Genome sequence of *Shigella flexneri* 2a: insights into pathogenicity through comparison with genomes of *Escherichia coli* K12 and O157

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ABSTRACT

We have sequenced the genome of Shigella flexneri serotype 2a, the most prevalent species and serotype that causes bacillary dysentery or shigellosis in man. The whole genome is composed of a 4607203 bp chromosome and a 221618 bp virulence plasmid, designated pCP301. While the plasmid shows minor divergence from that sequenced in serotype 5a, striking characteristics of the chromosome have been revealed. The S.flexneri chromosome has, astonishingly, 314 IS elements, more than 7-fold over those possessed by its close relatives, the non-pathogenic K12 strain and enterohemorrhagic O157:H7 strain of Escherichia coli. There are 13 translocations and inversions compared with the E.coli sequences, all involve a segment larger than 5 kb, and most are associated with deletions or acquired DNA sequences, of which several are likely to be bacteriophage-transmitted pathogenicity islands. Furthermore, S.flexneri, resembling another human-restricted enteric pathogen, Salmonella typhi, also has hundreds of pseudogenes compared with the E.coli strains. All of these could be subjected to investigations towards novel preventative and treatment strategies against shigellosis.

INTRODUCTION

Shigella species are Gram-negative, non-sporulating, facultative anaerobes causing bacillary dysentery or shigellosis in man with estimated annual episodes of 160 million and 1.1 million deaths, most of which are children under 5 years old in developing countries (1). In China, more than 10 million cases are estimated per annum, of which 50-70% are caused by Shigella flexneri serotype 2a and most are associated with epidemic and pandemic shigellosis (2). Shigella are highly invasive in the colon and the rectum, and are able to proliferate in the host cell cytoplasm, triggering an inflammatory reaction. The clinical manifestations of Shigella infection vary from short-lasting watery diarrhea to acute inflammatory bowel disease characterized by fever, intestinal cramp and bloody diarrhea with mucopurulent feces (1). Since the current preventive and treatment strategies are found to be inadequate, the World Health Organization has placed an anti-Shigella vaccine as a priority (3).

Shigella was recognized as the etiologic agent for bacillary dysentery in the 1890s, and was adopted as a genus in the 1950s and subgrouped into four species: *S.dysenteriae*, *S.flexneri*, *S.boydii* and *S.sonnei* (4). However, a recent genetic study argues that *Shigella* emerged from multiple independent origins of *Escherichia coli* 35 000–270 000 years ago and may not constitute a genus (5). Genes on a virulence plasmid encode the primary virulence determinants, including the invasion plasmid antigens (Ipa) and their devoted Mxi-Spa type III secretion apparatus, but many chromosomal loci are

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also virulence required (6). Thus, the defining point for Shigella to evolve from E.coli must be the acquisition of the precursor of the current-day virulence plasmid carrying genes necessary for the bacteria to invade and access the host cell cytoplasm. This is a niche unique amongst the enteric pathogens with the exception of enteroinvasive E.coli that also possesses the virulence plasmid causing similar pathogenic characteristics (4). Subsequent evolution of the chromosome, however, enables the full expression of virulence. Hence, despite the fact that plasmid sequences from serotype 5a have become available (7,8), we felt it necessary to sequence the whole genome of S.flexneri 2a, the most prevalent species and serotype. Particularly, the expression of virulence depends on a complex regulation mechanism that involves dialog between the chromosome and the virulence plasmid (9), and a better understanding of this requires the availability of the whole genome sequence. Indeed, though the virulence plasmid from serotype 2a has minor divergence from that of servtype 5a, we have revealed the highly volatile and dynamic nature of the Shigella chromosome in comparison with the genomes of the non-pathogenic K12 strain and the enterohemorrhagic O157:H7 strain of E.coli (10,11). Furthermore, we have uncovered many chromosomal loci that potentially contribute to virulence in addition to those identified by the classic genetic studies (6).

MATERIALS AND METHODS

Shigella flexneri 2a strain and growth conditions

Shigella flexneri strain 301 (abbreviated Sf301), which we sequenced, was isolated from a patient with severe acute clinical manifestations of shigellosis in the Changping District, Beijing, in 1984, and has since been used as a reference strain for *S.flexneri* in China. The strain was routinely grown at 37°C overnight on tryptic soy agar containing 0.01% Congo red. Red colonies were inoculated into tryptic soy broth and grown to stationary phase at 37°C for isolating plasmid and chromosomal DNAs.

