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Ribosomal RNA Gene Restriction Patterns Provide Increased Sensitivity for Typing *Salmonella typhi* Strains

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To date, epidemiologic associations among strains of *Salmonella typhi* are based exclusively on phage typing, which may be of limited value if a common phage type is involved. Analysis of ribosomal RNA gene restriction patterns allows separation of most independently isolated strains of identical phage types. The sensitivity of the method is dependent on the restriction enzymes used to digest chromosomal DNA. It was highest for *Pst*I, which separated 16 of 20 strains that belonged to 8 phage types including 3 untypable strains. Three strains differed in their phage types but had identical ribosomal RNA gene restriction patterns. Also, two pairs of strains indistinguishable by phage typing exhibited identical patterns; however, two of these strains were expected to be identical because they were isolated from two patients who were likely exposed to the same source. Ribosomal RNA gene restriction patterns appear to be stable. Thus, the method may complement phage typing and aid in further differentiation of strains.

Typhoid fever is an acute, febrile illness that frequently represents a major public health problem in developing countries and to travelers to these regions. Large outbreaks usually linked to asymptomatic carriers who have excreted *Salmonella typhi* for years, however, have been reported from developed countries [1]. Bacteriophage typing requiring the presence of the Vi antigen is the method of choice to demonstrate epidemiologic associations among strains of this species [2], but its use is limited to reference

laboratories that have all 100+ necessary phage suspensions available. Also, depending on the geographic location, only a few phage types represent the majority of strains isolated [3, 4]. For example, nearly 25% of the strains isolated in the USA belong to phage type E1 and up to 80% of the strains encountered in certain African countries are of phage type A [4]. Therefore, phage typing provides little additional epidemiologic information if the same phage type is found both in patients and in the suspected carrier [2, 3].

The specificity of Vi types is determined by the carriage of temperate phages and by the nonlysogenic precursor type [5]. This implies that loss or uptake of a lysogenic phage can result in a change of the phage type. This has been proven experimentally and has been shown to occur naturally, thus complicating epidemiologic analysis [6]. Attempts to de-

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velop other reliable, sensitive, and easy-to-perform typing methods that might complement phage typing have failed because *S. typhi* strains are homogeneous in their phenotype. Multilocus enzyme electrophoresis, although useful for other groups of bacteria such as *Legionella pneumophila* [7], *Escherichia coli* [8], or *Aeromonas* species [9], did not reveal a single difference among 26 strains of diverse geographic origin and phage types with 24 different enzymes that had been analyzed [10]. We describe the variation of *S. typhi* strains with regard to ribosomal RNA (rRNA) gene restriction patterns [11] and their potential as an epidemiologic tool.

Strains and Methods

Strains. *S. typhi* strains isolated from humans were selected from the culture collection at the Centers for Disease Control, Atlanta, mainly to represent groups of identical phage type but to be independent from each other otherwise, that is, they were isolated at different geographic locations at differ-

ent times (table 1). In addition, two strains were included that were isolated from two patients in the same hospital who had probably been exposed to the same source. *Serratia fonticola* 3965 was used to provide a molecular weight standard for rRNA gene restriction patterns [11].

Restriction endonuclease analysis and ribosomal RNA gene restriction patterns. For restriction endonuclease analysis and rRNA gene restriction patterns, the method developed by Grimont and Grimont [11] was used with some modifications. Briefly, high-molecular-weight chromosomal DNA was isolated from strains grown on blood agar for 48 h at 37°C by a modified, small-scale version of a published procedure [12]. DNA samples were digested with restriction endonucleases (New England Biolabs, Beverly, Mass), electrophoresed on either 0.8% or 1.0% agarose gels in Tris-acetate buffer (0.04 M Tris acetate, 0.002 M EDTA, pH 8.0), and stained in ethidium bromide (1 µg/ml) for 20 min. Gels were then photographed and blotted to nitrocellulose according to established procedures [13]. Plas-

Table 1. Typing of *Salmonella typhi* strains of various phage types by analysis of restriction and rDNA patterns.

