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# The Molecular Epidemiology of Resistance in Cefotaximase-Producing *Escherichia coli* Clinical Isolates from Dublin, Ireland.

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1 **Title:**

2 The molecular epidemiology of resistance in cefotaximase (CTX-M)-producing *Escherichia coli*  
3 clinical isolates from Dublin, Ireland

4

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## 21 **Introduction**

22 Extended-spectrum  $\beta$ -lactamases (ESBLs) are amongst the most important resistance determinants in  
23 Enterobacteriaceae, conferring resistance to oxyimino-cephalosporins. ESBLs have increased  
24 remarkably in diversity and range recently, predominantly due to the evolution of cefotaximase  
25 (CTX-M)-type enzymes, which are now the most common ESBLs worldwide, and are often found on  
26 widespread transferable plasmids in *Escherichia coli* responsible for infections in both the community  
27 and hospitals<sup>4</sup>. In addition to human-to-human transmission, the contribution of zoonotic transmission  
28 by direct contact and through the food chain, to the diversity and mobilisation of *bla*<sub>CTX-M</sub> genes is  
29 increasingly recognised.<sup>14,18,21</sup> CTX-Ms are organized into five clusters (groups 1, 2, 8, 9, and 25)  
30 based on their amino acid sequences ([www.lahey.org/studies/webt.stm](http://www.lahey.org/studies/webt.stm)), with group 1 (including  
31 CTX-M-1, -3 and -15) and group 9 enzymes (including CTX-M-9, -14 and -27) widespread in  
32 Europe. The insertion sequences (ISs), *ISEcp1* and *ISCR1* are responsible for the mobilization of  
33 *bla*<sub>CTX-M</sub> and drive their expression. CTX-M genes are frequently found downstream of *ISEcp1* within  
34 modular multidrug-resistance regions (MRRs) on ESBL plasmids<sup>9,29</sup> or are linked to *ISCR1* on  
35 complex class 1 integrons which also bear highly variable gene cassette regions encoding resistance to  
36 multiple antimicrobial classes. The surveillance of non *bla*<sub>CTX-M</sub> - associated integrons in nosocomial  
37 ESBL is important as ESBL genes co-localised on the same plasmid may be co-selected in the gut  
38 flora of animals where these antibiotics are used and may be a source of zoonotic transfer to humans  
39 through the food chain. Multiple copies of IS26 are also found on MRRs of ESBL plasmids and have  
40 contributed to the spread of *bla*<sub>CTX-M</sub> by facilitating genetic rearrangements of these regions between  
41 ESBL plasmids.<sup>22,29</sup> The staggering propensity for recombination and transposition, afforded by this  
42 genetic arrangement, has resulted in the evolution of mosaic ESBL plasmids of narrow and broad host  
43 ranges that confer multidrug-resistance, including to the carbapenems<sup>2</sup>. In the healthcare setting, these  
44 plasmids are likely to be vertically and horizontally disseminated, as the multi-drug resistant (MDR)  
45 phenotype they confer provides a survival advantage to bacteria in the antimicrobial laden  
46 environment. Their association with widespread successful *E. coli* clones, such as the pandemic  
47 O25b-ST131 clone, further facilitates their dissemination by clonal expansion<sup>2</sup>.

48 We previously investigated the genetic relatedness of 100 ESBL-producing *E. coli* (ESBL-  
49 EC) from North Dublin<sup>3</sup> revealing the widespread dissemination of ST131 and other clones within  
50 Beaumont Hospital, Dublin and the local community. The aims of the present study were: (1) to  
51 characterise the mobile genetic elements, including the ISs, integrons and plasmids to which *bla*<sub>CTX-M</sub>  
52 and other resistance genes were associated in this collection with reference to the published literature  
53 from human and animal *E. coli* and (2) to determine the potential for horizontal transfer of CTX-M  
54 plasmids from clinical *E. coli* to laboratory strains.

55

## 56 **Materials and Methods**

### 57 **Bacterial strains and culture conditions**

58 One hundred ESBL-EC clinical isolates were collected as part of a previous study between January  
59 2009 and December 2010 in Beaumont Hospital, Dublin, Ireland and were previously subjected to  
60 routine diagnostic antimicrobial susceptibility tests and PFGE, which identified 12 clusters A-L.<sup>3</sup>  
61 Appropriate control strains for beta-lactamase negative, *bla*<sub>TEM</sub>, *bla*<sub>SHV</sub> and each of the five *bla*<sub>CTX-M</sub>  
62 groups and epidemic UK strain A were sourced from the American Type Culture Collection (ATCC)  
63 and the National Collection of Type Cultures (NCTC). A sodium azide-resistant *E. coli* strain J53 was  
64 the recipient for conjugations and was a gift from Prof. Martin Cormican, Department of  
65 Bacteriology, National University of Ireland, Galway. *E. coli* ElectroMAX™ DH5α-E™ (Invitrogen)  
66 was the recipient for transformations. All isolates were routinely grown on Mueller-Hinton (MH)  
67 agar.

### 68 **Characterisation of resistance genes and their associated genetic elements**

69 DNA was prepared from overnight cultures using the Wizard® Genomic DNA Purification Kit  
70 (Promega, Madison, WI, USA). The carriage of *bla*<sub>TEM</sub>, *bla*<sub>SHV</sub> and the five *bla*<sub>CTX-M</sub> gene groups in  
71 the isolates was investigated by multiplex PCR using previously described primers and cycling  
72 conditions.<sup>17,30</sup> Specific *bla*<sub>CTX-M</sub> alleles and their upstream genetic environments were identified by

73 PCR mapping and were fully sequenced. The genes, *bla<sub>SHV</sub>* and *bla<sub>TEM</sub>* were sequenced in isolates that  
74 were *bla<sub>CTX-M</sub>*-negative. However, resource restraints prevented sequencing of the upstream genetic  
75 environments of *bla<sub>TEM</sub>* and *bla<sub>SHV</sub>* alleles in favour of the more widely disseminated *bla<sub>CTX</sub>* alleles.  
76 PCR and sequencing primers used to detect class 1 and 2 integron motifs, *ISEcp1*, *IS26*, and *ISCR1*  
77 and their arrangement in relation to *bla<sub>CTX-M</sub>* are listed in Table S1. Sequencing was performed by  
78 Source BioScience (Dublin, Ireland) and GATC Biotech (Köln, Germany) and sequence analysis was  
79 carried out using CLC Main Workbench 6.6.2 (CLC Bio, Aarhus, Denmark). Representative integron  
80 variable region amplicons of each different size were sequenced and used to perform BLAST searches  
81 on the National Centre for Biotechnology Information (NCBI) nucleotide databases.

