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Insertion Hot Spot for Horizontally Acquired DNA within a Bidirectional Small-RNA Locus in *Salmonella enterica*[∇]

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In *Escherichia coli* and *Salmonella enterica*, RyeA and RyeB RNAs are encoded on opposite DNA strands at the same locus. We present evidence indicating that the last 23 bp of the *ryeB* gene, corresponding to an internal portion of the *ryeA* gene, served repeatedly as the integration site for exogenous DNA during *Salmonella* evolution and still act as an attachment site for present-day bacteriophages. Interestingly, *ryeA* sequence and expression are modified upon lysogenization.

Lateral gene transfer is a prolific source of evolutionary changes in microorganisms and is thought to have had a major impact in the emergence of bacterial pathogens. In particular, the acquisition of so-called pathogenicity islands is regarded as being a key event in the conversion of ancestral extracellular bacteria into intracellular pathogens (2, 9, 11, 15). A common pathway to DNA acquisition involves the integrative recombination of circular DNA molecules into the host genome. The step is catalyzed by integrases, a class of site-specific recombinases encoded by temperate bacteriophages and plasmids. Integrases introduce staggered cuts at specific sequences on both donor and host DNAs and promote strand exchange and ligation (8, 14). As a result, the sequences recognized by the integrase are duplicated at each end of the inserted DNA. When conserved, such directed repeats allow a precise definition of the site of the original integration event.

Structure of the *Salmonella* CS 40 island. *Salmonella enterica* serovars harbor a panoply of genomic islands and islets. Some such elements are associated with prophage remnants, suggesting that they were incorporated as a result of lysogenization events. This is the case for a 16-kb insert lying at about centisome (CS) 40 on the chromosome map. The CS 40 island contains various loci linked to pathogenicity, such as *mig-3* (18) *pagK-pagO* (10), and *sopE2* (1, 16), interspersed with sequences reminiscent of phage genes. Among the latter is a presumptive integrase gene (STM1871) (12) located near one end of the island and oriented outwards. As in lambdoid prophage maps (*int* gene near *attL*), the STM1871-proximal end is hereafter designated the “left” end of the island (Fig. 1). Alignment with the *Escherichia coli* K-12 genome sequence reveals that the *Salmonella* CS 40 island is inserted into the intergenic region between the *pprA* gene (also named *pphA*) and the ortholog of the *yebY* locus (STM1873) (Fig. 1). Recently, the *pprA-yebY* intergenic region of *E. coli* was shown to contain two small-RNA (sRNA) genes with opposite polarities, *ryeA* and *ryeB*, transcribed from opposite DNA strands, with the *ryeB* sequence entirely contained within the larger

ryeA gene (19, 20). The core region of this locus, a 146-bp segment comprising the entirety of *ryeB*, is highly conserved in *Salmonella* and is located immediately to the left of the CS 40 island near the end of the putative integrase gene.

Since we wondered if the interval between *ryeB* and STM1871 contained the integrase recognition site, we examined whether a portion of the sequence was repeated on the opposite end of the island. No such repeat could be identified at the right end of the element; however, a sequence identical to the last 23 bp of the *ryeB* gene, and in the same orientation, was located inside the island, approximately 10 kb from the left end. Interestingly, this sequence lies adjacent to an open reading frame (STM1861) whose putative product shares 77% identity (89% similarity) with the C-terminal half of the STM1871-encoded integrase. This strongly suggests that the CS 40 island is in fact made of two separate islets lying side by side, one carrying *mig-3* and *pagKO* (left) and the other containing the *sopE2* gene (right) (Fig. 1). The lack of a recognizable attachment site at the right end of the insert, as well as the apparent defective nature of STM1861, suggests that the *sopE2* islet was acquired earlier and has since suffered extensive decay. Consistent with this idea, *Salmonella bongori*, a lineage sharing a common ancestor with *Salmonella enterica*, carries *sopE2* (13) but lacks the *mig-3-pagKO* islet (data not shown).

