 excision between intracellular bacteria and extracellular bacteria were determined (relative value in figure 5). As shown in Fig. 5A, excision rates in intracellular bacteria were increased at all time points, especially after 18 h of DCs infection (Fig. 5A) and after 2 h of macrophages infection (Fig. 5B). Similar results were obtained when the ratio of ROD21 excision between intracellular bacteria and bacteria grown in either LB or cell culture media was determined (data not shown). Finally, because similar intracellular bacterial loads were observed at all time points during infection of phagocytic cells (Fig. 5C and 5D), it is unlikely that our results could be due to variability in the amount of bacteria recovered after infection. These findings suggest that the excision of ROD21 might be induced by the environmental conditions found by S. Enteritidis inside phagocytic cells during infection.

Peroxide treatment increases ROD21 excision in vitro

Given that phagocytic cells produce reactive oxygen species upon phagocytosis of bacteria [24], whether oxidative stress could increase the frequency of ROD21 excision was evaluated. S. Enteritidis PT1 was grown in LB medium until OD600 equal to 0.6 and then 3.6 x 10⁹ bacteria were incubated in N medium, which mimics the intracellular conditions found inside eukaryotic cells (i.e. reduced magnesium concentration [25]). At 30 min before the end of the incubation in N medium, bacteria were treated with 0.25 mM hydrogen peroxide and the frequency of ROD21 excision was determined by qPCR. As shown in figure 6A, ROD21 excision was significantly higher in bacteria grown for 18 h in N medium and treated with peroxide, as compared to the same strain grown in N medium alone.

To explore a possible mechanism for the enhancement of excision induced by oxidative stress, the relative expression of the gene coding for the integrase (SEN1970) required for the integration and excision process was measured. We obtained mRNA from bacteria grown either on LB or N medium that were treated or not with hydrogen peroxide and evaluated the relative amount of the SEN1970 mRNA, using qPCR. As shown in Fig. 6B, the amount of SEN1970 mRNA was reduced when bacteria were exposed to N medium and hydrogen peroxide. These results suggest that intracellular conditions, such as reduced nutrients and oxidative stress may increase the excision rate of ROD21 due to a decreased expression of integrase, as it has been described previously for other bacteria [26].

In summary, our data suggest that the intracellular environment of dendritic cells and macrophages promotes excision of ROD21. It is likely that the survival of Salmonella inside these phagocytic cells might be increased by the excision of this pathogenicity island.

Discussion

The acquisition of genes through lateral transfer is a major source of variation and evolution for pathogenic bacteria. An example of mobile elements that can be transferred from one bacterium to another are pathogenicity islands, which can harbor several genes required by bacterial pathogenesis during infection [27]. Previous studies have described that some pathogenicity islands can excise from the bacterial chromosome, either spontaneously or in response to certain stimuli [28]. In this study, we have provided evidence suggesting that ROD21, a pathogenicity island of S. Enteritidis, is an unstable genetic element that can undergo spontaneous excision by two different recombination events. Our data show that recombination of attT-2 (attL) with the DRS (attR) results in the excision of ROD21 (excision type 1), which forms a circular episomal genetic element. In addition, recombination of attT-2 with attT-3 (excision type 2) also leads to the excision of ROD21 from the chromosome, but in this case the excision leads to a circular DNA fragment that includes ROD21 and the fragment located between DRS and attT-3. Both circular elements generated by excisions type 1 and 2 were detected in this study for S. Enteritidis. The presence of a gene coding for an integrase (SEN1970) in ROD21 suggests that this protein may catalyze excisions type 1 and 2. However, it remains to be determined whether this gene encode a functional protein. Further, it is also possible that other integrases encoded by genes located in different sites in the chromosome could catalyze the excision of this region, as it has been described for pathogenicity islands of E. coli [8]. Mechanisms of homologous recombination, known to participate in the excision of unstable genetic elements [29], could also contribute to the excision of ROD21. Given that a gene encoding the RecA protein is present in the S. Enteritidis genome [30] and that the sequences involved in the recombination events are 100% identical, it is likely that this protein may also contribute to catalyzing ROD21 excision. Moreover, the presence of three tRNA genes near ROD21, all of them identical in sequence, raises the possibility that 3 additional recombination events could take place: (1) recombination between attT-1 and the DRS, (2) recombination between attT-1 and attT-3, and (3) recombination between attT-3 and the DRS. To define whether these additional excisions occur further research would be required.

