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# Restriction Fragment Length Polymorphism Analysis of Some Flagellin Genes of Salmonella enterica

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Salmonellae often have the ability to express two different flagellar antigen specificities (phase 1 and phase 2). At the cell level, only one flagellar phase is expressed at a time. Two genes, fliC, encoding phase-1 flagellin, and fljB, encoding phase-2 flagellin, are alternatively expressed. Flagellin genes from 264 serovars of Salmonella enterica were amplified by two phase-specific PCR systems. Amplification products were subjected to restriction fragment length polymorphism (RFLP) analysis by using endonucleases HhaI and HphI. RFLP with HhaI and HphI yielded 64 and 42 different restriction profiles, respectively, among 329 flagellin genes coding for 26 antigens. The phase-1 gene showed 46 patterns with HhaI and 30 patterns with HphI. The phase-2 gene showed 23 patterns with HhaI and 17 patterns with HphI. When the data from both enzymes were combined, 116 patterns were obtained: 74 for fliC, 47 for fljB, and 5 shared by both genes. Of these combined patterns, 80% were specifically associated with one flagellar antigen and 20% were associated with more than one antigen. Each flagellar antigen was divided into 2 to 18 different combined patterns. In the sample of strains used, determination of the phase-1 and phase-2 flagellin gene RFLP, added to the knowledge of the O antigen, allowed identification of all diphasic serovars. Overall, the diversity uncovered by flagellin gene RFLP did not precisely match that evidenced by flagellar agglutination.

In developed countries, salmonellosis is a major economic problem for the food industry, as well as a public health hazard for the consumer. In developing countries, the death toll from salmonellosis (typhoid and diarrhea in children) is very high. Individualization of strains of the pathogen is essential to the study of the association between clinical cases and possible sources of infection.

The genus *Salmonella* is composed of two species, "*Salmonella enterica*" (quotes indicate pending nomenclatural status) and *S. bongori* (11, 17). The primary basis for the typing of "*S. enterica*" is a serotyping scheme (the White-Kauffmann-Le Minor [WKL] scheme) in which 2,375 serovars have been recognized on the basis of the antigenic properties of the cell wall lipopolysaccharide (O antigen), the phase-1 flagellar protein (H1), and the phase-2 flagellar protein (H2) (15, 16).

The flagellar protein or flagellin constitutes the subunit of the helical filament that forms the flagellar organelle. *Salmonella* flagellin consists of extremely conserved terminal regions and a variable central region (7, 23). This central region of the molecule carries the antigenic specificity (14). For the phase-1 flagellin, 63 antigens have been distinguished. For the phase-2 flagellin, 37 antigens have been described. Some of these antigens are defined by a single factor (antigen i, d, or r); others are defined by several subfactors (e.g., antigens l,v; l,w; g,m; and e,n,x) (15).

The antigenic specificities of phase-1 and phase-2 flagellins are encoded by flagellin genes fliC and fljB, respectively. These flagellar genes are found at two different locations on the chromosome. At one location is the gene fliC. At another location is an operon containing the genes hin, encoding the Hin recombinase; fljB, encoding phase-2 flagellin; and fljA, encoding a repressor for fliC. The Hin recombinase catalyzes the revers-

ible inversion of a 993-bp segment of the chromosome containing a promoter. In one orientation, the promoter directs transcription of the *fljB* and *fljA* genes. Phase-2 flagellin and the repressor are produced (thus repressing *fliC*). In the other orientation, repression of the *fliC* gene is relieved and phase-1 flagellin is expressed (6, 25).

Salmonella isolates expressing two antigenically distinct types of flagellin are biphasic. Monophasic Salmonella strains expressing only one type of flagellar antigen include many clinically and epidemiologically important salmonellae, e.g., serovar Typhi, the agent of typhoid fever, and serovar Enteritidis, a major foodborne pathogen associated with poultry and eggs (19). One serovar, Gallinarum, is always nonflagellated (12). Occasionally, nonmotile (nonflagellated) variants of normally motile serovars are isolated from specimens. These isolates cannot be identified with a known serovar by serotyping.

Molecular techniques such as restriction fragment length polymorphism (RFLP) could reflect the flagellar antigenic diversity of salmonellae at the genetic level (9). The purposes of this study were to determine whether (i) 26 flagellar antigens (carried by 237 serovars) could be differentiated by flagellin gene RFLP, (ii) genes coding for phase-1 and phase-2 antigens with the same designation could have identical RFLP patterns, (iii) flagellar antigens could be subdivided into RFLP patterns, and (iv) serovars could be identified by using flagellin gene RFLP. The results obtained showed these purposes to have been partially achieved.

#### MATERIALS AND METHODS

Collection of strains. The 237 reference strains from different serovars of *S. enterica* subsp. enterica (subsp. 1) and salamae (subsp. II) used in this study were from the World Health Organization Collaborating Center for Reference and Research on Salmonella (from M. Y. Popoff, Institut Pasteur, Paris, France). We also studied 27 strains received at the Centre National de Reference des Salmonella et Shigella or the Centre National de Reference pour le Typage Moleculaire Enterique (both centers are located in the Unité des Entérobactéries, Institut Pasteur). These included 7 isolates of serovar Typhi representing different ribotypes; 11 isolates of serovar Typhimurium, corresponding to six phage types (12 atypical, 29, 113, 114 atypical, 120, and 153); 1 isolate of serovar

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TABLE 1.	Flagellar	antigens	represented	in	this	study

Flagellar	No. of	No. of strains				
antigen(s) <sup>a</sup>	serovars	Subsp. I	Subsp. II			
Phase 1						
a	1	1				
b	2	1	1			
c	1	1				
d	22	24	1			
j	1	5				
e,h	3	3				
e,n,x	1		1			
g series	49	36	13			
i	25	36				
r	8	8				
r,i	8	9				
l,v	51	51				
l,w	17	17				
Z	1	1				
z6	1		1			
z10	1	1				
z35	1	1				
z39	1		1			
z42	1		1			
Phase 2						
l,w	70	70				
1,2	8	8				
1,5	17	17				
1,6	6	6				
1,7	7	7				
1,2,7	1	1				
e,n,x	6	6				
e,n,x,z15	1	1				
e,n,z15	10	10				
z6	6	6				
z35	2	2				
Total: 26	237	245	19			

<sup>&</sup>lt;sup>a</sup> Selected antigens are in boldface.

