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Identification and typing of Salmonella for public health surveillance using whole genome sequencing

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In April 2015, Public Health England implemented whole genome sequencing (WGS) as a routine typing tool for public health surveillance of Salmonella, adopting a multilocus sequence typing (MLST) approach as a replacement for traditional serotyping. The WGS derived sequence type (ST) was compared to the phenotypic serotype for 6887 isolates of S. enterica subspecies I, and of these, 6616 (96%) were concordant. Of the 4% (n=271) of isolates of subspecies I exhibiting a mismatch, 119 were due to a process error in the laboratory, 26 were likely caused by the serotype designation in the MLST database being incorrect and 126 occurred when two different serovars belonged to the same ST. The population structure of S. enterica subspecies II-IV differs markedly from that of subspecies I and, based on current data, defining the serovar from the clonal complex may be less appropriate for the classification of this group. Novel sequence types that were not present in the MLST database were identified in 8.6% of the total number of samples tested (including S. enterica subspecies I-IV and S. bongori) and these 654 isolates belonged to 326 novel STs. For S. enterica subspecies I, WGS MLST derived serotyping is a high throughput, accurate, robust, reliable typing method, well suited to routine public health surveillance. The combined output of ST and serovar supports the maintenance of traditional serovar nomenclature while providing additional insight on the true phylogenetic relationship between isolates.

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Running title: WGS derived MLST serotyping for Salmonella
Key words: whole genome sequencing, multilocus sequence typing, Salmonella

26 Abstract

27 In April 2015, Public Health England implemented whole genome sequencing (WGS) as a routine typing 28 tool for public health surveillance of Salmonella, adopting a multilocus sequence typing (MLST) 29 approach as a replacement for traditional serotyping. The WGS derived sequence type (ST) was 30 compared to the phenotypic serotype for 6887 isolates of S. enterica subspecies I, and of these, 6616 31 (96%) were concordant. Of the 4% (n=271) of isolates of subspecies I exhibiting a mismatch, 119 were 32 due to a process error in the laboratory, 26 were likely caused by the serotype designation in the MLST 33 database being incorrect and 126 occurred when two different serovars belonged to the same ST. The 34 population structure of S. enterica subspecies II-IV differs markedly from that of subspecies I and, based 35 on current data, defining the serovar from the clonal complex may be less appropriate for the 36 classification of this group. Novel sequence types that were not present in the MLST database were 37 identified in 8.6% of the total number of samples tested (including S. enterica subspecies I-IV and S. 38 bongori) and these 654 isolates belonged to 326 novel STs. For S. enterica subspecies I, WGS MLST 39 derived serotyping is a high throughput, accurate, robust, reliable typing method, well suited to routine 40 public health surveillance. The combined output of ST and serovar supports the maintenance of 41 traditional serovar nomenclature while providing additional insight on the true phylogenetic relationship 42 between isolates.

44 Introduction

45 The Salmonellae are major human pathogens and represent a significant global public health issue 46 causing morbidity and mortality resulting in a high social and economic burden worldwide (Majowicz et 47 al., 2010). The genus consists of 2 species; Salmonella enterica and S. bongori. There are six subspecies 48 of S. enterica differentiated by biochemical variations, namely subspecies enterica (I), salamae (II), 49 arizonae (IIIa), diarizonae (IIIb), houtenae (IV) and indica (VI) (Threlfall et al. 1999). Subspecies I, S. 50 enterica subsp. enterica cause 99% of human and animal infections. The two main pathologies 51 associated with S. enterica are gastroenteritis and typhoidal disease. The typhoidal Salmonellae include 52 S. Typhi and S. Paratyphi A, B and C. They are host restricted, monophyletic, rarely undergo 53 recombination events and exhibit convergent evolution driven by genome degradation (Wain et al. 54 2015). The majority of gastroenteritis in the UK is caused by the host generalist serovars, such as S. 55 Typhimurium and S. Enteritidis, and host adapted serovars that are adapted to a specific animal 56 reservoir but can infect man and include S. Dublin, S. Gallinarum S. Choleraesuis, and S. Bovismorbificans 57 (Langridge et al. 2015).