Here we have determined that ROD21 excision may result in the loss of this genomic region in a small percentage of the bacterial population, consistently to what has been observed for other bacteria possessing unstable pathogenicity islands [8]. By using a selection assay based on tetracycline sensitivity, bacteria that had lost ROD21 due to type 2 excision (recombination between attT-2 and attT-3 genes) were isolated. However, bacteria that lost ROD21 due to recombination between attT-2 and the DRS could not be isolated, even if both, attB and attP sequences generated by type 1 excision could be detected by PCR.

| ORF     | Size  | Location       | Hypothetical role                    | serU island of the UPEC | % identity |
|---------|-------|----------------|--------------------------------------|-------------------------|------------|
| SEN1999 | 876   | 2086401–2087276 | Hypothetical protein (S. Gallinarum and S. Enteritidis). Identities: 100%. | CFT073                  | 1999       |
|         |       |                |                                      |                         | 40         |

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rsD gene was quantified as an indication of the total amount of bacterial chromosomes per sample. Data were expressed as the number of attB copies/number of rpsD copies.
It is likely that type 1 excision leads to the formation of a stable episomal element, which is kept inside bacteria and probably re-integrated into the chromosome. Although the episomal element formed after an excision by recombination between \( \text{asnT-2} \) and \( \text{asnT-3} \) can be detected by PCR, this element might be unstable and eventually degraded. This possibility could explain why only bacteria that lost this genomic region only due to type 2 excision could be isolated.

We also observed that type 2 excision-mediated loss of ROD21 extended the time required by \( S. \) Enteritidis to cause a lethal disease in mice. Further, in vivo competitive assays at equivalent infection times showed that the \( \Delta \text{ROD21} \) strain was less capable of colonizing spleens and livers as compared to WT \( S. \) Enteritidis. These data are in agreement with a potential role for ROD21 as a cluster for virulence genes. This notion finds additional support on a previous report showing that lack of the ROD21-encoded \( \text{SEN1975} \) gene reduces the capacity of \( S. \) Enteritidis to cause lethal disease in mice [19]. Moreover, ROD21 also codes for \( \text{SEN1993} \), which is homologous to HnsT, a protein that promotes the expression of virulence genes in uropathogenic \( E. \) coli. It is likely then that \( \text{SEN1993} \) could work in \( S. \) Enteritidis as an inducer of the expression of virulence genes and that the lack of this protein would reduce pathogenicity in mutant strains. However, additional research is required to evaluate whether other ROD21 genes contribute to the virulence of \( S. \) Enteritidis in mice and other hosts.

We observed that the excision frequency of ROD21 increases when bacteria infect phagocytic cells, such as DCs and macrophages. The highest excision frequencies were obtained...
for bacteria recovered from the intracellular space of infected cells. It is possible that recognition of signals present inside host cells would lead to the activation of proteins involved in ROD21 excision. As a consequence of excision, transcription of genes contained in these unstable genetic elements could be enhanced, as described for unstable elements found in *Corynebacterium glutamicum* [22]. Increased expression of potential virulence genes contained in ROD21 when *Salmonella* locates inside host cells could be relevant for its ability to survive intracellularly.

The conditions present in the *Salmonella*-containing vacuole, such as low pH, high amounts of reactive oxygen species, nitric oxide, reduced Mg\(^{2+}\) concentration, among others, could act as signals that promote production or activation of proteins involved in the excision process. In agreement with this notion, our results show that the presence of hydrogen peroxide increases the frequency of ROD21 excision. Furthermore, the transcription of the integrase gene (*SEN1970*) is reduced when bacteria are grown in N medium and exposed to hydrogen peroxide. These data are consistent with the idea that the presence of hydrogen peroxide promotes the excision of ROD21.