82

### 83 **Transfer of CTX-M plasmids by conjugation and transformation**

84 Conjugation was attempted by broth mating following the method of Gray *et al.* with modifications<sup>10</sup>  
85 for all ESBL-EC strains except the 33 strains that clustered with the single reference UK strain A  
86 (PFGE cluster A), whose ESBL plasmids lack the necessary conjugation machinery.<sup>29</sup> Briefly, after 4  
87 h aerobic growth in tryptone soya broth (TSB) cultures of donor and recipient strains were mixed at  
88 1:1 ratio and incubated at 37°C overnight. Mating mixtures were pelleted, rinsed and serially diluted  
89 in PBS. Transconjugants were selected using MH agar containing 100 µg/ml sodium azide and 1  
90 µg/ml cefotaxime. Presumptive transconjugants were confirmed by PCR<sup>5</sup> and *bla* multiplex PCR<sup>17</sup>  
91 with reference to donor and recipient strains, using 10 µl of cleared cell lysate as the template. Where  
92 conjugation was unsuccessful, transformation of ESBL plasmids into *E. coli* DH5α was attempted by  
93 electroporation performed at 1700 V and at a time constant of 4.8-5.0 ms using an electroporator  
94 (Eporator®, Eppendorf, UK).

95

### 96 **Characterisation of CTX-M plasmids**

97 Cefotaxime (CTX) minimum inhibitory concentration (MIC) assays were performed and interpreted  
98 using the guidelines of the Clinical and Laboratory Standards Institute (CLSI) for all clinical isolates  
99 and their transconjugants/transformants using Etest® strips (bioMérieux).<sup>6</sup> High-level cefotaxime  
100 resistance was defined as MIC  $\geq$ 128  $\mu$ g/ml. Plasmid DNA was isolated from clinical isolates and their  
101 transconjugants/transformants using the phenol-chloroform extraction method of Kado and Liu.<sup>12</sup> The  
102 presence of *bla*<sub>CTX-M</sub> genes, associated IS elements and integrons in plasmid extracts of  
103 transconjugants/transformants was investigated by PCR as described for clinical isolates. CTX-M-  
104 containing plasmids were sized by S1 PFGE.<sup>1</sup> Plasmid incompatibility typing was carried out on  
105 transconjugant/transformant plasmid preparations and clinical isolates using PCR-based replicon  
106 typing (PBRT). Transferable IncF plasmids were further characterised by replicon sequence typing  
107 (RST) and IncN and IncI plasmids were further characterised by plasmid multi-locus sequence typing  
108 (pMLST). All typing schemes were performed and interpreted using the methods and databases  
109 available at <http://pubmlst.org/plasmid/>.

110

## 111 **Results**

### 112 **Genotypic characterisation of ESBL genes in *E. coli* clinical isolates**

113 Investigation of the carriage of *bla*<sub>CTX-M/TEM/SHV</sub> genes revealed that 94% of isolates were *bla*<sub>CTX-M</sub>  
114 gene positive. *Bla*<sub>TEM</sub> genes were the second most common (46%) and 5% of clinical isolates carried a  
115 *bla*<sub>SHV</sub> gene. Group 1 was the most common cefotaximase gene cluster (80/94; 85%) and comprised  
116 66 *bla*<sub>CTX-M-15</sub>, 7 *bla*<sub>CTX-M-1</sub>, 4 *bla*<sub>CTX-M-3</sub>, 2 *bla*<sub>CTX-M-55</sub> and 1 *bla*<sub>CTX-M-32</sub>. Thirteen group 9 (13/94; 14%)  
117 were identified, comprising 8 *bla*<sub>CTX-M-14</sub>, 3 *bla*<sub>CTX-M-9</sub> and 2 *bla*<sub>CTX-M-27</sub>. A single isolate contained a  
118 *bla*<sub>CTX-M-2</sub> gene, but no group 8 or group 25 CTX-M genes were detected.

119

### 120 **Specific *bla*<sub>CTX-M</sub> genes and their upstream genetic environments**

121 The arrangement of *bla*<sub>CTX-M</sub> genes and associated upstream IS elements are summarised in Table 1,  
122 with reference to identical previous GenBank entries. A number of common genetic arrangements  
123 were identified within and between the PFGE cluster groups and these arrangements were grouped  
124 together for comparison with arrangements previously described in the literature. An *ISEcp1* promoter  
125 was located upstream of *bla*<sub>CTX-M</sub> in 53 isolates (56%). In 35 (66%) of these isolates, PCR detected the  
126 full length *ISEcp1* element (1.7 kb) including the *tnpA* transposase gene (0.8 kb). However, PCR  
127 results suggested truncation within *ISEcp1* in the other 18 isolates (no *ISEcp1* amplicon was  
128 detected), which apparently occurred within *tnpA* for 10 isolates where no *tnpA* amplicon was  
129 detected either. Many of the genetic environments identified between *ISEcp1* and *bla*<sub>CTX-M</sub> matched  
130 the frequently described “W”, “X” and “V” common regions first described by Eckert *et al*<sup>9</sup> the most  
131 common of which was the 48 bp W spacer region typical of UK strains B to E<sup>31</sup>, which was present in  
132 32 isolates in combination with *bla*<sub>CTX-M-15</sub> and generally associated with high-level cefotaxime  
133 resistance (MICs  $\geq$ 128  $\mu$ g/ml). PCR and sequencing confirmed reversely oriented IS26 inserted  
134 within the terminal inverted repeat of *ISEcp1* (24 bp before the 3' end of *ISEcp1*) and upstream of  
135 *bla*<sub>CTX-M-15</sub>, as described previously for UK strain A<sup>31</sup> in all but one (33/34) PFGE cluster A isolates  
136 and an ST131 isolate from cluster J. These 34 isolates also contained the alternative ‘promoter X’  
137 region with -35 TTCATG and -10 GGGGATGAT sequences positioned 140 bp and 115 bp,  
138 respectively, upstream of the *bla*<sub>CTX-M-15</sub> start codon within IS26, conferring variable levels of  
139 cefotaxime resistance as described previously.<sup>7</sup> The remaining PFGE cluster A isolate contained  
140 *bla*<sub>CTX-M-14</sub> downstream of *ISEcp1*. The *ISCR1* element was located upstream of *bla*<sub>CTX-M-9</sub> as part of a  
141 complex class 1 integron in all three PFGE cluster G isolates, which upon sequencing resembled the  
142 *sul1*-type integron In60-D.<sup>20</sup> *ISCR1* was also detected upstream of a *bla*<sub>CTX-M-2</sub> gene as part of an In35-  
143 like complex class 1 integron.