58

59 Approximately 8,000 isolates are referred to the Salmonella Reference Service (SRS) at Public Health 60 England (PHE) each year from local and regional hospital laboratories. In April 2015, PHE implemented 61 whole genome sequencing (WGS) as the routine typing tool for public health surveillance of Salmonella 62 infections. Prior to April 2015, presumptive Salmonella isolates referred to SRS were speciated and sub-63 speciated using PCR (Hopkins et al. 2009, 2011) and grouped into serovars as described in the White-64 Kauffman-Le Minor scheme (Grimont & Weill 2007, Guibourdenche et al. 2010, Issenhuth-Jeanjean et al. 65 2014). This methodology is based on reactions of rabbit antisera to the lipopolysaccharide (O antigen 66 encoded by rfb genes) and flagellar antigens (phases 1 and 2 of H antigen encoded by fliC and fliB). The 67 scheme utilises this phenotypic variation, expressed as an antigenic formulae, to divide Salmonella into 68 more than 2600 serovars. Epidemiological investigations of Salmonella infecting humans and animals 69 have relied on serotyping for over 70 years; national and international governmental agencies base 70 guidelines and regulations on the serotyping method and the use of this nomenclature is a globally 71 recognised form of communication (Swaminathan et al. 2009, EFSA 2010). Furthermore, serovars have 72 often been shown to correlate with host range and disease sequelae (Gordon et al. 2011, Wain et al. 73 2015, Langridge et al. 2015).

75 There are, however, a number of issues with the serotyping approach; specifically, the expense and 76 expertise required to produce the antisera and , furthermore, serotyping does not reflect the genetic 77 relatedness between serovars, nor does it provide an evolutionary perspective. Alternative molecular 78 serotyping methods have been described previously including Pulsed-field gel electrophoresis. 79 ribotyping, repetitive extragenic palindromic sequence-based PCR (rep-PCR) and combined PCR- and 80 sequencing-based approach that directly targets O- and H-antigen-encoding genes (Ranieri et al. 2013, 81 Shi et al. 2015). In 2012, Achtman et al. proposed a sequenced based approach, multilocus sequence 82 typing (MLST), based on the sequences of multiple house-keeping genes. Isolates that possess identical 83 alleles for the seven gene fragments analysed are assigned a common sequence type (ST) and related 84 STs from clonal complexes are termed e-Burst Groups (eBGs). They showed that ST and eBGs strongly 85 correlated with serovar and so utilising this approach would facilitate backward compatibility with 86 historical data, minimise disruption for reference laboratory service users and facilitate data exchange 87 with other colleagues in the field.

88

89 Advances in whole genome sequencing (WGS) methodologies have resulted in the ability to perform 90 high throughput sequencing of bacterial genomes at low cost making WGS an economically viable 91 alternative to traditional typing methods for public health surveillance and outbreak detection (Koser et 92 al. 2012). Whilst WGS provides the opportunity to resolve bacterial strains to the single nucleotide 93 resolution needed for identifying cases linked to a common source of infection (Dallman et al. 2015), 94 grouping isolates into higher taxonomical clones (e.g. those defined by serotyping) is an important step. 95 The decision to adopt WGS as a routine typing method at PHE provided the opportunity to review our 96 approach to typing Salmonella and to implement the MLST approach in parallel with WGS. 97 98 The aim of this study was to evaluate MLST, as derived from WGS data, as a replacement for 99 conventional serotyping of Salmonella for routine public health surveillance and to provide insight into 100 the genetic population structure of all Salmonella species in England and Wales during a 12 month

101 period.

- 103 Methods
- 104 Bacterial strains
- 105 All isolates (n=7465) of Salmonella from human cases of gastrointestinal disease submitted to SRS from
- 106 local and regional hospital laboratories in England & Wales between 1st April 2014 and 31st March 2015

107 were sequenced in parallel with phenotypic serotyping (Supplementary Table). Of these, 7338 were 108 identified as subspecies I and included 263 different serovars. The ten most common serovars in this 109 dataset were Enteriditis (2310), Typhimurium (1407), Infantis (184), Typhi (184), Newport (173), Virchow 110 (162), Kentucky (160), Stanley (146), Paratyphi A (135) and Java (99). One hundred and twenty seven 111 isolates were identified as subspecies II-IV (S. enterica subspecies salamae n=28; S. enterica subspecies 112 arizonae n=25; S. enterica subspecies diarizonae n=49; S. enterica subspecies houtenae n=20) and there 113 was one isolate of *S. bongori*. No isolates belonging to subspecies VI (*S. enterica* subspecies *indica*) were 114 submitted to SRS during the study period. 115

$116 \quad \text{DNA extraction for WGS}$

117 DNA extraction of *Salmonella* isolates was carried out using a modified protocol of the Qiasymphony

118 DSP DNA midi kit (Qiagen). In brief, 0.7 ml of overnight *Salmonella* culture in a 96 deep well plate was

119 $\,$ harvested. Bacterial cells were pre-lysed in 220 μl of ATL buffer (Qiagen) and 20 μl Proteinase K $\,$