Shotgun sequencing and sequence assembly

The plasmid and the chromosomal libraries were separately constructed using pBluescript KS(-) (Strategene) as vectors. Approximately 48 000 clones were sequenced from both ends using the big-dye kit (ABI) and ABI377 or ABI3700 automated sequencers, giving rise to 10 times coverage of the genome.

Sequences were assembled initially using the phred/phrap program (12) when the sequence coverage was ~4-fold over the estimated size of the genome. The program was run with optimized parameters and the quality score was set to ≥ 20 . Further assembly was carried out repeatedly using the same program when more sequences were obtained. When 100 500 sequences were assembled into 318 contigs, the Consed program was used for sequence finishing (13). Gaps among contigs were closed either by primer walking on selected clones, which were identified by analysis on the forward and the reversed links between contigs using the perl/Tk algorithm, or by sequencing the DNA amplicons generated by polymerase chain reaction (PCR).

Prediction of open reading frames (ORFs) and identification of gene families

Glimmer 2.0, a program that searches for protein coding regions, was used to identify those ORFs possessing more than 30 consecutive codons (14). Overlapping and closely clustered ORFs were manually inspected. Predicted polypeptide sequences were used to search the non-redundant protein database with BLASTP, and the clusters of orthologous groups of proteins (COGs) database was used to identify families to which predicted proteins are related (15).

Mobile elements and repetitive sequences were identified using pair-wise comparison. tRNA sequences were identified by the program tRNAscan-SE (16). Repetitive regions were defined as those that have at least 200 bp with the significance of e^{-10} by BLASTN against the Sf301 genome itself and known IS databases. Sequence annotation and graphs of the circular and linear genomic maps were prepared using a newly developed Perl-Script tool kit (available at ftp:// ftp.chgb.org.cn/pub/).

Whole genomic comparison with *E.coli* K12 MG1655 (accession no. U00096) and O157 EDL933 (accession no. AE00517H) was performed using the GenomeComp program (J.Yang, J.Wang, Q.Jin, Y.Shen, Z.Yao and R.Chen, manuscript in preparation).

Accession of the genome sequence

The accession numbers for Sf301 chromosome and plasmid pCP301 are AE005674 and AF386526, respectively, in GenBank.

RESULTS AND DISCUSSION

General features of the genome

The primary features of the Sf301 genome are summarized in Table 1 and graphically viewed in Figures 1 and 2. The whole genome of Sf301 is composed of a 4 607 203 bp chromosome and a 221 618 bp virulence plasmid, designated pCP301. The chromosome shares a common 'backbone' sequence ~3.9 Mb with those of *E.coli* K12 (MG1655) (10) and O157 (EDL933) (11), which is essentially collinear. However, the backbone sequence is interrupted by numerous segments of K12-, O157and Shigella-specific DNA, designated 'K-islands' (KIs), 'O-islands' (OIs) and 'S-islands' (SIs), respectively (Fig. 1, circle 1). The co-linearity is also broken by numerous inversions and translocations compared with the E.coli sequences, 13 of which involve DNA segments >5 kb and are all bordered by IS elements and mostly associated with deletions or SIs (Fig. 2). All of these were confirmed by subsequent PCR sequencing of the junctions of each of the translocations and inversions. In the case of EDL933, there is only one inversion near the replication terminus with respect to K12 as noted previously (Fig. 2) (11). The dynamic gene shifts of the Sf301 chromosome are in contrast to the conserved genetic maps of E.coli K12, Salmonella typhimurium, other E.coli strains, other Salmonella spp., Klebsiella pneumoniae and many other enterics (17). However, there is no evidence for gene drift mediated by recombination between rRNA operons as observed in S.typhi (18) and in some Shigella strains (19). All the rRNA operons of Sf301 fall in approximately the same loci as those of E.coli

Table 1. General features of the Sf301 genome compared with genomes of *E.coli* K12 and 0157, and the virulence plasmid, pWR501, from *S.flexneri* M90T 5a