The above-described findings tentatively define the last 23 bp of the *ryeB* gene as the core region of the integration site. Since this sequence is reconstituted upon integration, the acquisition of the CS 40 islets is not expected to have affected *ryeB* gene structure or expression. In contrast, if the *ryeA* gene is positioned in *Salmonella* as in *E. coli*, the incorporation of the islets should have separated the structural portion of the gene from its original promoter. Thus, it seemed relevant to determine the *ryeA* transcriptional status in *Salmonella*. For this purpose, RNA extracted from *S. enterica* serovar Typhimurium strain LT2 was subjected to Northern hybridization analysis using oligonucleotides complementary to the predicted sequences of RyeA and RyeB RNAs as probes. As shown in Fig. 2, both probes gave positive signals. In the case of RyeA, the most intense band corresponded to an RNA of 250 to 300 nucleotides, while the RyeB analysis detected an RNA of approximately 100 nucleotides. These sizes are closely comparable to those of RyeA and RyeB RNAs in *E. coli* (19,

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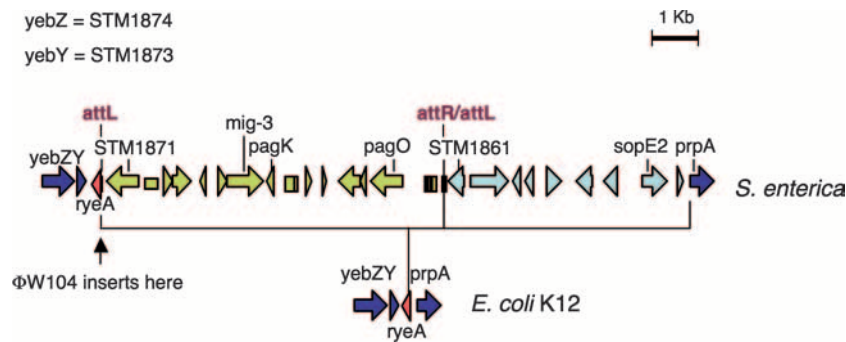


FIG. 1. Organization of the CS 40 island in *Salmonella enterica* serovar Typhimurium and of the corresponding region in the *E. coli* chromosome. Horizontal arrows represent open reading frames. The arrow clusters depicted in green and light blue correspond to segments proposed to originate from separate insertion events (see text). Phage Φ W104 inserts at the left boundary of the island. The diagram is based on data from references 10, 12, 16, 19, and 20).

20). As observed in *E. coli* (20), additional fainter signals were detected with both probes. These minor bands are likely to represent processing products.

A survey of the *Salmonella* genome sequences revealed the

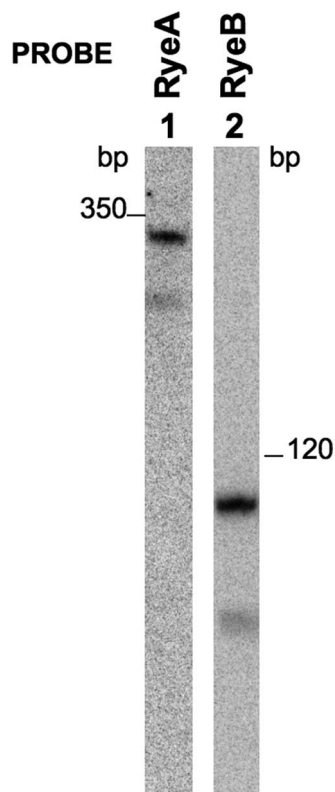


FIG. 2. Northern blot analysis of RyeA and RyeB RNAs in *Salmonella enterica* serovar Typhimurium LT2. Cultures grown overnight in LB were diluted 1:200 in fresh LB and grown to an optical density at 600 nm of 0.35. RNA was extracted as previously described (4), fractionated on a 6% polyacrylamide-8 M urea gel (lane 1) or on an 8% polyacrylamide-8 M urea gel (lane 2), transferred onto a Hybond-N⁺ membrane, and hybridized to ³²P-labeled oligonucleotides complementary to RyeA (pp925 [5'-GGAAAACCTGGCGTCGTCATCTATTCTTAAAGGGC AAGGCGA-3']) and RyeB (pB13 [5'-GATTCTGTATTCGGTCCAG GGAAATGGCTCTTGGGAGAGAG-3']). Sizes were estimated from migration distances of tmRNA and 5S RNA (not shown).

occurrence of yet another insertion event at the *ryeA/ryeB* locus. Some strains carry a prophage-related insert between the end of *ryeB* and the left boundary of the *mig-3-pagK* islet (Fig. 1). The length and structure of the *Salmonella ryeA/ryeB* locus insert vary considerably, from a full-length prophage in some isolates (e.g., *S. enterica* serovar Typhi strain CT18) (17) to a shortened and scrambled version in others (e.g., *S. enterica* serovar Enteritidis strain LK5) (7). In all instances, the terminal 23 to 26 bp of *ryeB* are found duplicated at the two ends of the element, indicating that this portion of the gene serves as the attachment site. Thus, in strains carrying the *ryeB*-linked prophage, *ryeA* transcription is expected to originate from within phage DNA.