120~ (Qiagen), and incubated shaking for 30 mins at 56°C. Four μl of RNase at 100 mg/ml (Qiagen) was added

121 to the lysed cells and re-incubated for a further 15 mins at 37°C. This step increases the purity of the

122 DNA for further downstream sequencing. Extraction of DNA from the treated cells was performed on

- 123 $\,$ the Qiasymphony SP platform (Qiagen) and eluted in 100 μl of water. DNA concentration using the
- 124 GloMax system (Promega) was determined for the following sequencing steps.
- 125

126 DNA sequencing

127 Extracted DNA was then processed using the NexteraXT sample preparation method and sequenced

- 128 with a standard 2x101 base protocol on a HiSeq 2500 Instrument in fast mode (Illumina, San Diego).
- 129

130 Bioinformatics Service Workflow

FASTQ reads were quality trimmed using Trimomatic (PMID 24695404) with bases removed from the trailing end that fell below a PHRED score of 30. If the read length post trimming was less than 50bp the read and its pair were discarded. A K-mer identification step was used to compare the sequenced reads with 1769 published genomes to identify the bacterial species (and *Salmonella* subspecies) and to detect cultures submitted by the local and regional hospital laboratories that contained more than one bacterial species (mixed cultures). ST assignment was performed using a modified version of SRST (PMID 25422674). Preliminary analysis was undertaken using the MLST database described in Achtman

138 *et al.* (2012).

139	
140	For isolates that had novel STs, or a ST but no associated serovar in the Achtman MLST database, the
141	serovar was determined by phenotypic serotyping at PHE. STs and corresponding serovars of isolates
142	serotyped and sequenced during this study were added to a modified version of the Achtman MLST
143	database, held and curated at PHE. These novel STs were assigned a preliminary ST (PST) and an
144	inferred serovar was determined. The PHE MLST database currently holds 7000 strains and 1,200
145	serovars and is up-dated every three months.
146	
147	Results
148	Achtman et al. (2012) described the population structure of Salmonella enterica as monophyletic
149	lineages of STs that have evolved from a single founder node and termed these discrete clusters eBGs.
150	The population structure of all the Salmonella species submitted to PHE between April 2014 and March
151	2015 is illustrated by the minimum spanning tree in Figure 1.
152	
153	Salmonella subspecies I
154	The ST and corresponding serovar designation obtained from the MLST database were used to compare
155	the WGS derived ST to the phenotypic serotype for 6887 (94%) of 7338 isolates of subspecies 1, and of
156	these, 6616 (96%) had the same result by both methods (Supplementary Table). It was not possible to
157	compare phenotypic serotyping with MLST-based serotyping for 451 (6%) subspecies I isolates because
158	either the phenotypic serotype could not be determined due to an incomplete antigenic structure (S.
159	Unnamed) (n=423) or the serovar could not be determined because the ST did not have a designated
160	serotype in the MLST database (n=70). Forty-two isolates were both S. Unnamed and had no MLST

- 161 designated serotype.
- 162

163 For the 423 (5.8%) subspecies I isolates reported as S. Unnamed, 318 (90%) were designated a serotype

- 164 from the WGS derived MLST data. The most common serovars typed in this way included S.
- 165 Typhimurium (118), S. Virchow (30), S. Stanley (17), S. Enteritidis (16), S. Infantis (14) and S. Thompson

166 (13). Of the 7338 strains tested, 70 (1%) had no serotype designation in the MLST database, of which 28

167 (40%) were serotyped phenotypically (Supplementary Table).

168

169 Subspecies I novel sequence types

170 Novel sequence types that were not present in the MLST database were identified in 8.6% (n=654) of 171 the strains (Supplementary Table). These 654 isolates belonged to a total of 326 novel STs, designated 172 PST; the modal number of isolates identified per PST was one (Figure 2a). There was no difference in 173 the distribution of number of isolates per PST depending on whether the PST had a known serovar or 174 belonged to an unnamed or ambiguous serotype. The rate at which PSTs were received throughout the 175 year was plotted and revealed a linear relationship ($R^2 = 0.98$, y = 1.04 * x, where x = number of days 176 since April 1st 2014) (Figure 2b).

