

## Distribution of molecular subtypes within *Salmonella enterica* serotype Enteritidis phage type 4 and *S. Typhimurium* definitive phage type 104 in nine European countries, 2000–2004: results of an international multi-centre study

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### SUMMARY

This study investigates the distribution of pulsed-field gel electrophoresis (PFGE) profiles within *Salmonella enterica* serotype Enteritidis phage type (PT) 4 and *S. Typhimurium* definitive phage type (DT) 104, from cases of human infection in nine European countries from 2000 to 2004. Isolates were subtyped using standardized methods and gel images submitted by each participating country to the coordinating centre (Health Protection Agency Centre for Infections, London, UK), where they were entered into a central database, developed within BioNumerics software, and designated using an agreed nomenclature. *S. Enteritidis* PT4 ( $n = 3637$ ) was differentiated into 38 different profiles. Simpson's index of diversity ( $D$ ) of profiles ranged from 0.2 to 0.4. Profile SENTXB.0001 represented at least 80% of all profiles in each country. *S. Typhimurium* DT104 ( $n = 1202$ ) was differentiated into 28 different profile types. Simpson's  $D$  was at least 0.6 in all countries except in Austria and Italy. In both these countries over 74% of *S. Typhimurium* DT104 profiles were STYMXB.0013. Profile STYMXB.0061, was predominant in Denmark, Spain, Finland and England & Wales where it represented between 36% and 45% of profiles. Profile STYMXB.0001 represented nearly half of all profiles in Scotland and 23% in England & Wales. PFGE is proving useful for further discrimination within *S. Enteritidis* PT4 and *S. Typhimurium* DT104. Ascertainment of international outbreaks involving common serotypes and phage types may be increased by the timely pooling of PFGE profiles within a central database readily accessible to all participating countries.

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## INTRODUCTION

The prompt ascertainment and investigation of outbreaks is important in the national and international prevention and control of salmonellosis. The International Surveillance Network for Enteric Infections (Enter-net), which encompasses 17 European and five non-European countries, was established to monitor human gastrointestinal infections caused by *Salmonella* and Vero cytotoxin-producing *Escherichia coli* (VTEC) O157, including their antimicrobial resistance [1]. For salmonellosis, the primary methods for differentiation of strains within Europe are serotyping, followed by phage typing for epidemiologically important serotypes [2], and antibiogram determination. These methods have been harmonized in all participating countries and the results pooled in a timely manner with other relevant information. These processes have been key to the success of Enter-net, which has detected and aided in the investigation of numerous international outbreaks [3].

*Salmonella enterica* serotypes Enteritidis and Typhimurium are the most commonly reported serotypes from cases of human infections in Europe. Between 2000 and 2002, these two serotypes represented 53% and 20% respectively of the 232 442 strains reported to Enter-net. Increasingly, subdivision within phage types is becoming necessary to elucidate outbreaks during investigations, whether these are national or international. Since the 1990s some phage types of epidemiological importance have predominated. Of all strains of *S. Enteritidis* phage typed during this period ( $n=87\,962$ ), 46% were phage type (PT) 4 and of the *S. Typhimurium* strains ( $n=18\,758$ ), 33% were definitive phage type (DT) 104 (Enter-net, unpublished data). For these phage types, any geographically or temporally scattered epidemiologically linked cases may go unnoticed or may be detected late. It is, therefore, necessary to consider further discrimination within common phage types for meaningful surveillance. Pulsed-field gel electrophoresis (PFGE) is currently the gold standard method for molecular subtyping of *Salmonella* [4, 5], and possibly also for the rapid identification of unusual *Salmonella* serotypes. The method involves cutting genomic DNA at specific sites to generate fragments of different sizes followed by determination of their molecular weights by running them through an electrically charged gel matrix.