Bovismorbificans; and 7 nonmotile isolates. Biochemical confirmation and serotyping were done by conventional methods.

Serovars were selected to include (i) most frequently encountered phase-1 flagellar antigens i, r, and the g series (f, g, m, p, q, s, t, u, and z51); (ii) most frequently encountered phase-2 flagellar antigens 1,2 and 1,5; (iii) flagellar antigen d, which is associated with medically important serovars; and (iv) flagellar antigens l,v and l,w, which are found in both phase 1 and phase 2. Since most strains were diphasic, a number of flagellar antigens other than the selected ones were de facto included in the study. Strains of *S. enterica* subsp. II were included in the study because they share the selected antigens with *S. enterica* subsp. I. Flagellar antigens are listed in Table 1.

Preparation of DNA. Isolates were grown in Trypto casein soy agar (Sanofi Diagnostics Pasteur, Marnes-la-Coquette, France). A single colony was grown in a shaking incubator for 18 h at 37°C in Trypto casein soy broth (Sanofi Diagnostics Pasteur). The culture was centrifuged at 10,000 rpm for 10 min. The pellet was suspended in 580 μl of lysis buffer (Tris-HCl at 0.05 M, EDTA at 0.05 M, NaCl at 0.1 M, pH 8) with 3 μl of a 20-mg/ml aqueous solution of pronase (Calbiochem, La Jolla, Calif.) and 32 μl of a 10% (wt/vol) sodium dodecyl sulfate solution and incubated for 1 h at 60°C to allow cell lysis. DNA was extracted with an AutoGen 540 automated DNA extraction system (AutoGen Instruments, Beverly, Mass.).

PCR amplification of fliC (phase 1). For amplification of the phase-1 flagellin gene, the primers used were CAAGTCATTAATACMAACAGCC (FSa1; M = A or C) and TTAACGCAGTAAAGAGAGGAC (rFSa1). Primer FSa1 was selected on the basis of fliC gene conservation among sequences of Salmonella, Escherichia coli, and Shigella flagellin genes (GenBank and EMBL accession no. M23773, M84972, M84973, M23772, M23774, X04505, M11332, M84976, M84978, M84979, Z15064, Z15065, Z15066, Z15069, Z15086, Z15070, Z15071, Z15072, L21912, L07387, D18821, and D16819). Primer rFSa1 was selected to amplify only the Salmonella fliC gene. The target of primer FSa1 was located at positions 18 to 40, and the target of primer rFSa1 was located at positions 1530 to 1510 of the serovar Muenchen fliC gene (accession no. M23774). The expected size of the amplified fragment was about 1.5 kbp, except for the H1-j flagellin gene of variant serovar Typhi, which contained a deletion of 261 bp (5).

DNA amplification by PCR was performed in a reaction volume of 100  $\mu l$  consisting of 10 mM Tris-HCl (pH 8.3); 50 mM KCl; 1.5 mM MgCl $_2$ , 0.01% (wt/vol) gelatin; 2.5 U of Hi-Taq DNA polymerase (Bioprobe, Montreuil-sous-Bois, France); 200  $\mu M$  each dATP, dTTP, dCTP, and dGTP; 50 pmol of each primer; and 1  $\mu l$  of sample DNA. The reaction mixture was overlaid with 50  $\mu l$  of mineral oil. Initial denaturation was carried out for 5 min at 94°C. Thirty-five cycles of amplification were performed in a PTC-100 thermal cycler (MJ Research, Watertown, Mass.). Each cycle consisted of three steps: denaturation for 1 min at 94°C, annealing for 1 min at 55°C, and extension for 1 min at 72°C. An additional step of extension for 5 min at 72°C was performed at the end of the amplification to complete extension of the primers. Amplification products were detected by electrophoresis in 0.8% (wt/vol) agarose gels in Tris-acetate buffer (0.04 M Tris-acetate, 0.002 M EDTA, pH 8.1), with the 1-kbp DNA Ladder (Gibco BRL, Gaithersburg, Md.) as a molecular size marker.

PCR amplification of *fljB* (phase 2). The primers designed for amplification of the phase-2 flagellin gene were CAAGTAATCAACACTAACAGTC (FSa2) and TTAACGTAACAGAGACAGCAC (rFSa2). The target of primer FSa2 was located at positions 7 to 28, and the target of primer rFSa2 was located at positions 1506 to 1486 of the serovar Abortusequi *fljB* gene (accession no. D13690). This PCR will be referred to as *fljB* amplification.

Because the GenBank and EMBL international databases contained only one fliB gene sequence, primer selectivity was assessed by the following procedure. A first amplification of DNA from serovars Abortusequi, Bloomsbury, Rubislaw, Typhimurium, Goldcoast, Anatum, Brandenburg, and Verona was done with primers ST-HIN-L and SA-FLJA-R (2). These primers selectively amplified a part of the Salmonella flagellar operon consisting of the hin, fljB, and fljA genes. This PCR will be referred to as hin-fljB-fljA amplification. The reaction required extracted DNA from bacteria expressing the phase-2 flagellar antigen. Amplification involved predenaturation at 94°C for 5 min, followed by 35 cycles of 94°C for 1 min, 47°C for 1 min, and 72°C for 1 min, with a final step of elongation at 72°C for 5 min. The 2,976-bp expected fragment was extracted from a 0.8% (wt/vol) agarose gel by using the JETsorb kit (Bioprobe) prior to fljB amplification (with primers FSa2 and rFSa2) to eliminate genomic DNA from the PCR mixture. The specific PCR with primers FSa2 and rFSa2 was performed by 35 cycles of 94°C for 1 min, 58°C for 1 min, and 72°C for 1 min. For elongation of the PCR product, a final step of 72°C for 5 min was included. The amplified products were detected by electrophoresis as described above.

fljB amplification was also done directly on the same strain DNAs with the same PCR parameters. The identities of amplified fragments were determined by RFLP as described below. After this primer specificity control, fljB amplification was done directly on Salmonella genomic DNA.