While tRNA and tmRNA genes constitute a favored target for temperate phages and other integrative elements (21), to our knowledge, only one example of the phage insert in an sRNA gene has been reported. Wassarman and colleagues found the previously mapped *attB* site for bacteriophage P2 in *E. coli* to correspond with the 3' end of the *ryeE* gene, which encodes an Hfq-binding sRNA of unknown function (20). In both the *ryeB* and *ryeE* genes, the attachment site lies within the sequence encoding the stem-loop of the transcription terminator, suggesting that the region of dyad symmetry participates in integrase recognition (14, 21).

Effect of phage integration on *ryeA/ryeB* expression. As part of a separate study, we examined the occupancy of the *ryeA/ryeB att* site in 84 *Salmonella enterica* serovar Typhimurium isolates of human or animal origin using a three-primer-based PCR assay. This analysis revealed the presence of a DNA insert in a fraction of the strains. Interestingly, all positive strains belonged to the DT104 phage type, suggesting that the acquisition of the insert occurred recently, possibly coinciding with the emergence of the virulent epidemic clone (3; N. Figueroa-Bossi, F. X. Weill, P. A. Grimont, and L. Bossi, unpublished data). Sequence data from the Sanger Institute website (<http://www.sanger.ac.uk/Projects/Salmonella/>) show the *ryeA/ryeB*-associated element to be a full-size lambdoid prophage. To assess its functional state, we deleted the prophage from the genome of a DT104 strain in our collection, MA6711 (5), and used the resulting strain (MA7860) as a host for monitoring the release of plaque-forming particles in cultures of the MA6711 parent. Tiny plaques from which active

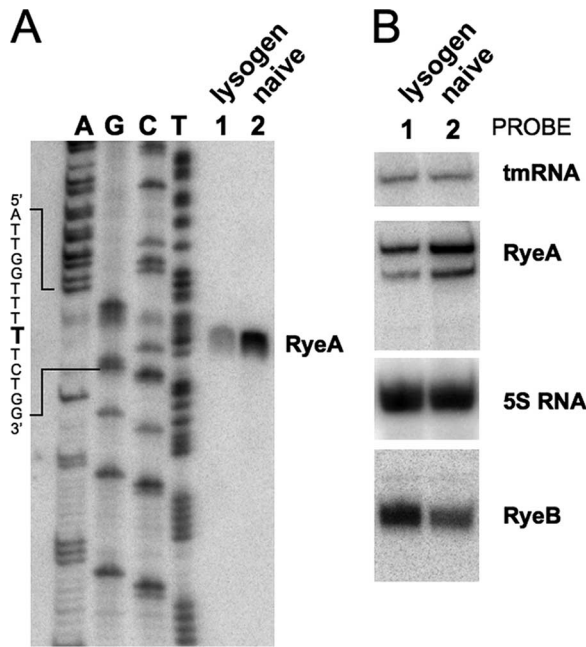


FIG. 3. Effect of phage Φ W104 integration on *ryeA/ryeB* expression. Bulk RNA was extracted from strains LT2 and MA7833 (Φ W104 lysogen) as described in the legend to Fig. 2. (A) Primer extension analysis of RyeA RNA from the lysogenic strain (lane 1) and from wild-type LT2 (lane 2). Reverse transcriptase reactions were carried out using primer ppB12 (5'-CCCTGGACCGAATACAGGA-3') as previously described (6). Sequencing reactions were performed with the *fmol* DNA cycle sequencing system from Promega according to the manufacturer's protocol. The template was a DNA fragment obtained by PCR amplification of chromosomal DNA from strain MA7833 with oligonucleotides pp490 (5'-TGGCGTCGCATCTATTC-3') and pp491 (5'-CAGGGACGCTATCACACA-3') as primers. (B) Northern blot quantification of RyeA and RyeB levels in strains MA7833 (lane 1) and wild-type LT2 (lane 2). Bulk RNA was fractionated on an 8% polyacrylamide-8 M urea gel. Membranes were probed with 32 P-labeled oligonucleotides pp925 and ppB13 (see the legend to Fig. 2). 5S RNA and tmRNA probed with ppB10 (5'-ACACTACCATCGGC GCTACG-3') and pp813 (5'-GCGGAGGCTAGGGAGAGAGG-3'), respectively, were used as internal controls. Due to the higher gel concentration, the two RyeA bands are less separated than in Fig. 2.

