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Title	Dissemination of pHK01-like incompatibility group IncFII plasmids encoding CTX-M-14 in Escherichia coli from human and animal sources
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Citation	Veterinary Microbiology, 2012, v. 158 n. 1-2, p. 172-179
Issued Date	2012
URL	http://hdl.handle.net/10722/152771
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1	Original research paper (VetMic-D-11-6397 revised)
2	Dissemination of pHK01-like incompatibility group IncFII plasmids encoding CTX-M-
3	14 in Escherichia coli from human and animal sources, 2002-2010
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19	Word counts: abstract (250 words), body text (3627 words)

Abstract

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22 Few studies have compared CTX-M encoding plasmids identified in different ecological 23 sources. This study aimed to analyze and compare the molecular epidemiology of plasmids 24 encoding CTX-M-14 among strains from humans and animals. The CTX-M-14 encoding 25 plasmids in 160 E. coli isolates from animal faecal (14 pigs, 16 chicken, 12 cats, 8 cattle, 5 26 dogs and 3 rodents), human faecal (45 adults and 20 children) and human urine (37 adults) 27 sources in 2002-2010 were characterized by molecular methods. The replicon types of the 28 CTX-M-14 encoding plasmids were IncFII (n=61), I1-I $\gamma$  (n=24), other F types (n=23), B/O 29 (n=10), K (n=6), N (n=3), A/C (n=1), HI1 (n=1), HI2 (n=1) and nontypeable (n=30). The 30 genetic environment, ISEcp1 - bla<sub>CTX-M-14</sub> - IS903 was found in 89.7% (52/58), 87.7% (57/65) 31 and 86.5% (32/37) of the animal faecal, human faecal and human urine isolates, respectively. 32 Subtyping of the 61 IncFII incompatibility group plasmids by replicon sequence typing, 33 plasmid PCR-restriction fragment length polymorphism and marker genes (vac, malB, 34 *eitA/eitC* and *parB/A*) profiles showed that 31% (18/58), 30.6% (20/65) and 37.8% (14/37) of 35 the plasmids originating from animal faecal, human faecal and human urine isolates, 36 respectively, were pHK01-like. These 52 pHK01-like plasmids originated from diverse 37 human (20 faecal isolates from 2002, 2007-2008, 14 urinary isolates from 2004) and animal 38 (all faecal, 1 cattle, 1 chicken, 5 pigs, 9 cats, 1 dogs, 1 rodents from 2008-2010) sources. In 39 conclusion, this study highlights the importance of the IncFII group, pHK01-like plasmids in 40 the dissemination of CTX-M-14 among isolates from diverse sources.

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42 Key words: pets, pigs, chicken, plasmid, extended-spectrum beta-lactamases

#### 44 Introduction

45 Asia is one of the epicentres of antimicrobial resistance (Hawkey, 2008; Ho et al., 2011c; Ho 46 et al., 2011a). In this part of the world, the CTX-M type extended-spectrum  $\beta$ -lactamases 47 (ESBLs) have spread extensively among bacteria in human, animals and the environment 48 (Hawkey, 2008; Ho et al., 2011a). Many studies have reported the overwhelming dominance 49 of the CTX-M-14 allele among clinical isolates of *Escherichia coli* and *Klebsiella* spp in 50 mainland China, Hong Kong and South Korea (Hawkey, 2008; Ho et al., 2007; Naseer and 51 Sundsfjord, 2011). In general, bacterial strains producing CTX-M-14 are genetically diverse 52 (Ho et al., 2007; Lo et al., 2010; Valverde et al., 2009). Hence, the dissemination of CTX-M-53 14 has been attributed to conjugative plasmids and other mobile genetic elements rather than 54 clonal expansion (Naseer and Sundsfjord, 2011).

55 In Spain, the spread of CTX-M-14 from 2000 to 2005 was largely due to pRYC105-56 like plasmids of the IncK incompatibility group disseminated among diverse E. coli lineages 57 (Valverde et al., 2009). It has further been shown that pRYC105 is related to the pCT plasmid 58 that has been found in CTX-M-14 producing strains from the United Kingdom, mainland 59 China and Australia (Cottell et al., 2011). Recently, we sequenced the IncFII incompatibility 60 group plasmid, pHK01 encoding CTX-M-14 and showed that it has disseminated widely 61 among E. coli isolates causing community-acquired urinary tract infections in women (Ho et 62 al., 2011b). Plasmids closely related to pHK01 have also been identified among Klebsiella 63 strains from mainland China (Yi et al., 2010). Besides plasmids, the acquisition and 64 horizontal transmission of *bla*<sub>CTX-M</sub> genes have been associated with insertion sequences (ISs), 65 putative transposons and class 1 integrons (Canton and Coque, 2006). These elements may 66 have mobilized the *bla*<sub>CTX-M</sub> gene from its progenitor and may drive the expression of the β-67 lactamase (Canton and Coque, 2006). In addition, these elements might accumulate 68 additional resistance genes to the *bla*<sub>CTX-M</sub> flanking regions (Canton and Coque, 2006). It has

also been suggested that IS*Ecp*,  $bla_{CTX-M}$  and IS*903* form a putative transposon and this block of genes could disseminate by transposition (Poirel et al., 2005).