Chromosome	Sf301	MG1655 ^a	EDL933 ^b
Total length (bp)	4 607 203	4 639 221	5 528 445
No. of total ORFs	4434	4289	5349
Average length of ORFs (bp)	891	954	905
Percentage of coding sequence (%)	80.4	87.8	87.1
G + C content			
Total genome (%)	50.89	50.79	50.40
Protein coding regions (%)	51.95	51.85	51.51
RNA genes (%)	54.79	54.84	54.88
Intergenic regions (%)	46.07	42.28	42.76
Ribosomal RNA			
No. of 16S	7	7	7
No. of 23S	7	7	7
No. of 5S	8	8	8
No. of transfer RNA	97	92	93
No. of tmRNA	1	1	1
No. of non-classical RNA	9	5	5
Translocations and inversions ^c	13	_	1
IS elements	314	39	40
Of which partial copies	67	7	19
Plasmid	pCP301	pWR501 ^d	
Total length (bp)	221 618	221 851	
No. of total ORFs	267	293	
Average length of ORFs (bp)	658	636	
Percentage of coding sequence	76.24	82.09	
G + C content			
Total (%)	45.77	46.36	
Coding regions (%)	46.13	46.95	
Intergenic regions (%)	44.59	43.69	
IS elements	88	92	
Of which partial copies	62	69	

^aData are from Blattner *et al.* (10).

^bData are from Perna *et al.* (11).

^cOnly those with DNA segments >5 kb are listed.

^dData are from Venkatesan *et al.* (8).

(10,11). Natural selection that optimizes all promoters has operated to conserve genetic maps among enterics (17). A gradient of gene dosage generated from rapid chromosomal replication constrains genes to certain locations relative to the replication origin, and actively transcribed genes have a strong bias to be transcribed away from the origin, whereas weakly transcribed genes are evenly orientated (20). Genetic rearrangements alter the locations, orientations, and the coding strand (in the cases of inversions) with respect to the origin of replication, possibly changing the amount of transcription of many genes. This in turn may affect their dosage, and in some cases impair growth (21,22). Hence, the changed genetic map suggests that *S.flexneri* may have reoptimized its promoters to cope with selection pressures in the unique intracellular or *ex vivo* environments.

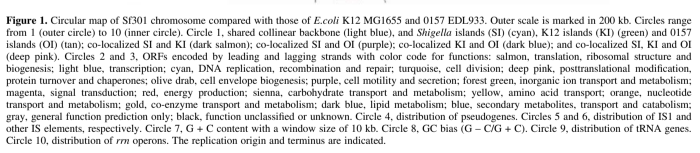
The Shigella islands

Sf301 has in total 64 SIs with sizes >1 kb, all of which are numbered and detailed in the 'linear map 1' (Supplementary Material). Among them, several, including the previously identified pathogenicity islands (PAIs) SHE-1 (23) and SHE-2 (24), have implications in virulence. Strikingly, there are

seven *ipaH* genes, five of which are located in five large SIs, designated as *ipaH* islands 1-5 (Fig. 2). All five *ipaH* genes in the islands are next to the genes that potentially encode proteins sharing 73-76% identity with a 188 amino acid hypothetical protein of unknown function from Salmonella bacteriophage P27 (accession no. NP_543109). The majority of the remaining genes in the *ipaH* islands share homologies with genes of different phages including those identified in the genome of O157 EDL933. But the overall gene contents and organizations in all 5 *ipaH* islands have little similarity. This suggests that the chromosomal *ipaH* genes were originally linked with phage P27 and subsequently transmitted to S.flexneri by different phages. The plasmid, pCP301, carries five *ipaH* genes, termed *ipaH*_{9.8}, *ipaH*_{7.8}, *ipaH*_{4.5}, *ipaH*_{2.5} and $ipaH_{14}$, at approximately the same loci as those in pWR100 and pWR501 from serotype 5a (7,8). None of these is next to the genes of the phage P27 paralogs, suggesting that they came from different sources or, alternatively, were transmitted to the plasmid via different vehicles. The pWR501-borne $ipaH_{7.8}$ is involved in the escape of Shigella from phagocytic vacuoles in the macrophages (25), but other *ipaH* genes have not been assigned a function. However, there is evidence that S.flexneri expresses more IpaH_{9.8} within host cells, and the proteins penetrate the host cell nuclei (26). This, and the fact that all IpaH proteins have a leucine-rich repeat region found in a diverse group of proteins from bacteria and eukaryotes (27), implies that IpaH might be involved in manipulating host gene expression. Alignment of all IpaH proteins indicates that they have identical C-terminal, but variable N-terminal, halves (Fig. 3). This suggests that they may interact with different host substrates, but exert similar functions.