**Figure 3. ROD21 excision can be generated by means of two different recombination events.** Amplification of *attB* and *attP* sequences generated after type 1 and type 2 excisions were detected by nested PCR in LK5, PT4, PT1 and PT21 strains of *S. Enteritidis*, using primer pairs described in Table 2 and in Figure 2. PCR products for *attB* (A) and *attP* (B) sequences for each type of excision were resolved in 1% agarose gels. The sequence of each PCR product was obtained (chromatograms in each Figure) and compared with the *attB* and *attP* sequences deduced for type 1 and type 2 excisions (labeled as theoretical), *attB* and *attP* sequences are highlighted in red in both alignments and chromatograms. Expected size for each PCR product: 591 bp for type 1 excision *attB*, 657 bp for type 2 excision *attB*, 995 bp for type 1 excision *attP* and 1050 bp for type 2 excision *attP*.

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in agreement with a recent report showing that Phage 4 integrase expression is downregulated to promote the excision of bacteria unstable genetic elements [26]. Then, it is possible that a reduction of integrase expression could promote ROD21 excision in response to phagocytic cell oxidative burst. It has been shown that oxidative burst starts early in macrophages [23], which is consistent with the early increase of ROD21 excision observed when S. Enteritidis infects J774.3 cells.

The excision of ROD21 in S. Enteritidis during infection of phagocytic cells is analogous to the observation that prophage excision is induced soon after infection. For instance, contact with pharyngeal epithelial cells promotes the excision of 5 prophages from the genome of group A Streptococcus [31]. Such a process induces the production of virulence factors encoded by these prophages. To the best of our knowledge, this is the first report describing the occurrence of the excision phenomenon when bacteria locate inside phagocytic cells, one of the most important stages of the infectious cycle of Salmonella. Additional studies are required to further evaluate if the excision of the ROD21 pathogenicity island contributes to the virulence of S. Enteritidis.
Materials and Methods

Ethics statement

All the 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 (07/06/2007). All animal work was performed according to the Guide for Care and Use of Laboratory Animals (National Institute of Health, USA) and Institutional guidelines were overseen by a Veterinarian.

Bacterial strains and growth conditions

*S. Enteritidis* strains used in this study are: *S. Enteritidis* LK5 (provided by Dr. Guido C. Mora, Universidad Andrés Bello, Chile), *S. Enteritidis* phagotype 1 (PT1), phagotype 4 (PT4) and phagotype 21 (PT21) (provided by Mrs. Alda Fernandez from the Public Health Institute of Chile, ISP). 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 and agitation. N-minimal medium was composed of 5 mM KCl, 7.5 mM (NH₄)₂SO₄, 0.5 mM K₂SO₄, 1 mM KH₂PO₄, 0.1 mM Tris-HCl pH 7.4, 30 μM MgCl₂, 0.2% glucose, 38 mM glycerol, and 0.1% casaminoacids; pH was adjusted to 5.0. When needed, hydrogen peroxide was added to the N-minimal medium, at a concentration of 0.25 mM. The strain *S. Enteritidis* ROD21::tetRA was generated by Lambda Red-mediated recombination, as described by Datsenko and Wanner [32]. Briefly, a PCR product encoding tetA and tetR genes was generated by PCR amplification from the mini Tn10 transposon T-POP [33]. The primers used were as follows:

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5' GGA ACT CTC CAT TGG AGA GAA TAC ATA TCA CTT GGG AAA Aaa tca tta agt taa ggt gga 3' 
5' ATG TTT GTG TTT AAA CAT TAT AAT AAA ATT TAA CTT TTA Ata tca taa tga tga tga tga 3'
```

The first 40 bp of these primers (capital) align with bp 2,066,174–2,066,213 and 2,066,321–2,066,360 of the *S. Enteritidis* chromosome, respectively. The last 20 bp of these primers (lowercase) align with the T-POP transposon. Competent *S. Enteritidis* PT1 harboring the thermosensitive plasmid pKD16 were prepared as described [32] and PCR products containing tetA and tetR genes were electrotransformed to these competent cells. After electrotransformation, bacteria were incubated for 1 h at 37°C with aeration in 1 ml of LB medium, and then seeded on solid LB medium supplemented with 25 μg/ml tetracycline. In order to verify the correct insertion of tetA and tetR genes, a PCR amplification was performed with genomic DNA of the mutant strains and primers SEN_1975_Fw (5' TTCTGATGAG-CAGCGTAAAGAGGC 3') and aslT_tetRA_(H2+P2). This PCR generates a product of 3,108 bp only if tetA and tetR genes are inserted in the correct position within ROD21.

Molecular biology techniques

The genomic DNA used in this work was prepared using the phenol-chloroform method described in [34]. PCR amplification was performed using standard PCR amplification cycles in a MaxiGene Gradient Thermocycler (Axygen). Approximately 1 ng/ml of DNA, 1 nmol/ml of each primer, 0.2 mM deoxynu-
gel-purified and cloned into the pCR-H specificitiy of the amplification reaction, some PCR products were bromide and visualized under UV light. To determine the attB-1 frequency of at a final concentration equal to 0.25 mM during the last 30 min of incubated for 2 or 18 additional hours. Hydrogen peroxide was added product as a template. PCR products were resolved by grown in LB medium and then approximately 3×10⁶ CFUs were sequenced by the Sequencing Facility at the Pontificia plasmid, according to the manufacturer’s instructions (Invitrogen) and sequenced by the Sequencing Facility at the Pontificia.Nested PCR was performed as described above using 1 µl of a first PCR product as a template. PCR products were resolved by electrophoresis in 1% agarose gels containing 0.5 µg/ml ethidium bromide and visualized under UV light. To determine the specificity of the amplification reaction, some PCR products were gel-purified and cloned into the pCR®-2.1®-TOPO cloning plasmid, according to the manufacturer’s instructions (Invitrogen) and sequenced by the Sequencing Facility at the Pontificia Universidad Católica de Chile.

Total RNA was obtained from each bacterial sample using TRIzol reagent (Invitrogen) and purified with RNeasy Mini kit (QIAGEN) according to the manufacturer instructions. After purification, RNA was treated with the DNA Free kit (Ambion) to remove contaminating genomic DNA. DNase-treated RNA was tested for DNA contamination amplifying rpoD gene by PCR. Samples with no positive amplification up to cycle 30 were considered to be DNA free. cDNA synthesis was performed using the ImPromII Reverse Transcription System (Promega) following the manufacturer instructions. One µg of RNA was used as a template and cDNA synthesis was performed using random hexamers. Reactions with no RNA or no reverse transcriptase were included to rule out gDNA contamination.

Isolation of tetracycline-sensitive S. Enteritidis S. Enteritidis PT1 ROD21::tetR4 was grown in LB medium supplemented with 25 µg/ml tetracycline at 37°C, with aeration provided by shaking, until an optical density at 600 nm (O.D₆₀₀) equal to 2.0 was reached. Serial dilutions were performed in sterile phosphate-buffered saline (PBS) to obtain 100 CFU in a volume of 100 µl. This volume was plated on LB agar supplemented with tetracycline and incubated at 37°C for 18 h. One colony was selected and grown in 2 ml of LB medium without antibiotics at 37°C for 48 h. Serial dilutions of this culture medium were prepared