Although the roles of plasmids and other genetic elements are recognized to be important in the dissemination of the CTX-M genes, few studies have compared the distribution of these mobile genetic elements among strains from humans and animals and their inter-relationship (Garcia-Fernandez et al., 2008). In this study, we investigated the molecular epidemiology of plasmids encoding CTX-M-14 and their  $bla_{CTX-M-14}$  genetic environment for a collection of *E. coli* isolates collected from diverse human and animal sources between 2002 and 2010.

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79 Methods

#### 80 Bacterial strains

81 One hundred and sixty CTX-M-14 producing E. coli isolates were studied. The strains were 82 chosen to provide representation from different time periods and animal and human sources. 83 All viable CTX-M-14 producing E. coli isolates recovered from four regional antimicrobial 84 resistance surveillances during 2002-2010 were eligible for inclusion (Ho et al., 2007; Ho et 85 al., 2008; Ho et al., 2011a; Lo et al., 2010). These included 37 isolates from female 86 outpatients with urinary tract infections, 65 human faecal isolates (45 adults and 20 children) 87 and 58 faecal isolates from animals (14 pigs, 16 chicken, 12 cats, 8 cattle, 5 dogs and 3 88 rodents) (Ho et al., 2007; Ho et al., 2008; Ho et al., 2011b; Ho et al., 2011a). The human 89 urine isolates were recovered in 2004 from female outpatients with urinary tract infections 90 (age range, 27 -80 years) and all CTX-M-14 isolates in the collection were included (Ho et 91 al., 2007). The 65 human faecal isolates include all of 62 CTX-M-14 isolates identified in a 92 study that examined faecal carriage in hospitalized children and their household contacts in 93 October 2007-September 2008 (Lo et al., 2010) and three (out of seven CTX-M-14 isolates

94 found) isolates from a study that assessed carriage in non-hospitalized individuals in 2002 95 (Ho et al., 2008). The remaining four isolates obtained in the 2002 faecal study were lost 96 during storage and therefore not included. All the animal isolates were obtained during 97 September 2008-August 2010 as part of an ongoing antimicrobial resistance by trained staff 98 in two government departments (Ho et al., 2011a). For cattle and pigs, rectal swabs were 99 obtained from fresh carcasses at a centralized slaughterhouse in Hong Kong (Ho et al., 100 2011a). Chicken samples were obtained while the animals were temporarily held for 101 inspection before sale at wet markets in Hong Kong. All the cattle were imported from 102 mainland China. The pigs and chicken included animals produced at local farms and those 103 imported from mainland China. Samples from the stray dogs, stray cats and rodents were 104 collected by trained staff at governmental animal management centres. These animals were 105 captured from urban areas from all over Hong Kong. The animals were sampled in batches: 106 chicken (20 animals per batch), cattle (10 animals per batch), pigs (2 to 7 animals per batch), 107 stray cats (1-10 animals batch), stray dogs (1-10 animals per batch) and urban rodents (2 to 108 23 animals per batch). In total, 2106 animals from 298 batches were tested. In 179 batches, 109 at least one animal was found to carry ESBL-producing E. coli. A subset of 132 isolates was 110 chosen randomly according to the collection period and batch number for CTX-M PCR and 111 sequencing. The isolates were selected randomly according to the collection period and batch 112 number. This subset included isolates recovered from 84 batches and covered the entire 113 specimen collection periods. Sequencing showed that 58 of the 132 isolates had CTX-M-14. 114 All the 58 CXT-M-14 producing isolates were included (Ho et al., 2011a).

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# 116 Susceptibility testing, conjugation and replicon typing

Antimicrobial susceptibility to amoxicillin-clavulanic acid, amikacin, cefotaxime,
ceftazidime, ciprofloxacin, chloramphenicol, cotrimoxazole, gentamicin, nalidixic acid,