Salm-gene, a EU research project, was established to evaluate the added value of using molecular subtyping

tools (PFGE essentially) for outbreak recognition of *S. Enteritidis* and *S. Typhimurium*, within the existing international surveillance of *Salmonella*. The project involved the participation of nine national reference laboratories in Europe: Austria, Denmark, Finland, Germany, Italy, The Netherlands, Scotland, Spain, England & Wales with France acting as software compatibility advisor.

We describe the distribution of pulsed-field profiles (PFPs) within isolates of *S. Enteritidis* PT4 and *S. Typhimurium* DT104 reported to Salm-gene from the nine participating European countries between January 2000 and September 2004.

## MATERIALS AND METHODS

### Strain source

All strains from cases of human salmonellosis in the nine countries that were referred to the national reference laboratory were eligible for subtyping using PFGE. Strains included any sporadic isolates and outbreak strains, with the proviso that only a single isolate from any outbreak within a country was represented.

### Selection of strains

At the start of the project all nine laboratories were asked to subtype retrospectively, using PFGE, a selection of strains of the most common *S. Enteritidis* and *S. Typhimurium* phage types from isolates that had been reported to them up to December 2001. This created a library of unique strain profiles and provided the background of profiles in strains in circulation prior to the study. Strains were selected in the order they had been reported, starting from the most recent, and working back temporally until a commonly agreed target number was reached.

'Real-time' subtyping was implemented from January 2003 and required each laboratory to submit to the Salm-gene coordinating centre (Health Protection Agency, England & Wales), up to three PFGE gels a week, representing 20–45 isolates in all. A protocol for selecting strains to be subtyped was sent to all laboratories. This involved all isolates of *S. Enteritidis* and *S. Typhimurium* if the total number was less than the PFGE typing capacity. If the number of strains reported nationally exceeded local subtyping capacity, a simple random sampling method was applied. The sample number varied by

serotype and was in proportion to the number of strains within each serotype. For example, assuming the typing capacity was 48 isolates, and 73 (70%) strains of *S. Enteritidis* and 31 (30%) strains of *S. Typhimurium* were reported to the national laboratory, 34 *S. Enteritidis* and 14 *S. Typhimurium* strains would be selected arbitrarily from these strains.

### PFGE subtyping

As described previously laboratory procedures for PFGE were standardized in all nine laboratories using the Bio-Rad CHEF<sup>®</sup> system [6]. The following harmonized protocol was used for the studies described:

After overnight growth on TSA or nutrient broth, cells were harvested by centrifugation and resuspended in 500  $\mu$ l cell suspension buffer [100 mM Tris, 100 mM EDTA (pH 8.0)]. The final cell density for plug preparation was 0.38–0.44 at 450 nm (McFarland No. 5). Proteinase K (0.5 mg/ml final concentration) was added to the cell suspension followed by mixing lysis of cell suspension 1:1 with agarose (Bio-Rad chromosomal grade). The resultant plugs were washed at least twice in distilled water and 2–3 times in TE buffer. Electrophoresis conditions were as follows: ramp initial 2 s; final 64 s; 6 V/cm; 14 °C, 22 h (Bio-Rad CHEF DRII<sup>®</sup>), 20 h (Bio-Rad CHEF DRIII<sup>®</sup>), 18 h (Bio-Rad CHEF Mapper<sup>®</sup>). DNA macrorestriction fragments were resolved on 1% agarose gels (Bio-Rad Pulsed-Field Certified<sup>®</sup> or Seakem Gold<sup>®</sup>) with a *S. Braenderup* strain H9812 (kindly supplied by PulseNet USA, CDC [4]) as a molecular reference marker. To ensure compatibility the electrophoresis conditions were identical to those used by PulseNet USA.

### Electronic submission of PFGEs

Gel images were transmitted in tag image file format (TIFF) by email together with additional microbiological and epidemiological information on each strain, including phage type when available, to the Salm-gene coordinating centre. All gels that were considered to be of acceptable quality (in terms of image clarity, band definition and use of Braenderup reference profile strains) were entered and analysed into the Salm-gene database, created within the BioNumerics software (v. 3.1, beta, Applied Maths, Sint-Martens-Latem, Belgium). This involved naming each pattern based on a comparison with the profiles of the library (see below).

