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Insight into Shiga toxin genes encoded by *Escherichia coli* 0157 from whole genome sequencing

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The ability of Shiga toxin-producing *Escherichia coli* (STEC) to cause severe illness in humans is determined by multiple host factors and bacterial characteristics, including Shiga toxin (Stx) subtype. Given the link between Stx2a subtype and disease severity, we sought to identify the *stx* subtypes present in whole genome sequences (WGS) of 444 isolates of STEC 0157. Difficulties in assembling the *stx* genes in some strains, were overcome by using two complementary bioinformatics methods; mapping and *de novo* assembly. We compared the WGS analysis with the results obtained using a PCR approach and investigated the diversity within and between the subtypes. All strains of STEC 0157 in this study had stx1a, stx2a or stx2c or a combination of these three genes. There was over 99% (442/444) concordance between PCR and WGS. When common source strains were excluded, 236/349 strains of STEC 0157 had multiple copies of different Stx subtypes and 54 had multiple copies of the same Stx subtype. Of those strains harbouring multiple copies of the same Stx subtype, 33 had variants between the alleles while 21 had identical copies. Strains harbouring Stx2a only were most commonly found to have multiple alleles of the same subtype (42%). Both the PCR and WGS approach to stx subtyping provided a good level of sensitivity and specificity. In addition, the WGS data also showed there were a significant proportion of strains harbouring multiple alleles of the same Stx subtype associated with clinical disease in England.

Insight into Shiga toxin genes encoded by Escherichia coli O157 from whole genome sequencing

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- 29 Conceived and designed the experiments: PA, TD, CJ
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- 34 Running title: Use of WGS for subtyping stx genes
- 35 Key words: STEC O157; WGS; stx subtyping

36 Abstract

37 The ability of Shiga toxin-producing Escherichia coli (STEC) to cause severe illness in humans is 38 determined by multiple host factors and bacterial characteristics, including Shiga toxin (Stx) 39 subtype. Given the link between Stx2a subtype and disease severity, we sought to identify the 40 stx subtypes present in whole genome sequences (WGS) of 444 isolates of STEC O157. 41 Difficulties in assembling the stx genes in some strains, were overcome by using two 42 complementary bioinformatics methods; mapping and *de novo* assembly. We compared the 43 WGS analysis with the results obtained using a PCR approach and investigated the diversity 44 within and between the subtypes. All strains of STEC O157 in this study had stx1a, stx2a or 45 stx2c or a combination of these three genes. There was over 99% (442/444) concordance 46 between PCR and WGS. When common source strains were excluded, 236/349 strains of STEC 47 O157 had multiple copies of different Stx subtypes and 54 had multiple copies of the same Stx 48 subtype. Of those strains harbouring multiple copies of the same Stx subtype, 33 had variants 49 between the alleles while 21 had identical copies. Strains harbouring Stx2a only were most 50 commonly found to have multiple alleles of the same subtype (42%). Both the PCR and WGS 51 approach to stx subtyping provided a good level of sensitivity and specificity. In addition, the 52 WGS data also showed there were a significant proportion of strains harbouring multiple alleles 53 of the same Stx subtype associated with clinical disease in England.

54 Introduction

55 Shiga toxin-producing Escherichia coli (STEC) are a rare but potentially fatal cause of 56 gastroenteritis. They are associated with a wide spectrum of disease ranging from mild to bloody 57 diarrhoea, through to haemorrhagic colitis and haemolytic uraemic syndrome (HUS) (1). The 58 main reservoir of STEC in England is cattle, although it is carried by other animals, mainly 59 ruminants. Transmission to humans occurs through direct or indirect contact with animals or their 60 environments; consumption of contaminated food or water, and through person-to-person 61 contact. Each year, there are approximately 900 cases of STEC O157 in England confirmed by 62 the Gastrointestinal Bacteria Reference Unit (GBRU) at Public Health England.

