The genome sequence of *Salmonella enterica* serovar Choleraesuis, a highly invasive and resistant zoonotic pathogen

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ABSTRACT

*Salmonella enterica* serovar Choleraesuis (*S*.Choleraesuis), a highly invasive serovar among non-typhoidal *Salmonella*, usually causes sepsis or extra-intestinal focal infections in humans. *S*.Choleraesuis infections have now become particularly difficult to treat because of the emergence of resistance to multiple antimicrobial agents. The 4.7 Mb genome sequence of a multidrug-resistant *S*.Choleraesuis strain SC-B67 was determined. Genome wide comparison of three sequenced *Salmonella* genomes revealed that more deletion events occurred in *S*.Choleraesuis SC-B67 and *S*.Typhimurium LT2. *S*.Choleraesuis has 151 pseudogenes, which, among the three *Salmonella* genomes, include the highest percentage of pseudogenes arising from the genes involved in bacterial chemotaxis signal-transduction pathways. Mutations in these genes may increase smooth swimming of the bacteria, potentially allowing more effective interactions with and invasion of host cells to occur. A key regulatory gene of TetR/AcrR family, *acrR*, was inactivated through the introduction of an internal stop codon resulting in overexpression of AcrAB that appears to be associated with ciprofloxacin resistance. While lateral gene transfer providing basic functions to allow niche expansion in the host and environment is maintained during the evolution of different serovars of *Salmonella*, genes providing little overall selective benefit may be lost rapidly. Our findings suggest that the formation of pseudogenes may provide a simple evolutionary pathway that complements gene acquisition to enhance virulence and antimicrobial resistance in *S*.Choleraesuis.

INTRODUCTION

*Salmonella* are important pathogens in humans and animals. Taxonomically, there are two species in the genus *Salmonella*: *S*.enterica (six subspecies) and *S*.bongori (one subspecies). Members of the seven subspecies can be serotyped into one of more than 2500 different serovars according to somatic (O) and flagellar (H) antigens. Some serovars such as *S*.Typhi and *S*.Paratyphi are highly adapted to humans without other known natural hosts, while others such as *S*.Typhimurium have a broad host range and infect a wide variety of animal hosts. A few serovars such as *S*.Choleraesuis have a narrow host range and occasionally infect humans. Nevertheless, when *S*.Choleraesuis infects humans, it usually causes invasive infections (1,2). It is one of the most common etiologic organisms for infective aneurysms, a devastating endovascular infection in humans (1,3). *S*.Choleraesuis also can cause swine paratyphoid, with clinical manifestations of enterocolitis and septicaemia (4). Furthermore, it is a common bacterial isolate from swine with porcine reproductive and respiratory syndrome (4–6), which is now found worldwide and leads to considerable economic loss in the swine industry (5).

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Antimicrobial therapy is essential in the treatment of \textit{S. Choleraesuis} infections in humans. The emergence of resistance to ampicillin, chloramphenicol, trimethoprim-sulfamethoxazole, and notably, fluoroquinolone antibiotics in \textit{S. Choleraesuis} has aroused concern on the use of these agents for the treatment of systemic infections caused by this organism (7). A multidrug-resistant strain of \textit{S. Choleraesuis} was recently isolated from a patient with sepsis (8). This isolate, SC-B67, was resistant to all antimicrobial agents commonly used to treat patients with invasive salmonellosis, including ceftiraxone [minimal inhibitory concentration (MIC), 16 \( \mu \)g/ml] and ciprofloxacin (MIC, 32 \( \mu \)g/ml) (8). To investigate the virulence factors, host-specificity and mechanism of antimicrobial resistance of \textit{S. Choleraesuis}, we sequenced the whole genome of \textit{S. Choleraesuis} SC-B67. This work enabled us to perform comparative analyses with two other sequenced serovars, \textit{S. Typhi} and \textit{S. Typhimurium} (9–11). The three \textit{Salmonella} serovars compared here are associated with three distinct human disease syndromes: typhoid fever caused by \textit{S. Typhi}, self-limited enterocolitis by \textit{S. Typhimurium}, and septicemic diseases with little involvement of the gastrointestinal tract by \textit{S. Choleraesuis}. In this study, we discovered significant differences in genomic composition and organization of these serovars, and we identified several mechanisms underlying the unique pathogenicity and antimicrobial resistance mechanism of \textit{S. Choleraesuis}.

