

## The SPI-3 Pathogenicity Island of *Salmonella enterica*

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**Pathogenicity islands are chromosomal clusters of pathogen-specific virulence genes often found at tRNA loci. We have determined the molecular genetic structure of SPI-3, a 17-kb pathogenicity island located at the *selC* tRNA locus of *Salmonella enterica* serovar Typhimurium. The G+C content of SPI-3 (47.5%) differs from that of the *Salmonella* genome (52%), consistent with the notion that these sequences have been horizontally acquired. SPI-3 harbors 10 open reading frames organized in six transcriptional units, which include the previously described *mgtCB* operon encoding the macrophage survival protein MgtC and the Mg<sup>2+</sup> transporter MgtB. Among the newly identified open reading frames, one exhibits sequence similarity to the ToxR regulatory protein of *Vibrio cholerae* and one is similar to the AIDA-I adhesin of enteropathogenic *Escherichia coli*. The distribution of SPI-3 sequences varies among the salmonellae: the right end of the island, which harbors the virulence gene *mgtC*, is present in all eight subspecies of *Salmonella*; however, a four-gene cluster at the center of SPI-3 is found in only some of the subspecies and is bracketed by remnants of insertion sequences, suggesting a multistep process in the evolution of SPI-3 sequences.**

The gram-negative bacterium *Salmonella enterica* is responsible for a variety of diseases, which include gastroenteritis and typhoid fever, depending on the nature of the infected host and on the serovar of the infecting bacteria. *Salmonella* has a complex life cycle in infected animals, and a large number of genes have been implicated in *Salmonella* virulence. Several of these virulence determinants are clustered within pathogenicity islands, i.e., large segments of horizontally acquired sequences present in pathogenic species but absent from closely related nonpathogenic species (17, 24). Pathogenicity islands constitute major elements in the evolution of bacterial pathogens, because their incorporation can, in a single step, transform a normally benign organism into a pathogen.

In addition to several small pathogenicity islets, five large pathogenicity islands have been identified in *Salmonella* (21, 51, 52). SPI-1, at 63 min on the *S. enterica* serovar Typhimurium chromosome, is a 40-kb island that governs the ability to invade epithelial cells (10, 38) and is required for *Salmonella*-induced macrophage apoptosis (9). The SPI-2 island is also 40 kb in length, maps downstream of a tRNA<sup>Val</sup> locus at 31 min (25), and harbors genes required for intramacrophage survival and systemic infection (40, 44). The SPI-3 island is located at 82 min, immediately behind *selC*, a tRNA locus that is the insertion site for distinct pathogenicity islands in enteropathogenic and uropathogenic strains of *E. coli* (3, 5, 33). Recently, a 27-kb *Salmonella*-specific DNA fragment at 92 min was designated the fourth *Salmonella* pathogenicity island because it includes a macrophage survival locus (34). A fifth pathogenicity island, containing genes mediating *Salmonella* enteropathogenesis, is located downstream of a tRNA<sup>Ser</sup> locus at 20 min in the chromosome (52).

The SPI-3 island harbors *mgtC*, a *Salmonella*-specific gene that is required for intramacrophage survival, virulence in mice,

and growth in low-Mg<sup>2+</sup> media (3). The *mgtC* gene is transcriptionally controlled by the PhoP-PhoQ regulatory system, which governs the adaptation to low-Mg<sup>2+</sup> environments (15, 46) and is the major regulator of virulence functions in *Salmonella* (14, 18). The *mgtC* gene is cotranscribed with *mgtB* (45), a Mg<sup>2+</sup> transporter gene dispensable for virulence in BALB/c mice (3). SPI-3 is 17 kb long and may contain additional genes that contribute to virulence or to other *Salmonella*-specific attributes.

In this study, we determined the molecular genetic structure of the SPI-3 island, examined the functions of the genes it carries, and investigated the distribution of SPI-3 sequences among salmonellae. We establish that at least 10 genes are encoded within SPI-3, some of which show similarity to known virulence factors from other bacterial species, and that the evolution of SPI-3 sequences occurred through a multistep process.

### MATERIALS AND METHODS

**Bacterial strains, plasmids, bacterial genetic techniques, and growth conditions.** Strains used in this study are listed in Table 1. The strains used in this study are derived from 14028s, except for TT10288 and AA3007, which are derived from LT2. Bacteria were grown at 37°C in Luria-Bertani broth (LB) (35). Ampicillin and kanamycin were used at 50 µg/ml, and chloramphenicol was used at 10 µg/ml. Phage P22-mediated transduction was carried out as described previously (12).

Construction of the *marT::cat* strain EG10207 was performed as follows. A 2.3-kb *Hind*III-*Bsp*MII fragment from plasmid pEG9106 (3) carrying *marT* was subcloned into pUC19 between the *Hind*III and *Xma*I sites to form plasmid pEG9109. A *cat*-containing 0.8-kb *Bam*HI fragment from plasmid pKRP10 (43) was introduced into the unique *Bg*II site in *marT* in plasmid pEG9109. The resulting plasmid (pEG9110) was used to transfer the *marT::cat* mutation into the *Salmonella* chromosome as described previously (23). The structure of the *marT* gene in the mutant strain was verified by Southern hybridization with both *marT*- and *cat*-specific probes (data not shown).