RFLP analysis. Endonucleases *Hha*I and *Hph*I were chosen after study of restriction maps of eight sequences of *fliC* and *fljB* from the EMBL and GenBank databases (accession no. M23774, M84973, D13690, Z15068, M23772, X04505, M11332, and L21912). Maps were obtained with the Mapdraw program of the Lasergene software (DNAstar, Madison, Wis.). The enzyme *Hha*I recognized and cleaved the sequence GCGC, while *Hph*I recognized the sequence TCACC or GGTGA and cleaved the sequence 8 or 9 bp further (10, 18). *Hha*I restriction sites were regularly distributed on the flagellin sequences, while *Hph*I preferentially cleaved the genes in the hypervariable region.

In a microtube, 10-µl portions of PCR mixtures containing amplified flagellin genes were digested. Digestion was done for 2 h at 37°C for both restriction enzymes. RFLPs were determined by electrophoresis of the digested DNA in 1% (wt/vol) agarose (Bioprobe) plus 1% (wt/vol) Nusieve agarose (FMC Bioproducts, Rockland, Maine) gels for 5 h at 4.8 V/cm. The 1-kbp DNA Ladder (Gibco BRL) was used as a molecular size marker. The restricted fragments were stained with ethidium bromide.

The RFLP patterns were scanned by using One-Scanner (Apple Computers, Cupertino, Calif.). Digitization and interpretation of RFLP profiles were done with the Taxotron package (Taxolab software; Institut Pasteur), including the programs RestrictoScan, RestrictoTyper, Adanson, and Dendrograf. Lanes and bands of the resulting TIFF images were detected with RestrictoScan. Fragment lengths were interpolated by using the Schaffer and Sederoff algorithm (20) implemented by RestrictoTyper. The program generated a normalized graph showing migration patterns. Fragments were considered identical if their sizes did not differ by more than 1% (percent tolerated error). The distance coefficient was calculated as the number of nonmatching fragments divided by the total number of bands in both patterns (complement of the Dice index [3]), and a distance matrix was built. The relationships between patterns were calculated by the average linkage (1), single linkage, and unweighted pair group using mathematical averages (UPGMA) (21) methods with the Adanson clustering program. Dendrograms were drawn by Dendrograf.

## RESULTS

**PCR amplification of** *fliC* **(phase 1).** A 1.5-kbp fragment was amplified from 259 strains and a 1.24-kbp fragment was amplified from five strains of variant serovar Typhi (H1:j), as expected. No other variation in *fliC* gene size was detectable on electrophoresis gels. Amplification occurred with strains from

nonmotile serovar Gallinarum, as well as other nonmotile isolates.

PCR amplification of fljB (phase 2). hin-fljB-fljA amplification generated 3-kbp amplification products from the eight serovars tested (Abortusequi, Bloomsbury, Rubislaw, Typhimurium, Goldcoast, Anatum, Brandenburg, and Verona). fljB amplification of total DNA and of hin-fljB-fljA amplification products yielded 1.5-kbp products. The fljB specificity of fljB PCR was demonstrated by HhaI and HphI restriction of fljB amplification products obtained from total DNA or from hin-fljB-fljA amplification products. For each strain tested, restriction profiles were identical in both cases (data not shown).

Thereafter, the specific fljB PCR was applied directly to Salmonella genomic DNA. For all of the diphasic serovars tested, the amplification product was invariably 1.5 kbp when amplification occurred. For the diphasic serovar Typhi strains tested (d:z66 and j:z66), fljB amplification failed to amplify a fragment. No fljB amplification was shown with the monophasic serovars Typhi, Paratyphi A, Enteritidis, Derby, Rissen, Agona, Borreze, Havana, Berta, Antarctica, Ona, Kingston, California, Congo, Giessen, Emek, Budapest, Dublin, Sylvania, Naestved, Essen, Gallinarum, Montevideo, Blegdam, Othmarshen, Rostock, Moscow, Senftenberg, Banana, Oranienburg, Hillingdon, Gateshead, Sangalkam, Ackwepe, and Keve.

**RFLP analysis of flagellin genes.** Restriction enzymes *Hha*I and *Hph*I were used on PCR products from the *fliC* and *fljB* genes of each of the 264 *Salmonella* strains studied and yielded profiles consisting of two to seven fragments sized between 62 and 1,310 bp (Fig. 1 and 2).

On repeated experiments, 42 strains tested with *Hha*I and 45 strains tested with *Hph*I could be unambiguously assigned to the same pattern as that generated by the first experiment.