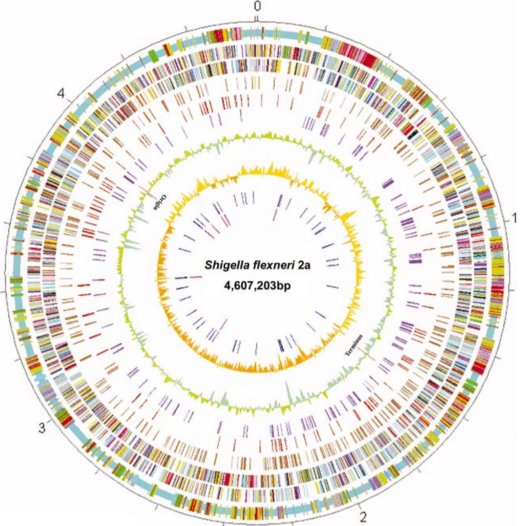
In *ipaH* island 2, four consecutive genes, similar to the Salmonella sitABCD and the Yersinea yfe operons (28), may encode proteins required for iron uptake. Since the Salmonella sitABCD can complement the growth in iron-restricted medium of an enterobactin-deficient E.coli, a role of the Salmonella sit-like (SSL) system in iron uptake is implicated. Iron uptake mechanisms have undergone complicated adjustments in S.flexneri. On the one hand, the enterobactin system is impaired due to the presence of stop codons in *fepE*, *fhuE* and *entC* genes (see Table 3), and on the other hand, the SSL system has been introduced, and additionally, SHE-2 encodes an aerobactin system (24). In some strains the E.coli fec enterobactin system is re-introduced along with so-called Shigella resistance locus PAI within a multiple resistance deletable element (MRDE) (29). However, MRDE is not present in Sf301.

Two other SIs are worthy of mention, the *sci* and the SfII islands (Fig. 2). The *sci* island is 22 789 bp in length and possesses a typical structure of PAI—inserted at an *asp*-tRNA and ends with an IS629 on the other side. It carries paralogs of the *Salmonella sciCDEFF* operon (accession no. AJ320483) of unknown function and of phage P22 and HK620, suggesting that it is possibly another phage-transmitted PAI. SfII has been demonstrated to be a lysogenic phage in which two genes, *bgt* encoding a bactoprenol glucosyl transferase and *gtrII* encoding a glucosyl transferase, are required for expression of the type II antigen (30). Thus, phage-mediated horizontal DNA transfer appears to be one of the major routes by which *S.flexneri* gains virulence determinants.



The IS elements

The IS elements identified in the Sf301 genome are listed in Table 2. In the chromosome, there are astonishingly 247 complete and 67 partial IS elements, which makes it the most IS-rich chromosome among enterics. The predominant species is IS1, followed by IS600, IS2 and IS4. They all are frequently associated with SIs, inversions and translocations, deletions and insertional gene inactivation (see 'linear map 1' in Supplementary Material). The IS elements are, therefore, probably the major cause of the dynamics of the Sf301 chromosome. Indeed, the presence of IS91 at two ends of MRDE (mentioned above) allows the precise acquisition and excision of the entire 99 kb segment (29). Furthermore, IS1 and other IS elements have also been shown to be able to mediate various genetic rearrangements (31,32), and IS1 in particular can cause inversions and deletions (32). It is plausible that the IS elements will mediate further evolution of the chromosome. Similarly, pCP301 has large numbers of IS elements, sharing similar composition with pWR501 from serotype 5a (Table 2), indicating that the virulence plasmids are also volatile and dynamic. One difference between pCP301 and pWR501 is that the former has two copies of iso-IS10R that may be transposed from the Sf301



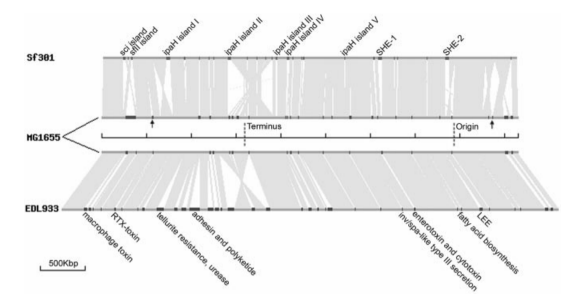


Figure 2. Schematic representation of translocations and inversions, and strain-specific islands (to scale). Chromosomes are represented by dark gray lines as indicated. Replication origin and terminus of MG1655 are indicated. The respective loci of Sf301 and EDL933 are at approximately the same positions. Regions in light gray indicate homologous sequences between paired chromosomes and triangular non-filling regions indicate the presence of inversions. Translocations are hardly visible because of the scale used. Regions in red, black and blue on the chromosomes represent SI, KI and OI, respectively. The known PAI among SIs and OIs are indicated. SIs with implications for a role in virulence are also indicated. Arrows indicate the KIs harboring *ompT* (left) and *cadA* (right) whose deletions are crucial for *Shigella* virulence.