**MATERIALS AND METHODS**

**Genome sequencing and analysis**

\textit{S. Choleraesuis} strain SC-B67 was isolated from a patient with sepsis in 2002. Whole-genome libraries in pUC18 were prepared from genomic DNA as described earlier (12). Random clones were sequenced by using dye-terminator chemistry and data collected on MegaBASE 1000 automated sequencers. The total number of reads (>200 bp) was 66,494, with an average read length of 540 bp. The Phred/Phrap/Consed software package was used for quality assessment and sequence assembly (13–15). Gaps were filled by sequencing opposite ends of linking clones and the use of several PCR-based techniques and primer walking methods. The final assembly was verified by comparison with restriction-enzyme digest patterns of I-CeuI, AvrII and XbaI. Furthermore, we performed PCR on genomic DNA to amplify regions covering the pseudogenes. All the sequence alterations were confirmed by re-sequencing the PCR products.

The potential CDS were established with GLIMMER 2.1 (16) and the annotated open reading frames (ORFs) of three known \textit{Salmonella} genomes (9–11); both predicted CDS and putative intergenic sequences were subjected to further manual inspections. ORFs that contain insertion, deletion or mutation to a stop codon compared with those known \textit{Salmonella} genes were selected as pseudogenes. To the 4445 determined ORFs, exhaustive BLAST searches with an incremental stringency against NCBI non-redundant protein database were performed to determine their homology (17). To further investigate the function of each protein we used the InterProScan to search against the InterPro protein family database, which included PROSITE, PRINTS, Pfam, ProDom, SMART, TIGRFAMs, PIR SuperFamily and SUPERFAMILY (18). The up-to-date KEGG pathway database (ftp://ftp.genome.ad.jp/pub/kegg/genomes) was used for pathway analysis. Genes from 127 bacteria were gathered and classified into 127 pathways, to which we searched the pseudogenes in the BLAST suite, and the best hit was selected. The BLAST search criteria were as follows: (1) ~30% of the subject sequence was aligned and (2) E-value = 10^{-10} or less. Putative aliens were detected by finding anomalous gene regions in genomic characterizations (19), and the pathogenic islands were further determined according to their functional annotation and comparison with other \textit{Salmonella} genomes (9,10). Unique regions of a genome were identified as sequences that have no counterpart in another compared genome with MegaBLAST search. Genome used for comparison are from GenBank and their accession numbers are: NC_003143, NC_004061, NC_002528, NC_004741, NC_004631, NC_004545, NC_000913, NC_004431, NC_004088, NC_003198, NC_002695, NC_002655, NC_004344, NC_004337 and NC_003197.

**Electron microscopy**

Scanning electron microscopy was performed to examine the flagella of \textit{S. Choleraesuis} SC-B67 and \textit{S. Typhimurium} LT2. In preparation, bacteria were collected from the culture plate and suspended directly to 3% glutaraldehyde–2% paraformaldehyde mixture in 0.1 mol/l cacodylate buffer (pH = 7.4) for fixation at room temperature for 1 h. After adhering to poly-L-lysine coated glass coverslips, they were then post-fixed in 1% osmium tetroxide in the same buffer at 4°C for 1 h. After dehydration with graded ethanol, the samples were critical point dried, mounted onto the specimen stub, coated with Pt/Pd alloy, and examined with Hitachi S-5000 electron microscope at 3 kV.

**Inhibition assay and western blot analysis**

MICs were determined by the standard agar dilution method on the Mueller-Hinton medium. The MIC was defined as the lowest concentration of the drug that completely inhibited visible growth after inoculation for 18 h at 37°C. MICs of ciprofloxacin were also determined in the presence of the efflux pump inhibitor, Phe-Arg-β-naphthylamide (Sigma), at the concentration of 80 \( \mu \)g/ml (22). The efflux pump protein, AcrA, expression in \textit{Salmonella} strains was analyzed by western blot hybridization. Fluoroquinolone-susceptible \textit{S. Typhimurium} BN18 and \textit{in vitro}-selected resistant mutants BN18/21, BN18/41 and BN18/71 were used as controls in these experiments (22). For comparison, a fluoroquinolone-susceptible \textit{S. Choleraesuis} isolate SC-B42, was also examined by these methods.