MudJ is a derivative of bacteriophage Mu that harbors a gene conferring resistance to kanamycin and a segment of the *lac* operon devoid of its promoter sequences (7). To isolate MudJ insertions in SPI-3, a P22 lysate grown in TT10288 was used to infect strain EG10207. A lysate grown on a pool of 25,000 kanamycin-resistant transductants was used to infect 14028s, with selection for both kanamycin and chloramphenicol resistance. To establish the orientation and approximate position of each MudJ insertion, PCR was performed with primers complementary to the ends of MudJ (i.e., *attL* or *attR*) and to known SPI-3 sequences.

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TABLE 1. *Salmonella* strains used in this study

| Strain  | Description                           | Reference or source              |
|---------|---------------------------------------|----------------------------------|
| 14028s  | Wild type                             | American Type Culture Collection |
| TT10288 | <i>hisD9953::MudJ hisA9944::MudI</i>  | 20                               |
| AA3007  | <i>polA2 ara-9</i>                    | 21                               |
| MS7953s | <i>phoP7953::Tn10</i>                 | 22                               |
| EG9527  | <i>mgtCB9232::MudJ</i>                | 15                               |
| EG9529  | <i>mgtCB9232::MudJ phoP7953::Tn10</i> | 15                               |
| EG10207 | <i>marT1::cat</i>                     | This work                        |
| EG10209 | <i>mgtCB9232::MudJ marT1::cat</i>     | This work                        |
| EG10277 | <i>sugR1::MudJ</i>                    | This work                        |
| EG10278 | <i>sugR1::MudJ marT1::cat</i>         | This work                        |
| EG10349 | <i>sugR2::MudJ</i>                    | This work                        |
| EG10389 | <i>sugR1::MudJ phoP7953::Tn10</i>     | This work                        |
| EG10755 | <i>misL1::MudJ</i>                    | This work                        |
| EG10756 | <i>misL1::MudJ marT1::cat</i>         | This work                        |
| EG10757 | <i>misL1::MudJ phoP7953::Tn10</i>     | This work                        |
| EG10759 | <i>rhuM1::MudJ</i>                    | This work                        |
| EG10760 | <i>rhuM1::MudJ marT1::cat</i>         | This work                        |
| EG10913 | <i>rhuM1::MudJ phoP7953::Tn10</i>     | This work                        |
| EG11086 | <i>marT2::MudJ</i>                    | This work                        |
| EG11087 | <i>marT2::MudJ phoP7953::Tn10</i>     | This work                        |

**Molecular biological techniques.** The nucleotide sequence of the 12-kb segment between the *selC* and *mgtB* genes was determined on both strands by using plasmid pEG9106 DNA as the template, starting with primers complementary to *selC* (*selC*-F) (11) and to the 3' end of *mgtB* (3' *mgtB*-F, 5'-ATCGTCGTGGTTAAACCGCGTCC-3') and walking with newly synthesized primers. DNA sequence analysis and protein sequence alignments were performed by using the GeneWorks (IntelliGenetics) and Genetics Computer Group (GCG) (University of Wisconsin) software packages.

PCRs were carried out on purified chromosomal DNA with *Taq* polymerase (Gibco BRL) according to the manufacturer's protocol. For amplification of long DNA fragments, we used the TaqPlus Long PCR system (Stratagene). To examine whether SPI-3 sequences are linked to *selC* in different *Salmonella* subspecies, we used primers *selC*-F (11), *selC*-1-25 (5'-GGAAGATCGTCGTCCGGTGAGGC-3'), and *slsA*-R (5'-TTGTACAAAATCGGCATTATCCCAGGC-3'). To determine whether *mgtC* and *orf307* are linked in different *Salmonella* subspecies, we used primers *mgtC*-R (5'-GCCCGCCCCAGAAAGCCAATCC-3') and E07-R (11).

Southern hybridization analysis was carried out with chromosomal DNA as described previously (11). To investigate the distribution of the *orf269* gene, a PCR-generated probe corresponding to the *Escherichia coli* K-12 *orf269* open reading frame (ORF) was used for hybridization to DNAs from *E. coli* K-12, *E. coli* D, *Shigella flexneri*, *S. enterica* serovar Typhimurium, *Citrobacter freundii*, *Enterobacter aerogenes*, *Enterobacter cloacae*, *Klebsiella pneumoniae*, *Serratia odif-*

*era*, *Yersinia enterocolitica*, *Yersinia pestis*, *Haemophilus influenzae*, *Mycobacterium avium*, and *Pseudomonas aeruginosa*. To investigate the distribution of SPI-3 sequences among salmonellae, probes were hybridized to DNAs from strains of the *Salmonella* Reference Collection C (6) and from *E. coli* K-12 strain MC1061. Probe 1 (410 bp) was generated from a PCR DNA fragment by using primers *selC*-415 (5'-AGATGATGTGGCTGGCG-3') and *selC*-R (11), probe 2 (780 bp) was generated by using primers described previously (3), and probes 3, 4, 6, 7, 8, and 9 were generated by using primers complementary to the 5' and 3' ends of the *sugR*, *rhuM*, *marT*, *slsA*, *mgtB*, and *mgtC* genes, respectively. Probe 5 was generated from a 4-kb *EcoRI*-*HindIII* restriction fragment (3).

