CHARACTERISATION AND TYPING OF BACTERIA

A framework for IS200, 16S rRNA gene and plasmid-profile analysis in Salmonella serogroup D1

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Summary. Chromosomal fingerprinting of the type strains of serotypes of Salmonella O-serogroup D1 with the DNA insertion sequence IS200 generated patterns which were either serotype-specific (e.g., Typhi), or conserved among groups of related serotypes (e.g., Dublin, Rostock and certain phage types of Enteritidis). The number of IS200 copies varied considerably, and the IS200 patterns of type strains of serotypes associated with systemic infections in man were specific and suitable for identifying strains within those serotypes. Polymorphism at 16S rRNA gene loci was examined among type strains and 11 16S rRNA gene profiles were characterised. The most prevalent of these was conserved among type strains of 11 serotypes, and the next most prevalent among type strains of nine serotypes; together, they encompassed 15 unique IS200 profiles. The distribution and mol. wts of plasmids carrying sprBC (virulence) genes could be directly related to certain chromosomal genotypes defined by IS200 patterns. The presence of virulence plasmids in serotypes Lomalinda, Antarctica and Wangata is reported for the first time. Comparison of 16S rRNA gene profiles and IS200 patterns provides a definition of genotype that is applicable to epidemiological studies of various group D1 serotypes and should prove particularly useful for those lacking plasmid DNA.

Introduction

The cell-surface antigen profiles of salmonellae are so diverse that serotyping has, historically, provided a reliable phenotypic method for their differentiation. In the diagnostic scheme of Kauffmann and White, > 2300 serotypes of Salmonella are distinguished.1 Clinically or epidemiologically important serotypes, such as Typhi or Enteritidis, can be subtyped by serotype-specific phage typing. However, for most other salmonella serotypes phenotypic subtyping is not available; hence, the development of chromosomal fingerprinting by a common methodology would prove enormously valuable. Ribosomal RNA genes have provided one such probe.2 The profile of the DNA insertion sequence IS2003 is also provides fingerprints of salmonella chromosomes that reproducibly discriminate epidemiological clonality in salmonella serotypes, if the copy number is sufficiently high.5

Certain serotypes of Salmonella contain high-mol.- wt plasmids, related by DNA homology and restriction endonuclease fingerprinting, that contain a common "virulence region".6,7 Six spr (salmonella plasmid virulence) genes have been characterised to the nucleotide sequence level; one, sprC, has a clearly demonstrable role in virulence for mice.8 However, virulence plasmids are absent from some other important serotypes of Salmonella.9,10

We have compared restriction fragment length polymorphism (RFLP) at IS200 and 16S rRNA gene loci, and examined the presence of virulence plasmids in type strains of serotypes of Salmonella of Kauffmann-White O-serogroup D1. The objective of this study was to provide a necessary background for the rapid molecular definition of genotype in epidemiological studies.

Materials and methods

Bacteria

Strains of Salmonella used in this study11,12 are listed in table I. Purity was checked on blood agar plates before growth in Luria-Bertani broth for isolation of DNA. Stock cultures were maintained on Dorset’s egg slopes.

DNA isolation, probes and hybridisation

Preparation of intra-genic probes for IS200 and the 16S rrn gene, labelling with 16-dUTP-biotin and hybridisation conditions have been described previously.3 Plasmid DNA was isolated by the method of Kado and Liu13 and genomic DNA by the method

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of Wilson. Small amounts (3 μg) of genomic DNA were digested with PstI, PvuII or BglII for determination of IS200 pattern, or with PvuII, PstI or Smal for 16S rRNA gene profile. These enzymes lack restriction sites within the corresponding DNA probe sequences. The spuBC virulence gene probe, a 4.1-kbp EcoRI fragment of plasmid IP1367, was separated by electrophoresis in low-melting-point agarose and labelled by random priming.

### Results

**IS200 patterns**

The distribution of IS200 was investigated principally in genomic Southern blots made with PstI, such as those shown in fig. 1, "IS200 bands", as discussed below, should correspond to insertion sites carried on PstI fragments. Simple conserved features were found in 10 strains, including the type strains of seven serotypes sharing the phase-1 flagellar antigen "g", and serotype Pullorum. Except for serotype Berta, these type strains possessed two IS200 bands and shared a 4.5-kbp IS200 band (see fig. 2). There were two subgroups, one of which contained IS200 bands of 4.5 and 5.2 kbp, and the other of 4.5 and 3.9 kbp. The former subgroup comprised the type strains of serotypes Enteritidis (PT4), Blegdam, Moscow and Antarctica, each of which showed the 16S rRNA gene profile D-R1 (see below and table II). It also included the type strain of the non-motile, non-flagellate serotype Pullorum. The latter subgroup comprised type strains of serotypes Enteritidis (PT11), Dublin and Rostock, each of which showed a 16S rRNA gene profile termed D-RII (see below and table II). The type strain of serotype Enteritidis PT8 had a distinct IS200 profile, but shared the 16S rRNA gene profile termed D-R1 (see below) of Enteritidis PT4.