177

- 178 The serovars with the highest number of new PSTs were *S*. Typhimurium (n=9), *S*. Stanley (n=9), *S*.
- 179 Enteritidis (n=9) and S. Newport (n=8), although the majority of these PSTs were single locus variants
- 180 (SLVs) of established STs, belonging to these serovars (S. Typhimurium 8/9, S. Stanley 7/9 and S.
- 181 Newport 7/8). There were also serovars for which a large number of PSTs were identified that were not
- 182 SLVs of established STs (S. Agama 5/5, S. Agbeni 5/5, S. Saint-Paul 5/5, S. Enteritidis 4/9) which may
- 183 represent new eBGs that share these serotypes.
- 184

185 Subspecies I mismatches

- 186 Four percent (n= 271) of the isolates tested exhibited a mismatch between the WGS MLST derived
- 187 serovar and the phenotypic serotyping results. Of the 271 mismatches, 119 were due to a process error
- 188 in the laboratory either in the phenotypic serotyping or the DNA extraction part of the WGS pipeline.
- 189 With respect to the phenotypic serotyping, common errors included mislabelling samples and
- 190 misinterpreting or incorrectly transcribing the antigenic structure, especially when the antigenic
- 191 structures were similar. For example, S. Agona (I 4, 12:f, g, s:-) and S. Derby (I 4, 12:f, g:-). DNA
- 192 extraction errors were associated with mislabelled samples.
- 193

194Twenty-six mismatches were potentially caused by the predicted serotype designation in the Achtman195MLST database being incorrect which may be attributed to single entries that had been misidentified at196the laboratory from which the MLST data was submitted. For example, in the original database ST1499197is represented by one entry identified by the submitter as *S*. Litchfield. Subsequently, phenotypic198serotyping at PHE identified this ST as *S*. Bovis-morbificans in more than five isolates. ST1499 belongs to199eBG34 which comprises two other STs both associated with S. Bovis-morbificans, indicating that the200original entry in the MLST database is likely to be incorrect.

The most common reason for mismatches occurring between the WGS MLST derived serotype and the phenotypic serotype (n=126) occurred when two different serovars belonged to the same eBG and the same ST (see Table 1 and discussed in more detail below).

205

206 Serovars Enteritidis and Dublin

207 Of the 2308 isolates of S. Enteritidis identified by both phenotypic serotyping and WGS MLST derived 208 serotyping, 2296 belonged to eBG4, including 2200 ST11 and 76 ST183 (Figure 1). There were five 209 additional SLVs of ST11, four of which were novel types. S. Gallinarum and S. Pulloram can be difficult to 210 distinguish from S. Enteritidis (Thomson et al. 2008) but neither of these serovars were identified in this 211 study. Serologically, S. Dublin ([1],9,12:g,p:-) has a similar antigenic structure to S. Enteritidis 212 ([1],9,12:g,m:-), and in Achtman et al. (2012), eBG32 (ST74) contained both S. Enteritidis and S. Dublin. 213 However, in this study both isolates belonging to ST74 eBG32 typed as S. Enteriditis. Of the 2308 214 isolates, 26 belonged to nine new PSTs. The most common was P3147, a previously undescribed SLV of

- 215 ST11, comprising 16 cases including 10 known to have travelled to Malaysia or Singapore.
- 216

217 Serovar Typhimurium

218 In this study, eBG1 contained 1392 isolates of S. Typhimurium and monophasic S. Typhimurium (rough

- and non-motile variants) (Hopkins et al. 2012). The monophasic variants also belong to eBG138
- 220 (primarily ST 36) and eBG243. In contrast to eBG1 described in Achtman *et al.* (2012), which was
- represented by a large central ST19 node with at least 27 SLV STs comprising much smaller numbers of
- strains, eBG1 in the PHE dataset shows a predominance of both ST19 and ST34 and less allelic variation.
- 223 Only nine SLVs to ST19 were identified including three undesignated STs (Figure 1).
- 224

225 Serovars Java/Paratyphi B data

226 Despite the different disease outcomes associated with S. Paratyphi B (most commonly associated with

- invasive disease and paratyphoid fever) and S. Java (most commonly associated with gastroenteritis) it is
- not possible to differentiate the two serotypes by serotyping alone. *S.* Java and *S.* Paratyphi B are
- therefore differentiated in the laboratory by their ability to ferment dextrorotatory tartrate (S. Java dTa+
- and S. Paratyphi B dTa-) (Malorny et al. 2003).
- 231
- 232 The 99 isolates identified by both phenotypic serotyping and WGS MLST derived serotyping as S. Java,
- 233 belonged to a diverse range of eBGs, STs and PSTs (Table 2 and Figure 1). Two of these 99 isolates

- (marked with * in Table 2) belonged to ST86 and the predicted serotype from the MLST database was S.
- Paratyphi B. One of these isolates was from a blood culture (associated with invasive disease) and,
- therefore, likely to have been misidentified phenotypically. All 12 isolates identified as S. Paratyphi B
- 237 phenotypically, were identified as *S*. Paratyphi B ST86 by WGS MLST.
- 238