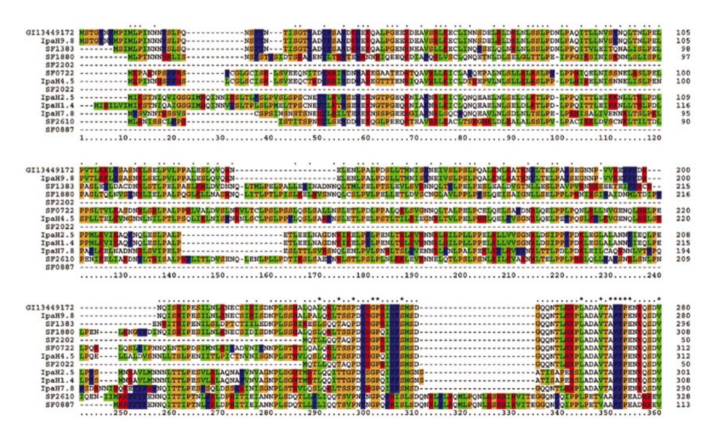


Figure 3. CLUSTALW amino acid sequence alignment of N-terminal halves of IpaH proteins identified in Sf301. IpaH_{9.8} of pWR501 (gi_13449172) serves as a reference on the top. The most homologous IpaH_{9.8} from pCP301 is placed in the second, and other IpaH family members are arranged in line with their homology to IpaH_{9.8}. The consensus line displayed above the aligned sequences depicts identical amino acids as asterisks, with conserved residues shown as dots.

Name Length No. of (bp) ORFs	No. of intact elements				No. of p	No. of partial elements						
	Sf301	K12	0157	pCP301	pW501	Sf301	K12	0157	pCP301	pWR501		
IS1	768	2	108	6	2	2	3	9	0	0	1	1
iso-IS1	803	2	0	0	0	0	0	1	0	3	5	5
IS2	1331	2	30	6	1	1	2	5	1	0	2	2
IS3	1258	2	5	5	0	0	0	3	0	2	7	8
IS4	1428	2	18	1	0	1	1	3	0	0	1	2
IS5	1198	1	0	10	0	0	0	0	1	0	0	0
iso-IS10R ^a	1329	1	13	0	0	2	0	0	0	0	0	0
IS21	2131	2	0	0	0	0	0	0	0	0	3	3
IS91	1830	1	3	0	0	0	0	2	0	0	6	6
IS100	1963	2	0	0	0	0	0	0	0	0	7	6
IS150	1443	3	0	1	0	0	0	5	0	0	2	2
IS186	1372	1	0	3	0	0	0	0	0	0	0	0
IS600	1264	2	35	0	0	3	2	17	1	6	10	13
IS629	1310	2	10	0	18	8	5	11	0	3	3	9
IS630	1164	1	0	0	0	1	1	0	0	4	2	2
IS911	1250	2	16	0	0	1	1	0	4	0	0	0
IS1294	1714	1	0	0	0	1	2	3	0	0	7	4
ISSfl1	929	1	0	0	0	1	1	0	0	0	2	3
ISSfl2	1374	1	6	0	0	2	2	0	0	0	1	0
ISSfl3	1302	1	0	0	0	1	1	1	0	0	1	1
ISSfl4	2754	3	3	Ó	Õ	2	2	7	0	1	2	2
Total		-	247	32	21	26	23	67	7	19	62	69

Table 2. IS elements identified in genomes of Sf301, MG1655 and EDL933, the virulence plasmid, and pWR501, from S.flexneri 5a

aiso-IS10R is a homolog of IS10R identified in Sf301 in this study.

chromosome (Table 2). But, it remains to be seen whether this IS element is present in the genome of serotype 5a. If not, it might be used as a marker for epidemiological studies.

The Escherichia coli islands (KIs and OIs)

With the respect to the Sf301 chromosome, MG1655 and EDL933 possess two kinds of islands. One kind is formed owing to the deletions of the corresponding *E.coli* DNA segments from the Sf301 chromosome, which is hardly surprising given the dynamics of the genome. These include the so called 'Black Hole' harboring *cadA* responsible for converting lysine to cardverine that adversely affects virulence (33) and the *kcp* locus harboring *ompT* that inhibits the induction of guinea pig keratoconjunctivitis (34) (arrows in Fig. 2). It remains to be investigated how many such 'Black Holes' have deletions of genes that would otherwise inhibit full expression of virulence.