### Profile designation

The designation involved assigning a six-letter code to the new pattern together with a four-digit numerical identifier [6]. The first pattern for *S. Enteritidis* digested with the enzyme *Xba*I was designated SENTXB.0001 and similarly for *S. Typhimurium* digested with the same enzyme, STYMXB.0001. A profile differing by at least one band to a pattern in the library was assigned a new name and added to the library.

### Data analysis

All strains of *S. Enteritidis* and *S. Typhimurium* entered into the database up to 30 September 2004 were included in the analysis. This represents a subset of isolates reported to the national laboratories between January 2000 and September 2004.

The diversity of PFGE profiles was calculated using Simpson's index of diversity (*D*) as described by Grundmann *et al.* [7]. The value of this index ranges between 0 and 1, with 0 indicating no diversity. The 95% confidence intervals were calculated based on the variance as suggested by Grundmann *et al.* Access 2000 (Microsoft, Washington, USA) was used for the extraction of data and Excel 2000 (Microsoft) was used for the calculation of Simpson's *D*, variance and confidence intervals.

## RESULTS

### *S. Enteritidis*

Of all strains of *S. Enteritidis* included in the Salm-gene database ( $n=11\,659$ ), 10 927 had a designated phage type, of which 3637 (33%) were PT4. Of the 3637 profiles, 3509 had been assigned a profile name by the database curator. Within the designated profiles *S. Enteritidis* PT4 isolates were differentiated into 38 different profile types, the 10 most common of which are illustrated in Figure 1. Profile SENTXB.0001 predominated in all countries representing at least 80% of all profiles (Table 1). Simpson's *D* of profile types within PT4 ranged from 0.2 to 0.4 and was lowest in Austria, Spain, England & Wales, Scotland and Germany. Some countries had country-specific profile types. Finland had the greatest number of unique profiles ( $n=6$ ), followed by England & Wales ( $n=3$ ), The Netherlands ( $n=2$ ), Denmark ( $n=1$ ), Spain ( $n=1$ ), Scotland ( $n=1$ ) and Italy ( $n=1$ ) (Table 1).

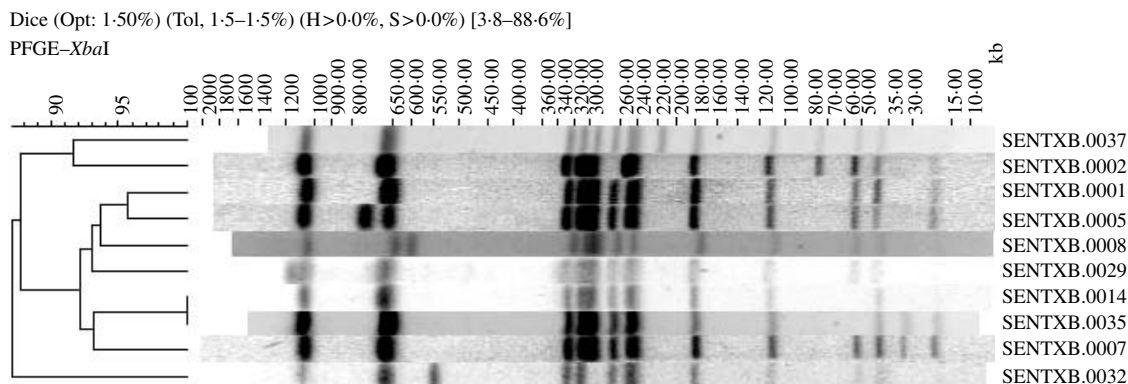


Fig. 1. Ten most common profile types within *S. Enteritidis* PT4.

