Genetic Variation in IncI1-ColIb Plasmids

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Abstract. Nucleotide sequences of portions of three plasmid genes *(cib, cir,* and *abi)* present in IncI1-ColIb colicin plasmids obtained from strains of *Salmonella typhimurium* isolated in either 1974 (Barker strains) or between 1935 and 1941 (Murray strains) were examined along with sequences of the chromosomal gene for 6 phosphogluconate dehydrogenase *(gnd).* Our principal findings were: (1) The plasmid genes were virtually identical to those in IncIl-ColIb plasmids from *E. coli,* suggesting that *Salmonella* and *E. coli* share overlapping pools of these plasmids. (2) The plasmid genes were much less polymorphic than *gnd* or any other known chromosomal gene from *Salmonella,* further suggesting horizontal transfer with rapid transmission and turnover. (3) No characteristic differences were found in either the plasmid genes or the chromosomal gene between the 1974 isolates and the Murray strains, indicating that these plasmids have been stable for at least several decades. (4) There was an excess of amino-acid replacement polymorphisms, relative to synonymous polymorphisms, in the plasmid genes, which is consistent with the hypothesis of diversifying selection among colicin-producing plasmid families. (5) The *abi* (abortive infection) gene present in each of the plasmids contained two single-nucleotide insertions relative to the published sequence. These result in a putative *abi* protein of 114 amino acids instead of 89.

Key words: Salmonella typhimurium -- Plasmid --Colicin Ib -- Abortive infection gene (abi) -- Diversifying selection

Introduction

The colicin plasmids comprise a diverse group of plasmids containing genes that code for proteins toxic to bacteria as well as genes conferring specific immunity to the host which produces those toxins (Konisky 1982). The distribution and abundance of colicin plasmids have been characterized in both *S. typhimurium* (Barker 1980) and *E. coli* (Riley and Gordon 1992). Approximately 12% of natural isolates of *S. typhimurium* (Barker 1980) and 35% of a reference collection of natural isolates of *E. coli* (Riley and Gordon 1992) produce colicins; among the most frequent colicin classes are ColIa and ColIb. Colicins specifying related bacterial toxins are identified by the presence of cross-immunity. At present, approximately 20 distinct types of colicins have been identified (Pugsley 1985; Riley 1993a).

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Colicins that are related physiologically are encoded in genes that are similar in nucleotide sequence. Comparisons of pairs of related colicin operons in three different colicin families (colicins E3 vs E6, E2 vs E9, and Iavs Ib) yielded nucleotide-sequence identities ranging from 85 to 96% (Riley 1993b). In comparing the rates of synonymous, nonsynonymous, and intergenic nucleotide substitution in these three pairs of closely related colicins, Riley (1993b) observed an excess of nonsynonymous substitutions in the immunity gene and in the immunity-binding domain of the colicin gene. The excess of amino acid replacements, relative to synonymous nucleotide substitutions, was hypothesized to result from selection for diversity in the immunity protein and in the immunity-binding region of the colicin (Riley 1993a,b).

The present analysis was carried out, in part, to determine whether independent plasmid isolates coding for a particular type of colicin also show an excess of

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amino-acid replacement polymorphisms, relative to synonymous polymorphisms. The study also addresses the issue of nucleotide diversity among genes present in plasmids. Most previous work on plasmid polymorphisms has been biased toward finding diversity because the studies have focused on isolates of plasmids that differ in one or more phenotypic attributes. For example, regions of several genes suspected to be responsible for the diversity of plasmid phenotypes have been sequenced and compared from a wide range of Frelated plasmids (Finlay et al. 1986a-c). The level of DNA sequence identity in these regions ranged from 68 to 100%. These data contrast with studies of several types of transposable insertion sequences (IS) indicating little or no genetic variation within species (Lawrence et al. 1992; Bisercic and Ochman 1993), a result attributed to rapid turnover and plasmid-mediated horizontal transmission of IS elements (Lawrence et al. 1992). Hence, a reduced level of genetic variation in plasmid genes, relative to chromosomal genes, might also be expected.