The primary STEC virulence factor responsible for the most serious outcomes of human infection is Shiga toxin (Stx), an AB₅ toxin that targets cells expressing the glycolipid globotriaosylceramide (Gb3), disrupting host protein synthesis and causing apoptotic cell death (2). Renal epithelial cell membranes are enriched for Gb3 resulting in the kidneys bearing the brunt of Stx toxicity and, in 5-10% of cases, this leads to the development of Hemolytic Uremic Syndrome (HUS) (1). There are two types of Stx; Stx1 and Stx2 and both have multiple subtypes. These subtypes can be differentiated using a PCR targeted at the encoring genes described by (3). In addition, a web-based tool, VirulenceFinder has been developed which uses a *de novo* assembly followed by BLAST approach to identify subtypes of Stx (4). This system was shown to have good, but not perfect, agreement with PCR, although how it handles strains that encode both *stx2a* and *stx2c* is uncertain as no strains that encoded both these subtypes were examined (4). The ability of STEC to cause severe illness in humans is determined by multiple bacterial factors (in addition to host factors), including Shiga toxin subtype. There is evidence that the Stx2a subtype is significantly associated with progression to HUS (5, 6).

77 As part of a project investigating the utility of whole genome sequencing (WGS) for public health 78 surveillance and outbreak investigation of foodborne pathogens, high throughput, short read 79 Ilumina GAII sequence data for 444 strains of STEC O157 isolated in England between 2009 80 and 2013 was obtained. We determined the presence, or absence of the Stx encoding genes 81 stx1 and/or stx2 genes in all 444 isolates of STEC O157 from the genome sequence data. Given 82 the link between Stx subtype and disease severity, we also sought to identify the stx subtypes 83 present using bioinformatics methods and to compare the results with those obtained using the 84 PCR scheme of (3).

85 WGS high throughput short read technologies are rapid and low cost compared to Sanger 86 sequencing but it was recognised early on that assembling short reads would be problematic (7). 87 A major difficulty in assembly is the presence of repeat sequences that are longer than the read 88 length. Furthermore the study by (3) clearly demonstrated that as well as a high level of similarity 89 between stx2a, stx2c and stx2d there is also considerable diversity within each of these 90 subtypes. The assembly of stx into one contig in strains of STEC O157 containing both stx2a 91 and *stx2c* is difficult because the regions of variation between these subtypes are concentrated 92 at the 5' and 3' ends of the coding DNA sequence (CDS), with a largely homogenous region in 93 the centre. Existing methods for subtyping stx from short read data have not been tested against 94 strains encoding stx2a and stx2c (4). This region of 100% identity is often longer than the typical 95 read length of short read sequencing technologies, so contiguous assembly of both subtypes 96 relies on information from the paired end reads, which has a limited ability to resolve repeats up 97 to the average fragment size (550-700 bp for Illumina Nextera mate-pair). The STEC O157 98 Sakai reference genome encodes 18 pro-phage that show a large degree of modularity and 99 similarity (8), this further complicates assembly of these regions (9). These difficulties have led 100 to a relative paucity of data on the presence of subtypes of Stx within the E. coli population 101 despite large WGS projects.

102 In this study a dual bioinformatic approach was taken, using both mapping and de novo

103 assembly to determine *stx* subtype. The results of the bioinformatic analysis were compared to 104 the results from the PCR typing method (3). In addition, the diversity within and between the *stx* 105 subtype genes were investigated and evidence that certain strains contained multiple copies of 106 the same *stx* subtype was assessed.

107 Methods

108 Strain selection

109 A total of 444 isolates of STEC O157 submitted to GBRU for confirmation and typing were 110 selected for sequencing, 365 from 2012 representing approximately one third of the culture 111 positive isolates (1002 total isolates) received by the reference laboratory that calendar year 112 from laboratories in England, Wales and Northern Ireland, 67 English historical isolates 113 submitted to GBRU between 1990 and 2011 and 12 isolates from 2013. The collection contained 114 strains from sporadic cases, known outbreaks, household clusters, and serial strains isolated 115 from the same patient. However, only sporadic strains and a single strain from any related cases 116 (e.g. household, outbreak) were included in the diversity and multiple allele analysis. A total of 18 117 phage types were represented.

118 Sequencing

Genomic DNA was fragmented and tagged for multiplexing with Nextera XT DNA Sample Preparation Kits (Illumina) and sequenced at the Animal Health Veterinary Laboratory Agency (Weybridge) using the Illumina GAII platform with 2x150bp reads. Multiplexing allowed 96 samples to be sequenced per run. Sequencing data with a phred score below 30 or a read length below 50 were removed from the data set using Trimmomatic (11). FASTQ data is available from the NCBI Short Read Archive, BioProject accession PRJNA248064.