Table 1. Number of *S. Enteritidis* PT4 profiles within each country

	Austria	Denmark	Spain	Finland	E&W*	Scotland	Italy	The Netherlands	Germany	Total
SENTXB.0001	819	40	277	200	908	194	78	173	405	3094
SENTXB.0014	62	5	16	19	39	11	4	5	23	184
SENTXB.0005	23		6	1	1		9	7	3	50
SENTXB.0037	11	1	2		14	2		1	10	41
SENTXB.0002	9	1			1		2	4	2	19
SENTXB.0008	3	1	3	1	6	2				16
SENTXB.0029	1			1	11			1		14
SENTXB.0032					12		1			13
SENTXB.0007	3		1		3	2		2	2	13
SENTXB.0035			1		9	1				11
SENTXB.0012			1	2	6				1	10
SENTXB.0031	4				3					7
SENTXB.0023	2		1			1		1	1	6
SENTXB.0013	3		1			1			1	6
SENTXB.0010	1			1	1	1				4
SENTXB.0059				1	2		1			4
SENTXB.0020			3		1					4
SENTXB.0045	1							2		3
SENTXB.0015			1					1	1	3
SENTXB.0021			2			1				3
SENTXB.0003					2			1		3
SENTXB.0004			1		1				1	3
SENTXB.0036	1				1					2
SENTXB.0038				2						2
SENTXB.0057				2						2
SENTXB.0044							1	1		2
SENTXB.0041				1	1					2
SENTXB.0046		1					1			2
SENTXB.0009			1							1
SENTXB.0011					1					1
SENTXB.0016					1					1
SENTXB.0027							1			1
SENTXB.0039								1		1
SENTXB.0040				1						1
SENTXB.0043								1		1
SENTXB.0048					1					1
SENTXB.0050				1						1
SENTXB.0026						1				1
Total	943	49	317	233	1025	217	98	201	450	3533
No designation	14	2	20	4	27	7	9	10	11	104

\* England & Wales.

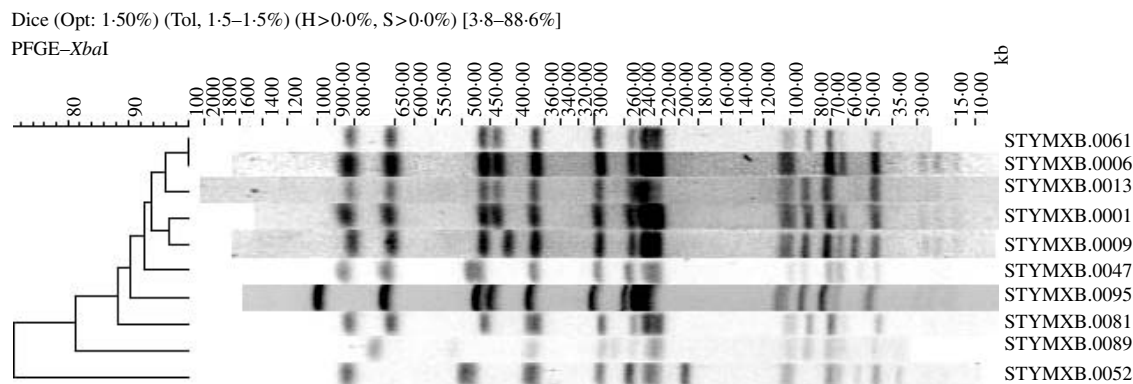


Fig. 2. Ten most common profile types within *S. Typhimurium* DT104.

Table 2. Diversity of profiles of *S. Enteritidis* PT4 and *S. Typhimurium* DT104 within each country

	Diversity index	95% CI
<i>S. Enteritidis</i> PT4		
Austria	0.24	0.20–0.28
Denmark	0.32	0.16–0.49
Spain	0.23	0.17–0.30
Finland	0.26	0.18–0.33
E&W*	0.21	0.18–0.25
Scotland	0.20	0.13–0.27
Italy	0.36	0.23–0.48
The Netherlands	0.26	0.17–0.34
Germany	0.19	0.14–0.24
<i>S. Typhimurium</i> DT104		
Austria	0.09	0.01–0.18
Denmark	0.72	0.63–0.81
Spain	0.74	0.68–0.80
Finland	0.61	0.53–0.69
E&W*	0.76	0.74–0.79
Scotland	0.72	0.64–0.81
Italy	0.42	0.23–0.60
The Netherlands	0.60	0.51–0.68
Germany	0.69	0.66–0.73

\* England & Wales.