119 nitrofurantoin, tetracycline was tested by the disc diffusion method (Clinical and Laboratory 120 Standards Institute, 2011). ESBL production was determined by the double-disc synergy test 121 using cefotaxime, amoxicillin-clavulanic acid and ceftazidime at inter-disc distances (centre 122 to centre) of 20 and 25 mm (Ho et al., 2000). Conjugation was carried out in filters with E. 123 *coli* J53Az<sup>r</sup> as the recipient. Donor and recipient cells were mixed at 1:10 ratio. 124 Transconjugants were selected on trypticase soy agar plates containing sodium azide (150 125 mg/L; Sigma Chemical Co.) for counterselection and cefotaxime (1 mg/L; Sigma Chemical 126 Co.) to select for plasmid-encoded resistance. For detecting plasmids, bacterial cells were 127 embedded in agarose plugs and disrupted by alkaline lysis. Subsequently, the plasmids were 128 converted to the linear forms by incubation with Aspergillus oryzae S1 nuclease (Sigma 129 Chemical Co., St Louis, MO, USA) and were sized by pulsed-field gel electrophoresis (PFGE) 130 as previously described (Ho et al., 2010; Ho et al., 2011b). The replicon types for E. coli 131 transconjugants with CTX-M encoding plasmids were determined by a scheme previously 132 described (Carattoli et al., 2005). Eighteen pairs of primers were tested in five multiplex and 133 three simplex PCR assays. The method allowed recognition of the following plasmid 134 incompatibility groups (Inc): FIA, FIB, FIC, FIIA, HI1, HI2, I1-I<sub>γ</sub>, L/M, N, P, W, T, A/C, K, 135 B/O, X, Y, F. An additional primer pair was used for identification IncFII replicons (Osborn 136 et al., 2000). The FII plasmids were further subtyped by the replicon sequence typing (RST) 137 scheme (Villa et al., 2010). This involved PCR and sequencing of DNA fragments containing 138 the copA region of the FII replicon, the iterons-*repE* region of the FIA replicon, the *repB* 139 gene of the FIB replicon and the *copA* region of the FIC replicon. Alleles were assigned to 140 each sequence and the replicon types were expressed according to the FAB formula (Villa et 141 al., 2010). Where a transconjugant could not be obtained, the replicon type for the plasmid 142 encoding CTX-M-14 was determined in the parent strains. In all the isolates, the replicon 143 location in the plasmids was confirmed by hybridization with probes specific for *bla*<sub>CTX-M</sub> and

144 rep amplified by PCR from different samples. For non-transconjugants, S1-PFGE was used 145 to separate all the plasmids and the plasmid carrying the  $bla_{CTX-M}$  gene was identified by 146 hybridization probe. PCR was used to determine all the plasmid replicons carried by the non-147 transconjugants. Afterwards, the non-transconjugants were tested by hybridization using 148 probes for all positive replicon types. The *rep* probes that hybridized to the *bla*<sub>CTX-M</sub> carrying 149 plasmid were used to define the replicon type. The four main phylogenetic groups (A, B1, B2, 150 and D) of the E. coli isolates were determined by a multiplex PCR assay (Clermont et al., 151 2000).

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# 153 PCR-RFLP analysis of IncFII plasmids

154 The IncFII plasmids were analysed further by a PCR-RFLP scheme previously described (Ho et al., 2011b). The PCR-RFLP scheme involved PCR amplifications of seven loci or regions 155 156 (designated A, B1, B2, and C to F) with sizes ranging from 1.7 kb to 7.8 kb. Following PCR 157 amplifications, the amplicons were digested with 5 U each of the following restriction 158 enzymes (New England BioLabs) according to the manufacturer's instructions: RsaI (locus 159 A); SmaI (locus B1); SfcI (locus B2); HhaI (locus C); BsmI (locus D); BstEII (locus E); and 160 DrdI (locus F). For each locus, the PCR-RFLP patterns were assigned numbers. Patterns with 161 one or more band differences were assigned different numbers. The profiles for the seven loci 162 were used to designate the pRFLP type. Primers for amplification of the A, B2, C to F loci 163 were those described previously (Ho et al., 2011b). Locus B1 was included in this study to 164 map the region upstream of *bla*<sub>CTX-M-14</sub>. The primer pair used to amplify locus B1 was: 165 repA1F (forward), 5'- CGCTCCTTCTGCGCATTGTAA -3' and CTX-M-9F (backward), 5'-166 CAAAGAGAGTGCAACGGATG -3'(Woodford et al., 2006). The content of the amplified 167 regions in pHK01 were (Ho et al., 2011b): locus A (finO, orf99-102, repA2, repA3 and 168 <u>repA1</u>, locus B1 (<u>repA1</u>, yacABC, yadA, malB, IS903, <u>bla</u><sub>CTX-M-14</sub>), locus B2, (<u>bla</u><sub>CTX-M-14</sub>),

169 ISECp1, eitD, eitC, eitB, eitA, orf18-19, parB), locus C (parB, parA), locus D (orf37, orf38-

40, ssb, orf42-43, psiB, psiA, orf45), locus E (traM, J, Y, A, L, E, K, B, P, trbD), and locus F
(traI, X, orf97, finO). The primers were designed within the underlined gene or
upstream/downstream of the gene in **boldface.** Plasmid with PCR-RFLP patterns identical to
the reference plasmid (pHK01) in three or more loci were considered to be pHK01-like. PCR
for the pHK01-associated malB, yac, eitA, eitC and parA/B genes was performed using
previously described primers (Ho et al., 2011b).