To ensure that no partial restriction occurred with the enzymes used, summation of fragment lengths for each profile was done. Surprisingly, the sum of the flagellin fragment lengths varied in accordance with the profiles studied and was often smaller than the size of the PCR product used for restriction. In studying profiles obtained from restriction maps of eight published sequences (accession no. M11332, M84973, Z15068, X03395-M23774, X03393-M23772, X04505, L21912, and D13690), we observed that only fragments larger than 60 bp were visible on the agarose gel. Product size, deduced from the sum of restriction fragment lengths, was reduced mostly with *Hha*I and slightly with *Hph*I (Table 2). In these cases, part of the difference between the size of the PCR product and the sum of the restriction fragment lengths was due to fragments smaller than 60 bp. In comparing fragment lengths (greater than 60 bp) obtained from restriction maps of published sequences and those deduced from migration distances on electrophoresis gels for the same genes, two situations occurred (Table 2). In situation 1, five *HhaI* profiles and five *HphI* profiles gave the same number of fragments as the profiles deduced from restriction maps. Differences between the sums of fragment lengths from restriction maps and the experimental gel estimates ranged from 0.8 to 3.1% of the total size (experimental error). In situation 2, four HhaI profiles and three HphI profiles included fragments of equal sizes which appeared as single bands on agarose gels (comigrating fragments). Thus, for such profiles, the difference between sums of fragment lengths from restriction maps and experimental gelbased estimates ranged from 4.3 to 12.9% of the total size (comigration bias).

Correspondence between patterns and antigens. Restriction of 329 (195 phase-1 plus 134 phase-2) flagellin genes yielded 64 *Hha*I profiles and 42 *Hph*I profiles (Table 3). Patterns were designated by a letter indicating the restriction enzyme used (A

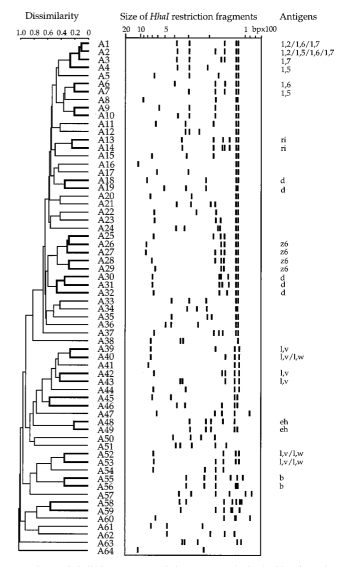


FIG. 1. Dissimilarity among restriction patterns obtained with *HhaI*. The dendrogram was generated by the UPGMA method. Clusters obtained with three clustering methods are indicated with thicker lines (robust clusters). Each branch of the tree faces each flagellin banding pattern. Antigenic specificities are indicated when typical of a cluster.

for *Hha*I or P for *Hph*I) followed by a number (Fig. 1 and 2). Phase-1 genes showed 46 patterns with *Hha*I and 30 patterns with *Hph*I. Phase-2 genes showed 23 patterns with *Hha*I and 17 patterns with *Hph*I. Forty-one *Hha*I patterns and 25 *Hph*I patterns were only associated with the *fliC* gene in this study. Eighteen *Hha*I patterns and 12 *Hph*I patterns were only associated with the *fljB* gene. Five patterns obtained with *Hha*I (A15, A39, A40, A52, and A53) and five obtained with *Hph*I (P1, P2, P8, P24, and P42) were associated with both the *fliC* and *fljB* genes.

Fifty *Hha*I and 20 *Hph*I patterns were each specifically associated with a single antigen. For example, the *Hha*I patterns found were A42 for the l,v flagellar antigen, A19 and A31 for d, A25 for i, A34 for j, and A54 for phase-2 l,w antigens; the *Hph*I patterns found were P38 and P41 for i, P17 for j, P18 for g,z51, and P4 and P5 for phase-2 l,w antigens. The other patterns were each associated with more than one antigen. Fig-

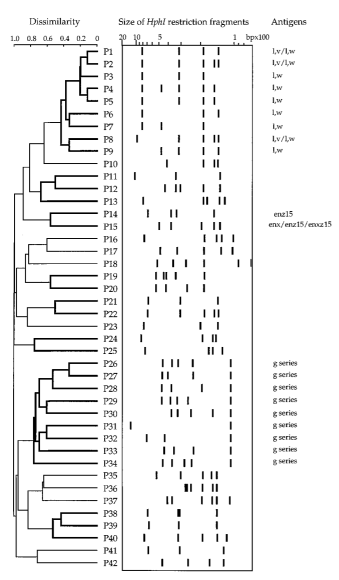


FIG. 2. Dissimilarity among restriction patterns obtained with *Hph*I. The dendrogram was generated by the UPGMA method. Clusters obtained with three clustering methods are indicated with thicker lines (robust clusters). Each branch of the tree faces each flagellin banding pattern. Antigenic specificities are indicated when typical of a cluster.

ures 1 and 2 show results of cluster analyses of restriction patterns. Dendrograms produced clusters of similar restriction patterns. It is striking that some clusters corresponded to related flagellar antigens.

In combining the data obtained with both restriction enzymes, 116 combined patterns (74 combined patterns for *fliC*, 47 combined patterns for *fljB*, and 5 combined patterns shared by both genes) were identified among the 26 antigens examined (195 phase-1 and 134 phase-2 genes were tested). *HhaI* divided 21 *HphI* patterns into 92 combined patterns. *HphI* divided 16 *HhaI* patterns into 66 combined patterns. Two to 18 different combined patterns were observed for each of the following flagellar antigens: d; e,n,x; e,n,z15; i; r,i; l,v; l,w; z6; 1,2; 1,5; 1,6; and 1,7 (Tables 3 and 4). The restriction profiles of the *fliC* and *fljB* genes revealed a molecular diversity greater than the antigenic diversity.