144

#### 145 **Integron content of ESBL *E. coli* clinical isolates**



146 Class 1 integrons were detected in 66% of isolates. Four isolates contained complex class 1 integrons  
147 bearing *bla*<sub>CTX-M</sub> as described above and these are indicated in Table 2. The remaining isolates  
148 contained class 1 integrons associated with resistance to agents such as trimethoprim, sulphonamides  
149 and aminoglycosides. Nine distinct class 1 integron variable region amplicons of different sizes were  
150 detected and their distribution amongst PFGE clusters is summarised in Table 2 and details of each  
151 individual isolate are given in supplementary Table 2. Seven isolates produced more than one  
152 amplicon, indicating multiple class 1 integrons were present. Six isolates contained class 2 integron  
153 variable regions of 2.2 kb (*dfrA1-sat1-aadA1*), five of which also contained a class 1 integron. The  
154 most common class 1 integron variable region was 1.7 kb (*dfrA17-aadA5*), present in 54% of isolates.  
155 Most integron variable regions identified in this study contained genes for trimethoprim (*dfr*) and  
156 streptomycin/spectinomycin (*aad*) resistance. The presence of dihydrofolate reductase-containing  
157 integrons correlated with resistance to trimethoprim and/or trimethoprim/sulfamethoxazole in all  
158 clinical *E. coli* isolates. However, streptomycin or spectinomycin susceptibility was unknown for  
159 ESBL-EC as they were not routinely tested for. Gene cassettes conferring resistance to gentamicin  
160 and tobramycin were detected infrequently, were co-localised with a chloramphenicol resistance gene:  
161 either *cmlA1*, *cmlA6* (sporadic isolates) or *catB8* (cluster G isolates). Although susceptibility patterns  
162 to chloramphenicol were untested, the presence of either *aacA4* or *aadB* correlated with phenotypic  
163 resistance to gentamicin in all five strains with these gene cassettes.

164

### 165 **Characterization of transferable CTX-M plasmids**

166 PBRT of all clinical isolates detected IncF plasmids in 90% of clinical isolates, IncII plasmids in  
167 30%, IncN plasmids in 3% and two isolates carrying L/M and B/O plasmids (Table S2). Transfer of  
168 ESBL plasmids was successful for 33 strains, 28 of which transferred a CTX-M-producing plasmid.  
169 The remaining 5 plasmids conferred an ESBL phenotype through expression of *bla*<sub>TEM</sub> or *bla*<sub>SHV</sub>. The  
170 CTX-M plasmids were transferred by conjugation (18) and transformation (10). The replicon types  
171 identified in these plasmids were IncF (12), IncII (11) and IncN (2). The remaining three CTX-M

172 plasmids were untypeable by PBRT. Characteristics of the transferable CTX-M plasmids are detailed  
173 in Table 3. Individual RSTs were indeterminable for two CTX-M bearing plasmids in recipient strains  
174 that harboured multiple IncF plasmids; pBHEC48 and pBHEC12. Six different RSTs were identified  
175 amongst the remaining ten IncF CTX-M plasmids. IncF plasmids carried group 1 CTX-M genes  
176 except for the 27.5 kb plasmid pBHEC91, which carried a group 9 CTX-M gene (CTX-M-27). The  
177 1.7 kb *dfrA17-aadA5* integron was commonly transferred by IncF plasmids. Among 11 IncI1 CTX-M  
178 plasmids identified, six different sequence type (ST)s were identified by IncI1 pMLST. These were  
179 ST3 (n=3, size range 107-112.5 kb), ST7 (n=3, size range 104.5-113 kb), ST 16 (n=2, 94 kb), ST31  
180 (n=1, 93 kb), ST 57 (n=1,84 kb) and the previously undefined ST159, 100 kb with  
181 *repI1/ardA/trbA/sogS/pilL* alleles 1/2/9/1/7 (pBHEC16). The 1.5 kb *dfrA1-aadA1* integron was  
182 transferred on two very similar IncI1-ST7 plasmids, pBHEC20 and pBHEC90. Two IncN type  
183 plasmids of ~30 kb were identified which belonged to the previously identified ST1 and ST6 types.  
184 CTX-M plasmids untypeable by PBRT comprised pBHEC66 (CTX-M-55), pBHEC54 (CTX-M-15,  
185 TEM) and pBHEC76 (CTX-M-15). Cefotaxime MICs for recipient strains bearing CTX-M plasmids  
186 were often less (by between 2 and 5 doubling dilutions) than those of their corresponding clinical  
187 isolate donor strains. Data for all CTX-M plasmids were deposited in the relevant plasmid MLST  
188 database at <http://pubmlst.org/plasmid/>.

189

## 190 Discussion

191 Similar to the situation reported in Europe and globally, our study identified *bla*<sub>CTX-M</sub> as the most  
192 common ESBL gene in clinical *E. coli* isolates collected during 2009-2010 with group 1 alleles the  
193 most common (80/94, 85%), especially *bla*<sub>CTX-M-15</sub>, which was found in two thirds (66%) of ESBL-  
194 EC. A previous nationwide 11 year study (1997-2007) reported a prevalence of 59% group 1 *bla*<sub>CTX-M</sub>  
195 among *bla*<sub>CTX-M</sub> producing *E. coli*. Group 9 CTX-M prevalence among *E. coli* was 144/348 (41.4%)  
196 compared to 13/94 (14%) in this study, however, this may reflect regional differences in prevalence or  
197 changes during the intervening period.<sup>19</sup> The increased prevalence of group 1 genes may be partly

198 due to *E. coli* ST131 dissemination in Ireland, associated with *bla*<sub>CTX-M-15</sub>.<sup>3,24</sup> The endemic nature of  
199 CTX-M-15-producing *E. coli* is evident in Dublin, as indicated by the detection of *bla*<sub>CTX-M-15</sub> in clonal  
200 and sporadically-occurring isolates. Isolates from the endemic PFGE cluster A were confirmed as  
201 genetically indistinguishable from UK strain A and contain *bla*<sub>CTX-M-15</sub> under the control of “promoter  
202 X”, as described previously.<sup>7</sup> Despite the local dominance of CTX-M-15, four other group 1 genes  
203 were detected amongst 14 isolates; *bla*<sub>CTX-M-1, -3, -32</sub> and *bla*<sub>CTX-M-55</sub>. To the best of our knowledge, this  
204 represents the first detection of the latter two genes, or indeed the group 2 gene *bla*<sub>CTX-M-2</sub>, amongst  
205 human *E. coli* isolates in Ireland. Molecular investigation of the promoter regions by PCR mapping  
206 enabled the identification of a number of allele-specific associations. Amongst group 9 CTX-M genes  
207 *bla*<sub>CTX-M-14</sub> was associated with *ISEcp1*, *bla*<sub>CTX-M-27</sub> with *IS26* and *bla*<sub>CTX-M-9</sub> with *ISCR1* as part of the  
208 In60-D integron.<sup>20</sup> Group 1 alleles were usually associated with *ISEcp1*, likely controlling *bla*  
209 expression and driving high-level cefotaxime resistance when associated with *bla*<sub>CTX-M-15</sub> as previously  
210 described.<sup>23</sup>