The other kind of island is apparently formed by laterally acquired DNA sequences, of which the large ones are evident in Figure 2 with the scales used. A FASTA query of these groups of OIs and KIs against the Sf301 genome reveals no significant homologous sequence, and a query of all the SIs against EDL933 and MG1655 genomes reveals no homologous sequence either. Thus, O157 and *S.flexneri* appear to have acquired their island DNA from different sources and have evolved from ancestral *E.coli* strains through unrelated paths. Furthermore, all the SIs, OIs and KIs have no duplicated copies, indicating that none of them is mobile.

We must point out that we do not define sequences shared by paired strains (EDL933 or MG1655 with Sf301) as islands, though these may appear to be 'islands' with respect to the third genome. These sequences may reflect genetic properties of the ancestral *E.coli* strain that Sf301 evolved from. An example of these are the *rfa/waa* genes involving LPS biogenesis (Fig. 4). Sf301 and EDL933 have identical numbers of genes that share 99% identity in each case, whereas MG1655 has an equivalent functional operon with more genes and poor homology with the former (Fig. 4). Studies into this type of shared sequence may shed more light on strain diversity and evolution.

Pseudogenes

Apart from deletions of corresponding *E.coli* DNA segments, the formation of pseudogenes through introduction of stop codons, frame shifts, truncations and insertions in the coding regions appears to also play a major part in losing unwanted genes in S.flexneri. Pseudogenes with known functions according to the *E.coli* protein database are listed in Table 3. Answers to many of the phenotypic characteristics of Shigella, such as the loss of motility and utilization of lactose, maltose and xylose, etc., can be found here. It is noted that 90% of these pseudogenes are intact in O157 EDL933. To this end, S.flexneri resembles S.typhi, another enteric pathogen restricted to humans. The presence of large numbers of pseudogenes has been postulated to be one of the main reasons that S.typhi evolved from the rest of the Salmonella species to become a solely human pathogen (35). Likewise, the originally closely linked O157 and Shigella have evolved in diverse directions. Strain O157 became a successful pathogen with broad host range mainly by acquiring DNA (Table 1 and Fig. 2), whereas Shigella also became a successful pathogen but restricted to humans only, by acquiring, as well as losing, DNA.

The virulence plasmid pCP301

Like previously sequenced virulence plasmids (pWR100 and pWR501) from serotype 5a strains (7,8), pCP301 is a mosaic of potential virulence-related genes, IS elements, maintenance