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#### 177 Analysis of *bla*<sub>CTX-M-14</sub> genetic environment

The genetic environment may be defined as the sequences flanking the open reading frame of the  $bla_{\text{CTX-M-14}}$  gene. This includes insertion elements that may be involved in the mobilization and expression of the  $bla_{\text{CTX-M-14}}$  gene. Based on published studies and sequences deposited in the GenBank database, eight different types of  $bla_{\text{CTX-M-14}}$  genetic environment have been reported (Table 1). For purpose of description, they were denoted types I to VIII. Monoplex PCRs using different primer pair combinations were used to map the genetic environment to one of the recognized types (Table 1).

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#### 186 Plasmid sequencing

The Illumina Genome Analyzer IIx was used for sequencing of plasmids at approximately 500-fold coverage, as described previously (Ho et al., 2011b). In brief, plasmids were extracted from the transconjugants by using the Qiagen Large Construct kit (Qiagen, Hong Kong). Purified plasmid DNA was fragmented by nebulization. The fragments were amplified and a library was constructed as described previously (Ho et al., 2011c). Based on the qPCR quantified concentration of the barcoded plasmid library, it was diluted to generate approximately 500,000 clusters and seeded with other samples in the same Solexa sample lane. Sequencing run of 76-base pair-end reads was performed according to the
manufacturer's recommendations. The Illumina Off-Line Basecaller (version v1.6), WebACT
and Geneious Pro (Version 5.0.1, Biomatters Limited, Auckland, New Zealand) softwares
were used for bioinformatics analysis (Ho et al., 2011c).

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199 **Results** 

In conjugation experiments, the plasmids harbouring CTX-M 14 could be transferred in 77.5% (124/160) of the isolates at frequencies of  $10^{-6}$  to  $10^{-1}$  per donor cells. In addition to cefotaxime resistance, the plasmids in 19 transconjugants encoded co-resistance to non- $\beta$ lactam antibiotics with profiles involving chloramphenicol, gentamicin, cotrimoxazole and/or tetracycline. These plasmids have origins from 17 faecal (chicken 8, pigs 3, dog 1, humans 5) and two human urine isolates (Supplementary Table S1).

206 Ten different replicons (IncFII, IncFIA, IncFIB, IncI1-Iy, IncB/O, IncK, IncN, IncA/C, 207 IncHI1, IncHI2), either alone or in combinations were found among the plasmids encoding 208 CTX-M-14 (Table 2). The two most common replicon types were IncFII (38.1%, 61/160) and 209 IncI1-I $\gamma$  (15%, 24/160). The frequencies of IncFII replicon were similar among isolates from 210 the three sources: animal faecal (36.2%), human faecal (38.5%) and human urine (40.5%, chi 211 square for trend, P = 0.7). Replicon IncI1-Iy was more common among animal faecal isolates 212 (20.7%) than in human faecal isolates (10.8%) and human urine isolates (13.5%), but the 213 difference was not statistically significant (P = 0.3). The replicons of the CTX-M-14 214 encoding plasmids for 19 transconjugants with co-resistance to non- $\beta$ -lactam antibiotics were 215 as follows: IncFII (n=1), IncFII, FIB (n=2), IncFIB (n=2), IncFII, I1 $\gamma$  (n=2), IncFII, FIA, FIB 216 (n=1), Inc I1 $\gamma$  (n=5), IncB/O (n=2), A/C (n=1), HI2 (n=1) and nontypeable (n=2)

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 Overall, 37.5% (60/160) isolates were phylogenetic group D, 22.5% (36/160) were

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 group B2, 20.6% (33/160) were group A and 19.4% (31/160) were group B1. Analysis of

plasmid replicon distribution showed that the frequency of IncFII replicon in the four phylogenetic groups were similar. The IncFII frequencies for the virulent groups B2 and D were 47.2% (17/36) and 40% (24/60), respectively; that for the commensal groups A and B1 were 39.4% (13/33) and 22.6% (7/31), respectively (virulent vs. commensal groups, P = 0.1).

224 PCR mapping and sequencing of representative products revealed six of the eight 225 reported types of *bla*<sub>CTX-M-14</sub> genetic environment (Table 3 and Supplementary Table S2). 226 One hundred forty-one (88.1%) isolates were found to have a genetic environment consisting 227 of an ISEcp1 element upstream and an IS903 element downstream of the  $bla_{CTX-M-14}$  gene 228 (type II). The proportions of isolates from animal faecal, human faecal and human urine 229 sources with type II CTX-M-14 genetic environment were 89.7%, 87.7% and 86.5%, 230 respectively. In addition to ISEcp1 and IS903, four other insertion sequences (IS10, IS26, 231 IS6100 and ISCR1) and class 1 integron were found in the other types of genetic environment 232 (III to V, VII and VIII) in different combinations. Type VII and VIII genetic environments 233 were found to have genes encoding resistance to aminoglycosides (amonoglycoside 234 acetyltransferases, *aacA4* and *aadA2*), chloramphenicol (*cml*), sulphonamides (*sul1*) and/or 235 trimethoprim (dihydrofolate reductase, dfrA12).