211 Class 1 integron carriage in our Dublin ESBL-EC collection (66%) was similar to that  
212 recorded in a longitudinal study of ESBL-EC isolates from Madrid (67%)<sup>16</sup> but lower than recorded in  
213 clinical *E. coli* isolates collected from 1998-2004 in Guangzhou, China (86%).<sup>28</sup> As previously  
214 reported, a low prevalence of complex class 1 integrons containing *bla*<sub>CTX-M</sub> was found<sup>16</sup>.  
215 Interestingly, *dfrA17-aadA5* was the most common integron array in the present study (54%) and the  
216 Chinese study (36%). This integron was also detected in Madrid and is globally disseminated in *E.*  
217 *coli*.<sup>11,16,22,28</sup> Class 1 integrons are frequently associated with Tn21-like transposons<sup>15</sup> or multiple  
218 copies of *IS26* on large conjugative plasmids in clinical Enterobacteriaceae, which facilitate the  
219 mobilisation of drug resistance elements among plasmids by homologous recombination.<sup>22,29</sup> This  
220 may explain the presence of the *dfrA17-aadA5* integron on at least four different IncF type plasmids  
221 in our *E. coli* collection. The epidemic distribution of host strains, as exemplified here by UK strain  
222 A, may account for its high prevalence.

223 Six of the nine distinct integron variable regions identified in the present study, including all  
224 those identified in multiple isolates, were documented previously in a study carried out at the

225 Veterinary Hospital, University College Dublin during 2007.<sup>13</sup> The authors characterised consecutive  
226 MDR *E. coli* isolates from predominantly faecal samples of horses (44), cattle (17), pigs (9), dogs (3)  
227 and a sheep, reporting high carriage rates of class 1 integron gene cassette regions (76%). None of the  
228 veterinary *E. coli* isolates belonged to the B2 phylogenetic lineage which is mainly associated with  
229 infections in humans.<sup>13</sup> Nonetheless, the identification of matching class 1 integron variable regions  
230 exemplifies a possible reservoir for these antimicrobial resistance determinants, which may evolve in  
231 the commensal *E. coli* of companion and food animals amidst the selective pressure of veterinary  
232 antimicrobial use and transfer to strains causing human disease.<sup>8,18</sup> The widespread and lengthy use of  
233 trimethoprim and sulphonamides in veterinary medicine has been implicated in the evolution and  
234 persistence of integron-bound resistance genes of the *dfr* and *sul* families.<sup>25</sup> Likewise, the *aad* genes  
235 for streptomycin resistance are positively selected for in animals, where it is used as a first-line drug  
236 for Gram-negative infections. One can speculate that co-carriage of ESBL genes on integrons or  
237 indeed plasmids containing these gene cassettes may drive their co-selection and propagation in the  
238 gut flora of animals treated with veterinary antimicrobials.

239 We noted a local dominance of multi-replicon IncF and IncI1 plasmids amongst CTX-M  
240 producing ESBL-EC in Dublin and identified four new RSTs. The range of IncF replicon sequences  
241 identified in this small cross section of ESBL-EC isolates from the same geographical location  
242 demonstrates the diversity in ESBL-bearing IncF plasmids. The diversity of STs amongst the ten  
243 transferred IncF CTX-M plasmids contrasts with the phylogenetic homogeneity of the host isolates,  
244 7/10 (70%) of which belonged to the ST131 pandemic clone. This reflects the plasticity of IncF  
245 plasmids, which contain multiple hotspots for genetic recombination.<sup>22</sup> The success of conjugation  
246 (18) and transformation (10) of CTX-M plasmids was limited. Low success of conjugation may be  
247 explained by assuming that the replicon type FII-FIA (pEK499-like) plasmids detected here in 41  
248 strains, including PFGE cluster A, lack the requisite *traW* to *traX* genes for conjugation, as has been  
249 shown previously.<sup>29</sup> Nonetheless, the importance of horizontal transfer in the dissemination of  
250 *bla*<sub>CTX-M</sub> and of trimethoprim and aminoglycoside resistance genes on class 1 integrons is evident,  
251 given their frequent location on conjugative plasmids.

252 Many of the 28 CTX-M plasmids characterised had RSTs and *bla*<sub>CTX-M</sub> alleles in common  
253 with those previously identified throughout Europe in both human and animal *E. coli* isolates.  
254 Identical ESBL genes, plasmids and strains of *E. coli* have been identified in Dutch poultry, chicken  
255 meat and humans.<sup>14,21</sup> There are relatively few studies on the prevalence of ESBLs in Irish animals  
256 and animal food products. However, it can be speculated based on the similarity of mobile genetic  
257 elements between commensal *E. coli* of animals and clinical ESBL-E from humans that they may be a  
258 reservoir for MDR plasmids. However, further investigations in this area are warranted. Ireland is a  
259 major exporter of animal meats with 75 % exported to UK and European markets and the remainder  
260 going to the rest of the world. Studies investigating epidemiological links between agricultural and  
261 human isolates of MDR Enterobacteriaceae, particularly in relation to antimicrobial resistance  
262 platforms should be investigated at least at a European level to provide an evidence base for informed  
263 policy in relation to antibiotic use in agriculture.

264 In conclusion, this study reveals the complex array of tools for the mobilization and  
265 expression of *bla*<sub>CTX-M</sub> and other antibiotic resistance genes within ESBL-EC circulating in Dublin  
266 and highlights the importance of group 1 and 9 CTX-M genes and specifically *bla*<sub>CTX-M-15</sub> and *bla*<sub>CTX-</sub>  
267 <sub>M-14</sub>. Our data supports significant roles for both horizontal transfer of ESBL and integron-bound  
268 resistance genes via conjugative IncF, I1 and N plasmids and vertical transfer via clonal spread of the  
269 pandemic ST131 clone. Zoonotic transfer of both integrons and ESBL plasmids to human-associated  
270 *E. coli* may occur through contact with animals or through the food chain.