and aliquots of 100 µl containing 10⁶ CFU were plated on solid Bochner-Maloy medium (5 g/l tryptone, 5 g/l yeast extract, 10 g/l NaCl, 10 g/l NaH₂PO₄ · H₂O, 50 mg/l chlorotetracycline hydrochloride, 12 mg/l fusaric acid, 0.1 mM ZnCl₂, 15 g/l agar) [21]. For each experiment, 100 plates were seeded with 1×10⁶ CFU each and the amount of tetracycline sensitive colonies was quantified. In addition, aliquots of 100 µl containing approximately 100 CFU were plated on LB agar to determine the exact CFU seeded in each BM plate. After 36 h of incubation at 37°C, tetracycline-sensitive colonies were selected and grown in liquid BM medium, at 37°C for 4 hours, then patched in LB agar and replica plated in LB agar containing tetracycline and incubated overnight at 37°C. Spontaneous loss of ROD21 in tetracycline-sensitive CFU was confirmed by PCR, using primers pairs indicated in Table 2 and in Fig. 2. The frequency of ROD21 loss was calculated dividing the number of tetracycline-sensitive colonies obtained by the total number of CFU seeded in each experiment.

Mouse infection assays Groups of 4 male C57BL/6 mice (5–6 week age) were used to evaluate the virulence of strain S. Enteritidis PT1 ARO21. Infections with WT or ARO21 S. Enteritidis PT1 strains were performed by growing these bacteria in LB medium at 37°C until an OD₆₀₀ equal to 0.6 was reached. The volume of bacterial culture containing 1×10⁶ CFUs was centrifuged in a refrigerated microcentrifuge (CT15RF, Hitachi) at 10,000 × g for 5 min. Bacterial pellets were thoroughly resuspended in 20 µl of PBS and used to orally infect mice. Infective doses were corroborated by seeding serial dilutions of the bacterial inoculum onto LB plates. After infection, survival rate was recorded on a daily basis. To perform competitive assays, a kanamycin or chloramphenicol resistance gene was introduced between genes rpoD and the copy number of S. Enteritidis chromosome.

Figure 6. Exposure to peroxide induces the excision of ROD21 from the S. Enteritidis chromosome. S. Enteritidis strain PT1 was grown in LB medium and then approximately 3×10⁶ CFUs were transferred to fresh LB medium (LB) or to N-minimal medium (N), and incubated for 2 or 18 additional hours. Hydrogen peroxide was added at a final concentration equal to 0.25 mM during the last 30 min of incubation (N+H) and either genomic DNA or RNA was isolated. (A) The frequency of attB-1 excision was quantified by qPCR using genomic DNA and was expressed as a relative value equal to the ratio between the copy number of attB-1 over the copy number of rpoD gene. (B) SEN1970 expression was determined by qPCR using cDNA and expressed as a relative value (the ratio between the copy number of SEN1970 and the copy number of rpoD). The results are the average of three independent experiments. **, p<0.01, one-way ANOVA and Tukey post test.

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cleside triphosphates, 1.5 mM MgCl₂ and 50 U/ml of Taq DNA polymerase (Invitrogen) were used in these amplifications. Nested PCR was performed as described above using 1 µl of a first PCR product as a template. PCR products were resolved by electrophoresis in 1% agarose gels containing 0.5 µg/ml ethidium bromide and visualized under UV light. To determine the specificity of the amplification reaction, some PCR products were gel-purified and cloned into the pCR®-2.1®-TOPO cloning plasmid, according to the manufacturer’s instructions (Invitrogen) and sequenced by the Sequencing Facility at the Pontificia Universidad Católica de Chile.
solid LB, LB/kanamycin (50 μg/ml) and LB/chloramphenicol (10 μg/ml) that were incubated for 12–16 h at 37°C. After 72 h of infection, mice were euthanized and livers and spleens were recovered and homogenized with two slides in a Petri dish, with 2 ml PBS. To determine the total amount of both kanamycin and chloramphenicol resistant bacteria, serial dilutions of the homogenized tissue were seeded in solid LB, LB/Kanamycin (50 μg/ml) and LB/Chloramphenicol (10 μg/ml) medium and incubated for 12–16 h at 37°C. Then, colonies were counted and the competitive index was calculated as follows: (CFU WT/CFU-D\_ROD21)\_Input/(CFUWT/CFU\_D\_ROD21)\_Output. The word “input” refers to the proportion of bacteria used to infect mice and “output” to the bacteria recovered from organs.