The other IS200 profiles found among type strains of serogroup D1 were diverse, and unique profiles were associated with many of them. Nonetheless, certain bands were conserved among serotypes. In ascending order of size, a 2.1-kbp EcoRI fragment of plasmid pIP1367, was separated by electrophoresis in low-melting-point agarose and labelled by random priming.

### Table I. Type strains of Salmonella of O-serogroup D1 used

<table>
<thead>
<tr>
<th>Serotype</th>
<th>Strain number*11</th>
<th>O antigens</th>
<th>H antigen phase 1</th>
<th>H antigen phase 2</th>
<th>Place and date of isolation*11</th>
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<tr>
<td>Miami</td>
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<td>1, 9, 12</td>
<td>a</td>
<td>l, 5</td>
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<td>a</td>
<td>l, 7</td>
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<td>9, 12</td>
<td>b</td>
<td></td>
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<tr>
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<td>1, 9, 12</td>
<td>b</td>
<td>e, n, z15</td>
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<td>NCTC 9868</td>
<td>9, 12</td>
<td>e</td>
<td></td>
<td></td>
</tr>
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<td>NCTC 8385</td>
<td>9, 12 [Vi]</td>
<td>c</td>
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<td>NCTC 8700</td>
<td>9, 12</td>
<td>d</td>
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<td>1, 5</td>
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<td>e, h</td>
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<td>f, g, t</td>
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<td>1, 9, 12</td>
<td>f, m, p</td>
<td>[1, 7]</td>
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<tr>
<td>Moscow</td>
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<td>g, q</td>
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<td>g, s, t</td>
<td></td>
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<td>g1, z23</td>
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<td>m, t</td>
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<td>Claibornei</td>
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<td>k</td>
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<td>l, v</td>
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<td>l, v</td>
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<td>l, w</td>
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<td>z10, z23</td>
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<td>z10</td>
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<td>Pullorum</td>
<td>NCTC 5776</td>
<td>1, 9, 12</td>
<td>z10</td>
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**16S rRNA gene profiles**

Restriction endonuclease *Pvu*II generated the most discriminatory 16S rRNA gene profiles; thus, 11 unique profiles were found among type strains of the 30 group D1 serotypes examined and were identified by the prefix D-R (I–XI). Characteristic examples are shown in fig. 3. With two exceptions, the number of homologous *Pvu*II fragments was six. Type strains of serotypes Typhi and Lomalinda, which display among the largest numbers (14) of chromosomal IS200 insertions, generated a five-band profile. A *Pvu*II fragment of 9.3 kbp was ubiquitous, whilst fragments of 3.4, 6.5, 9.5 and c. 15 kbp occurred in otherwise diverse profiles.

The commonest profile, D-R1, shared by type strains of 11 serotypes, was distributed throughout serogroup D1 without reference to phase-1 flagellar antigen composition; this group included the type strains of the epidemic phage types (PT4 and PT8) of serotype Enteritidis, as well as serotypes Moscow, Blegdam and Antarctica which also possess the phase-1 flagellar antigen "g". However, the type strains of serotypes Miami, Saarbruecken, Ndolo and Goettingen which have other diverse flagellar antigens (see table II) also showed the D-R1 profile.

The second commonest profile (D-RII) was conserved among the type strains of nine serotypes, including a phage-type strain (PT11) of serotype Enteritidis, serotype Dublin and serotype Berta. The D-RIII 16S rRNA-gene profile was shown by the type strains of serotypes Napoli and Portland (fig. 3, track 4) and D-RIV by the type strains of serotypes Daressalaam and Neasden, which have the same (c, n, x) phase-2 flagellar antigens (data not shown). Seven other profiles (D-RV–D-XI) were found in individual type strains only (e.g., fig. 3, tracks 2, 3, 6, 7, 8 and 9). Profile D-RX was observed in serotype Pullorum, which had the same IS200 profile (*SeCLI*: table II) as the Enteritidis (PT4) group. The overall occurrence of 16S rRNA gene profile in relation to IS200 pattern is documented in table II.

**Plasmid analysis**

Eighteen strains belonging to 16 serotypes carried...
plasmids, ranging in size from 1 to 126 MDa, the majority of which were > 30 MDa in size. The type strains of four serotypes carried two plasmids; those of 14 other serotypes carried none (table II). None of the plasmids showed any homology with IS200. Eleven plasmids ranging in size from 28 MDa (in serotype Lomalinda) to 59 MDa (in serotype Enteritidis) exhibited homology with the plasmid virulence genes \( spBC \) (fig. 4A versus B) of Typhimurium. These virulence plasmids are reported for the first time for serotypes Lomalinda, Antarctica and Wangata.