236 Table 4 summarized the plasmid subtyping result for the 61 CTX-M-14 encoding 237 IncFII plasmids. RST revealed that the IncFII plasmids had four allelic variants: F2 (n=42), 238 F35 (n=17), F43 (n=1) and F51 (n=1). According to the PCR-RFLP patterns for the seven 239 loci (Figure 1), 32 unique plasmid RFLP (pRFLP) profiles were obtained for the 61 plasmids 240 encoding *bla*<sub>CTX-M-14</sub> (Supplementary Table S3). These included 16 plasmids (group 1) with 241 all seven PCR-RFLP patterns identical to the reference plasmid (pHK01), 36 plasmids (group 242 2) with identity over three to six loci, and nine plasmids (group 3) without any identical PCR-243 RFLP pattern or identical patterns in one or two loci only. Hence, 85.2% (52/61) of the 244 IncFII plasmids (group 1 and 2) were considered to be pHK01-like. These 52 pHK01-like 245 plasmids had their origins from human faecal (n=20), human urine (n=14) and animal faecal 246 (n=18, including 1 cattle, 1 chicken, 5 pigs, 9 cats, 1 dog and 1 rodent) sources. There were 247 15 unique pRFLP profiles for plasmids with the F2 and F35 replicon allele each. The 248 presence of the pHK01-associated malB, yac, eitA, eitC and parA/B genes was assessed by 249 monoplex PCRs using the purified plasmids as template. The result showed that 90.4% 250 (47/52) of the pHK01-like plasmids were positive for all targets. In contrast, these genes were 251 variably found among group 3 plasmids (Supplementary Table S3). The proportions of 252 animal faecal, human faecal and human urine isolates with pHK01-like plasmids were 31% 253 (18/58), 30.6% (20/65) and 37.8% (14/37), respectively.

254 The CTX-M-14 encoding plasmids, pHK09 (E. coli strain C017e) and pHK17a (E. 255 coli strain P0014ST) originating from the faecal samples of a child aged one year in 2007 and 256 a pig in 2008, respectively were sequenced (Ho et al., 2011a; Lo et al., 2010). The sizes of 257 pHK09 (GenBank accession JN087528) and pHK17a (GenBank accession JF779678) were 258 70382 bp and 70060 bp, respectively. The sequences of the two plasmids were highly similar 259 (>95%) to pHK01 (Hong Kong, 2004, GenBank accession HM355591, pRFLP 1-1-1-1-1) 260 and the pHK01-like plasmid, pKF3-70 (China, 2006, GenBank accession FJ494913, in silico 261 pRFLP 1-1-1-1-2-4). Figure 1 showed an alignment of the four plasmids (pHK01, pHK09, 262 pHK17a and pKF3-70) according to the functional regions. Except for some sequence 263 variations and possible insertions/deletions, the four plasmids were virtually identical. With 264 reference to pHK01, sequence variations were found in the *repA4* gene (94.5% similarity, 265 pHK17a), the ssb gene (98.9% similarity, pHK17a) and the ISEcp1 element (99.8% similarity, 266 pKF3-70). The transfer region from *traM* to *traX* in pHK09 was more similar to pHK01 267 (99.3%) than to pHK17a (91.6%) or pKF3-70 (91.5%). The regions (traM to traX) in 268 pHK17a and pKF3-70 were almost identical to each other (99.6% sequence identity).

Sequence variations in the transfer region correlated with the observed difference in the restriction digestion pattern over locus E and F in pHK17a and pKF3-70. In pHK17a, sequence variations within orf37 (encoding an adenine-specific DNA methylase gene) correlated with negative PCR result for locus D. The *yfiA* gene (hypothetical protein), which was found between *traR* and *traC* in pHK01 and pHK09, was absent in pHK17a and pKF3-70. In contrast, an *artA* gene (hypothetical protein), absent in pHK01 and pHK09, was inserted between *traF* and *traQ* in pHK17a and pKF3-70.

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### 277 Discussion

278 This study demonstrated the widespread dissemination of pHK01-like plasmids among 279 isolates originating from multiple human and animal sources. The finding showed that 85% 280 (52/61) of the IncFII plasmids and 32.5% (52/160) of all the plasmids encoding CTX-M-14 281 were pHK01-like. Our previous work showed that pHK01-like plasmids play an important 282 role in the community emergence of CTX-M-14 among urinary E. coli in 2004 (Ho et al., 283 2007; Ho et al., 2011b). The present work extends the observation by showing that pHK01-284 like plasmids were found among human and animal faecal isolates collected in different time 285 periods. Of note, pHK01-like plasmids appeared to have approximately equal representation 286 among E. coli isolates of the four phylogenetic groups. This may partially explain its 287 widespread prevalence in bacteria of both animal and human origin. In Hong Kong, studies 288 have revealed that identical gentamicin resistance genes, integron and cassette arrays were 289 shared by isolates from animal faecal, human faecal and human urinary sources at similar 290 prevalence (Ho et al., 2009; Ho et al., 2010).