271

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281

282 **Disclosure Statement**

283 No competing financial interests exist.

284

285 **Table 1.** *bla*<sub>CTX-M</sub> genes and their genetic environments in 94 ESBL-*E. coli* clinical isolates

N	PFGE clusters	<i>bla</i> <sub>CTX-M</sub> allele	Upstream IS	Spacer Region <sup>a</sup>	CTX MIC <sup>b</sup>	GenBank ID	Integrans (size in kb)
34	A, J	15	IS26 (rev)	24 bp ISEcp1 + W	2-256 (8)	GU264003	32 x <i>dfrA17-aadA5</i> (1.7) 1 x <i>dfrA1-sat1-aadA1</i> (2.2)
32	B-E, I, K, L, sporadic	15	ISEcp1	W	16-256 (256)	AY463958	13 x <i>dfrA17-aadA5</i> (1.7) 2 x <i>dfrA1-sat1-aadA1</i> (2.2)
7	H, L, sporadic	1	ISEcp1	XW	12-256 (256)	AM003904	2 x <i>dfrA1-aadA1</i> (1.6)
3	E, sporadic	3	ISEcp1	W	12-256	HF549092	2 x <i>dfrA17-aadA5</i> (1.7)
1	sporadic	3	ISEcp1	VW	12	EU935740	none
1	sporadic	55	ISEcp1	45 bp	256	KC576516	none
1	sporadic	55	ISEcp1	W	256	GQ456159	none
1	sporadic	32	ND	NS	256	AJ557142	none
8	I, F, sporadic	14	ISEcp1	NS	32-256	AF252622	6 x <i>dfrA17-aadA5</i> (1.7) 1 x <i>aadA1</i> (1.0)
2	B	27	IS26	NS	48-128	AY156923	1 x <i>dfrA17-aadA5</i> (1.7)
3	G	9	ISCR1	326 bp	8	AM040708	3 x <i>dfrA12-orfF-aadA8b</i> (1.8) + [ <i>orf513-bla</i> <sub>CTX-M-9</sub> ]
1	sporadic	2	ISCR1	498 bp	256	EF592570	1 x <i>dfrA1-aadA1</i> (1.6) +

286 <sup>a</sup>Spacer region size or letter-coded description: W = 48 bp, X = 32 bp, V = 79 bp (see text and <sup>7</sup>). <sup>b</sup>Cefotaxime

287 MIC range in mg/L (mode, where distinguishable). ND: None Detected; NS: Not Sequenced.

288

289

290 **Table 2.** Characteristics of 28 transferable CTX-M plasmids

Plasmid ID	Replicon Sequence	Previousl y found	Size (kb)	Transfe r	Insertio n	ESBL s	Integron transferre	Dono r	Recipie nt CTX
<b><i>IncF plasmids</i></b>									
pBHEC9	F1:A:-B-	Novel	27.5	T	IS26	CTX-	No transfer	128	12
pBHEC8	F2:A:-B-	H/A WW	78.5	C	ISEcp1	CTX-	<i>dfrA17-</i>	>256	8
pBHEC8	F2:A1:B-	H UK	95	T	IS26	CTX-	<i>dfrA17-</i>	32	6
pBHEC8	F2:A1:B-	H UK	115	T	IS26	CTX-	<i>dfrA17-</i>	3	3
pBHEC3	F45:A1:B-	Novel	100	C	ISEcp1	CTX-	N/A	>256	>256
pBHEC9	F45:A4:B-	Novel	151	C	ISEcp1	CTX-	<i>dfrA17-</i>	>256	32 <sup>d</sup>
pBHEC4	FII, FIA	N/A	137	T	IS26	CTX-	<i>dfrA17-</i>	12	8 <sup>d</sup>
pBHEC1	FII, FIB	N/A	122	C	ISEcp1	CTX-	N/A	>256	32 <sup>d</sup>
pBHEC8	F22:A1:B2	H/A EU	157	C	ISEcp1	CTX-	<i>dfrA7</i>	>256	64 <sup>d</sup>
pBHEC3	F31:A4:B1	H/A EU	157	C	ISEcp1	CTX-	<i>dfrA17-</i>	192	32
pBHEC5	F31:A4:B1	H/A EU	162	T	ISEcp1	CTX-	<i>dfrA17-</i>	>256	32
pBHEC9	F31:A4:B1	H/A EU	146.	C	ISEcp1	CTX-	<i>dfrA17-</i>	>256	32
<b><i>IncII plasmids</i></b>									
pBHEC0	I1-ST3	H/A EU	107	T	ISEcp1	CTX-	N/A	>256	>256
pBHEC1	I1-ST3	H/A EU	112.	C	ISEcp1	CTX-	N/A	12	8
pBHEC2	I1-ST3	H/A EU	112.	C	ISEcp1	CTX-	N/A	>256	64
pBHEC5	I1-ST7	H/A EU	104.	T	ISEcp1	CTX-	No transfer	>256	128
pBHEC2	I1-ST7	H/A EU	113	C	ISEcp1	CTX-	<i>dfrA1-</i>	16	6
pBHEC9	I1-ST7	H/A EU	113	C	ISEcp1	CTX-	<i>dfrA1-</i>	48	6
pBHEC1	I1-ST16	H UK	94	C	ISEcp1	CTX-	No transfer	12	4
pBHEC5	I1-ST16	A UK	94	C	ISEcp1	CTX-	N/A	>256	64
pBHEC3	I1-ST31	H/A EU	93	C	ISEcp1	CTX-	No transfer	>256	64
pBHEC7	I1-ST57	H EU	84	C	ISEcp1	CTX-	N/A	64	24
pBHEC1	I1-ST159	Novel	100	T	ISEcp1	CTX-	N/A	48	32
<b><i>Other plasmids</i></b>									
pBHEC3	N-ST1	H/A EU	30	C	ND	CTX-	N/A	>256	>256
pBHEC7	N-ST6	H UK	31	C	ND	CTX-	No transfer	>256	16
pBHEC6	N/D	N/A	62	C	ISEcp1	CTX-	N/A	>256	>256
pBHEC5	N/D	N/A	116	T	ISEcp1	CTX-	N/A	>256	24
pBHEC7	N/D	N/A	115	T	ISEcp1	CTX-	N/A	>256	32

291 Abbreviations: ND = none detected; N/A = not applicable; UTD = unable to determine; T =  
 292 transformation; C = conjugation; sporadic = sporadically occurring strain; IS = insertion sequence;  
 293 CTX = cefotaxime; MIC = minimum inhibitory concentration (mg/L).<sup>a</sup> Plasmid with same RST and  
 294 CTX-M allele previously found in human (H) or animal (A) isolates worldwide (WW), in the UK  
 295 (UK) or in Europe (EU), see <http://pubmlst.org/plasmid/>; <sup>b</sup> Transfer Method = Transformation (T) or  
 296 Conjugation (C); <sup>c</sup> ESBLs: Specific CTX-M alleles detected. <sup>d</sup> Integrons transferred on plasmid <sup>d</sup>  
 297 recipient contains >1 β-lactamase plasmid