**Discussion**

A conceptually simple and satisfactory approach to ribotyping\(^2\) employs PCR-generated intragenic probes to obtain 16S rRNA gene-specific profiles.\(^5,17\) In this way, 11 distinct 16S rRNA gene profiles were observed among 32 strains in this study. Seven of these were specific to individual serotypes. Two conserved profiles (termed D-R1 and D-R1I) were found among strains with 15 distinct IS200 patterns, suggesting a common ancestry among these strains. The 16S rRNA gene profile D-RIV was shared by serotypes Dares-salaam and Neasden, which lacked IS200; this is consistent with their assignment to a different subspecies (subsp. salamae) of Salmonella enterica.

Both IS200 and 16S rRNA gene profiles indicate that there is indeed a close phylogenetic relationship between two subgroups of serotypes forming the “g” flagellar antigen group. Serotypes Enteritidis (PT11), Dublin and Rostock have one set of identical IS200 and 16S rRNA gene profiles whereas serotypes Enteritidis (PT4), Blegdam and Moscow have another. With respect to serotypes Enteritidis (PT11), Dublin and Rostock, multilocus enzyme electrophoresis (MLEE) has demonstrated that certain electrophoretic types (ETs) of serotype Dublin are virtually identical to the most common ET of serotype Enteritidis, and that serotype Rostock too is virtually identical to Dublin.\(^{18,19}\) The pattern of conservation of IS200 and 16S rRNA gene profiles in this study is consistent with the suggestion\(^20\) that the globally predominant ET of serotype Enteritidis resembles the ancestor of serotypes Dublin and Pullorum. Another point of evolutionary interest concerns serotypes Israel and Goettingen, whose identical IS200 profile indicates their common chromosomal genetic origin. These serotypes share the same phase-2, but have different phase-1, flagellar antigens. One, or other, may represent a case of gene-specific evolution to new serotype by horizontal transfer of the \( fliC \) (phase-1 flagellin) gene, a mechanism proposed by Smith et al.\(^{18}\)

Lam and Roth\(^4\) noted that each of the five serotypes of Salmonella shown to carry more than six IS200 copies was a mammalian pathogen. In the present study, the type strains of three serotypes with a large number of IS200 bands—Typhi, Lomalinda (NCTC 6705; isolate from a baby with meningitis) and Portland (NCTC 9907; isolate from woman with peritonitis)—were from systemic infections of man,
Table II. 16S rRNA-gene profiles, IS200 patterns and plasmids of type strains of serotypes of *Salmonella* of O-serogroup D1

<table>
<thead>
<tr>
<th>Serotype</th>
<th>16S rRNA gene profile (D-R)</th>
<th>IS200 pattern</th>
<th>number of IS200 bands</th>
<th>Plasmid sizes (MDa)</th>
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<tr>
<td>Miami</td>
<td>RI</td>
<td>u</td>
<td>4</td>
<td>60</td>
</tr>
<tr>
<td>Saarbruecken</td>
<td>RI</td>
<td>u</td>
<td>6</td>
<td>...</td>
</tr>
<tr>
<td>Lomalinda</td>
<td>RV1</td>
<td>u</td>
<td>14</td>
<td>28*</td>
</tr>
<tr>
<td>Durban</td>
<td>RI</td>
<td>u</td>
<td>5</td>
<td>...</td>
</tr>
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<td>Onarimon</td>
<td>RJII</td>
<td>u</td>
<td>6</td>
<td>...</td>
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<td>...</td>
<td>0</td>
<td>52</td>
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<td>u</td>
<td>14</td>
<td>...</td>
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<tr>
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<td>RI</td>
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<td>...</td>
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<td>RI</td>
<td>C</td>
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<td>Berta</td>
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<td>u</td>
<td>3</td>
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<td>Enteritidis (PT11)</td>
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<td>59*</td>
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<td>Enteritidis (PT8)</td>
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<td>C: ScCLI</td>
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<td>Enteritidis (PT4)</td>
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<td>2</td>
<td>36*</td>
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<tr>
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<td>C: ScCLIII</td>
<td>2</td>
<td>50*</td>
</tr>
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<td>Rostock</td>
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<td>2</td>
<td>50*</td>
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<tr>
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<td>C: ScCLI</td>
<td>2</td>
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<td>...</td>
<td>4†</td>
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<td>RII</td>
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<td>RIII</td>
<td>u</td>
<td>11</td>
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<td>RIII</td>
<td>u</td>
<td>9</td>
<td>54</td>
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C or u indicates conserved or unique IS200 pattern, respectively.
* Indicates plasmid hybridising to *spvBC* probe.
† DNA does not digest with *PstI* but four copies in digests with other enzymes.
‡ These serotypes belong to ssp. II (salamae).