Since CTX-M-14 is the only antibiotic resistance gene in most pHK01-like plasmids,
persistence and spread of the plasmids cannot be attributed to co-selection by non-β-lactam
antibiotics. This single resistance feature is shared by the group of epidemic pCT-like

294 plasmids (IncK) encoding CTX-M-14 that have been reported in bacteria from human and 295 animals (Cottell et al., 2011). Furthermore, most pHK01-like plasmids had the 1-1 restriction 296 pattern at locus B1 and B2, and were PCR positive for *vac*, *malB*, and *eitA/eitC*. The finding correlated with a conserved array of genes in the variable region (yac-malB-IS903-bla<sub>CTX-M</sub>-297 298 14-ISEcp1-eitABCD). It is possible that the genes in the variable region may be advantageous 299 to the host bacteria. The toxin-antitoxin plasmid stability system (yac) is recognized to play 300 roles in the stable maintenance of large plasmids while the other transport proteins (malB and 301 eitABCD operon) could provide the host bacteria with some metabolic or virulence 302 advantages (Cheng et al., 2009; Zhao et al., 2009). Interestingly, the vac system and eitABCD 303 operon combination were also found in two major avian pathogenic plasmids, pAPEC-O2-304 ColV (GenBank accession AY545598) and pAPEC-O1-ColBM (GenBank accession DQ381420) that are widespread among poultry isolates (Johnson et al., 2006). 305

306 This study subtyped the IncFII plasmids by several methods. The RST scheme 307 discerned related from unrelated plasmids according to the copA sequence, while the pRFLP 308 scheme analysed the plasmid scaffold (Ho et al., 2011b; Villa et al., 2010). Our experience 309 showed that the RST scheme was easy to apply and the discriminatory power was good. 310 However, the RST allele did not consistently predict the plasmid scaffolds. Despite the fact 311 that the two FII alleles (F2 and F35) cluster into two distinct replicon subgroups (Villa et al., 312 2010), plasmids of the two replicon alleles could share highly similar pRFLP profiles. 313 Complete sequencing of two other pHK01-like plasmids confirmed that variations in the 314 pRFLP profiles correlate with DNA insertion, deletion, point mutations and possible 315 homologous recombination. The finding suggests that multiple genetic processes are involved 316 in the evolution of the pHK01-like plasmids.

Among isolates of all sources, the predominant genetic environment of  $bla_{CTX-M-14}$ consisted of an upstream IS*Ecp1* and a downstream IS*903* (type II). This type of genetic environment was similarly prevalent among the IncFII and non-IncFII plasmids. This study also revealed the presence of several other mobile genetic elements (IS10, IS26, IS6100, ISCR1 and class 1 integron) in association with  $bla_{CTX-M-14}$  and/or ISEcp1/IS903 in six different combinations. Thus, many genetic elements might be involved in the mobilization of  $bla_{CTX-M-14}$  into different plasmids (Barlow et al., 2008). In the present study, almost all recognized genetic environment types were found. This could possibly be due to the diverse origins of animals.

326 This study has limitations. Since pigs and chicken from different farms had been 327 mixed at the time of sampling, the geographic origins of these samples could not be traced 328 individually. Therefore, the observed variations in CTX-M-14 genetic environment and 329 divergence in plasmid sequences could be partly related to mixed origins of the samples from 330 local farms and importation from mainland China. Secondly, only two plasmids were 331 completely sequenced. As a result, the full spectrum of sequence divergence in the plasmids 332 could not be comprehensively resolved. Nonetheless, multiple pRFLP profiles involving 333 variations in different loci were found for isolates that originated from the same animal and 334 human sources. The finding highlights plasticity among related plasmids in different modules 335 of the plasmid backbones and the *bla*<sub>CTX-M-14</sub> genetic environment. In the future, additional 336 plasmids with farm and geographical origins clearly defined should be investigated to clarify 337 the epidemiologic pattern of different plasmid sequences. Preferably, a larger number of 338 pHK01-like plasmids from different time periods should be completely sequenced to 339 delineate how plasmid sequence divergence correlates with the epidemiologic information 340 and how that changes over time.

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342 Conclusion

This study showed that the pHK01-like plasmids encoding CTX-M-14 were shared among *E. coli* isolates collected in different time periods from multiple human and animal sources. Public health authorities should adopt measures to reduce the direct and indirect transfer of resistant bacteria or resistance determinants within animal populations, from animals to human and vice versa (Codex Alimentarius Commission, 2005).

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### 350 Acknowledgements

This work was supported by grants from the Research Fund for the Control of Infectious Diseases (RFCID) of the Health, Welfare and Food Bureau of the Government of the HKSAR, and the Consultancy Service for Enhancing Laboratory Surveillance of Emerging Infectious Disease for the HKSAR Department of Health.