125 Subtyping of stx by assembly

126 High quality reads were assembled using Velvet v1.2.03 (10) with k-mer chosen using VelvetK 127 (http://bioinformatics.net.au/software.velvetk.shtml). The resulting contigs were then compared 128 against a set of stx reference genes (stx1a, L04539.1; 1c, Z36901.1; 1d, AY170851.1; 2a, 129 X07865.1; 2b, X65949.1; 2c, AB071845.1; 2d, AY095209.1; 2e, AJ249351.2; 2f, AB472687.1; 130 2g, AY286000.1) using BLASTn within the BioPython framework (12). Only matches with an E-131 value less than 1x10⁻²⁰ were included in further analysis. For each strain, the length of the best-132 matched sequence (in terms of the BLAST score) between the contigs and each stx reference 133 gene was calculated. For example where both stx2a and stx2c were present, there may be five 134 query sequences each of 600 bp. If three of them matched stx2a with the highest BLAST score, 135 and two of them matched stx2c with the best BLAST score, then stx2a would score 1200 and

137 Subtyping of stx by mapping

138 An alignment of stx1a, stx1c, 1d, 2a, 2b, 2c, 2d, 2e, 2f and 2g sequences (taken from Scheutz et 139 al., 2012) was generated using ClustalW within the MEGA 5 software package (13). Three bases 140 for each reference subtype that, when combined, had 100% sensitivity and specificity for each 141 subtype were identified. High quality sequencing reads were mapped to a set of reference stx 142 genes (same genes as BLAST approach described above) using BWA-MEM (http://bio-143 bwa.sourceforge.net/). Reads that mapped to more than one place in the reference set (i.e. 144 ambiguous reads) were removed from the resultant SAM file using Samtools (14). If at least 10 145 reads and 90% of the total reads concordantly mapped to all three discriminatory positions for a 146 specific subtype, then a positive match was returned for that subtype.

147 Determination of the presence of multiple alleles of the same *stx* subtype by mapping148 depth

149 Multiple copies of the same stx allele could be identified using two complementary approaches. 150 In the first approach, reads were mapped to the stx reference genes, with ambiguous mapping 151 allowed. Then the coverage of each stx allele, which had been identified by the mapping and 152 assembly methods described above, was calculated using the Samtools 'depth' option. A 153 distribution of mapping depth in all the strains that were positive for one particular Stx subtype 154 was plotted revealing a bimodal distribution with the higher mode approximately twice the lower 155 mode. The lower mode represented strains with only one copy of stx and the higher mode 156 represented strains with multiple alleles of stx. There was no bimodal distribution of mapping 157 depth for strains that encoded both stx2a and stx2c, due to the redundant mapping between 158 these two strains. For example, if a strain encoded stx2a only and mapped to an stx2c reference 159 gene, it showed approximately one third of the average coverage compared to if it were mapped 160 to an stx2a gene. This cross-mapping meant that multiple alleles of the same stx subtype could 161 not be detected in strains that encoded both *stx2a* and *stx2c*.

162 In the second approach, the bam file resulting from the mapping of the reads to the *stx* reference 163 set was parsed for mixed positions with the minority variant present in at least 25% of reads i.e. 164 one position in the reference gene was mapped by two different bases. Only strains that were 165 known to encode only one of stx2a or stx2c from the subtyping results were analysed, as the 166 high similarity between stx2a and stx2c can result in pseudo-mixed bases when compared with 167 *stx* reference genes. If there were mixed bases present in an alignment (where the depth was 168 greater than 20x and minority variant present in greater than 15% of reads), from a strain

- 169 encoding only one of stx2a or stx2c, the presence of multiple alleles of a specific stx subtype
- 170 that vary by at least one base was assumed to be present (supplementary materials Figure 1).