**RESULTS AND DISCUSSION**

**General features of the genome**

The principal features of \textit{S. Choleraesuis} SC-B67 genome is composed of a 4,755,700 bp chromosome, a 138,742 bp large plasmid, designated pSC138, and a 49,558 bp virulence plasmid, designated pSCV50 (Table 1 and Figure 1A–C). The principal features of \textit{S. Choleraesuis} genome is classified into 127 pathways, to which we searched the pseudogenes in the BLAST suite, and the best hit was selected. The BLAST search criteria were as follows: (1) ~30% of the subject sequence was aligned and (2) E-value = 10^{-10} or less. Putative aliens were detected by finding anomalous gene regions in genomic characterizations (19), and the pathogenic islands were further determined according to their functional annotation and comparison with other \textit{Salmonella} genomes (9,10). Unique regions of a genome were identified as sequences that have no counterpart in another compared genome with MegaBLAST search. Genome used for comparison are from GenBank and their accession numbers are: NC_003143, NC_004061, NC_002528, NC_004741, NC_004631, NC_004545, NC_000913, NC_004431, NC_004088, NC_003198, NC_002695, NC_002655, NC_004344, NC_004337 and NC_003197.

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Table 1. Features of the Salmonella genomes

| Serovar       | S.Choleraesuis SC-B67 | S.Typhimurium LT2 | S.Typhi CT18 |
|--------------|-----------------------|-------------------|-------------|
| Chromosome (size, bp) | 4,755,700             | 4,857,432         | 4,809,037   |
| G+C content (%)      | 52.11                 | 53                | 52.09       |
| Coding sequences (excluding pseudogenes) | 4445                  | 4450              | 4359        |
| Pseudogenes          | 151                   | 39                | 204         |
| Ribosomal RNAs       | 6 × (16S-23S-5S), 1 × (16S-23S-5S-5S) | 6 × (16S-23S-5S), 1 × (16S-23S-5S-5S) | 6 × (16S-23S-5S-5S), 1 × (16S-23S-5S-5S) |
| tRNAs                | 85                    | 85                | 78          |
| Average gene length (bp) | 898                  | 946               | 958         |
| Plasmid              | pSC138                | pSCV50            | pSLT        |
| Size (bp)            | 138,742               | 49,558            | 93,939      |
| G+C content (%)      | 51.30                 | 52.10             | 52.09       |
| Coding sequences (excluding pseudogenes) | 178                  | 51                | 108         |
| Pseudogenes          | 2                     | 0                 | 6           |

aFrom ref. (9).
bFrom ref. (10).

Genome comparison

The overall comparison of S.Choleraesuis SC-B67 genome to two other sequenced Salmonella genomes is shown in Table 1 (9,10). The S.Choleraesuis SC-B67 chromosome is 101,732 bp shorter than S.Typhimurium LT2, and 53,337 bp shorter than S.Typhi CT18; however, S.Typhi CT18 genome contains a unique 134 kb DNA segment coding for Vi capsule (9,10). The overall similarity of SC-B67 genome to those of LT2 and CT18 (see Supplementary Figure S1) is demonstrated by CROSS_MATCH comparison of the genomes (P. Green, unpublished data) with NUCmer (24). They are generally collinear and share >98% homology at the DNA level; however, such co-linearity between SC-B67 and CT18 is in part broken by two major inverted segments. Rearrangements between the rRNA operons are common in S.Typhi (25), but are not found in S.Typhimurium LT2 and S.Choleraesuis SC-B67. A comparison of the insertions and deletions (indels) among the three sequenced serovars indicates that such events generally involve a small number of genes (Figure 2). The three Salmonella genomes are similar in size, indicating that the acquisition of genes has been counterbalanced by deletions; however, more deletion events occurred in S.Choleraesuis and S.Typhi (Figure 2). There are three major differences between S.Typhi and the two non-typhoidal Salmonella. S.Typhi contains Vi capsule, many more deletions, and a different profile of adherence- and motility-related genes, for which it lacks sti, stf, lpf, pef and pilA, but harbors genes like sta, stg, sef and pil. Vi is important in the pathogenesis of typhoid fever because of its role in antiphagocytosis and complement evasion. It is, however, difficult to address the relatedness between the different fimbrial patterns and pathogenicity, due to the redundancy of fimbrial operons in Salmonella.