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### 356 Transparency Declaration

# 357 Conflicts of interest: Nothing to declare.

358

# 359 Appendix A

360 Supplementary data

#### 361 Table 1

Oligonucleotide primers for PCR mapping of CTX-M-14 genetic environment 362 363 \_\_\_\_\_

					Amplicon size	
Primer name <sup>a, b</sup>	Direction	Nucleotide sequence (5' to 3')	Position	Accession No.	( <b>bp</b> )	Source
ISEcp1U2	F	AATACTACCTTGCTTTCTGA	7650-7669	HM355591	1464	(Saladin et al., 2002)
M9L	В	CCCTTCGGCGATGATTCTC	6206-6224			(Saladin et al., 2002)
ISEcp1U1	F	AAAAATGATTGAAAGGTGGT	7210-7229	HM355591	366	(Saladin et al., 2002)
СТХ-М-9В	В	ATTGGAAAGCGTTCATCACC	6864-6883			(Woodford et al., 2006)
ISCR1F	F	GCCACCAACCCGACCAGAC	4001-4019	EF450247	2019	This study
СТХ-М-9В	В	ATTGGAAAGCGTTCATCACC	6000-6019			(Woodford et al., 2006)
tnpIS26-R	F	AACTCTGCTTACCAGGCG	1413-1430	GQ385314	1507	This study
СТХ-М-9В	В	ATTGGAAAGCGTTCATCACC	2900-2919	-		(Woodford et al., 2006)
ISEcp_uw	F	AACATCAAACGAATCGACCG	799-818	EU136400	1468	This study
M9L_dw	В	CACCTGCGTATTATCTGCGG	2247-2266			This study
Int-F	F	GCCACTGCGCCGTTACCACC	322-341	EF450247		(Ho et al., 2010)
Int1-285B	В	GCACAGCACCTTGCCGTAGAA	66-86		276	This study
CTX-M-9B	В	ATTGGAAAGCGTTCATCACC	6000-6019		5698	(Woodford et al., 2006)

	M9U	F	ATGGTGACAAAGAGAGTGCA	7056-7075	HM355591	1265	(Saladin et
							al., 2002)
	IS903D-5811R	В	TAACCGACTTTGCCCGCCTG	5811-5830			This study
364	<sup>a</sup> Based on publishe	d studies	and sequences deposited in the GenBank of	database, eight diffe	erent types of <i>bla</i> <sub>CT</sub>	<sub>K-M-14</sub> genetic en	vironment have
365	been reported. Type	e I (ISEcp	$1 - bla_{\text{CTX-M-14}}$ ) consisted of an upstream IS	<i>Ecp1</i> but the down	stream IS903 was n	ot detected by P	CR (Kim et al.,
366	2011). However, th	e presence	e of a truncated IS903 has not been confirmed	med by plasmid se	quencing. The GenH	Bank accession	numbers for the
367	other types were HM	M355591 (	(type II, IS <i>Ecp1 - bla</i> <sub>CTX-M-14</sub> - IS903), EU13	36400 (type III, ΔΙS	$SEcp1 - IS10 - bla_{CT}$	<sub>X-M-14</sub> - IS903), (	GQ385314 (type
368	IV, IS26 - $\Delta$ ISEcp1	- bla <sub>CTX-M</sub>	<sub>A-14</sub> - IS903), AB180674 (type V, IS26 - $\Delta$ I	S10 - bla <sub>CTX-M-14</sub> - I	S <i>903</i> ), EU056266 (t	ype VI, Class 1	integron- orf5 -
369	IS6100 -ISCR1 - ΔΙ	SEcp1 - b	<i>la</i> <sub>CTX-M-14</sub> - IS903), FQ482074 (type VII, Cl	lass 1 integron- orf.	l -ISEcp1 - bla <sub>CTX-M</sub>	-14 - IS903) and	EF450247 (type
370	VIII, Class 1 integro	on- ISCR1	-bla <sub>CTX-M-14</sub> - IS903). The class 1 integron i	include type VI and	VIII contain intl1-d	frA12-orfF-aad	42-qacE∆1-sul1
371	(type VI and VIII) of	or ⊿intl1-a	acA4-cml-qacE∆1-sul1 (type VII).				
372	<sup>b</sup> Monoplex PCRs u	sing differ	ent primer pair combination was used to ma	ap the genetic envir	conment to one of the	e eight recognize	ed types. All the
373	assigned types yield	led amplic	ons of the expected sizes. Representative an	nplicons were seque	enced for confirmation	on.	
374							
375							