171 Diversity of *stx* associated with STEC O157 in the England, Wales and Northern Ireland

172 The stx genes that were successfully assembled into a single contig were extracted from the de 173 novo genome assemblies using BLAST and aligned. Only strains that subtyping had shown to 174 encode one of stx2a or stx2c were included in this part of the study. Where the complete 175 sequence of both stxA and stxB, including the intergenic region, was assembled into a single 176 contig, the CDSs were aligned and represented in minimum spanning trees generated using 177 Bionumerics v6 (http://www.applied-maths.com/bionumerics). Strains where the stxA and stxB 178 subunits could not be assembled into a single contig (e.g. due to the presence of multiple copies 179 of the same stx subtype with sequence variation between them), were aligned against a 180 reference gene and the resulting Sam file was parsed using custom python scripts to identify 181 variant positions. The sequences of stx1a, stx2a and stx2c present in the strains investigated 182 here were compared with a representative sample of stx subtype sequences in the National 183 Centre for Biotechnology Information (NCBI, <u>http://www.ncbi.nlm.nih.gov/</u>) nucleotide database to 184 assess diversity and identify novel alleles.

185 Stx real-time qPCR and block-based subtyping PCR

DNA was prepared by inoculating single colonies into 490ul distilled water and boiled in a water bath for 10 mins. The real-time qPCR described by the European Union Reference Laboratory (EURL) for *stx1* and *stx2* was performed as previously described (15). For the block-based subtyping PCR, DNA was amplified on a block-based DNA Engine platform using the stx subtyping primers and amplification parameters described by (3). Amplified DNA was electrophoresed on a 2% gel, stained with ethidium bromide and visualised with UV light.

192 **Results**

193 Stx subtyping of 444 STEC O157 in the UK – comparison of NGS and PCR

Subtyping results from PCR and WGS were identical in 422/444 strains (Table 1), there was agreement for 85 *stx2a* encoding strains, 153 *stx2a/stx2c* strains and 187 *stx2c* strains. When the subtyping PCR was repeated for the 22 discordant strains, results for 442/444 strains were identical. Of the two strains where PCR and sequencing were discordant, one strain was positive for *stx2c* by PCR but no *stx2c* was identified in the sequencing data by the bioinformatics algorithms described here, and one strain was positive for *stx2a* by sequencing that was not detected by PCR. The strain that had a positive PCR result for *stx2c* but no corresponding result in the WGS data had a very low level of mapping (54 reads, <7x average coverage) to *stx2c*. This was not enough to definitively identify *stx2c* by either the mapping or assembly algorithms, although is indicative of its presence. The *stx2a* gene sequence of the strain that was PCR negative but that had *stx2a* reads identified in the WGS data, was analysed for mutations in the primer binding sites, but none were identified.

206 **Detection of multiple alleles of** *stx*

A subset of 349 sporadic strains of STEC O157 (i.e. not from same person, household or outbreak) was investigated for the presence of multiple alleles of the same subtype of *stx*. The detection of multiple copies of the same *stx* subtype was performed using two complimentary methods (i) mapping and determining the short read coverage of a particular *stx* subtype relative to the coverage of the whole genome and (ii) the detection of mixed bases (coverage >20x, minority variant >15%, see supplementary material Figure 1) in an alignment to a single reference gene.

214 The stx1a gene was detected, in 6 different combinations with other subtypes/alleles, in 100 215 strains from independent sources (Table 2). For clarity, the relative coverage of stx1a in three of 216 the observed combinations (totalling 77 strains) is presented (Figure 1). The relative coverage of 217 stx1a in all 6 combinations observed in the 100 stx1a strains can be seen in supplementary 218 material Figure 2. In Figure 1, a bimodal distribution was clear, with the higher mode being 219 approximately twice as high as the lower mode. There were 11 strains in the higher mode 220 (Figure 1). When the *stx1a* alignments were examined for the presence of mixed bases, there 221 were 97 strains with no mixed bases and three strains that had at least one mixed base position. 222 The relative coverage was examined in the context of the presence of mixed bases and strains 223 with no mixed bases had a median relative coverage of 1.7x whereas strains with at least one 224 mixed base had a mean coverage of 2.8x (Figure 1). There were nine strains without mixed 225 bases that had relative stx1a coverage closer to the average of mixed base position strains than 226 the average of no mixed bases suggesting that two identical copies of the stx1a gene were 227 present.