Pseudogenes

A remarkable difference among the three serovars is the possession of 151 pseudogenes in S.Choleraesuis SC-B67 in contrast to the numbers that have been observed, 39 in S.Typhimurium LT2 and 204 in S.Typhi CT18 (9,10). It appears that S.Choleraesuis has joined those few host range serovars of Salmonella, such as S.Typhi and S.Paratyphi A, that have undergone genome degradation by point mutations (10,26). Most of the 151 pseudogenes in S.Choleraesuis are intact in S.Typhimurium and S.Typhi: only two shared by S.Typhimurium and 17 by S.Typhi. These pseudogenes have been inactivated through the introduction of a stop codon (68 out of 151) or a frameshift mutation (83 out of 151). The latter case involved 1- or 2-bp shift (insertion or deletion) in 76 pseudogenes, truncation in six, and transposon insertion in one. A significant number of the pseudogenes in S.Choleraesuis (64 out of 151) and S.Typhi (75 out of 204) were predicted to be metabolism-related, according to the clusters of orthologous groups of proteins (COGs) database (Table 2). The presence of such pseudogenes may be the reason that S.Choleraesuis evolved from the rest of the non-typhoidal Salmonella to become a narrow host range pathogen. Results of the cross-genome comparison of the pseudogenes of the three Salmonella genomes can be found in the website http://saldb.cgu.edu.tw.

Notably, in S.Choleraesuis we found five pseudogenes derived from genes of fimbriae synthesis or from flagellar or chaperone-usher operons. Two other pseudogenes were derived from genes that are likely to contribute to enteropathology of S.Typhimurium: icsA encoding a Peyer’s patch colonization and shedding factor and STM1669 encoding a protein homologous to invasin C of Yersinia. The mutation in these genes may have implications for bacterial adhesion to intestinal epithelial cells and may explain why S.Choleraesuis produces diarrhea more rarely than S.Typhimurium. Similar mutations in fimbriae clusters were also found in S.Typhi and S.Paratyphi A (10,26).

Furthermore, several pseudogenes (13 out of 151) arose from genes that are involved in the regulation of the virulence or metabolism-related genes. We searched the 151 pseudogenes in the KEGG database. Using the search criteria described earlier, we were able to identify appropriate pathways for 6 of the 13 regulatory genes (Table 3). S.Choleraesuis is a motile organism. We observed that it possesses a number of long, peritrichous flagella using electron microscopy (data not shown). Such flagella were also present in S.Typhimurium LT2. Motility is believed to increase the probability that bacteria reach suitable sites for invasion; chemotaxis, on the other hand, allows bacteria to sense the environmental cues, thereby modulating bacterial movement. S.Choleraesuis apparently has several pseudogenes derived from genes involved in the bacterial chemotaxis signal-transduction pathways (Table 4).
In an *in vitro* assay, *S*. *Choleraesuis* SC-B67 showed a relatively decreased chemotactic activity, compared with *S*. *Typhimurium* LT2. In the latter case, the 24-h swarm rate increased significantly in the presence of glucose 10 mM with or without the addition of PQQ (Figure 3). Previous studies have indicated that mutations, which disrupt the normal functioning of chemotaxis, enhance the invasive phenotype of *S*. *Typhimurium* (27,28). Such invasive phenotype was also observed *in vivo* in a murine infection model (28). Subsequent work revealed that defects in modulatory genes of the chemotaxis response (*cheA*, *cheR*, *cheW* and *cheY*) are responsible for the hyperinvasiveness of the mutants (28). These data are most consistent with the idea that motility affects the ability of *Salmonella* to enter host cells. As described previously, during smooth swimming the flagella operate as a single concerted bundle to propel the bacterium forward (28,29). The mutations in chemotaxis receptors may increase smooth swimming, thus allowing more effective bacterium–epithelial cell interactions and leading to cellular entry to occur at a higher frequency. As suggested for *S*. *Paratyphi* A (26), our data also suggest that loss of the genes involved in chemotaxis pathways may play a role in hyperinvasiveness of *S*. *Choleraesuis*.