Table 3.	Pseudogenes	with	known	functions	identified	in	Sf301	genome
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Pathway	Mutation	Description
Carbohydrate metabolism		
araA	Stop codon	L-Arabinose isomerase; arabinose catabolism
ugd	Stop codon	UDP-glucose 6-dehydrogenase; colanic acid synthesis
fucK	Stop codon	L-Funulokinase, fucose catabolism
glcD	Stop codon	Glycolate oxidase subunit D
xylA and B	Stop codon	D-Xylose isomerase; D-xylose catabolism and D-glucose conversion
aceB daoA	Stop codon Stop codon	Malate synthetase A; glyoxylate bypass D-Galactonate hydro-lyase; galactonate catabolism
dgoA fdhF ^a	Stop codon	Formate dehydrogenase-H; anaerobic respiration
zwf	Stop codon	G6PD; oxidative branch of pentose phosphate pathway
Energy metabolism	Stop codon	Sol D, oxiduate of allen of periode phosphate participation
cyoB	Stop codon	Cytochrome o ubiquinol oxidase subunit I; active under high oxygen growth conditions
cyoA	Truncation	Cytochrome o ubiquinol oxidase subunit II; as <i>cuoB</i>
acs	Stop codon	Acetyl-CoA synthetase; scavenging acetate
hyfB	Stop codon	Hydrogenase 4 subunit; anaerobic respiration
narZ	Stop codon	NRZ; anaerobic terminal electron acceptor
torA	Stop codon	Trimethylamine N-oxide reductase subunit; electron acceptor (anaerobic respiration)
torD	Insertion	Chaperone of TorA; preventing TorA degradation
Lipid metabolism		
hcaD	Stop codon	Ferredoxin reductase; utilization of aromatic acids
Amino acid metabolism	C4- 1	Omitting december in an in the interview of the interview
speF	Stop codon	Ornithine decarboxylase isozyme; putrescine synthesis
speG madP	Frame shift	Spermidine acetyltransferase; polyamine synthesis
nadB	Stop codon	Quinolinate thynthetase B; pyridine synthesis Succinate-semialdehyde dehydrogenase; aminobutyrate catabolism
gabD	Stop codon Frame shift	Peptidoglycan enzyme; cell wall formation
mtgA metA	Truncation	Homoserine transsuccinylase; methionine synthesis
cstC	Stop codon	Acetylornithine transaminase; arginine catabolism
Cofactors and vitamins	Stop codoli	Activitinine transammase, arginine catabolism
nfnB	Insertion	Dihydropteridine reductase; recycling the quinoid dihydrobiopterin cofactor by reducing it
lhr	Stop codon	ATP-dependent helicase, dispensable
lplA	Frame shift	Lipoate-protein ligase A; ligation of lipoyl to apoprotein
Complex lipids		
gldA	Stop codon	Glycerol dehydrogenase; glycerol dissimilation
Complex carbohydrates	1	
ycjM	Insertion	Putative polysaccharide hydrolase
otsA	Truncation	Trehalose-6-phosphate synthase; response to high osmolarity
aceK	Stop codon	Isocitrate dehydrogenase kinase/phosphatase; control flux between the TCA cycle and the glyoxylate bypass
Translation		
_ prfB	Stop codon	Peptide chain release factor RF-2
Transport	a 1	
araF	Stop codon	L-Arabinose-binding periplasmic protein
cysW	Stop codon	Sulfate transport system permease W protein
yhdX	Truncation	Permease; putative amino acid ABC transporter
ugpC	Insertion	ATP-transporter; glycerol-3-phosphate uptake
rbsA rbsB	Insertion Stop codon	ATP-biding component; D-ribose transport ABC transporter; D-ribose periplasmic binding protein
rusb glvG	Frame shift	6-Phospho-β-glucosidase; arbutin fermentation
ptsA	Stop codon	PEP-protein phosphotransferase system enzyme I
yphF	Stop codon	ABC transporter; periplasmic binding
Signal transduction	Stop todon	
citB	Truncation	Regulator (paired with <i>citR</i>); citrate fermentation
kdpE	Stop codon	Regulator of the kdp operon; potassium transport
kdpD	Stop codon	Sensor of the <i>kdpDE</i> system; potassium transport
narQ	Stop codon	Nitrate/nitrite sensor protein; acts on NarL/NarP
arp	Stop codon	Regulator of acetyl CoA synthetase
malT	Stop codon	Positive regulator of mal operon
Cell motility		
fliA	Frame shift	σ^{28} for flagellar operons
flgF	Stop codon	Cell-proximal portion of basal-body rod
flgK	Stop codon	Hook-filament junction protein 1
flgL	Stop codon	Hook-filament junction protein
fliF	Stop codon	Basal-body MS-ring and collar protein
fliJ	Truncation	FliJ protein
flhA Unaccional angumas	Stop codon	Export of flagellar proteins
Unassigned enzymes	Stop	And Co A thissetsman I by dealy and shain this
tesA	Stop codon	Acyl-CoA thioesterase I; hydrolyzes long chain acyl thioesters
pphA	Stop codon	Protein phosphatase 1; modulates phosphoproteins signaling protein misfolding

Table	3.	Continue	a
Table	3.	Continue	

Pathway	Mutation	Description
pphB	Stop codon	Removal of a phosphate group attached to serine or threonine residue; signaling protein misfolding through cpxRA system
Unassigned non-enzyme	s	
yaaJ	Stop codon	Transport protein; sodium/alanine symporter
nfrA	Stop codon	Omp; bacteriophage N4 receptor
csgG	Stop codon	Transporter; curli assembly
csgA	Insertion	Curlin major subunit; coiled surface structures
fepE	Stop codon	Transporter; ferric enterobactin (enterochelin)
fhuE	Stop codon	Omp; receptor for ferric iron uptake
entC	Stop codon	Isochorismate synthase; enterobactin biosynthesis
hlyE	Stop codon	Hemolysin E; hemolytic to sheep blood
hslJ	Truncation	Heat shock protein HslJ
uidB	Truncation	Transporter; specific to α - and β -glucuronides
celD	Insertion	Negative regulator of <i>cel</i> operon (cryptic); ferment cellobiose, arbutin and salicin
molR	Insertion	Molybdate metabolism regulator, first fragment
molR_2	Stop codon	Molybdate metabolism regulator, fragment 2
cirA	Stop codon	Porin and receptor; colicin I uptake
focB	Frame shift	Formate transporter (formate channel 2)
emrA	Stop codon	Multidrug resistance secretion protein
ppdA	Frame shift	Prepilin peptidase dependent protein A
glcF	Frame shift	Glycolate oxidase iron-sulfur subunit; ferridoxin related
aer	Stop codon	Aerotaxis sensor receptor; transducing signals for aerotaxis
ompG	Truncation	Outer membrane protein; forms large channels
yaeG	Stop codon	Regulator of D-galactarate, D-glucarate and D-glycerate metabolism
nagD	Stop codon	N-Acetyleglucosamine metabolism
fimD	Insertion	Export and assembly of type 1 fimbriae