We have examined nucleotide-sequence variation in three genes in conjugative plasmids of the incompatibility group IncI1 containing the ColIb colicin operon. The plasmids were present in strains of *Salmonella typhimurium* obtained as natural isolates either from Scotland in 1974 (Barker 1980; Barker et al. 1980) or from geographically diverse locations and collected by E.D.G. Murray in 1935-1941 (Hughes and Datta 1983; Jones and Stanley 1992). The latter strains represent those of the preantibiotic era and so provide an opportunity to detect evolutionary changes in plasmid genes over a period of approximately 40 years. The plasmids were chosen solely because they were IncI1-Collb and so have not been preselected for phenotypic or genetic diversity. The loci analyzed were *cib, cir,* and *abi.* The *cib* and *cir* loci are tightly linked and code for the colicin Ib toxin and immunity proteins, respectively (Mankovich et al. 1986); the *abi* locus codes for the abortive-infection protein that is responsible for abortive infection by the bacteriophages BF23 or T5 (Gupta and McCorquodale 1988). We also examined sequences of the chromosomal gene, *gnd,* coding for 6-phosphogluconate dehydrogenase, from the same bacterial strains.

Materials and Methods

Bacterial Strains and Plasmids. S. typhimurium strains M102, M140, M303, M496, M498, and M501 from the Murray collection were obtained from Dr. John Stanley and the National Collection of Type Cultures, London. M140 was added to the Murray collection in 1935; the other strains were added in 1941. The Murray strains are sometimes referred to as preantibiotic era (PAE) strains (Hughes and Datta 1983; Jones and Stanley 1992). The IncI1-ColIb plasmids in the strains were identified by Jones and Stanley (1992). More recent *S. typhimurium* strains 205/74 (biotype 17g) and 375/74 (biotype 9f) were collected in 1974 from water in Glasgow and from a cattle farm in

Wigtownshire, respectively, and characterized by Barker et al. (1980). These strains were provided by Dr. Jordan Konisky.

DNA Amplification. Oligonucleotide primers for the polymerase chain reaction (PCR) and for DNA sequencing were designed from published sequences of *cib* and *cir* (Mankovicb et al. 1986), *abi* (Gupta and McCorquodale 1988), and *gnd* (Reeves and Stevenson 1989) using the OLIGO 4.0 program for Macintosh (Rboads and Rychlik 1989). The following PCR primers were used: for the region containing parts of the *cib* and *cir* loci, 5'-AAGCAGTTAATTC-ATTTGTTT-3' and 5'-ACAGGAGCACCTCCGTTCCAT-3'; for the *abi* locus, 5'-ACAGGAGGATGCGAATGAACA-3' and 5'-GCGT-TGCGGCTACTTACTTTA-3'; and for the *grid* locus, 5'-GAGTAT-GTAATGTCCAAGCAA-3' and 5'-CTCAGCTCGCCGTTATTC-CAC-3'.

The optimal annealing temperature of each primer pair was estimated using the OLIGO-4.0 program. Following amplification (Saiki et al. 1988), the PCR products were purified by diluting the 30 - μ l reaction volume to a final concentration of 100 μ l with distilled and dionized H₂O, followed by phenol extraction, precipitation in the presence of 2.5 M sodium acetate and one volume of absolute ethanol, and washing with 70% ethanol (Sambrook et al. 1989).

DNA Sequencing. Sequencing of PCR products was performed with an Applied Biosystems model 373A automated DNA sequencing system using the *Taq* DyeDeoxy terminator cycle-sequencing kit using reaction conditions and other methods recommended by the manufacturer. Organic contaminants were removed by means of the phenol-chloroform extraction protocol provided with the sequencing kit. Both strands of all regions were sequenced in their entirety. Sequences were aligned and analyzed by hand.

Nucleotide sequences from the following regions (numbered according to the published sequences) were determined: for the *cib* locus, sites 1399-1519, 1640-1716, 2244-2520, 2568-2598, and 2601-2621 (total 527 bp); for the *cir* locus, sites 3057-3163 and 3201-3260 (total 167 bp); for the *abi* locus, sites 1343-1508 (total 166 bp); and for the *gnd* locus, sites 235-385 (total 151 bp). All strains were sequenced in these regions except strain 205/74, in which the *abi* and *gnd* regions were not analyzed. Nucleotide sequence was also determined for the *abi* locus between sites 1509 and 1617 in strain M140 and sites 1509-1596 in strain M496.

Results

Nucleotide-Sequence Variation

Analysis of nucleotide variation in the IncI1-ColIb plasmids was based on approximately 7 kb of sequence from three plasmid genes in IncI1 plasmids present in two strains isolated in 1974 as well as in Incll plasmids present in six Murray strains isolated in 1935-1941. The sequenced regions included 527 nucleotides of *cib* (the colicin Ib structural gene), 167 nucleotides of *cir* (the colicin Ib immunity gene), and 166 nucleotides of *abi* (abortive infection gene). For comparison, a region of approximately 150 nucleotides was determined for the chromosomal gene *gnd,* coding for 6-phosphogluconate dehydrogenase, in the same strains.