**von Willebrand factor**

The genome of *S*. *Choleraesuis* encodes a protein bearing homology to von Willebrand factor type A domain of humans (SC2319, *vwa*). The von Willebrand factor is a multimeric glycoprotein that is synthesized in megakaryocytes and endothelial cells (30). During normal hemostasis, it adheres to the subendothelial matrix following vascular damage. This *vwa* gene was also found in *S*. *Typhimurium*, but not in *S*. *Typhi*. Because a gene (SC1603) encoding microbial collagenase has been identified, it is reasonable to propose that collagenase-mediated vascular injury during infection and subsequent combined effect of the Vwa could lead to...
the development of septic vascular thrombi. This may explain the observed severe vascular sequelae of non-typhoidal Salmonella infections, namely infective aneurysm.

Pathogenicity islands

Genomic islands are regions of DNA on the chromosome of bacteria acquired via horizontal gene transfer. They are named pathogenicity, metabolic or resistance islands depending on functions of the genes on the islands. Salmonella have acquired a large number of virulence genes required during various stages of pathogenesis. In S. Choleraesuis, we have identified the five Salmonella pathogenicity islands (SPIs) shared by S. Typhimurium and S. Typhi. The average G+C content of the five SPIs in S. Choleraesuis genome is 46.62% (range, 43.55–50.33%). Of the five, we found variations in SPI-1, -3 and -5 (9,10,31). SPI-3 of S. Typhimurium harbors 10 genes, including the mgtCB operon that is required for survival in macrophages and growth in low-Mg\(^{2+}\) environment. Two genes, sugR and rhuM, were not found in SPI-3 of S. Choleraesuis. In addition, the high level of variability of sequence adjacent to selC among different Salmonella serovars implied that SPI-3 may be still evolving through gene acquisitions (31). SPI-1 and SPI-5 encode the most important invasion and enteropathology phenotypes of Salmonella. A 321 bp orf (SC1043) was found between pipB and pipC of SPI-5 in S. Choleraesuis, instead of the 431 bp orf (STM1089) in S. Typhimurium. Two genes (STM1092 and STM1093) between sopB and pipD of SPI-5 were absent in S. Choleraesuis. Within SPI-1 of S. Choleraesuis, while avrA and three other genes of unknown function (STM2901-STM2903) were not identified, three other unique genes (SC2841-SC2843) were present between pphB and STM2908. Two genes within SPI-1 of S. Typhimurium encoding a putative permease and a LysR transcriptional regulator are pseudogenes in S. Choleraesuis. The variation in SPI-1 and -5 may be another reason to explain why S. Choleraesuis produces fewer diarrheas when infecting humans. In addition to the SPIs 1-5, we also found SPI-6 and -9 (Figure 1A) that were previously identified in S. Typhi in the genome of S. Choleraesuis (10). We further discovered two SPIs (designated SPI-11 and -12) that had not been described previously and two metabolic islands in S. Choleraesuis (Table 5). These DNA segments have abnormal G+C contents that deviate from the genome average and are in proximity to tRNA genes or mobile elements. All the major SPIs, including SPIs 1-5 and the newly described SPIs 11-12 are highly conserved among the three Salmonella genomes (9–11,31).