**Virulence and  $\beta$ -galactosidase assays.** Macrophage survival assays with the macrophage-like cell line J774 and invasion assays with canine kidney epithelial (MDCK) cells were conducted as described previously (28). Virulence assays were performed with 7- to 8-week-old female BALB/c mice (10 mice per mutant) inoculated orally with 100  $\mu$ l of bacteria diluted in phosphate-buffered saline.  $\beta$ -Galactosidase assays were carried out in triplicate with bacteria grown exponentially in LB as described previously (35).

**Nucleotide sequence accession number.** The sequence reported in this paper has been deposited in the GenBank database (accession no. AF106566).

## RESULTS AND DISCUSSION

### Molecular analysis of SPI-3 genes and encoded proteins.

We have previously identified a pathogenicity island downstream of the *selC* gene in the *S. enterica* serovar Typhimurium chromosome (3). This island includes the *mgtCB* operon, which codes for the virulence protein MgtC and the Mg<sup>2+</sup> transporter MgtB. To further examine the role of SPI-3 in *Salmonella*, we determined the molecular genetic structure of the DNA region between the *selC* and *mgtB* genes (Fig. 1).

The SPI-3 island is 17 kb long, and in addition to the *mgtCB* operon, it harbors eight ORFs, all of which contain potential Shine-Dalgarno sequences (Fig. 1). (Additional small ORFs [encoding <120 amino acids] lacking a clear translation start site and similarity to proteins in the sequence databases are not reported here). The 10 genes carried within SPI-3 appear to be organized in six transcriptional units. Characteristics of the SPI-3 ORFs, as well as features of DNA stretches that have identity with sequences in the databases, are described in Table 2. Except for the *rmbA* gene, which is part of region with a very low G+C content, the codon usages of SPI-3 ORFs do not appear to be significantly different from those of highly expressed *E. coli* and *Salmonella* genes.

The first gene of the island, *sugR*, encodes a protein that exhibits closest similarity to the PgaA antigen of the periodontopathogen *Porphyromonas gingivalis* (43a) and to a putative

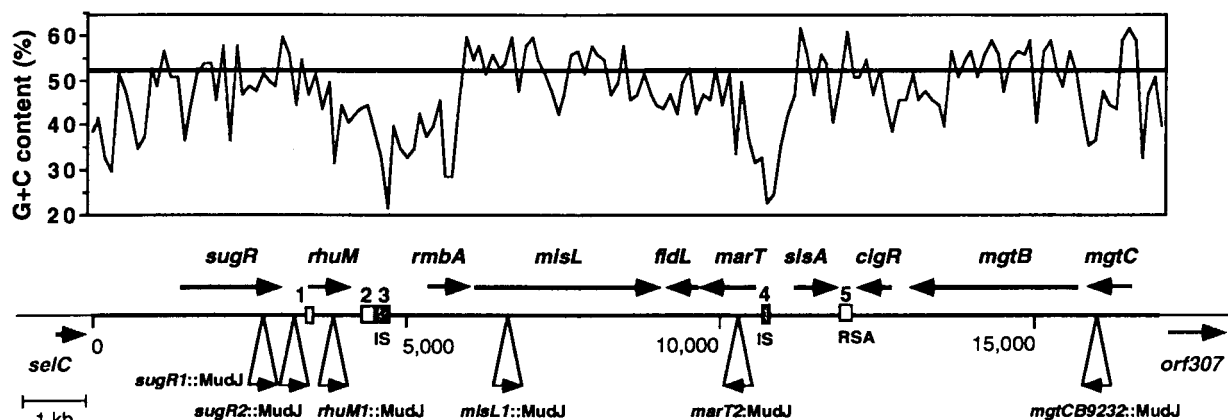


FIG. 1. Physical and genetic maps of the SPI-3 pathogenicity island. (Top) G+C content of the SPI-3 island. The graph was created by using the program Cricket Graph with data generated by the program Windows (GCG) (window, 100 bp; sliding increment, 100 bp). The line at 52% indicates the overall G+C content estimated for the *S. enterica* serovar Typhimurium chromosome. (Bottom) Positions and orientations of ORFs encoding products larger than 120 amino acids and containing potential Shine-Dalgarno sequences. DNA sequences reported in Table 2 are indicated by numbers (IS-like sequences are represented by gray squares). The map positions of MudJ insertions in the SPI-3 region are indicated by triangles. RSA refers to a family of protected sequences present in the genomes of members of the family *Enterobacteriaceae*.