There were 210 isolates that encoded stx2a, either alone or in combination with other subtypes (Table 2). For clarity, only the relative coverage of stx2a from the 73 strains that encoded only stx2a were presented in Figure 2 (the relative coverage of stx2a in all strains which encoded this subtype can be seen in supplementary material Figure 3). Inspection of the distribution of coverage of the short reads in stx2a revealed at least two modes within the relative coverage of

233 stx2a, with the upper mode (1.8x) being twice that of the lower (0.9x) (Figure 2). Of the 70 234 strains that encoded stx2a but not stx2c, 31 (42%) had short read coverage in the upper mode 235 (1.8x), of which all 31 had mixed base positions in their alignments, indicating the presence of 236 two alleles of stx2a. When the mixed position data was compared with the relative coverage 237 distribution, the mean relative coverage of the strains with mixed positions of stx2a was 1.9x 238 while the coverage in strains with no mixed positions was 0.75x (Figure 2). There were 1 (n = 239 29), 2 (n = 1) or 3 (n = 1) positions with mixed bases between the alleles in the 31 strains with 240 multiple copies.

The relative coverage of the 279 isolates that encoded stx2c was calculated. For clarity, only the relative coverage of the 139 strains that encoded stx2c but not stx2a are presented in Figure 3. The relative coverage of stx2c in all 279 strains can be seen in the supplementary materials and analysis of the distribution of relative stx2c coverage showed that the majority of these strains fell into an approximately normal distribution around 1x relative coverage (Figure 3). Twelve (8.6%) of the 139 strains had a relative coverage > 1.5x but no mixed base positions were found.

248 Diversity of *stx* associated with STEC O157 in the UK

249 The diversity of stx found in a subset of 349 sporadic strains of STEC O157 (i.e. not from same 250 person, household or outbreak) was investigated. Ninety-seven complete stx1a genes from this 251 study were compared with nine stx1a alleles from NCBI, and a total of 16 variant positions were 252 identified along the 1392 bp length of the gene. Of the five different alleles present in the strains 253 investigated here, three were not present in the NCBI database (as of 06/23/14, Figure 4). The 254 most frequently observed allele accounted for 76 (78.3%) of the 97 assembled stx1a genes from 255 this study, while the second most frequently observed allele accounted for 16 (16.5%) stx1a256 genes. Both the most frequently observed alleles had been previously identified in E. coli 257 O103:H2 (BAI33872.1) and E. coli O157:H7 (EF079675.1), respectively. The five remaining 258 stx1a genes comprised three different alleles, none of which had been previously submitted to 259 the NCBI database ((as of 06/23/14), although they were all within a single variant of previously 260 observed alleles (Figure 4).

The 38 fully assembled *stx2a* genes from this study were compared with 21 *stx2a* alleles from the NCBI nucleotide database. There were a total of 48 variant positions in a 1442 bp alignment of the 59 *stx2a* genes that included 25 different alleles (Figure 5). Of these 25 alleles, six were present in the strains investigated here. The most frequently observed allele was a single variant from a *stx2a* allele observed in *E. coli* O157 (AF524944.1), *E. coli* O111 and *E. coli* O145 and was present in 18 (47.3%) of the strains in this study. The second most frequently observed allele was present in 11 (28.5%) strains and was widely distributed, including in Bacteriophage 933W (X07865). The other nine strains represented four alleles, two of which had been identified before. The remaining allele (from strain H124840173) was highly divergent from the other *stx2a* alleles, with six SNPs compared to any previously identified *stx2a* gene and 11 variants compared to the closest *stx2a* observed in this study. Interestingly, this strain was a sorbitolfermenting (SF) STEC O157, the only SF strains to be included in this study.

273 There were 132 fully assembled stx2c genes from this study that were compared with 18 274 previously identified *stx2c* alleles from NCBI. There was a total of 59 variant positions along the 275 1441 bp gene alignment of the 150 stx2c sequences, comprising 22 unique alleles, of which 276 seven were identified in the strains analysed here (Figure 6). The most frequently observed 277 allele accounted for 115 (87.1%) of the 132 fully assembled stx2c genes. This allele had been 278 previously observed in a single E. coli O157:H- strain (AB015057.1). The 17 other stx2c genes 279 represented six distinct alleles that, with one exception, were within two variants of the most 280 frequently observed allele (Figure 6). There were two strains encoding the most divergent 281 observed stx2c allele, with six variant positions compared with the most frequently observed 282 allele. This divergent allele had been previously identified in *E. coli* RM10648 (KF932369.1).