Strain-year	Location of isolation	Source	Phage type	Restriction patterns			Ribosomal DNA patterns		
				<i>Pst</i> I	<i>Eco</i> RI	<i>Sma</i> I	<i>Pst</i> I	<i>Eco</i> RI	<i>Sma</i> I
0486-78	Massachusetts	Blood	A	A		A	G		A
1081-79	California	Blood	A	A		A	H		C
3018-79	New York	Stool	A	A		A	H		C
2082-86	North Carolina	Blood	A	A		A	D		B
2051-87	New York	Stool	A	A		A	I		B
2835-79	Massachusetts	Bile	A (VIII-)	A		A	K		D
2003-88	Alabama	Unknown	A (VIII-)	A	B*	A	C	B*	E
1207-84	New York	Blood	A (III-)	A		A	L		F
0571-78	Texas	Blood	E1	A		A	N		C
0048-80	New York	Stool	E1	A			M		B
2030-87	Florida	Blood	E1	A			E		
3036-77	Texas	Blood	Degraded Vi/A† (mex ^r)	B			P		
0891-79	New York	Stool	Degraded Vi/A (mex ^s)	A			O		
2115-87	New York	Unknown	Degraded Vi/A (mex ^s)	A	A		E	A	
2001-88	California	Bile	Untypable	A	A		A	A	
2004-88	California	Bile	Untypable	A	A		D	A	
2005-88	California	Blood	Untypable	A	A		E	A	
2006-88	Connecticut	Bile	Degraded Vi (45-, 51-)	A	A		F	A	
2007-88	Connecticut	Bile	Degraded Vi (45-, 51-)	A	A		F	A	
2002-88	Pennsylvania	CSF	46 (atypical)	A	A		B	A	

NOTE. Isolates were epidemiologically related except strains 2006-88 and 2007-88, which were isolated from two patients in the same hospital ward. CSF = cerebrospinal fluid.

* DNA was not cut by *Eco*RI.

† Degraded Vi/A, degraded Vi approaching A. These strains are sometimes called the "Mexican strain" and occur in the two phenotypes mex^r (antibiotic resistant) or mex^s (antibiotic sensitive).

mid pKK 3535 was used as a probe. This plasmid is a pBR322-derived plasmid containing essentially a ribosomal RNA operon consisting of one copy each of the genes coding for 5S RNA, 16S RNA, 23S RNA, and tRNA^{Glu} [14]. Plasmid DNA was labeled using [³²P]dCTP (DuPont NEN Research Products, Boston) and a commercial nick translation kit (BRL, Gaithersburg, Md), heat-denatured and hybridized to the filter-bound restriction fragments as described by Maniatis et al. [13] with the following modifications: filters were prehybridized in 6× standard saline citrate (SSC), 10× Denhardt solution for 60 min, and in hybridization solution (2× SSC, 5× Denhardt solution, 0.02 M Tris hydrochloride, pH 7.4, 0.1% SDS, and 41% formamide) at room temperature for at least 60 min before addition of the radioactive probe. Hybridization was done overnight at 37°C, equivalent to an effective hybridization temperature of 66.5°C. Filters were then washed as described [9], dried, and autoradiographed for 3–24 h at –70°C using Kodak X-OMAT film and two intensifying screens (Kodak, Rochester, NY).

Results

Restriction endonuclease analysis. Among the 20 strains (17 strains belonging to 7 different phage types and 3 untypable strains), only slight variability could be demonstrated (figure 1 top; table 1). All nine strains digested with restriction endonuclease *Sma*I exhibited the same banding pattern. For the two enzymes *Eco*RI and *Pst*I the results were similar. Only one strain of each differed in its pattern. Strain 2003–88 was repeatedly resistant to the action of *Eco*RI, although the DNA was cut by other enzymes, and strain 3036–77, the only strain of that phenotype (mex^r, table 1), differed from all other strains by the presence of two high-molecular-weight bands (figure 1 top).