TABLE 2. Properties of SPI-3 DNA sequences and SPI-3-encoded proteins

| Sequence               | Length              | Characteristics <sup>a</sup>  |
|------------------------|---------------------|---|
| <b>DNA<sup>b</sup></b> |                     |   |
| 1                      | 150 bp              | 83% identical over 125 nucleotides to sequences downstream of <i>E. coli selC</i> (AE000443)  |
| 2                      | 190 bp              | 91% identical to sequences downstream of <i>fimU</i> in <i>S. typhimurium</i> (L19338) and 87% identical to sequences upstream of <i>insA</i> in <i>E. coli</i> (L20943)  |
| 3                      | 200 bp              | 82% identical to IS1351 of <i>S. enteritidis</i> (Z83734)   |
| 4                      | 100 bp              | 95% identical to repetitive sequence 1-3 in <i>E. cloacae</i> (D00952); 76% identical to IS911 of <i>S. dysenteriae</i> (X17613)  |
| 5                      | 190 bp              | RSA repetitive sequence (interrupted by an additional 60 bp)  |
| <b>Proteins</b>        |                     |   |
| SugR                   | 519 aa <sup>c</sup> | Predicted cytoplasmic protein; 44% identical over 153 aa to PgaA antigen of <i>P. gingivalis</i> (X95938) and 34% identical over 249 aa to putative ATP binding protein from <i>E. coli</i> clinical isolate (S28007)   |
| RhuM                   | 215 aa              | Predicted cytoplasmic protein   |
| RmbA                   | 205 aa              | Predicted cytoplasmic protein; 39% identical over 190 aa to 230-aa <i>E. coli</i> ORF product (U29581)  |
| MisL                   | 955 aa              | Probable autotransported protein; putative N-terminal signal sequence; C-terminal domain is 43% identical to C-terminal domain of <i>E. coli</i> AIDA-I (Q03155) and 38% identical to C-terminal domain of <i>S. flexneri</i> VirG (A32247)   |
| FidL                   | 154 aa              | Predicted inner membrane protein; putative N-terminal signal sequence; 44% identical over 124 aa to 164-aa <i>E. coli</i> ORF product (U29581)  |
| MarT                   | 285 aa              | Predicted inner membrane protein (putative transmembrane domain comprises residues 172–188); N-terminal domain is 68% identical to 99 aa of 269-aa <i>E. coli</i> ORF product (U29581) and 32% identical to 98 aa of <i>V. cholerae</i> ToxR protein (P15795); C-terminal domain is 31% identical to 116 aa of 269-aa <i>E. coli</i> ORF product (U29581) |
| SlsA                   | 226 aa              | Predicted inner membrane protein (putative transmembrane domain comprises residues 114–130); 48% identical over 197 aa to <i>E. coli</i> ORF product (P21367)   |
| CigR                   | 159 aa              | Predicted membrane protein (putative transmembrane domain comprises residues 134–158); putative cleavable N-terminal signal sequence  |
| MgtB                   | 908 aa              | Mg <sup>2+</sup> transporter  |
| MgtC                   | 231 aa              | Predicted membrane protein; virulence determinant   |

<sup>a</sup> Similarities with other sequences as revealed by BLAST searches of sequence databases (<http://www.ncbi.nlm.nih.gov/BLAST>). Accession numbers in the EMBL database are indicated in parentheses. Prediction of protein localization sites was performed by using the PSORT WWW server (<http://psort.nibb.ac.jp/>).

<sup>b</sup> For DNA sequences 1 through 5, see Fig. 1.

<sup>c</sup> aa, amino acids.

ATP binding protein encoded in the genome of a clinical isolate of *E. coli* (29). The SugR protein contains an imperfect nucleotide-binding Walker A motif (APNGAGKT) that is missing the first conserved G of the consensus Walker sequence (GXXGXGKS/T) (48).

The MisL (for membrane insertion and secretion) protein exhibits similarity to the immunoglobulin A1 protease family of autotransported proteins, which have been found only in pathogenic bacteria (26, 32). These proteins consist of an N-terminal effector domain and a C-terminal conserved domain that forms a pore in the outer membrane through which the N-terminal domain is translocated. The similarity between MisL and the AIDA-I protein from enteropathogenic *E. coli* and the VirG protein from *S. flexneri* is limited to the C-terminal region (Fig. 2A), suggesting a similar autotransporter function rather than specific functional similarities with these two proteins, which have been implicated in diffuse adherence to HeLa cells (2) and cell-to-cell spreading (16), respectively. Neither the 955-amino-acid MisL protein nor the 1,286-amino-acid AIDA-I protein contains cysteine residues, a feature that precludes the formation of disulfide bonds and is believed to be crucial for membrane translocation (26). MisL also contains a predicted N-terminal signal sequence required for the translocation of the protein across the inner membrane.

The MarT (for membrane-associated regulator) protein has homology with a protein from *E. coli* K-12 (ORF269) and exhibits similarity in its N-terminal domain to the ToxR protein from *Vibrio cholerae* (Fig. 2B). ToxR is a transmembrane regulatory protein that is required for the synthesis of cholera toxin in *V. cholerae* (36). It consists of an N-terminal cytoplasmic domain, which is homologous to the OmpR family of transcription factors and probably involved in DNA binding, and a C-terminal domain which is thought to be involved in sensing environmental signals (30, 37). Like ToxR, MarT con-

tains a potential transmembrane domain in its central region and exhibits similarity with the putative DNA binding domain of the CadC transcriptional activator of *E. coli* K-12 (49), another member of the OmpR family, suggesting that the *marT* gene encodes a regulatory protein.

Finally, the *rhuM*, *rmbA*, *fidL*, *slsA*, and *cigR* gene products do not exhibit sequence similarity to proteins with known functions in the sequence databases. FidL and the glycine- and asparagine-rich CigR contain putative signal sequences and might be exported proteins.