239 Subspecies II-IV and S. bongori

- 240 Isolates from subspecies II, III, IV, VI and S. bongori were not well represented in Achtman's MLST
- 241 database and thus the majority of isolates from these sub-species sequenced in this study did not
- 242 belong to a previously designated eBG or ST. The population structure of the 127 non-subspecies I
- 243 isolates differs markedly from that of *Salmonella enterica* (subspecies 1) (Figure 1) and shows some
- similarity to the population structure of lineage 3 in being a connected network of STs.
- 245

Sixteen of the 28 isolates belonging to subspecies II were previously designated *S*. Unnamed and the 28 strains belonged to 20 different STs. There were 25 isolates classed as subspecies IIIa (belonging to 10 different STs) and 49 in subspecies IIIb (belonging to 27 different STs). Of the 20 isolates identified as subspecies IV, 10 were designated *S*. Wassenaar (P3029) by phenotypic serotyping and the 20 isolates belonged to five different STs. All isolates of subspecies II-IV and *S. bongori* were correctly speciated using the k-mer ID approach.

252

253 *Population structure*

As highlighted by Achtman *et al.* (2012), the majority of isolates in the dataset belong to eBGs that have a one-to-one relationship with a specific serovar including *S*. Typhi, *S*. Paratyphi A and *S*. Heidelberg. In this study, of the serovars comprising more than 25 isolates, there were 17 serovar specific eBGs and 10 examples of a single serovar being associated with multiple eBGs (Figure 1). There were at least six examples of more than one serovar belonging to the same eBG but different STs, for example *S*. Hadar (ST33) and *S*. Kottbus (ST582) both belong to eBG22 and *S*. Bredeney (ST306) and *S*. Schwarzengrund (ST96) both belong to eBG33 (Supplementary Table).

261

262 There were seven examples where two serovars belonged to the same eBG and the same ST (Table 1).

- 263 In all of these examples, the antigenic structures of the two serovars were similar with only one antigen
- 264 differentiating the two serovars. Further analysis was carried out on two examples to determine
- 265 whether this difference in antigenic structure represented a true difference in strain relatedness or a

- random change that is not reflected in phylogeny (for example, the insertion of phage encoded antigen).
- 267 The analysis showed that the change in antigenic structure in S. Richmond (I 6,7:y:1,2) and S. Bareilly (I
- 268 6, 7:y:1,5), both ST 909, and in S. Saintpaul (I 4,5,12: e,h: 1,2) and S. Haifa (I 4,5,12: z,10: 1,2), both ST49,
- 269 reflected a true phylogenetic difference (Figures 3a and 3b).
- 270

271 The same higher strata population structure referred to as lineage 3 for S. enterica subspecies I, as 272 described by Achtman et al. (2012), was observed in this dataset (Figure 4). Genomes of these 273 Salmonellae are in constant flux and homologous recombination among unrelated eBGs is frequent 274 (Achtman et al. 2012, Didelot et al. 2011). Serovars in this lineage mainly consists of multiple eBGs and 275 are polyphyletic by nature. Achtman et al. (2012) suggested that the population structure of lineage 3 276 does not comprise of independent startbursts, as observed with other serovars of subspecies I, but 277 rather a connected network (Figure 4). The five most common examples of this in the current study, 278 were S. Oranienburg, S. Montevideo, S. Chester, S. Poona and S. Bredeney (Figure 4 and Supplementary 279 Table). These five serovars are not represented in the top 10 serovars submitted to SRS during this 280 surveillance period.

281

282 K-mer identification

There were 249 cultures submitted to SRS by the local hospital and regional laboratories for *Salmonella* typing that were a mix of *Salmonella* and non-*Salmonella* species. These were identified by the k-mer identification step and included 138 *Escherichia coli*, 40 *Morganella morganii*, 11 *Citrobacter species* and four *Escherichia albertii*.