Plasmids and antimicrobial resistance

S. Choleraesuis SC-B67 has two plasmids, pSC138 and pSCV50 (Figure 1B and C). The sequence of pSCV50 is almost identical to that of pKDSC50 (32). Compared with the virulence plasmid (pSLT) of S. Typhimurium LT2 (9,33), pSCV50 contains two major deletions in pef-RepFI1A and traT-sama. The plasmid pSCV50 is the smallest, among the seven known Salmonella virulence plasmids (32,33).
The result of such reductive evolution is that pSCV50 becomes non-mobilizable but more stable (34). The plasmid pSC138 is composed of two distinguishable regions similar in size (~68 kb and 70 kb). The 68 kb segment is a mosaic of multiple antimicrobial resistance genes, insertion sequences and transposons, suggesting that the resistance genes were acquired through multiple lateral gene transfer events under selective pressure. The other segment contains replication and partial transfer regions of IncI1 plasmid ColIb-P9 (35,36). Many clinically relevant resistance genes were identified on this segment, including *dhfr* (trimethoprim), *sulI* and *sulII* (sulfonamide), *catI* and *cmlA* (chloramphenicol), *bla* (TEM-1; ampicillin), *strA* and *aadA2* (streptomycin), *tetRA* (tetracycline), *aph* (kanamycin), *sat* (streptothricin), *mef* (macrolide), *mer* (mercury), *ehr* (ethidium bromide) and *qac* (quaternary ammonium compound) (see Supplementary Figure S2). A β-lactamase inhibitor-resistant β-lactamase gene, *bla*<sub>TEM-67</sub>, was found on this plasmid (see Supplementary Figure S2). This gene, originally identified on the plasmid pANG-1 of *Proteus mirabilis*, codes for an enzyme similar to TEM-2 type β-lactamase that confers the organism resistance to ampicillin as well as β-lactamase inhibitors, such as clavulanic acid (37). On the other hand, the *ampC* (*bla<sub>CMY-2</sub>*) that mediates resistance to extended-spectrum cephalosporins was found located on a transposon-like element inserted in *finQ* of the ColIb-P9 plasmid (8). There was no *ampR*, previously described in resistant *Salmonella* that produced AmpC β-lactamase, in the upstream region of the element (8,38). The *ampR* is a regulatory gene of LysR family to *ampC* (38). The loss of *ampR*, which results in high-level AmpC expression, may be associated with cephalosporin resistance phenotype of the organism (8). The pSC138 is non-conjugative, but contains *oriT* sequence and a defective transfer system; therefore, it is probably mobilizable (39). All of the antimicrobial resistance genes may be disseminated to other organisms through the spread of this potentially transmissible plasmid.

On the other hand, we found that ciprofloxacin resistance of *S*.Choleraesuis SC-B67 is conferred through mutations of *gyrA* and *parC* in the chromosome. We further verified this finding by checking many more resistant strains of *S*.Choleraesuis using PCR and sequencing (7), and found only when both *parC* and *gyrA* mutations are present is ciprofloxacin resistance reliably detected. Furthermore, recent reports have indicated that the AcrAB-ToLC efflux system and its regulatory gene, *acrR*, may participate in the resistance to fluoroquinolones in *Salmonella* (22). It was noticed that when the *acrB* gene is inactivated, the resistance level to fluoroquinolones significantly reduces even when multiple target gene mutations

### Table 2. Functional classes of the corresponding genes of pseudogenes according to the clusters of orthologous groups of proteins (COGs) database in the three sequenced *Salmonella* genomes

| Functional category                              | S.Choleraesuis SC-B67 | S.Typhimurium LT2 | S.Typhi CT18 |
|-------------------------------------------------|-----------------------|-------------------|-------------|
| Information storage and processing               | 1                     | 1                 | 1           |
| Translation, ribosomal structure and biogenesis  | 10                    | 1                 | 13          |
| Transcription                                    | 7                     | 6                 | 29          |
| Replication, recombination and repair            | 11                    | 1                 | 13          |
| Cellular processes and signaling                 | 5                     | 0                 | 9           |
| Cell cycle control, cell division, chromosome partitioning | 10            | 0                 | 3           |
| Defense mechanisms                               | 1                     | 0                 | 4           |
| Signal transduction mechanisms                   | 1                     | 0                 | 8           |
| Cell wall/membrane/envelope biogenesis           | 11                    | 1                 | 13          |
| Cell motility                                    | 5                     | 0                 | 9           |
| Intracellular trafficking, secretion and vesicular transport | 0   | 0                 | 1           |
| Posttranslational modification, protein turnover, chaperones | 10        | 0                 | 3           |
| Metabolism                                       | 15                    | 1                 | 11          |
| Energy production and conversion                 | 18                    | 7                 | 27          |
| Carbohydrate transport and metabolism            | 13                    | 3                 | 13          |
| Amino acid transport and metabolism              | 2                     | 0                 | 6           |
| Nucleotide transport and metabolism              | 5                     | 0                 | 7           |
| Coenzyme transport and metabolism                | 3                     | 0                 | 4           |
| Lipid transport and metabolism                   | 6                     | 0                 | 6           |
| Inorganic ion transport and metabolism           | 2                     | 0                 | 1           |
| Secondary metabolites biosynthesis, transport and catabolism | 23         | 2                 | 16          |
| Poorly characterized                             | 9                     | 2                 | 22          |
| General function prediction only                  | 3                     | 1                 | 12          |