**Expression of SPI-3-encoded genes.** To examine the expression of the genes encoded within SPI-3, MudJ transposon insertions were isolated in this region of the *Salmonella* genome, and the  $\beta$ -galactosidase activities produced by the resulting strains were determined (see Materials and Methods) (Fig. 1). When the strains were grown in LB broth,  $\beta$ -galactosidase activity was produced by *lac* gene fusions to the *sugR*, *rhuM*, and *marT* genes but not by the *misL-lac* fusion (Table 3), suggesting that the *misL* gene may respond to signals not present in laboratory media. A *lac* fusion to the intergenic region between *sugR* and *rhuM*, located approximately 200 to 250 bp upstream of *rhuM*, produced a level of  $\beta$ -galactosidase activity comparable to that of the *rhuM-lac* fusion, suggesting that *sugR* and *rhuM* may constitute an operon even though the distance between *sugR* and *rhuM* is 580 bp.

Expression of horizontally acquired genes is often controlled by regulatory proteins encoded by linked genes within the acquired sequences. For example, several genes in the *Salmonella* SPI-1 island are regulated by the HilA and InvF proteins, which are also encoded within SPI-1 (1, 27). Likewise, a two-component system encoded in the *Salmonella* SPI-2 island governs expression of several SPI-2 genes (46–47). Despite its similarity to the regulatory protein ToxR, MarT does not appear to control expression of the *sugR*, *rhuM*, *misL*, and *mgtC*