287

288 Discussion

289 In their seminal 2012 paper Achtman and colleagues argued convincingly for replacing serotyping with a 290 MLST approach based on genetic population groupings for typing S. enterica (Achtman et al. 2012). The 291 key aspects of this approach that led PHE to adopt this strategy were (i) the robustness of the 292 population structure as defined by the natural eBG clusters (ii) the fact that eBG designation provides an 293 accurate representation of strain relatedness and (iii) that this approach lends itself to automation. At 294 the same time, it was necessary for PHE to maintain serovar nomenclature in order to facilitate data 295 exchange with other colleagues in the field and maintain backward compatibility with historical data. It 296 was suggested that by using the MLST approach to infer serovar, and by reporting both inferred serovar

- and ST, it would be possible to utilise the advantages of both methods and implement a state-of-the-art
 typing system while keeping disruption for reference laboratory service users to a minimum.
- 299

300 The PHE dataset of 6887 subspecies I isolates that were serotyped using both traditional phenotypic 301 methods and a derived serotype based on MLST data extracted from the genome during a 12 month 302 time frame, provided further evidence of the robustness of the ST/eBG approach to typing. The 96% 303 concordance between the two techniques in a reference laboratory setting is evidence of the validity 304 and suitability of this approach. There were 451 isolates that had to be excluded from the comparison 305 because both types of data (phenotypic and genotypic) were not available. Of these, for 94% of the 306 isolates, it was the phenotypic serotype that could not be determined indicating that WGS MLST derived 307 serotyping is more robust.

308

309 The PHE dataset included single serovars associated with multiple eBG, for example S. Typhimurium and 310 S. Newport (Sangal et al. 2010, Achtman et al. 2012) and multiple serovars belonging to the same eBG 311 but with different STs, for example S. Java (ST43) and S. Paratyphi (ST86) both belong to eBG5 (Achtman 312 et al. 2012). In both these scenarios, the correct serovar was determined from the MLST WGS data and 313 the combination of serovar and ST/eBG provided insight into the true phylogenetic relationship between 314 isolates. This data clearly supports Achtman and colleagues argument that eBG and ST designation 315 provides a more accurate representation of strain relatedness than the traditional serovar designation. 316 The phenomenon of multiple serovars belonging to the same ST (for example S. Richmond/S. Bareilly 317 and S. Haifa/S. Saintpaul) was a rare but important example of serotyping providing a higher level of 318 strain discrimination within a ST. These strains could be differentiated in silico using a tool to infer 319 serovar from the genes that determine antigenic struture, such as seqsero (Zhang et al. 2015).

320

321 Despite the implementation of WGS, a limited phenotypic serotyping facility continues to be maintained 322 at PHE in order to serotype isolates that cannot be matched to a serovar; either because the ST in the 323 MLST database has no serovar designation or the ST is a novel type. Additionally, it ensures that we 324 maintain the ability to perform the standard reference method for serotyping Salmonella. The PHE MLST 325 database is regularly up-dated to include STs recently matched to a serotype by linking the ST to PHE 326 phenotypic serotyping data and novel PSTs. This approach was adopted because at the time of analysis, 327 the Achtman MLST database was not accepting submissions generated by WGS. There was no decrease 328 in the rate at which PSTs were observed during the 12 month study period and the majority of PSTs

329 were only sampled once in that time frame. Many PSTs were SLVs of known STs, indicating that we have

- 330 not yet sampled the full diversity of known eBGs. New PSTs, not part of any previously identified eBG,
- 331 were also observed and further diversity was found within *S. enterica* subspecies II-IV and the lineage 3
- population. This suggests that there is a large amount of previously unidentified diversity within the
- 333 species *Salmonellae* associated with both domestically acquired and travel related gastrointestinal
- disease in human cases resident in England and Wales.
- 335
- 336 Isolates exhibiting monophasic properties that could not be fully serotyped phenotypically because they 337 had an incomplete antigenic structure were matched to a ST derived serotype. The monophasic variants 338 in this study mainly belonged to eBG1, eBG138 and eBG243 and previous studies have also shown that 339 monophasic variants of S. Typhimurium have emerged as a result of multiple independent genetic 340 events (Soyer et al. 2009, Switt et al. 2009, Tennant et al. 2010). Strains with monophasic properties are 341 reportable to European Centre for Disease Prevention and Control (ECDC) but cannot be determined 342 using the ST approach. Alternative strategies for determining monophasic characteristics by PCR are 343 available (Prendergast et al. 2013) and methods for extracting this information from the genome 344 sequencing data have been developed at PHE (Personal communication: Philip Ashton & Anna Lewis, 345 publication in preparation).
- 346

347 In contrast to S. Typhurmurium, where ST could not be used to determine monophasic characteristics, in

348 this study ST was able to differentiate the complex relationship between S. Java (Hazard Group (HG) 2

organism) and S. Paratyphi B (HG3) with the latter belonging to either ST42 or ST86. If this ST

 $350 \qquad \text{designation proves to be robust, MLST will facilitate the diagnosis of invasive disease and life}$

351 threatening paratyphoid fever.