298



299 **Table S1.** Primers for analysis of *bla*<sub>CTX-M</sub> genetic environment

Primer name	Sequence (5'-3')	Target	Reference
<i>tmpA</i> ISEcp1	AATACTACCTTGCTTTCTGA	<i>tmpA</i> of ISEcp1	2
ISEcp1 5'	TTCAAAAAGCATAATCAAAGCC	ISEcp1 5'-3'	2
ISEcp1 reverse	CAACCACCTTTCAATCATTTTT	ISEcp1	2
Orf513	TGGAAGAGGGCGAAGACGAT	Orf513 of ISCR1	2
Orf513 rev	GCGTTTTATCGGTAGTCGTC	Orf513 of ISCR1	2
<i>Int1</i> -F	GGTCAAGGATCTGGATTTTCG	<i>Int1</i>	3
<i>Int1</i> -R	ACATGCGTGTAATCATCGTC	<i>Int1</i>	3
5'CS	GGCATCCAAGCAGCAAG	Class 1 integron	3
3'CS	AAGCAGACTTGACCTGA	Class 1 integron	3
<i>Int2</i> -F	CACGGATATGCGACAAAAAGGT	<i>Int2</i>	3
<i>Int2</i> -R	GTAGCAAACGAGTGACGAAATG	<i>Int2</i>	3
<i>attI2</i> -F	GACGGCATGCACGATTTGTA	Class 2 integron	3
<i>orfX</i> -R	GATGCCATCGCAAGTACGAG	Class 2 integron	3
PROM+	TGCTCTGTGGATAACTTGC	<i>bla</i> <sub>CTX-M</sub> +	4
PRE-CTX-M-3B	CCGTTTCCGCTATTACAAAC	<i>bla</i> <sub>CTX-M-1</sub> +	4
IS26	AGCGGTAAATGCTGGAGTGA	IS26	5
CTX15 rev	ATTCGGCAAGTTTTTGCTGT	IS26 + <i>bla</i> <sub>CTX-M-1/3</sub>	5
M9 upper	ATGGTGACAAAGAGAGTGCA	<i>bla</i> <sub>CTX-M-9</sub>	1
M9 lower	CCCTTCGGCGATGATTCTC	ISEcp1/IS26 +	1

300

301

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321

322



324 **Table S2.** Summary of antimicrobial resistance platforms for 100 ESBL-producing *E. coli* collected in Dublin

CG <sup>a</sup>	Strain	PG	IS <sup>b</sup>	ESBL	β-L <sup>c</sup>	Integron gene cassette arrays	CTX MIC	Replicons	Transfer Method	Plasmid ST (size kb) <sup>d</sup>
A	BHEC44	B2-ST131	IS26	CTX-M-15	ND	<i>dfrA17-aadA5</i>	3	F2, F1A	N/A	N/A
A	BHEC45	B2-ST131	IS26	CTX-M-15	ND	<i>dfrA17-aadA5</i>	16	F2, F1A	N/A	N/A
A	BHEC48	B2-ST131	IS26	CTX-M-15	ND	<i>dfrA17-aadA5</i>	12	F2, F1A	Trans	F-UT (137)
A	BHEC26	B2-ST131	IS26	CTX-M-15	ND	<i>dfrA17-aadA5</i>	4	F2, F1A	N/A	N/A
A	BHEC27	B2-ST131	IS26	CTX-M-15	ND	<i>dfrA17-aadA5</i>	3	F2, F1A, I1	N/A	N/A
A	BHEC28	B2-ST131	IS26	CTX-M-15	ND	<i>dfrA17-aadA5</i>	4	F2, F1A	N/A	N/A
A	BHEC34	B2-ST131	IS26	CTX-M-15	ND	<i>dfrA17-aadA5</i>	4	F2, F1A	N/A	N/A
A	BHEC38	B2-ST131	IS26	CTX-M-15	ND	<i>dfrA17-aadA5</i>	64	F2, F1A	N/A	N/A
A	BHEC40	B2-ST131	IS26	CTX-M-15	ND	<i>dfrA17-aadA5</i>	2	F2, F1A	N/A	N/A
A	BHEC74	B2-ST131	IS26	CTX-M-15	ND	<i>dfrA17-aadA5</i>	24	F2, F1A	N/A	N/A
A	BHEC85	B2-ST131	IS26	CTX-M-15	ND	<i>dfrA17-aadA5</i>	>256	F2, F1A	No	N/A
A	BHEC96	B2-ST131	IS26	CTX-M-15	ND	<i>dfrA17-aadA5</i>	6	F2, F1A	No	N/A
A	BHEC86	B2-ST131	IS26	CTX-M-15	ND	<i>dfrA17-aadA5</i>	8	F2, F1A	No	N/A
A	BHEC63	B2-ST131	IS26	CTX-M-15	ND	<i>dfrA17-aadA5</i>	48	F2, F1A	No	N/A
A	BHEC92	B2-ST131	IS26	CTX-M-15	ND	<i>dfrA17-aadA5</i>	16	F2, F1A	N/A	N/A
A	BHEC94	B2-ST131	IS26	CTX-M-15	ND	<i>dfrA17-aadA5</i>	8	F2, F1A	N/A	N/A
A	BHEC65	B2-ST131	IS26	CTX-M-15	ND	<i>dfrA17-aadA5</i>	6	F2, F1A	N/A	N/A
A	BHEC67	B2-ST131	IS26	CTX-M-15	ND	<i>dfrA17-aadA5</i>	4	F2, F1A, I1	N/A	N/A
A	BHEC21	B2-ST131	IS26	CTX-M-15	ND	<i>dfrA17-aadA5</i>	96	F2, F1A, I1	N/A	N/A