283 Although the complete gene sequence could not be determined for strains that had more than 284 one copy of a stx subtype, an alignment of the reads against a reference was analysed to 285 identify variant positions. Of the three strains with multiple alleles of stx1a, all three had the 286 same four variant positions. There was one SNP in all three multiple-stx1a strains that was not 287 previously identified in the stx1a sequences described above or in the NCBI reference 288 sequences. Of the 30 strains with multiple copies of stx2a, 28 had only a single variant position 289 that was the same in all 28 strains and that had been previously identified. Of the other two 290 strains, one had the same SNP as the 28 other mixed position strains and an additional SNP 291 that had not been previously observed in the strains described in this study above or in the NCBI 292 reference strains. The final strain had three unique mixed positions, all of which had been 293 previously observed in this study.

294 Discussion

In this study we have developed novel, robust and highly accurate methods for subtyping of *stx*

- from short read sequence data, validating this method against PCR for 444 STEC O157 isolates.
- 297 Furthermore, we have mined the WGS data to show that a significant proportion of strains
- 298 encode multiple copies of the same subtype of Shiga-toxin gene. The diversity of *stx* genes from

300 There was over 95% initial agreement (422 of 444 strains) between WGS subtyping and PCR 301 subtyping in determining subtypes of stx2 which shows that WGS is an acceptable method for 302 subtyping stx in O157. The strains where there were discrepancies between WGS and PCR 303 were subjected to a repeat subtyping PCR, after which all but two of the discrepancies became 304 concordant. One possible reason for the discrepancy between the initial and repeat PCR results 305 is the high stringency of the subtyping PCR. During a multi-centre evaluation of the subtyping 306 PCR, there were differences observed in the subtyping results obtained between different 307 laboratories and these were ascribed to the use of different reagents and thermocyclers, with the 308 main source of variability thought to be the use of different polymerase. While the taq 309 polymerase recommended by (3) was used here, variations in other laboratory reagents and 310 equipment may have resulted in the discrepancies. The excellent concordance between the 311 PCR and WGS results, even despite the problems associated with analysis of homologous 312 genes using short read data, provides evidence of the accuracy of the bioinformatics algorithm 313 showing that WGS could replace PCR for subtyping.

314 Using mapping coverage to detect multiple copies of the Stx phage has been described 315 previously using more challenging metagenome data (16). The novelty of this work is to use the 316 mapping coverage of stx relative to the average coverage of the whole genome to identify 317 strains encoding multiple alleles of the same stx subtype. There are stx sequences in the NCBI 318 database that indicate that multiple alleles of the same subtype encoded by the same strain 319 have been previously observed i.e. these sequences contain ambiguous bases. However, the 320 studies associated with these sequences make no mention of the possibility of multiple alleles 321 (17, 18, 19, 20). The presence of multiple alleles of the same stx type has been previously 322 identified by WGS (21), however this is the first study to present a large sale comparison of this 323 method with PCR subtyping. Some of the ambiguous positions in sequences in the NCBI 324 database were the same positions in stx as the mixed positions observed in this study, 325 supporting the evidence that multiple alleles exist and are present in the same strain. While 326 mapping of short reads has been successful at detecting multiple copies of the same subtype, it 327 has not been possible in strains that encode stx2a and stx2c due to ambiguous mapping 328 between these types (see supplementary materials, Figures 2-4). For characterisation of these 329 stx2a/stx2c strains, and full characterisation of the insertion sites and genomic context of the stx 330 alleles in strains encoding multiple copies of the same subtype, longer sequencing reads from 331 e.g. PacBio, are needed. There was also evidence that some strains of STEC O157 encoded 332 multiple alleles of both stx1a and stx2c, further characterisation of these strains to determine whether they had a genetic determinant that made Stx phage acquisition more likely would beinteresting.