The polymorphic nucleotides in each of the three plasmid loci, as well as in the *gnd* locus, are summarized in Fig. 1, in which the nucleotides are classified according to position, type of nucleotide change, and ef-

| | | | <u>Strain</u> | | | | | | | | |
|-------|------------|----------------------|---------------|----------------|----------------------|-------------|-----------|-----------|----------------|------|--|
| | | | | M | M | M | M | Μ | M | | |
| | | 2 | 3 | $\mathbf{1}$ | $\mathbf{1}$ | 3 | 4 | 4 | 5 | | |
| | | 0 | 7 | 0 | 4 | 0 | 9 | 9 | 0 | | |
| Locus | Site | 5 | 5 | \overline{c} | Ō | 3 | 6 | 8 | $\overline{1}$ | Type | |
| cib | 1404 C | | | | | , | | | A | Rep | |
| | 1655 G | | \bullet | A | | \bullet | ٠ | ٠ | | Syn | |
| | 2420 T | | ۰ | | ٠ | $\mathbf C$ | ۰ | ٠ | ٠ | Syn | |
| | 2428 T | ä | ۰ | | | ٠ | C | | ۰ | Rep | |
| | 2432 T | $\ddot{}$ | C | | | C | Ċ | | C | Syn | |
| | 2455 A | G | | | ۰ | ٠ | ۰ | | | Rep | |
| | 2484 G | | , | | T | ٠ | | | | Rep | |
| | 2514 G | | | | A | ٠ | | | | Rep | |
| | 2570 A | т | T | T | T | Τ | T | T | T | Rep | |
| | 2597 A | | ٠ | | C | | | | | Rep | |
| cir | 3063 A | | | | | | G | | | Rep | |
| | 3084 T | Ċ | | | | | | | | Rep | |
| | 3103 A | | | C | | | | C | C | Rep | |
| | 3126 A | C | | | | | | \bullet | | Rep | |
| | 3139 T | Ċ | | | | | | | | Syn | |
| abi | >1354 - | | Т | T | т | Т | Т | T | Т | | |
| | 1366 T | | ٠ | ٠ | A | | | | | Rep | |
| | 1442 T | | G | Ġ | Ġ | | | | | Rep | |
| gnd | 237 Т | | ۰ | | ۰ | | ۰ | ٠ | с | Syn | |
| | ŕ 240 | | | | | | ۰ | ٠ | Ċ | Syn | |
| | 252 C | | ٠ | ٠ | ٠ | | | T | | Syn | |
| | 261 G | | ă | | Ċ | ٠ | \bullet | Ċ | ٠ | Syn | |
| | 276 Α | | ۰ | ٠ | $\ddot{}$ | T | T | | | Rep | |
| | 282 C | | Т | | ۰ | \bullet | T | \bullet | | Syn | |
| | 300 C | | T | T | | Ţ | | | | Syn | |
| | 330 Т | | C | | | | ۰ | | ٠ | Syn | |
| | 353 A | | | | | | | | G | Rep | |
| | 371 G | | | | | | | C | | Rep | |
| | 378 T | | Ġ | | | | | | | Syn | |
| | 381 T | | ٠ | G | G | | | | | Syn | |

Fig. 1. **Site refers to the numbering of the nucleotides in the published sequences of Mankovich et** al. (1986) **for** *cib* **and** *cir,* **Gupta and McCorquodale (1988) for** *abi,* **and Nasoff et** al. (1984) for *gnd.* Nucleotides **identical to the published sequences are indicated by** *dots.* **Type (replacement of synonymous) refers to the effect on the amino acid sequence. The symbol** *>1354* **in the** *abi* **gene indicates a single nucleotide insertion following site** 1354.

fect on amino acid sequence (replacement difference vs synonymous difference). The proportion of polymorphic nucleotides (K) in each gene is summarized in the second column in Table 1. All three plasmid genes are polymorphic: taken together, the three plasmid genes have $K = 0.021 \pm 0.005$. The third column in Table 1 **summarizes the polymorphism data in terms of the nu**cleotide diversity, π , which is the average number of nu**cleotide differences per site between two randomly chosen alleles. The nucleotide diversity has the convenient property that it is independent of both sample size and total number of sites (Nei 1987, p 256). The weighted** average π for the three plasmid loci (weighted by the reciprocal of the estimated variance) is 0.0071 .