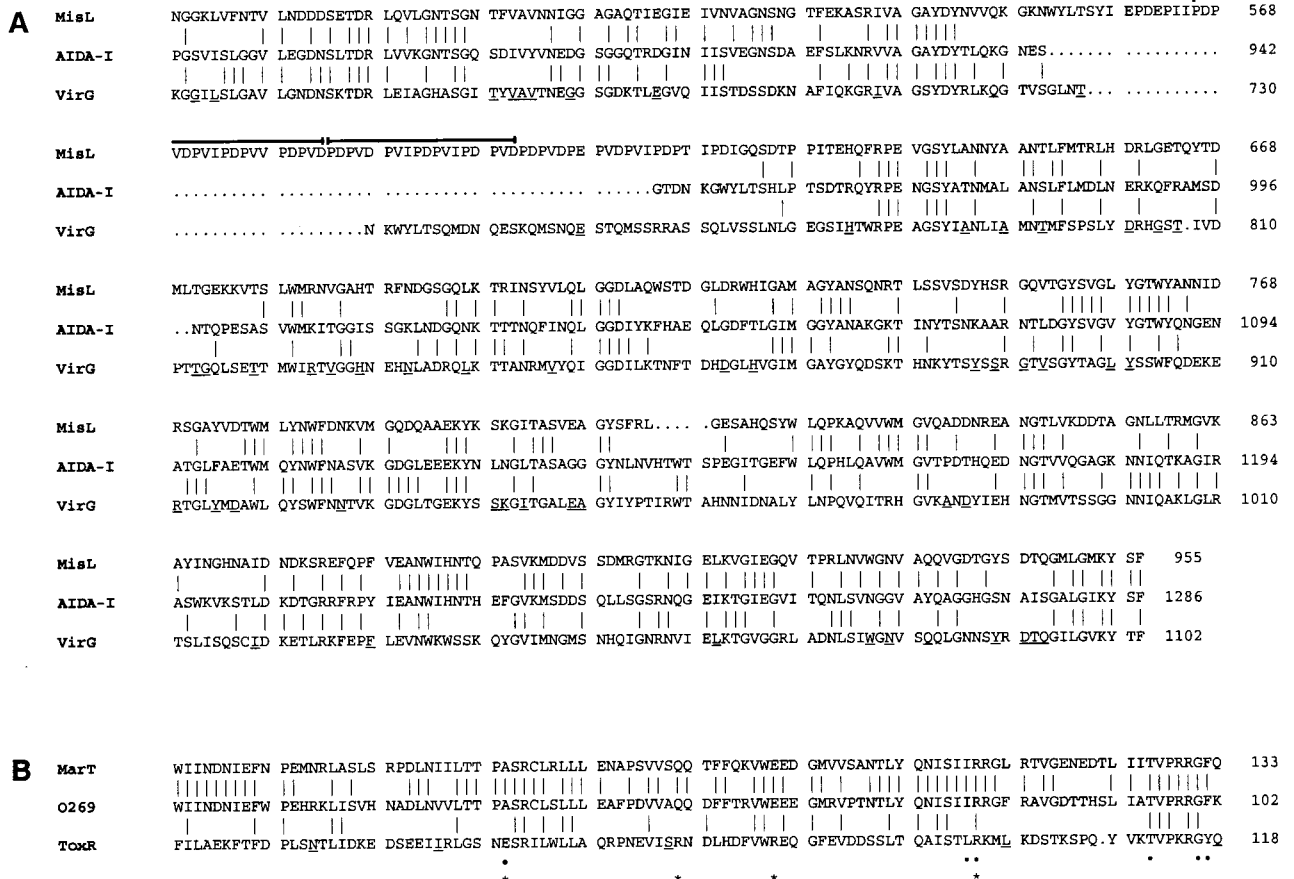


FIG. 2. (A) Alignment of the C-terminal domains of the *S. enterica* MisL protein, the plasmid-encoded AIDA-1 protein from enteropathogenic *E. coli*, and the VirG protein of *S. flexneri*. An 18-amino-acid duplicated region rich in Pro, Asp, and Val within MisL is indicated by a horizontal line. Amino acids that are identical between the MisL and AIDA-1 proteins or between the AIDA-1 and VirG proteins are linked by vertical lines. Amino acids that are identical between the MisL and VirG proteins are indicated by a short underline in the VirG residue. (B) Alignment of the N-terminal domains of the MarT protein, ORF269 (O269) of *E. coli*, and the ToxR regulator from *V. cholerae*. Highly conserved residues among OmpR homologs are indicated by dots (30). Amino acids that have been shown to be important for ToxR function (41) are marked by asterisks. Amino acids that are identical between the MarT and ORF269 proteins or between the ORF269 and ToxR proteins are linked by vertical lines. Amino acids that are identical between the MarT and ToxR proteins are indicated by a short underline in the ToxR residue. Alignments were performed by using the PILEUP program (GCG).

genes, because similar levels of  $\beta$ -galactosidase were displayed by isogenic *marT*<sup>+</sup> and *marT* mutant strains when bacteria were grown in LB broth (Table 3). MarT does not appear to regulate its own expression either, because the wild-type *marT* gene on a multicopy plasmid did not modify the  $\beta$ -galactosidase activity of a *marT2*::MudJ strain (data not shown). The MarT protein has different amino acids than the ToxR protein at three of four positions shown to be important for ToxR regulatory function (Fig. 2B) (41), which raises the possibility of MarT being involved in a function other than transcriptional regulation. However, it is also possible that the MarT protein governs transcription of other genes within SPI-3 or under different growth conditions.

To coordinate their expression with that of the rest of the genome, foreign sequences often recruit host regulators in addition to those encoded within the acquired sequences. One striking example is provided by the regulatory protein PhoP, which is present in both pathogenic and nonpathogenic bacterial species (19) and controls expression of several horizontally acquired sequences involved in *Salmonella* virulence, including the SPI-3-carried *mgtC* gene (18). However, expression of *sugR*, *rhuM*, *misL*, and *marT* is not dependent on the PhoP regulatory protein, since similar levels of  $\beta$ -galactosidase were

displayed by isogenic *phoP*<sup>+</sup> and *phoP* mutant strains (Table 3).

**Virulence properties of SPI-3 mutants.** Because the *misL* and *marT* genes encode proteins with similarity to known virulence factors, we investigated whether these genes were required for *Salmonella* virulence. Strains with a *misL1*::MudJ

TABLE 3. Transcriptional activities of SPI-3-encoded genes in wild-type, *marT*, and *phoP* strains

| MudJ insertion                   | $\beta$ -Galactosidase activity <sup>a</sup> |                            |                        |
|----------------------------------|--|----------------------------|------------------------|
|                                  | Wild type                                    | <i>marT1</i> :: <i>cat</i> | <i>phoP7953</i> ::Tn10 |
| <i>sugR1</i> ::MudJ              | 116  | 114                        | 118                    |
| <i>sugR2</i> ::MudJ <sup>b</sup> | 175  | ND <sup>c</sup>            | ND                     |
| <i>rhuM1</i> ::MudJ              | 200  | 205                        | 192                    |
| <i>misL1</i> ::MudJ              | No <sup>d</sup>                              | No                         | No                     |
| <i>marT2</i> ::MudJ              | 71   | ND                         | 74                     |
| <i>mgtCB9232</i> ::MudJ          | 18   | 18                         | No                     |

<sup>a</sup> In Miller units.  
<sup>b</sup> Insertion in the intergenic region between *sugR* and *rhuM*.  
<sup>c</sup> ND, not determined.  
<sup>d</sup> No, nondetectable activity.



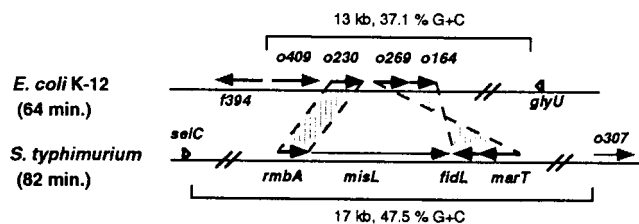


FIG. 3. Organization of the *rmbA*, *misL*, *fidL*, and *marT* genes in *S. enterica* serovar Typhimurium SPI-3 in comparison to the *orf230* (*o230*), *orf269* (*o269*), and *orf164* (*o164*) genes of *E. coli* K-12. The deduced amino acid sequences of *rmbA*, *fidL*, and *marT* are about 40% identical to the deduced proteins encoded by *o230*, *o164*, and *o269*, respectively (Table 2), suggesting that these proteins are orthologues rather than homologues.

mutation (EG10755) or with a *marT1::cat* mutation (EG10207) exhibited wild-type levels of survival within macrophages and invasion of epithelial cells (data not shown). Moreover, their ability to cause a lethal infection in mice was as efficient as that of the wild-type parent when tested orally on BALB/c mice at doses of  $2 \times 10^6$  and  $1.5 \times 10^7$  CFU (the 50% lethal dose of the wild-type strain is  $6 \times 10^5$  CFU) (22). These results indicate that the *misL* and *marT* genes are not essential for virulence under the conditions investigated. However, these genes could be involved in other aspects of pathogenesis, such as chronic infection and host specificity, or they could play a role in processes specific to *Salmonella* that are unrelated to virulence.