Phagocytic cells assays

The monocyte/macrophage J774.3 cell line used in this study was kindly provided by Dr. María Inés Becker (Biosonda S.A., Chile). J774.3 cells were routinely grown in high-glucose DMEM medium (GIBCO, Invitrogen) supplemented with 10% Fetal Bovine Serum (HyClone) and 1 mM HEPES (GIBCO, Invitrogen) in T75 bottles. Cells were incubated at 37°C and 5% CO₂ until 95% of confluence. Before infection assays, cells were treated with 0.1 mg/ml trypsine (HyClone) for 5 min, recovered in 50 ml polypropylene tubes, and centrifuged at 1,800 × g for 5 min at room temperature. After three washes with supplemented DMEM medium, cell number and viability was determined in a haemocytometer, using the trypan blue staining (1 mg/ml, Invitrogen). 5 × 10⁵ cells/ml were seeded in 24 well-plates and incubated overnight at 37°C and 5% CO₂. DCs were prepared from bone marrow precursors of C57BL/6 mice. Cells were incubated in complete RPMI 1640 medium supplemented with 5% FCS (Hyclone), 2 mM glutamine, 1 mM non-essential amino acids, 1 mM pyruvate, 1 mM HEPES, and 10 ng/ml of recombinant murine GM-CSF (Peprotech). All cell culture media were acquired from GIBCO (Invitrogen). Culture media was replaced every 2 days. After 6 days, the phenotype of DCs was analyzed by flow cytometry for the expression of the surface markers CD11c, CD86 and CD40, which revealed over 70% CD11c\_+ with an immature phenotype. Before the infection assays, DCs were washed three times with PBS and then culture media was replaced with complete RPMI medium without antibiotics. DCs and macrophages were infected with S. Enteritidis PT1 at a multiplicity of infection (MOI) equal to 25. The MOI was confirmed by plating serial dilutions of bacterial cultures on LB agar. After 1 h of incubation at 37°C and 5% CO₂, the supernatant of infected cells was recovered and stored for genomic DNA preparation. Cells were washed two times with PBS and 1 ml of appropriate medium supplemented with gentamicin.

### Table 2. Primers used in this study.

| Primer | Nucleotide sequence | Coordinates* |
|--------|---------------------|--------------|
| 1      | GAGGGAAATCTTTTCGCCGTG | 2060716-2060735 |
| 2      | GGTCATGACCTGCGTTGATCA | 2060768-2060787 |
| 3      | CGGGTAACTTCTTGCACCAT | 2061726-2061707 |
| 3'     | TTGGCCGCAGACACGGGAAAGG | 2061683-2061662 |
| 4      | TACGGGGATCCCTTCCGAGCT | 2062007-2061988 |
| 5      | CAGCAGAACGGCGCAAGGACT | 2087049-2087068 |
| 6      | GCTGATGGGATTGGGATCGGG | 2087203-2087222 |
| 7      | GTTAATGACCTGGTAAAGGCAG | 2087380-2087411 |
| 8      | AGTGGGCTTATGGGATCCG | 2087962-2087943 |
| 9      | TCATAATCACCAGGGAGCTG | 2098867-2098886 |
| 10     | CCCAGGCTTAAAAGGGCAACCAC | 2098923-2098942 |
| 11     | CAGGCCCTGGCCTTTAATAATCCT | 2099721-2099699 |
| 12     | GTTAATGAGGTGCTTGGAGG | 2099774-2099755 |
| SEN1975 Fw | TTCTGTGACAGGCGTAAAGGCC | 2065343-2065366 |
| SEN1975 Rev | GCAAGGGACGGCAGAAAATCTAC | 2065807-2065784 |
| asfT\_tetRA\_(H1\_P1) | GGAACCTCTTCATGGAGAATACACATCCTCGGGAAAAAACCATTAAGTAAAGTGGGA** | 2066174-2066213 |
| asfT\_tetRA\_(H2\_P2) | ATGTTTTGTGTTTTAACATTTAAATTTATACTTTAACTTTAATCAGAAATCCTGGTAAAGTGGGA** | 2066360-2066321 |
| SEN\_1075\_Fw | TCTGACGAGGACGGAGCAGG | 2065343-2065366 |
| rpoD-Fw-2 | TGGTCTGAAATCTGGCTGAG | 3264619-3264640 |
| rpoD-Rev | TTCGGGGAATACGCTGAGCAAT | 3266113-3266094 |
| rpoD-RT-Fw | GTGGACCCGGGAAGGGCAACCA | 3266488-3266470 |
| rpoD-RT-Rev | CAGAACCACGGTGGTGTCGG | 3264908-3264889 |
| SEN1970-RT-Fw | CGATGCTGTGAGGAGGCGCT | 2061725-2061745 |
| SEN1970-RT-Rev | GGCACCGCTCTTTCATCATC | 2061954-2061934 |
| attB1nested-RT-Fw | GTTAATGACCTGGTAAAGGCAG | 2061082-2061104 |
| attB1nested-RT-Rev | CCGATTAAAGCCCTAAAATAGTATG | 2087781-2087759 |