There is a difference in character of the polymorphisms, as well as in the level, between the plasmid and chromosomal genes. In particular, the proportion of replacement polymorphisms is significantly greater in the colicin genes *cib* **and** *cir* **than in the chromosomal gene** *gnd* (Fig. 1; $\chi^2 = 6.2$, $P < 0.02$). This difference sug**gests either relaxed selective constraints on the amino**

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^a Proportion of polymorphic nucleotides in the sample \pm binomial **standard error**

b **Average number of nucleotide differences per site between randomly chosen pairs of alleles _+ standard error** (Nei 1987, p 276)

acid sequence of the colicin genes or positive selection for amino acid diversity.

Plasmids from 1935-1941 Isolates vs 1974 Isolates

There are some pairs of duplicates in the Murray strains (Hughes and Datta 1983), and so a serious consideration a priori was whether the strains added to the collection at the same time and with similar numbers might be duplicates of the same isolate. This consideration would especially include strains M496, M498, and M501, which were all added to the collection in 1941. However, the DNA sequencing results clearly indicate that the strains are different. Not only does the *gnd* **gene differ in all three strains, but so does** *cib (cir* **also differs between M496 and M498/M501).**

There are no characteristic differences in any of the plasmid genes between the 1974 Scotland isolates and the Murray strains. For example, dendrograms of the strains constructed from the plasmid genes by means of the unweighted pair-group method using arithmetic averages (UPGMA), as implemented in the Clustal V program (Higgins et al. 1992), showed no grouping of the 1974 isolates apart from the Murray isolates (data not shown).

The UPGMA dendrograms also showed no tendency to group the plasmid gene sequences determined for IncI1-ColIb plasmids derived from *E. coli* **apart from the plasmid-gene sequences determined for IncI1-ColIb plasmids derived from** *S. typhimurium* **(data not shown). This observation provides evidence that the Incll-ColIb plasmid pools in** *E. coli* **and** *S. typhimurium* **are not distinct, genetically divergent plasmid subpopulations.**

Extended Open Reading Frame in the abi Gene

Unexpectedly, all of the *abi* **sequences determined in this study differed in a significant way from the published sequence, and also from a** *Salmonella* **IncI1- ColIb plasmid (Gupta and McCorquodale 1988). In particular, all sequences contain a single-nucleotide insertion relative to the published sequence, following nucleotide site 1354. The resulting frameshift generates a termination codon 33 amino acid residues from the** published putative initiation codon. Downstream sequence determined for *abi* in the Murray strains M140 and M496 revealed another single-nucleotide insertion—a guanosine following nucleotide site 1570 which extends the published open reading frame by 42 codons. The structure of the *abi* gene proposed by Gupta and McCorquodale (1988) was determined on the basis of analyses of open reading frames and the location of appropriate promoter signals within a 0.8-kb restriction fragment known from deletion-mapping experiments to overlap the *abi* locus in an IncI1-Collb plasmid. Based on our data indicating an additional nucleotide following sites 1354 and 1570, we propose that the *abi* coding region extends from the methionine at site 1357 to the UAA stop codon at site 1698 (numbered as in Gupta and McCorquodale 1988). This open reading frame codes for a putative *abi* protein about 30% longer than that suggested by Gupta and McCorquodale (1988), yielding 114 amino acids vs 89. It is preceded by a Shine-Dalgarno sequence (GGAGG) five nucleotides upstream from the putative initiating methionine and overlaps, in frame, the proposed reading frame of Gupta and McCorquodale (1988) by 71 codons. Computer searches of both protein and nucleic acid databases performed at the National Center for Biotechnology Information using the BLAST network service failed to reveal any significant homologies with the proposed alternative coding sequence for the *abi* gene.

Discussion

An analysis of genetic variation among IncI1-ColIb plasmids was undertaken to elucidate factors affecting plasmid evolution as well as the evolutionary relationships between plasmids in strains collected in 1935- 1941 vs 1974 and those between IncI1-ColIb plasmids in *Salmonella* and *E. coli.* To assess the levels of nucleotide polymorphism in plasmid strains, we determined the DNA sequences of three *loci--cib, cir* and *abi--present* in IncI1-Collb plasmids in isolates of S. ty*phimurium* from 1974 and from the Murray strains.