A	BHEC60	B2-ST131	IS26	CTX-M-15	TEM	ND	6	F2, F1A	N/A	N/A
A	BHEC46	B2-ST131	IS26	CTX-M-15	TEM	<i>dfrA17-aadA5</i>	8	F2, F1A	N/A	N/A
A	BHEC4	B2-ST131	IS26	CTX-M-15	ND	<i>dfrA17-aadA5</i>	8	F2, F1A	N/A	N/A
A	BHEC9	B2-ST131	IS26	CTX-M-15	ND	<i>dfrA17-aadA5</i>	16	F2, F1A	N/A	N/A
A	BHEC50	B2-ST131	IS26	CTX-M-15	ND	<i>dfrA17-aadA5</i>	24	F2, F1A	N/A	N/A
A	BHEC61	B2-ST131	IS26	CTX-M-15	ND	<i>dfrA17-aadA5</i>	8	F2, F1A	N/A	N/A
A	BHEC77	B2-ST131	IS26	CTX-M-15	ND	<i>dfrA17-aadA5</i>	>256	F2, F1A	N/A	N/A
A	BHEC50	B2-ST131	IS26	CTX-M-15	TEM	<i>dfrA17-aadA5</i>	1.5	F2, F1A	N/A	N/A
A	BHEC55	B2-ST131	IS26	CTX-M-15	ND	<i>dfrA7, dfrA17-aadA5</i>	6	F2, F1A, I1	N/A	N/A
A	BHEC35	B2-ST131	IS26	CTX-M-15	TEM	<i>dfrA17-aadA5</i>	2	F2, F1A, I1	No	N/A
A	BHEC7	B2-ST131	IS26	CTX-M-15	ND	<i>dfrA17-aadA5</i>	8	F2, F1A	N/A	N/A
A	BHEC18	B2-ST131	IS26	CTX-M-15	ND	<i>dfrA17-aadA5</i>	2	F2, F1A, I1	N/A	N/A
A	BHEC23	B2-ST131	IS26	CTX-M-15	ND	<i>dfrA17-aadA5</i>	64	F2, F1A	N/A	N/A
A	BHEC82	B2-ST131	IS26	CTX-M-15	ND	<i>dfrA17-aadA5</i>	32	F2, F1A	Trans	F2:A1:B- (95)
A	BHEC16	B2-ST131	ISEcp1	CTX-M-14	TEM	ND	48	F2, F1A, F1B, I1	Trans	I1-ST159 (100)
B	BHEC43	B2-ST131	ISEcp1	CTX-M-15	TEM	<i>dfrA17-aadA5</i>	>256	F1A, F1B, I1	No	N/A
B	BHEC57	B2-ST131	ISEcp1	CTX-M-15	TEM	ND	>256	F1A, F1B, I1	No	N/A
B	BHEC56	B2-ST131	ISEcp1	CTX-M-15	ND	<i>dfrA17-aadA5</i>	>256	F2, F1A, F1B	Trans	F31:A4:B1 (162)
B	BHEC15	B2-ST131	IS26	CTX-M-27	ND	ND	48	F2, F1A, F1B	No	N/A
B	BHEC91	B2-ST131	IS26	CTX-M-27	ND	<i>dfrA17-aadA5</i>	128	F2, F1A, F1B	Trans	F1:A-B- (27.5)
C	BHEC32	B2-ST131	ISEcp1	CTX-M-15	TEM	ND	>256	F2, F1A	Conj	F45:A1:B- (100)
C	BHEC47	B2-ST131	ISEcp1	CTX-M-15	TEM	ND	>256	F2, F1A	No	N/A

C	BHEC84	B2-ST131	ISEcp1	CTX-M-15	TEM	ND	>256	F2, F1A, F1B	No	N/A
C	BHEC1	B2-ST131	ISEcp1	CTX-M-15	ND	ND	>256	F2, F1A	No	N/A
C	BHEC33	B2-ST131	ISEcp1	CTX-M-15	TEM	<i>dfrA17-aadA5</i>	>256	F2, F1A, F1B	No	N/A
C	BHEC10	B2-ST131	ISEcp1	CTX-M-15	TEM	<i>dfrA17-aadA5</i>	>256	F2, F1A, F1B, B/O	No	N/A
D	BHEC58	B2-ST131	ISEcp1	CTX-M-15	ND	ND	>256	F2, F1A	No	N/A
D	BHEC88	B2-ST131	ISEcp1	CTX-M-15	TEM	<i>dfrA7</i>	>256	F2, F1A, F1B, I1	Conj	F22:A1:B20 (157)
E	BHEC3	B2	ISEcp1	CTX-M-3	ND	ND	12	F2, F1A, F1B	No	N/A
E	BHEC62	B2	ISEcp1	CTX-M-15	TEM	ND	>256	F2, F1A, F1B	No	N/A
F	BHEC41	B2	ISEcp1	CTX-M-14	TEM	<i>dfrA17-aadA5</i>	64	F2, F1A, F1B, L/M	No	N/A
F	BHEC42	B2	ISEcp1	CTX-M-14	TEM	<i>dfrA17-aadA5</i>	32	F2, F1A, F1B, L/M	No	N/A
G	BHEC2	D	ISCR1	CTX-M-9	TEM	<i>drfA12-orfF-aadA8b,</i> <i>orfD-aacA4-orf105-catB8</i>	8	F2, F1B	No	N/A
G	BHEC11	D	ISCR1	CTX-M-9	TEM	<i>drfA12-orfF-aadA8b,</i> <i>orfD-aacA4-orf105-catB8</i>	8	F2, F1B	No	N/A
G	BHEC13	D	ISCR1	CTX-M-9	TEM	<i>drfA12-orfF-aadA8b,</i> <i>orfD-aacA4-orf105-catB8</i>	8	F2, F1B	No	N/A
H	BHEC31	D	N/A	SHV	SHV	ND	0.75	I1	Conj	ND (32.5)
H	BHEC90	D	ISEcp1	CTX-M-1	TEM	<i>dfrA1-aadA1,</i> <i>dfrA1-sat1-aadA1</i>	48	F2, F1A, F1B, I1	Conj	I1-ST7 (113)