When 14 antigens from 19 serovars of subsp. II were tested,

gels for the same sequences FABLE 2. Comparison of fragment sizes from restriction maps and fragment sizes interpolated from migration distances on

				Map-, gel-bas	Map-, gel-based sizes $(bp)^a$			
Enzyme	Dublin g,p antigen	Enteritidis g,m antigen	Rubislaw r antigen	Muenchen d antigen	Paratyphi A a antigen	Typhimurium i antigen	Typhi d antigen	Abortusequi e,n,x antigen
Hhal	706, 681 431, 425 115, 122 104, 117	706, 707 431, 441 115, 125 104, 119	676, 672 306, 317 174, 184 115, 122 104, 117	709, 705 309, 315 174, 179 115, 122 104, 118	484, 484 195, 199 154, 151 146, — <sup>b</sup> 113, 125 104, 118	667, 683 306, 309 174, 122 104, 118	508, 520 309, 320 204, 218 115, 122 113, —	399, 417 273, 289 174, 182 115, 122 113, —
Sums (SD [%])	1,356, 1,345 (0.8)	1,356, 1,392 (2.5)	1,375, 1,412 (2.6)	1,411, 1,439 (1.9)	1,196, 1,063 (12.5)	1,251, 1,232 (1.5)	1,353, 1,297 (4.3)	1,178, 1,127 (4.5)
<i>Нрh</i> I	463, 465 365, 373 318, 324 224, 239 93, 110	463, 457 365, 368 318, 323 224, 240 93, 110	831, 806 195, 199 186, 188 147, 151 130, 135	756, 756 186, 194 186, — 147, 154 130, 139 107, 112	830, 817 194, 193 186, — 147, 161 130, 146	706, 698 333, 332 306, 316 130, 144	759, 771 333, 337 183, 196 130, 138 107, 112	488, 498 363, 379 192, 201 186, — 147, 156 130, 138
Sums (SD [%])	1,463, 1,511 (3.1)	1,463, 1,498 (2.3)	1,489, 1,479 (0.7)	1,512, 1,355 (11.6)	1,487, 1,317 (12.9)	1,475, 1,490 (1)	1,512, 1,554 (2.7)	1,506, 1,372 (9.8)
a The vertex of The	a The motivation of the factor of the section of the	40.						

The values in boldface are significantly high —, band not seen.

TABLE 3. Diversity of flagellin gene restriction patterns obtained with *Hha*I and *Hph*I

Flagellar antigen(s)		patterns n by:	No. of combined patterns given by	n by No. of	
antigen(s)	HhaI	HphI	HhaI + HphI		
H:1					
a	1	1	1	1	
b	2	2	2	2	
c	1	1	1	1	
d	8	6	12	22	
j	1	2	2	1	
e,h	2	1	2	3	
e,n,x	1	1	1	1	
g series	8	12	17	49	
i	9	7	13	25	
r	4	2	4	8	
r,i	5	7	8	8	
1,v	8	4	11	51	
l,w	2	3	5	17	
Z	1	1	1	1	
z6	1	1	1	1	
z10	1	1	1	1	
z35	1	1	1	1	
z39	1	1	1	1	
z42	1	1	1	1	
Total	46	30	74	195	
H:2					
l,w	4	10	18	70	
1,2	2	3	3	8	
1,5	8	4	10	17	
1,6	3	3	4	6	
1,7	4	3	5	7	
1,2,7	1	1	1	1	
e,n,x	1	3	3	6	
e,n,z15	3	4	5	10	
e,n,x,z15	1	1	1	1	
z6	5	4	5	6	
z35	1	1	1	2	
Total	23	17	47	134	

13 different combined patterns were obtained (Table 4). Of these combined patterns, 12 were specifically associated with subsp. II. The exception was a serovar of subsp. II (O30; g,m,s; e,n,x) which shared *fliC* pattern A45P26 with serovars Giessen and Emek of subsp. I (Table 4).

Of the 116 combined patterns, 93 (80%) were each specifically associated with a single antigen and 23 (20%) each corresponded to several antigens (Table 4). For 71 combined patterns associated with one antigen (among the antigens tested), one enzyme provided specificity. In the 22 other combined patterns associated with only one antigen, specificity was given by the combination. The combination increased the number of specific RFLPs (combined patterns) associated specifically with one antigen but did not differentiate all of the antigens. Of the 13, 4, 8, and 12 combined patterns obtained for the i; r; r,i; and d antigens, 5 corresponded to several antigens (Table 4). Pattern A15P40 corresponded to both the d and r,i antigens. Pattern A44P22 corresponded to both the i and r,i antigens. Patterns A8P13 and A44P13 corresponded to both the i and r antigens.

When antigens shared antigenic subfactors, RFLP did not differentiate among them. Phase-2 antigens 1,2; 1,5; 1,6; 1,7; and 1,2,7 yielded four combined patterns (A1P1, A1P24, A2P3,

and A15P1) without correspondence between antigenic factors and patterns. Antigens e,n,x and e,n,z15 shared three combined patterns (A10P12, A10P15, and A10P19). The sensitivity of the RFLP method for recognition of the g series of antigens was as high as 97.9%. Only one strain tested (serovar Borreze) shared pattern A44P13 with other antigens (r and i). However, the discrimination power of the RFLP method within the g series was very low. For 19 different antigens, only 17 combined patterns were obtained. Each of the combined patterns A16P33, A36P34, A45P26, A45P32, and A45P33 corresponded to two to eight antigens of the g series. For l,v and l,w antigens, 22 of 23 combined patterns corresponded to both antigens. Pattern A39P24 was shared by the l,v (serovar Coromandel) and z35 antigens. Of the 23 combined patterns obtained, only 6 seemed to be specifically associated with the phase-1 l,v antigen flagellin gene (A40P2, A41P2, A42P2, A43P1, A44P8, and A53P24) and 13 were specifically associated with the phase-2 l,w antigen flagellin gene (A40P24, A52P3, A52P4, A52P5, A52P6, A52P8, A52P9, A53P3, A53P4, A53P5, A53P7, A54P1, and A54P7). Some genes encoding l,v or l,w antigens shared the same comined patterns (A52P1, A52P2, A52P24, and A53P2). All of the combined patterns of genes encoding phase-1 l,w antigen (A52P1, A52P2, A52P24, A53P1, and A53P2) were found for phase-2 l,w antigen. RFLP analysis showed that phase-1 and phase-2 genes sharing l,w antigens were not differentiated by the endonucleases used.