### *S. Typhimurium*

Overall, 5817 strains of *S. Typhimurium* were entered into the Salm-gene database. DT104 represented 26% ( $n=1202$ ) of all strains with a designated phage type ( $n=4631$ ). Of these DT104 strains, 1060 were assigned a PFGE profile type. Twenty-eight different profile types were identified, the 10 most common of which are shown in Figure 2. There were differences between countries in the distribution and diversity of profile types. Simpson's  $D$  was at least 0.6 except in Austria and Italy (0.09 and 0.42 respectively) (Table 2). In

both these countries the majority of profiles were STYMXB.0013. This profile was found in all other countries but more frequently in Germany and The Netherlands, where it represented 45% and 74% respectively of all profiles in these countries. Profile STYMXB.0061, was predominant in Denmark, Spain, Finland and England & Wales where it represented between 36% and 45% of profiles. Profile STYMXB.0001 represented nearly half of all profiles in Scotland and a high proportion of profiles (23%) in England & Wales (Table 3).

A remaining 104 profiles of *S. Enteritidis* and 142 profiles of *S. Typhimurium* had no matching patterns in the library at the time of analysis. These strains could not be designated due to late submission and time constraints.

### DISCUSSION

Phage typing, based on common protocols, is vital for improving outbreak ascertainment and for outbreak investigations of *S. Enteritidis* and *S. Typhimurium*. For common phage types within these serovars, such as PT4 and DT104 respectively, further discriminatory methods are necessary to differentiate these strains. Molecular typing methods are commonly used as a tool to help with outbreak investigations, PFGE being currently the gold standard for *Salmonella*. This method has been implemented through standardized protocols in the United States [4] and several countries in Europe, with the view of enhancing *Salmonella* outbreak ascertainment internationally.

Both PT4 and DT104 are recognized as highly clonal organisms [8]. As the present study has demonstrated, these phage types have been differentiated into several PFGE subtypes in all nine countries.

Table 3. Number of *S. Typhimurium* DT104 profiles within each country

	Austria	Denmark	Spain	Finland	E&W*	Scotland	Italy	The Netherlands	Germany	Total
STYMXB.0013	77	3	21	2	40	4	26	23	146	342
STYMXB.0061	3	23	40	19	104	13	1	37	84	324
STYMXB.0001		7	6	2	67	33		1	59	175
STYMXB.0006	1	6	15	22	50	12			11	117
STYMXB.0095		10	6			3		1	7	27
STYMXB.0047				1	10	1				12
STYMXB.0081		1		1	4	1				7
STYMXB.0052							6			6
STYMXB.0089									5	5
STYMXB.0009					1	1			3	5
STYMXB.0002			1		3					4
STYMXB.0005			1		2	1				4
STYMXB.0014		1	2		1					4
STYMXB.0051								4		4
STYMXB.0084					1	1	1		1	4
STYMXB.0069									3	3
STYMXB.0086								1	2	3
STYMXB.0050			1		1					2
STYMXB.0003					1				1	2
STYMXB.0056					1				1	2
STYMXB.0062							1			1
STYMXB.0066						1				1
STYMXB.0074						1				1
STYMXB.0011								1		1
STYMXB.0075			1							1
STYMXB.0008					1					1
STYMXB.0091			1							1
STYMXB.0093								1		1
Total	81	51	95	47	287	72	35	69	323	1060
No designation	3	9	36	7	26	3	5	10	43	142

\* England & Wales.