*Coordinates are those of the S. enterica serovar Enteritidis PT4 NTCT NCTC13349 sequence.
**Italics indicate the region that anneals to the 5’ or 3’ end of a mini Tn10 transposon.

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50 mg/ml was added to cell cultures to kill the remaining extracellular bacteria. After 2, 18 and 24 h of infection, cells were removed from the wells and centrifuged at 783 x g for 5 min. No significant changes on cell viability were observed after infection with S. Enteritidis at the time points used on the experiments (data not shown). To recover intracellular bacteria, phagocytic cells were treated with 1 ml of lysis solution (19% ethanol, 0.1% SDS, 1% saturated basic phenol) for 30 min on ice. After the incubation period, the cell lysate was centrifuged at 7.043 x 10^6 for 2 h and after that time bacteria were recovered by incubation in 1 ml of cell medium. The incubation was performed for 2 h and after that time bacteria were recovered by centrifugation at 7.043 x g for 5 min, and genomic DNA was prepared as described above.

**Peroxide treatment**

To evaluate whether oxidative stress has a role in the excision of ROD21, we used hydrogen peroxide (H₂O₂) as an oxidative agent and analyzed its effects in bacterial cultures grown in LB, N-minimal medium, and N-minimal medium+H₂O₂. S. Enteritidis strain PT1 was grown in LB until OD₆₀₀ equal to 0.6 was reached. Then, the culture was split in 6 samples, each containing approximately 3 x 10⁹ bacteria. The samples were pelleted and then two of them were resuspended in fresh LB medium while the other four were resuspended in N medium. From these 6 new cultures, two of the N cultures were incubated for an additional 1.5 hour. Next, 0.25 mM H₂O₂ (Merck) was added and cultures were incubated for additional 30 min. The other two N samples were incubated for 17.5 hours, then 0.25 mM H₂O₂ (Merck) was added and cultures were incubated for additional 30 minutes. After the H₂O₂ treatment, all samples were pelleted and stored at −80°C until analyzed. Genomic DNA or total RNA was prepared as described above.

**Quantitative real time PCR assays**

Quantitative real time PCR for the quantification of ROD21 excisions was performed using Brilliant SYBR Green QPCR kit (STRAGENE), following the manufacturer’s instructions. The reaction mixture contained 2 µl of genomic DNA as a template and 0.12 pmol/µl of each primer. Standard curves for attB-1 and rpoD were generated using serial dilutions of a plasmid containing the corresponding PCR fragment for attB-1 and rpoD. Thermal cycling conditions were: segment 1; one cycle at 50°C for 2 min followed by initial denaturation at 95°C for 10 min; segment 2, 40 cycles of 30 s at 95°C, 1 min at 50°C, 1 min at 72°C and 15 s at 90°C; segment 3, 1 cycle of 1 min at 95°C, 30 s at 55°C and 30 s at 95°C. The copy number of attB-1 and rpoD was calculated as follows: The number of attB-1 and rpoD copy number 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 that allows us to know the number of copies of the sample, which has a specific Ct [35]. The results were expressed as the ratio of attB-1 copy number/rpoD copy number (relative value).

Quantitative real time for the quantification of SEN1970 expression was performed as described above, but using as a template 2 µl of a 10⁻¹ dilution of the RT-PCR reaction. rpoD was used as reference gene to normalize the copy numbers of SEN1970. Standard curves were made of serial dilutions of purified PCR products.

**Statistical analyses**

Statistics significance was determined using the analyses of variance (ANOVA) test with the Prism Graphpad software.

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**Author Contributions**

Conceived and designed the experiments: SMB AMK. Performed the experiments: TSQ HET RJL PAN CPQ DVA. Analyzed the data: TSQ PAN AMK SMB. Contributed reagents/materials/analysis tools: SMB CAR CAS AMK. Wrote the paper: SMB PAN AMK.

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