virus could be isolated and characterized were detected. The phage, hereafter named Φ W104, proved capable of infecting and lysogenizing a variety of serovar Typhimurium strains including LT2, ATCC 14028, and SL1344. The isolation of such lysogenic derivatives provided a system for studying how Φ W104 integration affected the expression of the *ryeA/ryeB* locus. To identify the *ryeA* promoter, RNA preparations from exponential cultures of strain LT2 and its Φ W104-lysogenized derivative MA7833 were subjected to primer extension analysis (Fig. 3A). These experiments located the 5' end of RyeA RNA at identical positions in the two strains, approximately 80 bp to the right of the *ryeB* gene (Fig. 4). The presence of sequences resembling the -10 and -35 consensus motifs of σ^{70} -dependent promoters immediately upstream from the 5'-end position is consistent with this being the *ryeA* transcription start site. Overall, the sequences around this region in naive and lysogenic strains are highly conserved (Fig. 4), with the identity extending to the Φ W104 putative *int* gene that strongly resembles STM1871 (see above). Nonetheless, the difference in the intensities of the primer extension bands shown in Fig. 3A suggested that RyeA RNA might be more abundant in strain LT2 than in its lysogenic derivative. The difference was confirmed by Northern blot hybridization analysis (Fig. 3B). Interestingly, RyeB followed an opposite trend, being synthesized at a lower level in LT2 than in MA7833 (Fig. 3B). Since the *ryeB* sequence is unaffected by the lysogenization event, the observed difference might reflect the change in *ryeA* transcription associated with such an event. Conceivably, RNA polymerases transcribing the *ryeA* gene could dislodge polymerases bound to the *ryeB* promoter, causing the activity of the latter to negatively correlate with that of the *ryeA* promoter. Some variability in the sequence upstream from the *ryeA* promoter, particularly a 9-bp deletion/insertion at position -49 (Fig. 4), might account for the difference in *ryeA* transcription rates.

Biological significance of prophage insertion at the *ryeA/ryeB* locus. The above-described data suggest that the integration of the Φ W104 prophage "resets" the levels of RyeA and RyeB RNAs in the cell. In addition, the 5' portion of RyeA RNA is changed upon lysogenization. It is tempting to specu-

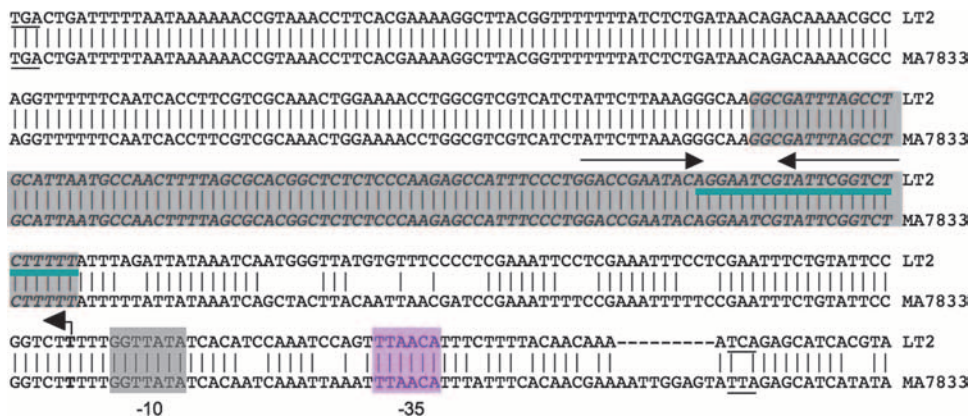


FIG. 4. Sequence alignment of *ryeA/ryeB* chromosomal regions in LT2 and Φ W104 lysogen. Underlined triplets indicate the translation termination codon of *yebY* and the complements of termination codons of STM1871 (LT2) (12) and the Φ W104 *int* gene (MA7833). The sequence of the *ryeB* gene is in italics and shaded. Gene boundaries are inferred from the 96% identical *E. coli* sequence (19, 20). Facing arrows indicate the transcription terminator stem sequence. Green underlining indicates the sequence found duplicated at the two ends of the Φ W104 prophage. The left-pointing arrow defines the *ryeA* transcription start site. Purple boxes indicate *ryeA* promoter elements.

late that these modifications might have physiological consequences. Unfortunately, the lack of information on the physiological roles of RyeA and RyeB does not offer much grounds for such speculation. RyeB RNA was shown to strongly bind the Hfq protein (20), suggesting its involvement in some Hfq-mediated regulatory mechanism. In contrast, RyeA bound Hfq with low affinity (20). Possibly, the role of this RNA is limited to regulating RyeB levels through transcriptional interference, as suggested above, or by a direct RNA-RNA interaction (19).

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