335 The functional implication of encoding multiple alleles of the same stx subtype remains unclear. 336 Three hypotheses to explain the 9% prevalence of strains encoding multiple alleles of the same 337 subtype are (i) these strains are more likely to cause symptomatic human disease (ii) these 338 strains have an fitness advantage which increases their chance of being present in the 339 environment (iii) carrying multiple alleles of the same subtype is 'merely' a side effect of the 340 recombinogenic capacity of Stx phage, which confers no phenotype. It is interesting that while 341 the multiple alleles of stx2a and stx2c always seem to have nucleotide differences, this is a 342 minority in the strains that encode multiple copies of stx1a. The close sequence relationship 343 between the multiple copies of the same subtypes raises the question whether they are derived 344 from multiple insertions by different Stx phage, or a phage or stx gene duplication.

345 This study reports on the diversity of stx observed in STEC O157 in the UK (except Scotland) 346 between 1990 and 2013, with a focus on 2012. Although there are 10 described subtypes of 347 stx1 and stx2 combined (3), in an examination of 444 strains covering a wide temporal spread 348 and range of phage types, only three subtypes (*stx2a, stx2c* and *stx1a*) were observed. Previous 349 studies examining strains from cattle and humans similarly found only stx2a, stx2c and stx1a 350 (22). The most diverse stx identified here was stx2a, followed by stx1a and then stx2c (Figures 351 4-6). The majority of stx2c were of a single genotype, and all the novel alleles identified were 352 within a single SNP of the majority genotype. This difference in diversity observed between 353 stx2a and stx2c is interesting considering that the background diversity of these two subtypes is 354 largely similar (3). Further studies in this laboratory aim to determine the phylogenetic context of 355 isolates encoding these two subtypes. This study also described 10 novel alleles of stx, with the 356 most diverse being an stx2a sequence 6 SNPs from any previously described stx2a. The fact 357 that this diverse stx was observed in a sorbitol fermenting strain indicates that there may be a 358 significant reservoir of stx diversity other serotypes of STEC. The majority of novel alleles had 359 sequences that were single nucleotide variants to previously described sequences.

This study is the first to describe stx subtyping by both PCR and WGS methods in a large number of strains of STEC O157. Both the PCR and WGS approaches to *stx* subtyping provided a good level of sensitivity and specificity. The WGS data also showed that a significant proportion of strains of STEC O157 harbour multiple alleles of the same Stx subtype. The functional significance of multiple alleles of the same subtype remains unclear, although this is the subject of on going work. Furthermore, the WGS analysis highlighted 10 novel alleles of *stx*

- 366 identified in this study and enabled us to study the diversity of stx sequences in a population of
- 367 STEC O157 associated with clinical disease in England.
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 438 Escherichia coli O157 isolates from Australia and the United States provides evidence of
 439 geographic divergence. Appl. Environ. Microbiol. **79**:5050–8.

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440 Tables and Figures

441 Table 1: Comparison of stx2 subtyping of 444 strains by sequencing and PCR. Strains that had

442 discrepant results between sequencing and PCR were subjected to a 'second pass' PCR.

Subtype	Sequencing Results	Subtyping PCR results - 1st pass	Subtyping PCR results - 2nd pass
2a	82	89	82
2c	194	196	196
2a/2c	167	155	166
No result	1	4	0
Total	444	444	444

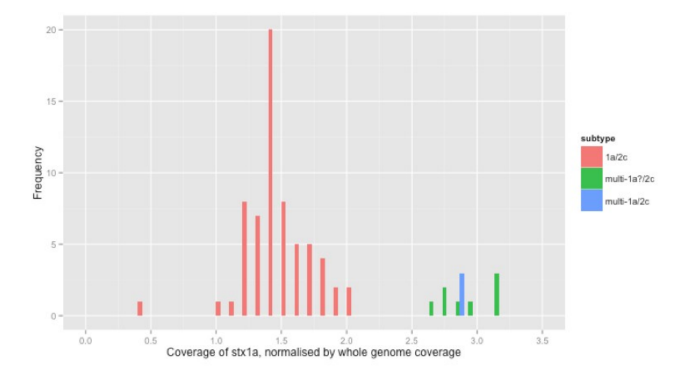
Table 2: Frequency of stx subtype profiles including stx1, derived from WGS analysis, not