I	BHEC14	D	ISEcp1	CTX-M-14	TEM	<i>aadA1, dfrA17-aadA5</i>	>256	F2, F1A, F1B, I1	No	N/A
I	BHEC37	D	ISEcp1	CTX-M-15	TEM	<i>aadA1, dfrA17-aadA5</i>	>256	F2, F1B, I1	Conj	I1-ST31 (93)
J	BHEC24	B2	N/A	TEM	TEM	<i>aadA1, dfrA1-sat1-aadA1</i>	2	F2	Conj	F-UT (15), F-UT (48.5), F-UT (66)
J	BHEC87	B2-ST131	IS26	CTX-M-15	TEM	<i>aadA1, dfrA1-sat1-aadA1</i>	3	F2, F1A	Trans	F2:A1:B- (115)
K	BHEC36	A	ISEcp1	CTX-M-15	ND	<i>dfrA17-aadA5</i>	192	F2, F1A, F1B, I1	Conj	F31:A4:B1 (157)
K	BHEC78	A	ND	CTX-M-15	ND	<i>dfrA17-aadA5</i>	96	F2, F1A, F1B	No	N/A
L	BHEC22	B1	ISEcp1	CTX-M-1	ND	ND	>256	F2, F1B, I1	Conj	I1-ST3 (112.5)
L	BHEC76	B1	ISEcp1	CTX-M-15	ND	ND	>256	ND	Trans	ND (115)
none	BHEC25	B2-ST131	ISCR1	CTX-M-2	TEM	<i>dfrA1-aadA1</i>	>256	F2, F1B, I1	No	N/A
none	BHEC72	A	ISEcp1	CTX-M-14	TEM	<i>dfrA17-aadA5, aacA4-cmlA1</i>	96	F2, F1B	No	N/A
none	BHEC81	A	ISEcp1	CTX-M-14	TEM	<i>dfrA12-orfF-aadA2</i>	>256	F2, F1B	No	N/A
none	BHEC64	B2-ST131	ISEcp1	CTX-M-14	TEM	<i>dfrA17-aadA5</i>	48	F2, F1A	No	N/A
none	BHEC49	D	ISEcp1	CTX-M-14	TEM	<i>dfrA17-aadA5</i>	96	F2, F1A, F1B	No	N/A
none	BHEC51	A	ISEcp1	CTX-M-15	ND	<i>dfrA17-aadA5</i>	>256	F2, F1A, F1B	No	N/A
none	BHEC80	A	ISEcp1	CTX-M-15	TEM	<i>dfrA17-aadA5, dfrA1-sat1-aadA1</i>	>256	F2	Conj	F2:A-B- (78.5)
none	BHEC99	A	ISEcp1	CTX-M-15	TEM	<i>dfrA17-aadA5, dfrA1-sat1-aadA1</i>	>256	F2, F1A, F1B, B/O	Conj	F45:A4:B- (151)
none	BHEC17	B1	ISEcp1	CTX-M-1	ND	ND	12	I1	Conj	I1-ST3 (112.5)
none	BHEC83	B1	ISEcp1	CTX-M-15	ND	ND	>256	ND	No	N/A

none	BHEC95	B1	ISEcp1	CTX-M-15	TEM	ND	>256	F1A	No	N/A
none	BHEC6	B2	ISEcp1	CTX-M-1	ND	ND	>256	F2, F1B, I1	Trans	I1-ST3 (107)
none	BHEC66	B2	ISEcp1	CTX-M-55	ND	ND	>256	ND	Conj	ND (62)
none	BHEC70	B2	ISEcp1	CTX-M-3	ND	ND	64	F2, F1B, I1	Conj	I1-ST57 (84)
none	BHEC93	B2-ST131	ISEcp1	CTX-M-15	ND	ND	>256	F2, F1A, F1B	No	N/A
none	BHEC97	B2-ST131	ISEcp1	CTX-M-15	ND	<i>dfrA17-aadA5</i>	>256	F2, F1A, F1B	Conj	F31:A4:B1 (146.5)
none	BHEC100	B2-ST131	ISEcp1	CTX-M-15	TEM	<i>dfrA17-aadA5</i>	192	F2, F1A	No	N/A
none	BHEC8	D	ISEcp1	CTX-M-15	TEM	ND	64	F2, F1A, F1B	No	N/A
none	BHEC12	D	ISEcp1	CTX-M-55	TEM	ND	>256	F2, F1B	Conj	F-UT (122)
none	BHEC19	D	ISEcp1	CTX-M-3	TEM	<i>dfrA17-aadA5</i>	12	I1	Conj	I1-ST16 (94)
none	BHEC29	D	ISEcp1	CTX-M-1	TEM	ND	64	F2, F1B, I1	No	N/A
none	BHEC30	D	ISEcp1	CTX-M-15	TEM	ND	>256	F2, F1A, F1B	No	N/A
none	BHEC53	D	ISEcp1	CTX-M-1	ND	<i>dfrA1-sat1-aadA1</i>	>256	I1	Trans	I1-ST7 (104.5)
none	BHEC54	D	ISEcp1	CTX-M-15	TEM	ND	>256	F2, F1B	Trans	ND (116)
none	BHEC98	D	ISEcp1	CTX-M-15	ND	<i>dfrA17-aadA5</i>	>256	F2, F1A, F1B	No	N/A
none	BHEC20	U	ISEcp1	CTX-M-1	ND	<i>dfrA1-aadA1</i>	16	I1	Conj	I1-ST7 (113)
none	BHEC52	U	ISEcp1	CTX-M-15	TEM	ND	>256	I1	Conj	I1-ST16 (94)
none	BHEC89	U	ISEcp1	CTX-M-15	SHV	ND	16	ND	No	N/A
none	BHEC59	A	N/A	SHV	TEM	ND	1.5	F2, F1B, I1	Conj	I1-ST3 (118)
none	BHEC69	A	N/A	SHV	TEM	<i>dfrA1-aadA1</i>	4	F2, I1, N	No	N/A
none	BHEC73	A	N/A	SHV	TEM	ND	4	F1B, I1	Conj	I1-ST3 (101.5)



none	BHEC68	B1	N/A	TEM	TEM	ND	32	F2, F1B, I1	Trans	ND (31.5), ND (21.5)
none	BHEC39	B1	ND	CTX-M-32	ND	ND	>256	F2, F1B, N	Conj	N-ST1 (30)
none	BHEC79	B2-ST131	ISEcp1 <sup>b</sup>	CTX-M-15	TEM	ND	>256	F2, F1A, F1B, I1	No	N/A
none	BHEC71	D	ND	CTX-M-3	TEM	<i>dfrA17-aadA5</i>	>256	F2, F1B	Conj	N-ST6 (31)
none	BHEC75	D	ISEcp1 <sup>b</sup>	CTX-M-15	TEM	<i>aadB-aadA1-cmlA6</i>	>256	F2, F1A	No	N/A

325 Abbreviations: ND = none detected; N/A = not applicable; UTD = unable to determine; Trans = transformation; Conj = conjugation; PG = phylogenetic  
326 group; B2-ST131 = O25B-ST131 pandemic clone; CG = clonal group, none = sporadically occurring strain; IS = insertion sequence; CTX = cefotaxime; MIC  
327 = minimum inhibitory concentration (mg/L). <sup>a</sup> Clonal groups as identified previously <sup>1</sup>. <sup>b</sup> IS not linked directly to CTX-M gene in indicated isolates. <sup>c</sup> Beta-  
328 lactamase gene families detected by PCR <sup>d</sup> Plasmid ST = plasmid sequence type; F:A:B formulae given for IncF plasmids; pMLST formulae given for I1 (I1-  
329 STx) and N (N-STx) plasmids; F-UT = F plasmid untypable by RST; approximate plasmid size (kb) is given in parentheses.

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