^afdhF has a stop codon (UAA) in addition to the stop codon UGA used for introducing selenocysteine.

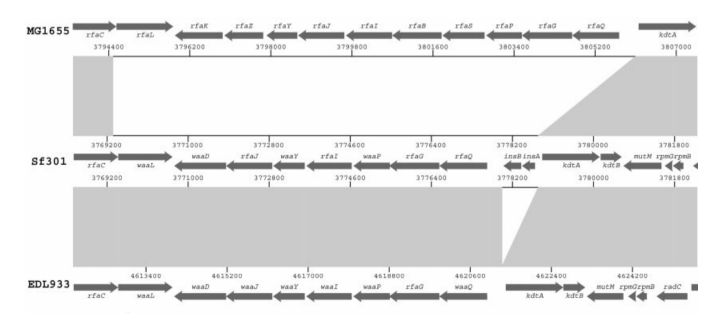


Figure 4. Comparison of the *rfa/waa* region (to scale). Arrows indicate predicted ORFs in both strands. Regions in gray indicate identical sequences among strains and the non-filling areas indicate sequences with non or low homology.

genes and functionally unknown ORFs. All the previously identified virulence genes are present in pCP301. These include the primary invasion genes *ipa* and *mxi-spa* (encoding the invasion plasmid antigens and the type III secretion system, respectively), *virG/IcsA* (required for polymerizing host actin to provide propelling force for intra- and intercellular spread) and *virF* (necessary for regulating virulence

gene expression). The replication origin (R100-like) *ori* and G site (single-strand initiation site) in pCP301 are identical to those of pWR501 and pWR100. pCP301 also has maintenance genes, *repA*, *copA* and *copB*, for replication; *parA* and *parB* for partitioning; and *ccdA* and *ccdB* for post-segregation killing. The noticeable difference between pCP301 and the plasmids from serotype 5a is the presence of more

IS-related DNA in pCP301, making its size close to pWR501 (221 851 bp) which is larger than pWR100 because of a *Tn*501 (8360 bp) insertion (8). So, both *Shigella* serotypes most likely acquired the ancestral virulence plasmid from the same source. One other minor divergence is that the *ipa-mxi-spa* loci in pWR501 and pCP301 are in the same orientation, whereas in pMYSH6000, the virulence plasmid from another 2a strain, they are in inverse orders (36). This indicates that the divergence of the plasmids does not necessarily correlate with serotypes. A detailed comparison of pCP301 with pWR501 is available in the Supplementary Material ('linear map 2').

CONCLUSION

Comparison of the *S.flexneri* genome with that of *E.coli* supports the previous genetic study (5) that *S.flexneri* is closely related to *E.coli* and may turn out to belong to the same genus. The global gene content (Table 1) and alignments (Fig. 2) indicate that *S.flexneri* is more closely related to the non-pathogenic K12 strain MG1655 rather than the pathogenic O157 strain EDL933. This is in agreement with the suggestions that O157 and K12 last shared an ancestor ~4.5 million years ago (37), whereas *Shigella* evolved from multiple *E.coli* strains much later, correlating with the appearance of early man in the paleolithic (5). All these studies call strongly to reclassify *Shigella* species as members of *E.coli*.

To meet the demand of its unique pathogenic lifestyle, the *S.flexneri* chromosome has evolved distinctive characteristics after acquisition of the large virulence plasmid. Most importantly, there are several potential bacteriophage-transmitted PAIs, many translocations, inversions and deletions of the corresponding *E.coli* DNA segments, and numerous pseudogenes. These findings provide an invaluable genetic basis for future studies into understanding bacterial evolution, as well as pathogenicity, and the development of novel preventive and treatment strategies against shigellosis.

SUPPLEMENTARY MATERIAL

Supplementary Material is available at NAR Online.

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