444 including outbreak strains. When a multi subtype result has a '?', it indicates that the only

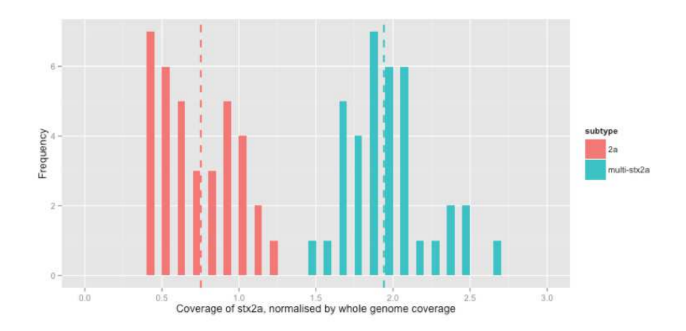
evidence suggesting the presence of multiple copies was the relative coverage (as opposed to

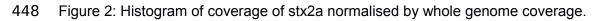
446 having mixed positions as well).

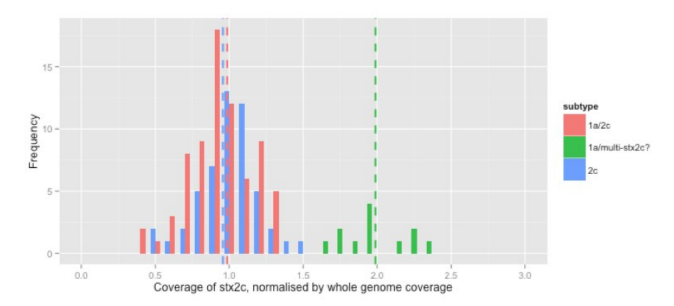
	Frequenc
<i>stx</i> profile	У
1a/2a	9
1a/2a/2c	3
1a/2c	64
1a/multi-stx2c?	10
2a	30
2a/2c	136
2c	51
multi-1a?/2c	9
multi-1a/2c	3
multi-stx2a	31
multi-stx2c?/multi-	
1a?	2
No stx detected	1



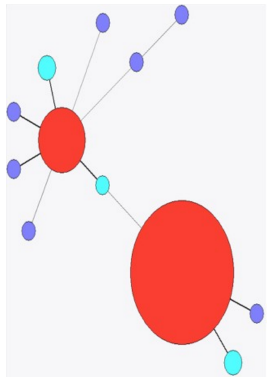
447 Figure 1: Histogram of coverage of stx1a normalised by whole genome coverage



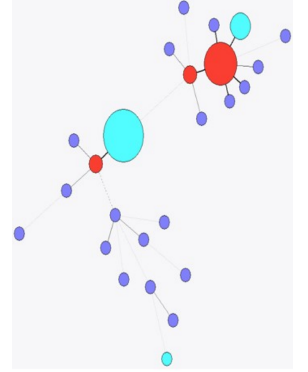




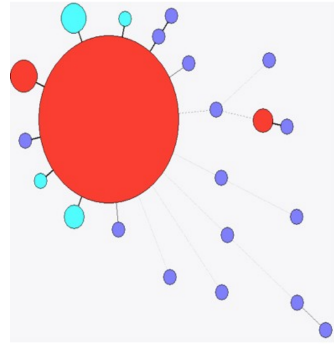
449 Figure 3: Histogram of coverage of stx2c normalised by whole genome coverage



450 Figure 4: Minimum spanning tree stx1a. Red = previously identified and observed in this study, 451 purple = previously identified but not observed in this study, light blue = novel allele.



452 Figure 5: minimum spanning tree *stx2a*. Colour as in Figure 4



453 Figure 6: Minimum spanning tree of *stx2c*. Colour as in Figure 4.

454 Supplementary figures

- 455 Supplementary Figure 1: Mixed position in an *stx2a* gene. Variant from reference highlighted.
- 456 Supplementary Figure 2: Histogram of coverage of stx1a normalised by whole genome coverage
- 457 Supplementary Figure 3: Histogram of coverage of stx1a normalised by whole genome coverage
- 458 Supplementary Figure 4: Histogram of coverage of stx1a normalised by whole genome coverage