Ribosomal RNA gene restriction patterns. More variability among strains could be demonstrated when restriction fragments were hybridized with radiolabeled DNA complementary to the genes coding for rRNA. Some sequences within these genes are conserved among a wide range of taxonomic groups (including gram-positive and gram-negative organisms) and thus allow hybridization with rRNA isolated from *E. coli* or, as done in this study, with *E. coli* DNA (cloned in plasmid pBR322) coding for rRNA. Multiple bands are expected because rRNA

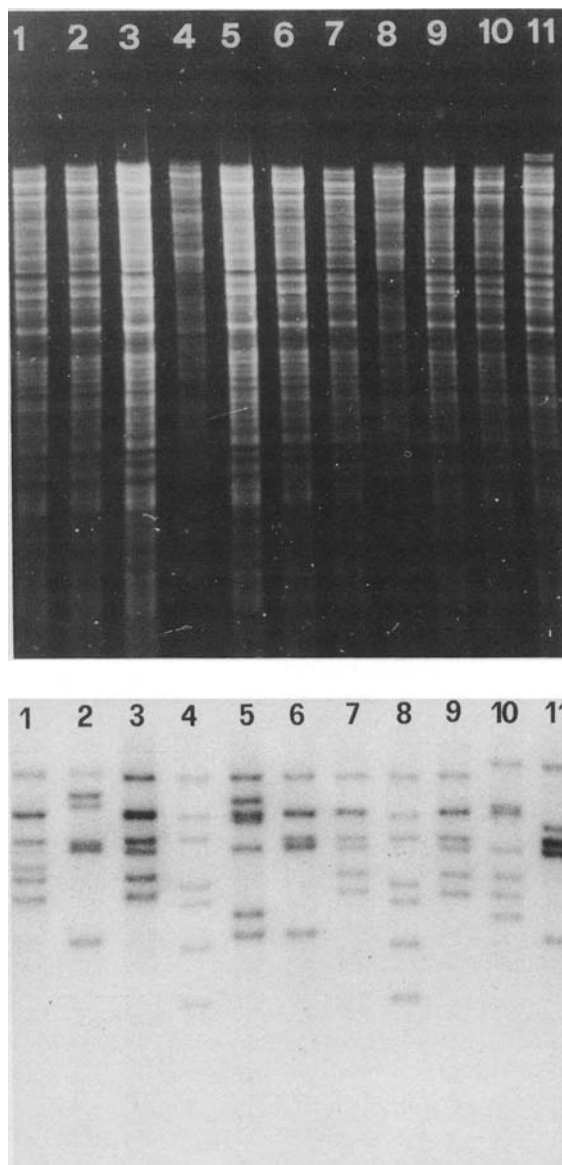


Figure 1. Restriction endonuclease patterns (top) and ribosomal RNA gene restriction patterns (bottom) of *S. typhi* DNA digested with *Pst*I. Lanes 4 and 8, *Serratia fonticola* DNA digested with *Hind*III (standard); lane 1, strain 0048-80; lane 2, strain 0571-78; lane 3, strain 2030-87; lane 5, strain 2001-88; lane 6, strain 2004-88; lane 7, strain 2005-88; lane 9, strain 2115-87; lane 10, strain 0891-79; lane 11, strain 3036-77. Strains belong to the following phage types: lanes 1-3, type E1; lanes 5-7, untypable; lanes 9-11, degraded Vi approaching A (Mexican strain). Sizes (in base pairs) of fragments generated by digestion of *S. fonticola* with *Hind*III (lanes 4 and 8) are 14,596, 11,853, 10,601, 8520, 7892, 6634, and 5419 (from top to bottom).

operons are redundant in most bacteria [11]. The variability of patterns is highly dependent on the restriction endonucleases used. *EcoRI* patterns were determined for five strains belonging to four different phage types and three untypable strains. All were identical except for the strain that was resistant to the action of this enzyme (table 1).