As can be seen from Figures 1 and 2, differences between profile types are mainly in the number and/or position of bands of high (450–1135 kb) and low (20–100 kb) molecular mass. In some cases the differences are minor (1–2 band difference). Although previous studies have concluded subtypes based on minor band differences may not be sufficiently discriminatory [9], minor band differences might be key in identifying outbreak strains and in their differentiation from sporadic cases [10, 11]. In this respect discrimination within phage type for investigating *Salmonella* outbreaks may be dependent on changes in a single band. This is contrary to the recommendations of Tenover *et al.* for other organisms [12]. Furthermore, the influence of plasmids may have some effect on PFGE profiles and whenever possible this should also be taken into account [11].

There was little variance with the genetic diversity of PT4 between the nine countries (Simpson's

$D=0.2-0.4$ ). This was in contrast to DT104. Countries with the highest diversity were Denmark, Finland, Germany, England & Wales, Scotland, The Netherlands and Spain (Simpson's  $D=0.6-0.8$ ) for DT104. The lower diversity observed in Austria (Simpson's  $D=0.2$ ) and Italy (Simpson's  $D=0.4$ ) may be due to fewer imported infections and/or fewer imports of contaminated food.

Certain profile types predominated within some countries. Within The Netherlands, Austria, Finland, Denmark, Germany, Italy, England & Wales, Scotland and Spain, the majority of profiles of *S. Enteritidis* PT4 were SENTXB.0001 although several other less common subtypes could also be observed. A similar distribution of profile patterns has been reported previously, although the restriction endonuclease used for PFGE was not always *XbaI* [13–15]. The predominant subtype SENTXB.0001 may be identical to those recognized in these studies.

This is not unlikely, considering the rapidity by which this phage type is known to have spread internationally [8, 16]. Further comparisons using the same enzyme and similar gel running times are required before this can be verified.

Some neighbouring countries shared the same predominant profile of *S. Typhimurium* DT104; for example in Austria and Italy, profile STYMXB.0067 represented the majority of profiles (74 and 90% respectively). The profiles of DT104 shared by neighbouring countries, suggests they are sharing local reservoirs or sources of multi-drug resistant DT104. In contrast Denmark, Spain, Finland, England & Wales and The Netherlands STYMXB.0061 predominated representing at least 36% of all profiles within each country. This profile was less frequently found in Austria, Scotland, Italy and Germany. Multi-drug resistant DT104 is known to have disseminated globally [17] and it is highly probable that this clone is represented in these subtypes.

Similarities in the occurrence and distribution of subtypes were apparent between countries. For example Austria and Germany shared eight subtypes of *S. Enteritidis* PT4 out of the 14 and 11 profiles observed in these countries respectively. Twenty-six of the PT4 subtypes were shared by at least two countries. The other 38 were unique to the reporting country. Because of the very low number of these profiles (in many cases only one isolate of this type was reported), we are unable to make conclusive remarks regarding their specificity. The present study has provided a timely analysis, at molecular level, of a representative sample of human *Salmonella* isolates across Europe. The laboratory protocols used in Salm-gene have been developed in line with those of PulseNet USA. This is a national subtyping network for the USA for foodborne pathogens using PFGE [4] and similar networks have been extended in Canada, and recently in Asia and Latin America. The use of the *S. Braenderup* control strain H9812 in this study has ensured compatibility between the Salm-gene and PulseNet network. As part of the PulseNet Europe ring-trial, the PulseNet network has been extended to 28 institutes in 19 European countries within and out of Salm-gene (PulseNet Europe) in late 2004, and the Salm-gene database is providing the foundation of this network for *Salmonella*. Combined with case and phenotypic information, the rapid pooling of PFGE profiles into a central database is readily accessible to Enter-net, Salm-gene and PulseNet Europe participating countries. This will facilitate the timely ascertainment

of international outbreaks involving common serotypes and phage types within Europe, and for countries with phage typing expertise, will also negate the expensive and time-consuming transfer of isolates through the postal system for subsequent molecular typing.

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## DECLARATION OF INTEREST

None.

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