Correspondence between patterns and serovars. The discrimination power of flagellin gene RFLP analysis alone was insufficient to distinguish all of the serovars tested. Among the 237 serovars studied, 112 combined patterns were each assigned to 1 serovar (Table 4). Each combined pattern was given by 1 to 22 serovars. For 51 serovars, combined patterns of the flagellin phase-1 gene were serovar specific. For 28 serovars, specificity involved only combined patterns from the fljB gene. For 33 serovars, combined patterns obtained from both genes were required. The specificity of the fliC gene combined pattern was given by HhaI for 27 serovars, by HphI for 11 serovars, and by both enzymes for 13 serovars. The specificity of the *fljB* gene combined pattern was given by *Hha*I for 12 serovars, by HphI for 7 serovars, and by both enzymes for 9 serovars. For r,i antigens, the discrimination of RFLP was so high that each serovar studied had its own specific combined pattern.

Specific patterns characterized some important serovars. Regular serovar Typhi (H1:d) showed specific pattern A19 with *Hha*I. Variant serovar Typhi (H1:j) showed specific patterns A34 with *Hha*I and P17 with *Hph*I. This result was tested on nine strains, seven of which showed different ribotypes. Strains from serovar Typhi harbored three types of specific combined patterns: A19P40 (regular Typhi), A34P17, and A34P22 (variant Typhi).

Serovar Typhimurium showed specific pattern P38 (H1:i) with *Hph*I. This result was confirmed for 11 strains, 6 of which had different phage types. Strains from serovar Typhimurium consistently showed combined pattern A44P38-A1P24 (phase-1 and phase-2 genes).

Some other epidemiologically important serovars were also differentiated (phase-1/phase-2 genes): Hadar (A46P35-A10P19), Heidelberg (A44P13-A1P24), Indiana (A60P10-A3P1), Newport (A49P24-A2P7), Choleraesuis (A47P24-A2P1), Saint-Paul (A48P24-A1P1), Goldcoast (A44P13-A52P24), Paratyphi A (A62P24/-), and Bovismorbificans H1: r,i (A15P13-A33P24).

Thus, among 170 strains studied in both phases, 134 strains (112 serovars) could be identified at the serovar level with the flagellin gene combined patterns. Lack of pattern specificity was observed for 36 serovars. For 25 serovars, the phase-2 gene

TABLE 4. Combined patterns and corresponding serovars

TABLE 4—Continued

	Anti	gen(s)			Anti	gen(s)	
Pattern	Phase 1	Phase 2	Serovar(s)	Pattern	Phase 1	Phase 2	Serovar(s)
A1P1		1,2	Saint Paul	A16P28	f,g,t		1,4,12:f,g,t:z6,z42 <sup>a</sup>
		1,7	Schwarzengrund, Banco	A16P30	f,g,t		Eingedi
1P24		1,2 1,6	Typhimurium, Heidelberg Quentin	A16P31	g,s,t		Gateshead
.2P1		1,7 1,5	Strasbourg, Dieuppeul Choleraesuis	A16P33	f,g f,g,s		Derby, Rissen, Havana Agona
2P3		1,6	Agama, Togo, Westeinde		f,g,t g,s,t		Berta Ona, Kingston Bloomsbury, Budapest, Senftenberg
2P7		1,7 1,2	Lika Newport, Muenchen, Stanley, Gabon,	A17P23	g,t i		Potto
			Overchurch	A18P1	d		Strasbourg, Olten, Plymouth
.2P12		e,n,z15	Tsevie	A18P25	d		Quentin
.2P20		1,6	Anatum	A18P40	d		Ontario
2P24		1,5	Fulda	A19P40	d		Typhi
.3P1		1,7	Indiana	A20P22	i		Hoboken
A4P24		1,5	Mono	A21P1		1,7	Truro
5P13	r		Brive	A22P39	z35		Pasing
A5P22	i		Douala, Bandia	A23P17	d		6,7:d:z42 <sup>a</sup>
.6P20		1,6	Dumfries	A24P35	g,m,s,t		Congo
7P24		1,5	Ontario	A25P16	i		Soerenga, Verona
.8P2		z35	Coromandel	A26P1		z6	Potto
.8P13	r i		Massakory Veneziana	A27P42		z6	Kentucky
9P3		1,5	Colorado	A28P1		z6	Tumodi
.10P12		e,n,x	Veneziana, Rubislaw	A29P24		z6	Assine, Plymouth
		e,n,z15	Olten, Toulon	A30P16	d		Tilburg
A10P15		e,n,x e,n,z15	Farsta, Brooklyn, Abortusequi Duisburg, Avonmouth, Surat, Branden-	A31P16	d		Livingstone, Mons, Putten, Cullingworth Dembe
		e,n,x,z15	burg Euston	A31P40	d		Birmingham
10P19		e,n,x	Hadar	A32P16	d		Niamey
		e,n,z15	Moussoro, Lomnava	A33P24		1,5	Bovismorbificans
11P21	e,n,x		9,12:e,n,x:-a	A34P17	j		Typhi
12P11		1,5	Lagos	A34P22	j		Typhi
.13P24	r,i		Euston	A35P1		1,5	Bloomsbury
14P23	r,i		Fareham	A36P27	m,t		Sangalkam
15P1		1,5	Alamo, Maricopa, Bovismorbificans, Victoria	A36P34	g,m,t m,t		California Banana, Oranienburg
45D12		1,2,7	Eingedi	A37P35	d		Isangi
.15P12		z6	Verona	A38P41	i		Agama, Farsta, Tsevie
.15P13	r,i		Bovismorbificans	A39P24	1,v		Coromandel
.15P16	d		Muenchen, Stanley, Duisburg, Manhattan, Caen			z35	Kotte
	i r,i		Moussoro, Dieuppeul Drogana	A40P2	l,v		India, Toronto
.15P24		1,5	Isangi, Manhattan, Gligji, Zigong, Pasing	A40P24		l,w	Brikama, Soerenga
15P40	d <sub>.</sub>		Schwarzengrund	A41P2	l,v		Sinchew
	r,i		Dumfries	A42P2	l,v		Nchanga, Horsham, Geraldton
A15P41	i		Lagos, Tumodie	A43P1	1,v		Burgas