Ten strains of four phage types were analyzed with restriction endonuclease *SmaI*, and only two patterns were not restricted to a single strain (table 1). Strains 2082-86 and 2051-87 (both of phage type A) and strain 0048-80 (phage type E1) were identical (pattern B, table 1), as were three strains with pattern C that belonged to either phage type A (two strains) or phage type E1 (one strain). The two type A strains were isolated in the same year in two different states (California and New York).

The best discrimination among strains provided restriction endonuclease *PstI*. For 20 strains analyzed, 16 different patterns were found (table 1, figure 1). Only three groups of strains could not be differentiated. Among these were the two strains of phage type A that were also identical using *SmaI* (see above) and three strains of phage type E1, degraded Vi/A (mex^s, table 1), and one that was not typable. Also, the two strains of the same phage type that had been isolated from two patients in the same hospital ward in Connecticut were identical.

Discussion

Phage typing has been the method of choice for decades to demonstrate epidemiologic associations among isolates of *S. typhi*. However, because this method frequently fails to provide additional information [3], some researchers have attempted to develop methods that would complement phage typing and allow differentiation of nonrelated strains that are of identical phage types [10]. Our results clearly show that the determination of rRNA gene restriction patterns, but not the comparison of restriction patterns, is a technique with the desired sensitivity.

The usefulness of the method depends on the enzyme(s) chosen for the analysis. *EcoRI* was the least useful of the enzymes tested in this study. All patterns were identical with the exception of a single strain whose DNA was not cut at all. The reason for this phenomenon is unknown. However, it was reproducible and probably not due to impurities in the

DNA preparation because the activity of other enzymes was not affected.

SmaI was used with only 10 strains. It was of intermediate sensitivity; two groups of three strains each were not distinguishable. Identical strains did not necessarily belong to the same phage type (table 1).

For the 20 strains used in this study, *PstI* proved to be by far the most useful enzyme. Most strains of identical phage types were easily differentiated and differed in one or several bands. Two pairs of strains that were not distinguishable are of special interest. The first two strains, both of phage type A, were isolated in 1979 in two states (California and New York). Neither was distinguishable from the other by *SmaI* patterns. Because no more epidemiologic information regarding these two infections is available, it can be speculated that the two patients had been exposed to the same source. It is more likely, however, that the methods evaluated here do not allow differentiation of the strains. For the second pair of identical strains, the epidemiologic background is better known. The strains were isolated from two related patients treated in the same hospital, who likely acquired the infection by exposure to a common source.

These two strains were chosen for our analysis to provide some data on the stability of the patterns. The results are consistent with the earlier finding that the patterns for a strain of *E. coli* remained constant after 16 serial subcultures in vitro [15]. The same authors had also shown that all eight colonies isolated from the primary sputum culture of a *Pseudomonas cepacia*-colonized patient were identical, although colonies of different morphology had been selected. This relative stability is important for typing methods if they are to be applicable in epidemics. The three strains of *PstI* pattern E could be differentiated by phage typing but not by rRNA gene restriction patterns, indicating the complementary nature of the two methods.

The specificity of the DNA probe compares well with that using rRNA since the pattern of *S. fonticola*-DNA is identical to that described [11]. In addition, rDNA patterns in *Aeromonas* [9] are also identical regardless of the protocol used (unpublished data). Therefore, the presence of the entire rRNA operon (plasmid insert) and the two antibiotic-resistant genes (vector pBR322) do not seem to cause any problems under the relatively stringent conditions used for hybridization.

The method described for the typing of *S. typhi*

strains offers an increased sensitivity over phage typing. It will be especially useful to analyze epidemics caused by common phage types and for the typing of strains that change phage type due to the loss or uptake of a temperate phage. The option of using the method on a nonradioactive basis (commercial kits for biotinylation of DNA and detection of hybrids are available) makes this method especially promising for use in nonreference laboratories and in developing countries.

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