### Table 3. Pseudogenes with predicted regulatory functions and pathways involved

| Pseudogene | Classification                                      | Pathway involved       | Source                | EC number or protein description |
|------------|-----------------------------------------------------|------------------------|-----------------------|----------------------------------|
| Pseudogene 20 | Putative transcriptional regulator, LysR family | Pyrimidine metabolism | Borelia burgdorferi | 3.5.4.5                           |
| Pseudogene 42 | Sensory histidine kinase of a two-component regulatory system (TarP) | Two-component system | E.coli | 2.7.3.–                            |
| Pseudogene 61 | Putative transcriptional regulator, LysR family | Purine metabolism      | Borelia burgdorferi | 2.7.7.7                           |
| Pseudogene 66 | Membrane protein, regulator of *shlT* expression | Two-component system   | E.coli | Regulatory protein UhpC             |
| Pseudogene 68 | Periplasmic sensor of a multi-component regulatory system (TonS) | ABC transporters, prokaryotic | *E.coli* | D-ribose transport system           |
| Pseudogene 134 | Transcriptional regulator, LysR family | ABC transporters, prokaryotic | Borelia burgdorferi | Oligopeptide transport system       |

*Oligopeptide transport system substrate-binding protein OppA-1*
are present (22). In *S. Choleraesuis* SC-B67, a pseudogene was formed through the introduction of an internal stop codon in *acrR*, which acts as a transcriptional suppressor to down-regulate the expression of *acrAB*. An efflux pump inhibitor, Phe-Arg-naphthylamide (22), at the concentration of 80 μg/ml reduced the ciprofloxacin MIC from 32 to 8 μg/ml, suggesting the involvement of efflux pumping mechanism in fluoroquinolone resistance (Figure 4). Moreover, western blotting analysis with anti-AcrA antibody showed that SC-B67 exhibited a detectable increased level of AcrA expression, compared to a ciprofloxacin-susceptible *S. Choleraesuis* isolate that contains an intact *acrR* gene (Figure 4). The data

| Pseudogene | Corresponding gene in the chemotaxis pathways | Source | EC number or protein description |
|------------|-----------------------------------------------|--------|--------------------------------|
| Pseudogene 48 | *cheY* | *Clostridium acetobutylicum* | Chemotaxis response regulator |
| Pseudogene 68 | *rbsB* | *E.coli* | n-ribose transport system substrate-binding protein |
| Pseudogene 112 | *cheA* | *Borrelia burgdorferi* | 2.7.3.– |
| Pseudogene 122 | *tar* | *Thermoanaerobacter tengcongensis* | Methyl-accepting chemotaxis protein |
| Pseudogene 123 | *cheR* | *Borrelia burgdorferi* | 2.1.1.80 |
| Pseudogene 129 | *motB* | *Clostridium acetobutylicum* | Flagellar motor protein |

Table 4. Pseudogenes with their predicted functions involved in the bacterial chemotaxis pathways

Table 5. Newly described pathogenicity and metabolic islands of *Salmonella*

**Figure 3.** Effect of chemotactic attractants on the swarming of *S. Choleraesuis* SC-B67 and *S. Typhimurium* LT2 by using tryptone swarm tubes. (A) Swarm distance for *S. Choleraesuis* SC-B67 and *S. Typhimurium* LT2 over the time. Control indicates no addition of chemotactic attractants at the bottom of the tube. Each point represents mean ± SD. (B) The swarm rate in the presence of chemotactic attractants of each strain. *S. Typhimurium* LT2 swarm rate was significantly higher (*P < 0.05 by Student’s t-test) in the presence of glucose (with or without PQ). This situation was not observed in *S. Choleraesuis* SC-B67.
suggested that fluoroquinolone resistance in *S*. Choleraesuis should be due to a combined effect of multiple target gene mutations and overexpression of AcrAB.

In conclusion, genome comparisons of the closely related *Salmonella* emphasize the insights that can only be gleaned from sequencing multiple genomes of a single species. Lateral gene transfer providing basic functions to allow niche expansion in the host and environment is maintained in the evolution of different serovars of *Salmonella*. On the other hand, genes providing little overall selective benefit may get lost rapidly. Our study reveals that *S*. Choleraesuis has evolved through gene deletion and sequence alternations to become a very efficient and successful pathogen among non-typhoidal *Salmonella*, particularly with regard to its pathogenicity and antimicrobial resistance.

**SUPPLEMENTARY MATERIAL**

Supplementary Material is available at NAR online.

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