A52P5

A52P6

Zuilen

Cullingworth

l,w

1,w

TABLE 4—Continued

TABLE 4—Continued

TABLE 4—Continued			TABLE 4—Continued					
Pattern Antigen(s)			Serovar(s)	Pattern		gen(s)	Serovar(s)	
	Phase 1	Phase 2			Phase 1	Phase 2		
A44P1	i		Avonmouth	A52P8		l,w	Cyprus	
A44P8	l,v		Sinstorf	A52P9		l,w	Alexanderpolder	
A44P13	i r		Lika, Kedougou Bovismorbificans, Heidelberg, Goldcoast,	A52P10	r		Surat	
	f,g,s		Grampian, Bochum Borreze	A52P24	1,v 1,w		Irumu, Bonn, Clackamas, Concord Mono, Togo, Colorado, Toulon, Over- church	
A44P22	i		Kentucky, Gloucester, Norton, Magherafelt, Pisa, Kaneshie			l,w	Bruck, Wyldegreen, Goldcoast, Abidjan	
	r,i		Zuilen	A53P1	l,w	l,w	Ackwepe Hallflod, Ohio, Bukuru, Langensalza,	
A44P23	r,i		Africana			2,11	Broughton, Niamey, Douala, Fairfield, Moroto, Dembe, Massakory, Niarembe	
A44P38	i		Typhimurium	A53P2	l,v		Mkamba, Kortrijk, Shanghai, Salford	
A45P26	g,m g,p		Essen, Enteritidis, Gallinarum, Othmarshen Dublin, Sylvania	A331 2	1,w	1	Kewe, Ramsey Mannheim	
	g,p,u		Rostock			l,w		
	g,m,p,s g,q		Montevideo Moscow	A53P3		l,w	Kedougou	
	g,m,q g,m,s g,p,s		Blegdam 30:g,m,s:e,n,x <sup>a</sup> , Giessen, Emek Naestved	A53P4		l,w	Wien, Mura, Bochum, Nantes, Birming- ham, Hoboken, Alkmaar, Carno, Kaneshie	
A45P28	g,m,s,t		1,13,23:g,m,s,t:1,5 <sup>a</sup>	A53P5		1,w	Grampian	
A45P29	g,z63		Antarctica	A53P7		l,w	Kuru, Epicartes, Newrochelle, Putten, Vridi, Demerara, Bandia	
A45P32	g,m g,m,s		Hillingdon Macclesfield	A53P24	l,v		Potsdam, Pakistan, Litchfield	
A45P33	f,g,m,t		6,8:f,g,m,t:- <i>a</i>	A54P1		1,w	Huettwillen, Mons, Preston	
	g,m,s g,m,s,t		40:g,m,s:e,n,x <sup>a</sup> 9,12:g,s,t:e,n,x <sup>a</sup> ; 1,9,12:g,m,s,t:e,n,x <sup>a</sup>	A54P7		l,w	Coleypark	
	g,s,t		$28:g,s,t:e,n,x^a$	A55P36	b		Kotte	
A45P34	g,m,t		4,12:g,m,t:z39 <sup>a</sup>	A56P42	b		6,7:b:z39 <sup>a</sup>	
A46P35	z10		Hadar	A57P24	z6		6,7:z6:1,7 <sup>a</sup>	
A47P24	c		Choleraesuis	A58P13	i		Truro	
A48P24	e,h		Saint Paul	A58P24	z39		6,7:z39:1,5,7 <sup>a</sup>	
A49P24	e,h		Newport, Anatum	A59P24	z42		6,7:z42:1,7 <sup>a</sup>	
A50P14		e,n,z15	Drogana	A60P10	z		Indiana	
A51P1	r,i		Banco	A61P33	g,t		52:g,t:-a; 3,10:g,t:-a	
A52P1	l,v		Koessen, Fyris, Azteca, Give, Gdansk, Zaiman, Goettingen, Svedvi, Fann	A62P24	a		Paratyphi A	
	l,w	1,w	Victoria, Assine Sindelfingen, Meleagridis, Tilburg, Calabar,	A63P18	g,z51		Alamo, Maricopa	
		1,W	Ordonez, Worthington, Minna, Vegesack, Yoruba, Caen, Fareham, Brive, Shannon	A64P33	g,z62		1,9,12:g,z62:-a; 4,12:g,z62:-a	
A52P2	l,v	l,w	Kimuenza, Loanda, Amherstiana, Manchester, Holcomb, Edmonton, Mendoza, Kapemba, London, Brandenburg, Bredeney, Ruzizi, Ngor, Parkroyal, Maracaibo, Bullbay, Lovelace, Borbeck, Nanga, Boecker Welikade, Los Angeles Gabon, Fulda, Gligji, Zigong, Westeinde, Brooklyn, Lomnava, Africana, Norton, Hillsborough, Magherafelt, Dabou, Bamboye, Anderlecht, Livingstone, Uhlenhorst	was mis phase-1 for sero and bot	was missing (monophasic strains) and for 11 diphasic serove phase-1 and phase-2 gene combined patterns were insufficitor serotype identification. The knowledge of the O antique and both flagellin gene RFLPs contributed to the identification.			
A52P3		l,w	Clerkenwell	of these 11 diphasic serovars (in the sample of serovars studied). For example, serovars Brooklyn and Brandenburg share the A52P2-A10P15 flagellin gene combined pattern but hat the O16 and O4 antigens, respectively.				
A52P4		l,w	Gloucester, Krefeld					

he identification of serovars studndenburg shared pattern but had the O16 and O4 antigens, respectively.