The level of genetic variation among the plasmid genes averaged $K = 0.021$ for the proportion of polymorphic nucleotides and $\pi = 0.0071$ for the pairwise number of nucleotide differences per site. These values are substantially smaller than those for chromosomal genes in *Salmonella*: estimates of π and K for the 6phosphogluconate dehydrogenase gene are $\pi = 0.030$, $K = 0.080$ (present study); those for the glyceraldehyde-3-phosphate dehydrogenase gene from 16 isolates are π $= 0.037$ and $K = 0.128$ (Nelson et al. 1991); and those for the proline permease from 16 isolates are $\pi = 0.046$ and $K = 0.147$ (Nelson and Selander 1992). A reduced level of nucleotide polymorphism in plasmid genes, relative to chromosomal genes, would be expected from a high rate of plasmid turnover and horizontal transmissibility. This effect is observed dramatically in the remarkable homogeneity of insertion sequences in E. *coli* (Lawrence et al. 1992). On the other hand, the reduction may be smaller than first appears because the chromosomal genes that have been sequenced are among the most polymorphic. In general, isolates of S. *typhimurium* have limited genetic variation compared with *E. coli* (Beltran et al. 1991; Ochman and Selander 1984), as evidenced by the fact that 76% of isolates of *S. typhimurium* have an identical electrophoretic pattern for each of 20 polymorphic enzyme loci (Beltran et al. 1991).

The average level of nucleotide-sequence divergence between homologous chromosomal genes in *Salmonella* and *E. coli* is 15.6% (Bisercic and Ochman 1993); however, the genes that were sequenced from the IncI1- Collb plasmids from *Salmonella* are not more similar to each other than they are to those from IncI1-ColIb plasmids from *E. coli* (Isaacson and Konisky 1974; Mankovich et al. 1986). This result would be expected if *Salmonella* and *E. coli* shared overlapping pools of IncI1-Collb plasmids. It should also be noted that some chromosomal copies of the transposable element IS 1 found in *Salmonella* are virtually identical to the IS1 found in *E. coli* (Bisercic and Ochman 1993), and it is thought that plasmids play a major role in the horizontal transmission of such elements among strains (Sawyer et al. 1987; Hartl and Sawyer 1988). Furthermore, no differences distinguishing the 1974 isolates from the Murray isolates were found in either the plasmid genes or the chromosomal genes. This result is consistent with the finding that many of the same plasmid incompatibility groups are present among plasmids of 1935-1941 as among those of 1974 (Jones and Stanley 1992; Hughes and Datta 1983). On the other hand, because there can be considerable sequence variation among genes in plasmids of the same incompatibility group (Finlay et al. 1986a-c), some differences in nucleotide sequence might have been expected. The lack of such differences implies that IncI1-Collb plasmids have been widespread in distribution and stable for at least a number of decades. Although our data support intergeneric exchange of IncI1-ColIb plasmids, it should be noted that this is not necessarily true of all plasmids. For example, Mercer et al. (1984) found no evidence that common plasmids are transferred among *E. coli* lineages in nature, although such plasmids can readily be transferred in the laboratory.

The levels of nucleotide polymorphism in the IncI1-ColIb plasmids reported here are lower than might be expected from results of Riley and Gordon (1992). These authors used restriction enzyme digests to study polymorphisms among a set of either ColE1 plasmids and five ColIa plasmids isolated from a reference collection of *E. coll.* Based on the number of restrictionfragment differences pooled over four restriction endonucleases, the nucleotide diversity among the ColE1

plasmids were estimated as $\pi = 0.204 \pm 0.059$ (to be compared with an estimated $\pi = 0.0071$ for the IncI1-Collb plasmids). The divergence among the ColIa plasmids was even greater: each had a unique pattern of restriction fragments, and no common set of fragments was shared among all ColIa plasmids. However, the data reported here and those of Riley and Gordon (1992) are not strictly comparable because, in our data, the plasmids were selected a priori for being IncI1-ColIb whereas, in their study, the plasmids were selected solely based on colicin type. Plasmids selected solely by colicin type may very well be more diverse than those selected by more stringent criteria. Alternatively, ColE1 and Colla plasmids from E. *coli* may have more genetic variation at the DNA sequence level than ColIb plasmids from *S. typhimurium*.

The higher proportion of replacement polymorphisms, relative to synonymous polymorphisms, observed in the colicin and immunity genes as compared with *gnd* suggests either relaxed selective constraints on the amino acid sequences or diversifying selection for variation in the colicin and immunity genes. Because the same pattern was observed in three pairs of related colicin operons, Riley (1993b) suggested positive selection for genetic diversity. The excess of replacement substitutions in our data is consistent with the postulated process of diversification among the allelic operons of a single colicin type present in IncI1-ColIb plasmids.

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