**Similarity of SPI-3 proteins to *E. coli* proteins encoded by horizontally acquired sequences.** Consistent with the notion the SPI-3 island was acquired by horizontal gene transfer, its overall G+C content is 47.5%, which is much lower than that of the *Salmonella* chromosome (52%) (39). Moreover, SPI-3 is located next to the *selC* tRNA gene, and tRNA genes are preferential sites of insertion of foreign sequences, including phages, plasmids, and pathogenicity islands (8, 20, 24). Furthermore, several SPI-3 gene products exhibit similarity with proteins encoded by horizontally acquired DNA sequences in other bacterial species. For example, the *sugR* gene product exhibits similarity with a protein from a clinical isolate of *E. coli* encoded by a gene that was probably acquired by lateral gene transfer, because it is part of a region with a low G+C content, located downstream of the *thrW* tRNA locus (29), and is absent from the *E. coli* K-12 genome (4).

The central region of SPI-3 includes four genes, i.e., *rmbA*, *misL*, *fidL*, and *marT*, three of which code for proteins with sequence similarity to *E. coli* K-12 ORF products encoded at 64 min in the chromosome (*rmbA*, *fidL*, and *marT* are similar to *orf230*, *orf164*, and *orf269*, respectively) (4). However, the genetic organizations of these genes are different in *E. coli* and *Salmonella* (Fig. 3). The *orf230*, *orf269*, and *orf164* genes appear to have been acquired horizontally into *E. coli* K-12, because they have an atypical codon usage and are part of a 13-kb region with a very low G+C content (37.1%) that is located downstream of the *glyU* tRNA gene (4). Consistent with the notion that this region is not ancestral to enteric bacteria but rather that it was incorporated into *E. coli* K-12 by horizontal gene transfer, Southern hybridization experiments revealed that *orf269*-hybridizing sequences are absent from 12 bacterial species, including *S. flexneri*, which is considered to be part of the *E. coli* species (see Materials and Methods) (data not shown).

**Evolution of SPI-3 sequences.** The G+C content is not uniform along SPI-3 (Fig. 1): genes with the lowest G+C content (*rmbA*, *fidL*, and *marT*, with 37.7, 45.2, and 47.3%, respective-

ly) are located in the central region of SPI-3. However, this region, which is homologous to an *E. coli* K-12 gene cluster (see above), is interrupted by the *misL* gene, which has a G+C content of 53%, suggesting that the incorporation of *misL* was a genetic event separate from that mediating the acquisition of *rmbA*, *fidL*, and *marT*. The central region of SPI-3 is surrounded by DNA segments with a very low G+C content that contain remnants of insertion sequences (Fig. 1): between the *rhuM* and *rmbA* genes, a 200-bp segment is homologous to the left inverted repeat and the 40 first residues of the transposase gene of the IS1351 element of *Salmonella enteritidis* (6a), and the region that separates the *marT* and *slsA* genes harbors a 100-bp sequence similar to a repetitive element from *E. cloacae* that is related to IS10 (31) and to the left inverted repeat of the IS911 element of *Shigella dysenteriae* (42). Taken together, these data suggest that SPI-3 has a composite structure and that the central region might have an independent origin.

To further examine the evolution of the SPI-3 island, we investigated the *Salmonella* Reference Collection C, which includes strains that encompass the eight subspecies of the genus *Salmonella* (6), for the presence of SPI-3 sequences. Southern hybridization experiments established that sequences hybridizing to the 5.3 kb at the right end of the island (including the *slsA* gene and the *mgtCB* operon) are present in all eight subspecies of *S. enterica* (Fig. 4). In contrast, the 5.5-kb central region did not hybridize to DNAs from strains of groups IIIa, IV, and VII and from one representative of group IIIb. Surprisingly, sequences hybridizing to this region were detected in *Salmonella bongori* (group V), the more divergent form of *Salmonella*. These results are consistent with the hypothesis that the central region of SPI-3 was incorporated as a separate genetic event, as suggested by the G+C composition and presence of surrounding insertion sequences (ISs). Thus, this portion of SPI-3 might have been acquired independently in *S. bongori*, or, alternatively, it may have been introduced into the *Salmonella* lineage or in the donor chromosome from which SPI-3 originated and then have been deleted in a subset of *Salmonella* subspecies. The left end of SPI-3 (including *sugR* and *rhuM*) appears to be less conserved than the central region and might have been the subject of deletions. As expected, a probe complementary to a region outside SPI-3, which included most of the *selC* gene and 350 bp of upstream sequences, hybridized to DNAs from all *Salmonella* subspecies and from *E. coli* K-12.

The *selC* tRNA locus is the site of insertion of SPI-3 in *Salmonella* and of the PAI-1 and LEE islands in pathogenic strains of *E. coli* (3, 5, 33). While this suggests a common mechanism for the acquisition of these foreign sequences, the SPI-3 island does not encode an integrase-like protein or harbor long repeated sequences. This is in contrast to PAI-1, which contains a cryptic integrase gene in its left end and is flanked by short direct repeats (5, 24), and to LEE, which harbors remnants of a transposase gene in its right end (13). The sequences of the *Salmonella* SPI-1, SPI-2, and SPI-4 islands and their boundaries have, thus far, not revealed long sequence repeats, phage attachment sites, or remnants of integrase genes, which could be responsible for the stability of these regions in the *Salmonella* genome.