Nonmotile isolates. A strain of serovar Gallinarum, a serovar failing to express flagella (lack of flagellar antigens), was assigned to fliC combined pattern A45P26. Other nonmotile isolates were studied. Two isolates (O9:H-:Vi-) had com-

bined pattern A45P26, which was observed for the flagellar antigen g series. These two isolates originated from poultry.

Four of the other isolates tested matched the A19P40 combined pattern that was observed for the H1:d antigen of serovar Typhi. Three of these isolates were O9,12:H-:Vi, and one was rough. No amplification of *fljB* was obtained.

One isolate (O4,5:H-) had a unique phase-1 flagellin gene combined pattern not described in this report.

#### DISCUSSION

This work shows that exploring the genetic diversity of *Salmonella* flagellin genes by RFLP is far more complicated than originally thought (9).

Subsp. II was the only subspecies other than subsp. I included in this study. This is because subsp. II strains share both O and H antigens with subsp. I strains. With a single exception, patterns of subsp. II flagellin genes were different from that of subsp. I. RFLP patterns suggest that flagellin genes from both subsp. I and II form separate evolutionary groups and are in support of the subspecies concept.

Phase-1 and phase-2 flagellin genes can be amplified separately. The phase-2 specificity of the *fljB* PCR system was verified by amplification of the *hin-fljB-fljA* region. It is remarkable that assignment of flagellin antigens to either phase was generally correct. There is still a question about serovars Kotte and Coromandel. Coromandel *fliC* and Kotte *fljB* have the same RFLP pattern (A39P24). However, the Coromandel phase-1 antigen is l,v whereas the Kotte phase-2 antigen is z35. Since the Coromandel phase-2 antigen is z35 (the Kotte phase-1 antigen is b), there is a possibility that the antigen phase assignment is wrong for Coromandel. More strains with the z35 and b antigens need to be studied to strengthen this hypothesis, and gene sequencing should be done.

Five patterns associated with antigen l,w were given by *fliC* and *fljB* amplified from different strains. In these cases, in spite of the lack of RFLP gene differentiation, some sequence difference must occur since the primers for amplification were different.

No fljB gene could be amplified from known monophasic serovars. This is in agreement with previous studies which showed that serovars Enteritidis, Typhi (monophasic), Berta, Agona, and Montevideo do not possess the fljA, hin, or fljB gene (2). It is striking that we were unable to amplify fljB from the (rarely occurring) serotype Typhi strains expressing second-phase z66. In contrast with the lack of a phase-2 gene in monophasic serotypes, the fliC gene could always be amplified from nonmotile isolates.

The high diversity of restriction profiles was attributed to variability within an internal region of the flagellin genes, whereas regions at the 5' and 3' ends are more conserved (6). In most cases, diversity highlighted by flagellin gene RFLP exceeded the diversity showed by antigens. This finding on the genetic variation of flagellin genes agrees with recently reported observations within populations of related strains of E. coli or Pseudomonas aeruginosa (4, 24). However, the flagellin gene sequence information was reduced by comigrating fragments. In addition, the choice of an endonuclease with restriction sites preferentially located in the variable region (HphI) did not yield better discrimination. Endonuclease *Hph*I showed fewer profiles than HhaI and fewer antigen- or serovar-specific patterns than *HhaI*. The use of a restriction enzyme which targets a highly variable region seems to increase the probability of generating falsely identical fragments, i.e., unrelated fragments with similar sizes in different patterns.

The correlation between flagellar antigens and flagellin

RFLP patterns is difficult to assess. It should be noted that building the WKL scheme has involved many historical and arbitrary decisions. The choice of strains for immunization and absorption was determining. Since antigenic factors could often be split further into subfactors, decisions had to be made about when to stop splitting. The WKL scheme is only a simplified summary of antigenic relationships among Salmonella serovars. When two antigens have the same designation, it indicates which antisera are likely to react, not that these antigens are identical. For example, serovars with H1:d may have different subfactors, such as d,d1 (Typhi), d,d3 (Stanley), and d,d3,d4 (Muenchen) (8). Flagellins of the g series have even more complex structures, such as g,o,m,z1,z2 for Enteritidis (summarized as g,m), g,o,m,q,z1 for Blegdam (summarized as g,m,q), or g,o,q,z3 for Moscow (summarized as g,q) (8). Phase-2 flagellins 1,5 and 1,7 are also very complex (8). On the other hand, i and r antigens have known antigenic relationships. Thus, it seems that RFLP patterns often split flagellar types in a different way than serotyping.

In several cases, RFLP was unable to discriminate among *fliC* genes encoding antigens with different designations such as d, i, or r,i antigens. This may be due to an unfortunate choice of endonucleases, although in these cases, sequences are not available for comparison. In other cases, lack of discrimination was due to excessive genetic similarity. The discrimination among antigens g,m (as in serovar Enteritidis) and g,p (as in serovar Dublin), which is essential in epidemiology, could not be achieved by RFLP analysis, since the corresponding genes differ by only six nucleotides (13). Genes encoding flagellin antigens 1,2; 1,5; and 1,6 are more than 96.2% related (22).

Although serotyping is the "gold standard" of Salmonella typing, all absorbed sera that are necessary for complete serotyping are not commercially available. This limits complete serotyping to National Reference Centers. A number of sera have to be prepared by Reference Centers. Since this preparation and serotyping itself (with phase inversion) are expensive and labor intensive, alternative methods which could be applied by clinical laboratories need to be sought. The dream that flagellar antigens could be deduced from flagellin gene RFLP has not been realized by this work. However, some important flagellin antigens, and even some serovars, can be deduced from RFLP patterns. When this is confirmed with more strains and combined with some PCR approach to Oantigen typing, then some major Salmonella serotypes could be identified by PCR and restriction, leaving serotyping for less common serotypes.

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