Kbp

<table>
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Fig. 3. Examples of 16S rRNA gene profiles showing genomic Southern blot (*PvuII*) hybridised with 550-bp PCR amplicon internal to the 16S rRNA gene of serotype Dublin. Lanes contained: 1, Miami; 2, Panama; 3, Lomalinda; 4, Napoli; 5, Onarimon; 6, Pensacola; 7, Pullorum; 8, Eastbourne; 9, Alabama.
whilst serotype Wangata has recently been implicated in outbreaks of human salmonellosis in England and Wales.\textsuperscript{21} Like serotype Typhi, serotypes Lomalinda (dulcitol- and d-tartrate-negative), Portland (dulcitol-negative and raffinose-positive) are Wangata (negative in glycerofuscin broth) and biochemically atypical. If the transposition rate of insertion sequences and their consequent mutagenic activity increase with copy number, the multiple chromosomal IS200 insertions in these serotypes might have compromised their survival.

Fig. 4. Virulence plasmids in serogroup D1. A. Plasmid mini-preparations of \textit{E. coli} 39R861, a plasmid mol. wt-marker strain, (lane 1) and of type strains of 15 \textit{Salmonella} serotypes. Lane 2, Miami; 3, Lomalinda; 4, Alabama; 5, Eastbourne; 6, Israel; 7, Enteritidis PT4-E2187; 8, Enteritidis PT11-E2019; 9, Bledgdam; 10 and 11, Dublin; 12, Rostock; 13, Moscow; 14, Antarctica; 15, Clairborne; 16, Wangata; 17, Portland; 18, Pullorum. B. Southern blots of gel in A, probed with \textit{speBC}: homology with 11 plasmids was observed.
abilities, a situation reflected by the above metabolic deficiencies. They may have evolved either towards obligate host adaptation (e.g., Typhi) or towards rarity (e.g., Lomalinda). Again, serotypes Portland and Lomalinda may be host adapted, albeit to hosts not yet described, and, hence, as a result appear "rare".

Plasmids carrying spec:BC-homologous regions (cf. table II) and of very similar size to that of serotype Enteritidis (PT4) were detected in Antarctica, Blegdam and Moscow, serotypes which share its chromosomal IS200 profile, SeCLI12 (table II). On the other hand, Rostock contained a 50-MDa virulence plasmid, the same size as that of Dublin, and these two serotypes shared IS200 pattern SeCLIIII, as found in Enteritidis (PT11), which is one of a number of Enteritidis phage types containing plasmids of this approximate size (59 MDa). The only serotype type strain sharing IS200 profile SeCLI, but which carried a virulence plasmid of an unusual size, was Pullorum. The virulence plasmid of serotype Pullorum has been shown to be genetically distinct, belonging to an incompatibility group different from the 38-MDa plasmid of serotype Enteritidis and the 50-MDa plasmid of serotype Dublin.22 In serotypes Antarctica, Lomalinda and Wangata, spe-homologous plasmids were detected for the first time. The virulence plasmid of Wangata was of the same size (52 MDa) as that of Pullorum. Serotype Wangata has been reported to be a causative agent of poultry-related outbreaks of human gastro-enteritis.23 The plasmid found in serotype Lomalinda is possibly the smallest virulence plasmid so far detected; only serotype Choleraesuis has been reported to carry one of such small size (33 MDa).23

Almost half of the type strains of group D1 serotypes in this study contained neither virulence nor other plasmids, confirming that virulence plasmids are by no means ubiquitous in salmonellae, and that the plasmid-profiling approach to subtyping is not applicable to many serotypes of Salmonella. Since many of these plasmid-free type strains, e.g., serotypes Typhi, Saarbruecken, Durban, Onarimon, Pensacola and Panama, contained > 5 IS200 bands, chromosomal typing with IS200 should give a useful level of inter-strain discrimination within the serotype, as was shown for Typhimurium,2 providing a unified approach to genotypic subtyping. Except for Typhi, which can be phage-typed, subtyping schemes do not presently exist for any of these serotypes.

In this study we have addressed the general framework of IS200 patterns for a major serogroup by examining type-strain profiles. It has been shown that a given serotype tends to have a specific, related range of profiles and copy numbers of IS200.24 Thus, the profiles reported here do indicate what range of profiling can be photographically or diagrammatically represented in a band-matching database, facilitating the comparison of data between laboratories—an important consideration for epidemiology. Where the number of IS200 bands is moderate to high, the chromosomal types so defined are consistent with the theoretical considerations defining clonality,24 and may be more definitive than phenotypic types for medico-legal aspects of salmonellosis. Results obtained with IS6110 typing of Mycobacterium tuberculosis, recently adopted internationally as a standard method,25 confirm the applicability and reliability of this approach in epidemiological typing.

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References


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