352

The MLST derived serovar correlated well with the traditional serovar designation and demonstrated many advantages over traditional phenotypic serotyping. Monophasic strains with incomplete antigenic structures were accurately assigned to serotypes. Phenotypic serotyping errors, such as misinterpreting or incorrectly transcribing the antigenic structure, were avoided. Novel types were identified, confirmed and given a PST designation. Finally, this approach lends itself to automation and rapid, high-throughput processing.

360 Two main issues arose during the evaluation of the MLST approach: (i) a number of STs did not have a 361 serovar designation in the MLST database (including subspecies II to IV) and (ii) the unexpectedly large 362 number of novel STs identified. Traditional phenotypic serotyping was required to type these isolates 363 and the MLST database was modified and up-dated to incorporate the new data. Clearly, as we move 364 forward the PHE MLST database will be constantly evolving and this data will be shared with colleagues 365 in the field via existing MLST databases and their WGS compliant successors e.g. EnteroBase & BIGSdb. 366 While it is difficult to draw conclusions based on our small sample size, MLST may not currently be an 367 appropriate tool for the classification of Salmonella sub-species II-IV, due to the lack of a discrete 368 population structure of EBGs. However, non-subspecies I isolates which are mainly adapted to cold 369 blooded animals and/or reptiles contributed to less than 1.7% of the workload during the time frame of 370 the study. Although MLST approach is generally more discriminatory than serovar, it does not always 371 provide the fine resolution required for public health surveillance. Further analysis based on single 372 nucleotide polymorphisms in the core genome compared to a type strain representing the most 373 common eBGs is performed for outbreak detection and investigation (Ashton et al. 2014). 374

375 In conclusion, WGS MLST derived serotyping is an accurate, robust, reliable, high throughput typing

376 method that is well suited to routine public health surveillance of Salmonella. This approach supports

377 the maintenance of traditional serovar nomenclature and provides further insight on the true

378 evolutionary relationship between isolates, as well as a framework for fine level typing within eGBs for

379 surveillance, outbreak detection and source attribution.

380

381

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388

389 **Data Deposition**

390 All data from the Salmonella surveillance project are deposited in the BioProject of the SRA

391 PRJNA248792.

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510 *Microbiol* 53(5):1685-92. doi: 10.1128/JCM.00323-15.

- 512 Tables
- 513 Table 1. Examples where two serovars belonged to the same eBG and the same ST
- 514

Serotype	Antigenic structure	ST
Bareilly	l 6,7: y: 1,2	909
Richmond	l 6,7: y: 1,5	
Saintpaul	I 4,5,12: e,h: 1,2	49
Haifa	I 4,5,12: z,10: 1,2	
Sandiego	l 4,12: l,v: e,n,z15	20
Brandenburg	l 4,12: e,h: e,n,z15	
Uganda	3, 10: ,z13: 1,5	684
Sinstorf	l 3, 10: l,v: 1,5	
Agona	I 4,12: f,g,s:-	13
Essen	I 4,12: f,g,m:-	
Napoli	l 1,9,12: l,z13: enx	P3141
Zaiman	l 1,9,12: l,v: enx	

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516

- 518 Table 2. S. Java isolates in this study belonged to a diverse range of eBGs and STs associated with S. Java
- 519 whereas S. Paratyphi B belonged to ST86 only

	eBG5				eBG 9	eBG59	eE	3G32	eBG95	
Phenotypic	ST	ST149	ST307	ST1577	ST8	ST88	ST28	ST423	ST682	1583
serovar	43				6	/127			/1588	
Java	45	7	4	3	2*	18	6	5	2	1
Paratyphi B	0	0	0	0	12	0	0	0	0	0

521	Figures
522	Figure 1. Population structure of all Salmonella enterica isolates submitted to PHE from local and
523	regional hospital laboratories in England and Wales between April 2014 and March 2015 (see
524	Supplementary Table for details)
525	
526	Figure 2a. Novel, preliminary STs (PST) and the modal number of isolates identified per PST
527	
528	Figure 2b. The rate at which PSTs were identified throughout the time frame of the study
529	
530	Figure 3a. Phylogenetic relationship of S. Richmond and S. Bareilly (ST909) (Figure 3a) and S. Saintpaul
531	and S. Haifa (ST49) (Figure 3b)
532	
533	Figure 4. Serovars in lineage 3 mainly consist of multiple eBGs and are polyphyletic by nature
FQ 4	

Figure 1(on next page)

Population Structure of Salmonella enterica submitted to PHE

Population structure of all *Salmonella enterica* isolates submitted to PHE from local and regional hospital laboratories in England and Wales between April 2014 and March 2015 (see Supplementary Table for details)



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Trends in Preliminary Sequence Types

(A) Novel, preliminary STs (PST) and the modal number of isolates identified per PST (B) The rate at which PSTs were identified throughout the time frame of the study.