The LEE pathogenicity island was originally identified at the *selC* locus of enteropathogenic strains of *E. coli* (33), but recent work indicates that LEE can be found at locations other than *selC* (50). We investigated whether SPI-3 is located at the *selC* locus in the different *S. enterica* subspecies by carrying out PCRs with primers complementary to the *selC* and *slsA* genes (these genes are 12 kb apart in *S. enterica* serovar Typhimurium, and *selC*- and *slsA*-hybridizing sequences have been

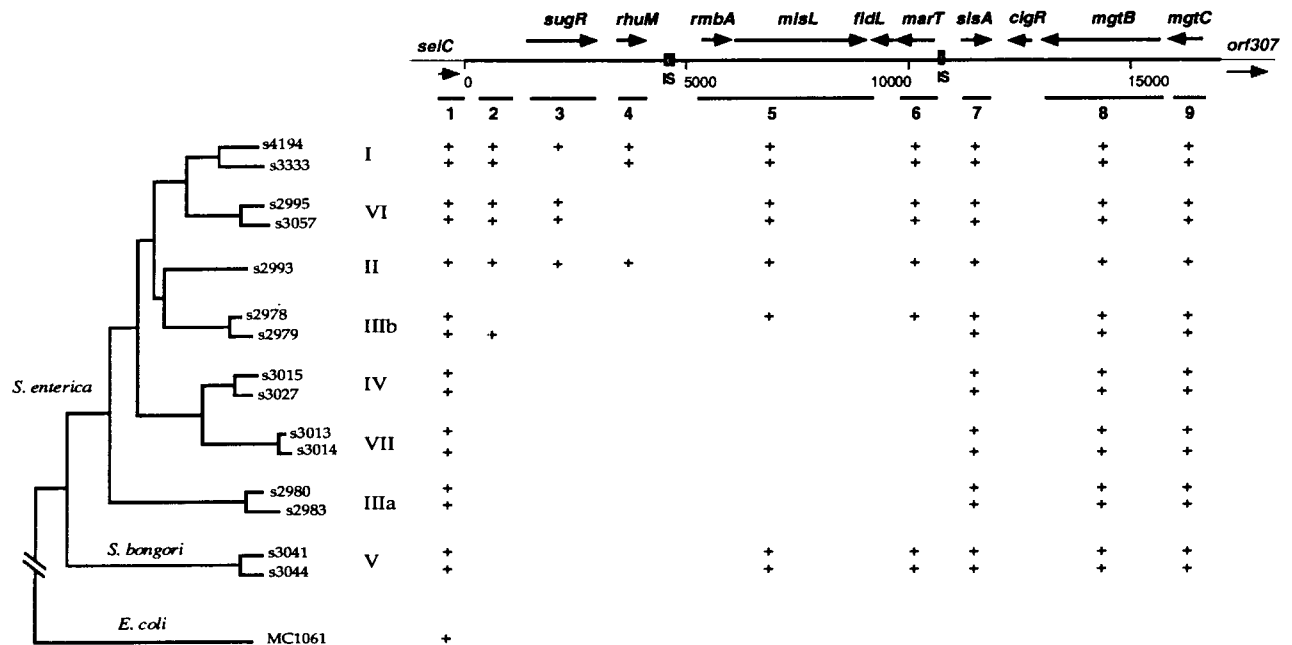


FIG. 4. Phylogenetic distribution of SPI-3 sequences among salmonellae based on Southern blot experiments carried out as described in Materials and Methods. Positions of the remnants of ISs are indicated. +, presence of a positive hybridization signal with the designated strain. Evolutionary relationships of *Salmonella* Reference Collection C strains are based on variation in the nucleotide sequences of five housekeeping genes (6). The roman numerals indicate the eight *Salmonella* subspecific groups.

detected in all *Salmonella* subspecies). The *slsA* gene is linked to the *selC* gene in most subspecies, because a 12-kb fragment was obtained when DNAs from *Salmonella* subspecies I, VI, and II were used as templates, and fragments smaller than 12 kb were amplified from at least one representative of each of the other subspecies (two different primers within *selC* were used to exclude nonspecific amplifications). At the right end of the island, the *mgtC* gene appears to be located next to *orf307* in all eight subspecies of *S. enterica*, because PCR experiments with primers complementary to *mgtC* and *orf307* resulted in the amplification of the same 0.76-kb fragment.

**Conclusion.** The molecular analysis of SPI-3 sequences and their phylogenetic distribution among the different subspecies that comprise *S. enterica* indicate that SPI-3 has a mosaic structure, most likely the result of a multistep evolutionary process, and encodes proteins that are not obviously functionally related. This is in contrast to the *Salmonella* SPI-1 and SPI-2 pathogenicity islands, which were likely acquired through single horizontal gene transfer events and encode functionally related proteins, which include type III export systems and secreted effector proteins (38, 40, 44). The different distribution of SPI-3 sequences may reflect the functional role of encoded genes: sequences present in only a subset of *Salmonella* subspecies could be involved in host specificity, tissue tropism, and disease manifestation, as recently proposed for a group of *Salmonella*-specific sequences of atypical base composition recovered by the in vivo expression technology procedure (11).

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