Figure 3(on next page)

Phylogenetic relationship of S. Richmond and S. Bareilly (ST909) (Figure 3a) and S. Saintpaul and S. Haifa (ST49) (Figure 3b)



Figure 4(on next page)

Phylogenetic relationship of S. Richmond and S. Bareilly (ST909) (Figure 3a) and S. Saintpaul and S. Haifa (ST49) (Figure 3b)

	<mark>52966_H14404061901-2 ['Salmonella Ha</mark> ifa', '1.1.7.7.14.14.15']
Feer Preprints	53998_H14398076401-1 ['Salmonella Haifa', '1.1.7.7.14.14.15']
	57607_H14402070601-2 ['Salmonella Haifa', '1.1.7.7.14.14.15']
	65426_H14416047001-1 ['Salmonella Haifa', '1.1.7.7.14.14.20']
	63546_H14418059101-2 ['Salmonella Haifa', '1.1.7.7.16.16.17']
	34208_H14312047901-2 ['Salmonella Haifa', '1.1.7.7.7.7.8']
	- 14533_H14204042501-1 ['Salmonella Haifa', '1.1.1.1.1.1.1']
	29937_H14302040905-1 ['Salmonella Haifa', '1.3.6.6.6.6.7']
	37784_H14322069001-2 ['Salmonella Haifa', '1.3.6.6.8.8.9']
	49729_H14392079401-1 ['Salmonella Haifa', '1.3.6.6.11.11.12']
	75923_H14160008201-1 ['Salmonella Haifa', '1.3.6.17.23.23.27']
	23476_H14258042801-1 ['Salmonella Haifa', '1.3.4.4.4.5']
	78648_H15012063801-1 ['Salmonella Haifa', '1.1.13.13.18.24.28']
	6503_H14180051001-1 ['Salmonella Haifa', '1.1.13.13.18.18.19']
	68878_H14436019101-1 ['Salmonella Haifa', '1.1.13.15.20.20.22']
	51250_H14384049601-2 ['Salmonella Haifa', '1.1.10.10.13.13.14']
	1510_H14142040401-1 ['Salmonella Haifa', '1.1.2.2.2.2.2']
	75922_H14154053101-1 ['Salmonella Haifa', '1.1.2.2.2.2.26']
	- 83324_H15046061505-1 ['Salmonella Haifa', '1.1.2.19.26.27.31']
	67557_H14462069401-2 ['Salmonella Haifa', '1.1.2.14.19.19.21']
	23469_H14258038805-1 ['Salmonella Saint-paul', '2.2.3.3.3.3.4']
	40358_H14356044201-2 ['Salmonella Saint-paul', '2.2.3.3.3.3.4']
	27788_H14274062301-1 ['Salmonella Saint-paul', '2.2.3.3.3.3.4']
	36382_H14328061401-1 ['Salmonella Saint-paul', '2.2.3.3.3.3.4']
	25243_H14282040701-1 ['Salmonella Saint-paul', '2.2.3.3.3.3']
	22242_H14260065705-2 ['Salmonella Saint-paul', '2.2.3.3.3.3']
	75666_H14512061301-1 ['Salmonella Saint-paul', '2.2.3.3.22.22.25']
	78746_H15014058001-2 ['Salmonella Saint-paul', '2.2.3.3.24.25.29']
	50317_H14374071701-2 ['Salmonella Saint-paul', '2.2.3.3.12.12.13']
	74300_H14508041701-1 ['Salmonella Saint-paul', '2.2.3.16.21.21.24']
	_ ref
	45593_H14372075905-1 ['Salmonella Saint-paul', '4.5.9.9.10.10.11']
	80547_H15030054801-1 ['Salmonella Saint-paul', '4.5.14.18.25.26.30']
	25237_H14282040101-1 ['Salmonella Saint-paul', '3.4.5.5.5.6']
	41952_H14354084901-1 ['Salmonella Saint-paul', '3.4.8.8.9.9.10']
	72306_H14434070401-2 ['Salmonella Saint-paul', '3.6.11.11.15.15.23']
	☐ 60035_H14424060601-2 ['Salmonella Saint-paul', '3.6.11.11.15.15.16']

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Salmonella lineage 3 population structure

Serovars in lineage 3 mainly consist of multiple eBGs and are polyphyletic by nature