# **Table 2** 377

Replicon	No. of	Plasmid sizes,	No. (%) according to source			
type <sup>a</sup>	isolates	kb				
		-	Animal	Human	Human UTI	
			faecal	faecal		
FII	61	55 100	21 (36 2)	25 (28 5)	15 (40.5)	
1.11	01	55-100	21 (30.2)	23 (38.3)	15 (40.5)	
FII, FIB	7	100-150	3	3	1	
FIB	6	55-190	1	3	2	
FII, I1 <b>-</b> Ιγ	3	70-100	1	2	-	
FIA	3	55-100	-	1	2	
FII, FIA	2	100-130	-	2	-	
FIA, FIB	1	80	1	-	-	
FII, FIA, FIB	1	150	-	1	-	
Ι1-Ιγ	24	40-120	12 (20.7)	7 (10.8)	5 (13.5)	
B/O	10	60-140	5	5	-	
K	6	80-100	-	5	1	
Others <sup>b</sup>	36	50-250	14	11	11	
Total	160	-	58	65	37	

378 Replicon types for plasmids encoding CTX-M-14 analysed in this study

<sup>a</sup> The replicon types were determined by probe hybridization in the transconjugants (n=84) or

380 the parent strains (n=76).

<sup>b</sup>These included N (n=3), A/C (n=1), HI1 (n=1), HI2 (n=1) and nontypeable (n=30).

382

383

#### **Table 3**

386 CTX-M-14 genetic environment of *E. coli* isolates from humans (*n*=102) and animals (*n*=58),

387 Hong Kong.

Type <sup>a</sup>	Genetic environment of <i>bla</i> <sub>CTX-M-14</sub>		No (%)	
		Animal-	Human-	Human-
		faecal	faecal	urine
II	ISEcp1 - bla <sub>CTX-M-14</sub> - IS903	52	57	32
		(89.7)	(87.7)	(86.5)
III	$\Delta ISEcp1 - IS10 - bla_{CTX-M-14} - IS903^{b}$	-	4	1
IV	IS26 - $\Delta$ ISEcp1 - $bla_{CTX-M-14}$ - IS903	3	1	-
V	IS26 - ΔIS10 - ΔIS $Ecp1$ - $bla_{CTX-M-14}$ - IS903 <sup>c</sup>	1	-	1
VII	Class 1 integron- orf1 -ISEcp1 - bla <sub>CTX-M-14</sub> - IS903	2	1	3
VIII	Class 1 integron- ISCR1-bla <sub>CTX-M-14</sub> - IS903	-	2	-
Subtotal		58	65	37

<sup>a</sup> The Genbank accession numbers were as follows: HM355591 (type II), EU136400 (type III),

389 GQ385314 (type IV), AB180674 (type V), EU056266 (type VI, Class 1 integron- orf5 -

390 IS6100 - ISCR1 -  $\Delta$ ISEcp1 -  $bla_{CTX-M-14}$  - IS903), FQ482074 (type VII) and EF450247 (type

391 VIII). The class 1 integron includes type VI and VIII containing intl1-dfrA12-orfF-aadA2-

392  $qacE\Delta 1$ -sul1 (type VI and VIII) or  $\Delta intl1$ -aacA4-cml-qacE $\Delta 1$ -sul1 (type VII).

393 <sup>b</sup> The  $\Delta$ IS*Ecp1* was interrupted by an IS*10* element.

<sup>c</sup> The  $\Delta ISEcp1$  element was very short and limited to a putative ISEcp1 promoter.

395

**397 Table 4** 

398 Characteristics of the IncFII plasmids encoding *bla*<sub>CTX-M-14</sub> in 61 *Escherichia coli* isolates

- No of plasmids<sup>a</sup> Group 2 Group 3 Group 1 All 36 61 Number of plasmids 9 16 FAB formula F2:A-:B-16 26 42 -F35:A-:B-10 7 17 \_ F43:A-:B-1 1 -\_ F51:A-:B-1 1 -\_ PCR profile malB-yac-eitA-eitC-parAB 16 31 2 49 Others<sup>b</sup> 1 5 6 -None 4 2 6 \_ Source 9 Human UTI 5 1 15 Human faecal 4 16 5 25 Chicken 1 1 --Cattle 1 1 \_ -Pig 1 4 1 6 Cats 3 6 1 10 Dogs 0 1 1 \_ Rodents 2 1 1 -
- 399 from humans and animals.

400	<sup>a</sup> The plasmids were categorised according to their similarity to the reference plasmid, pHK01:
401	group 1, all seven PCR-RFLP patterns were identical to reference; group 2, three to six
402	restriction patterns were identical; and group 3, less than three restriction patterns were
403	identical.
404	<sup>b</sup> Including the following profiles: <i>yac-eitA-eitC-parAB</i> (n=2), <i>yac-eitC-parAB</i> (n=1), <i>mal-</i>
405	<i>eitA-eitC</i> ( <i>n</i> =1), <i>parAB</i> ( <i>n</i> =2).
406	

# FIGURE 1. Schematic representation of pHK09 and pHK17a and comparison with two other CTX-M-14 encoding plasmids, pHK01 and pKF3-70.

The sequence of pHK01 was used as the reference for all comparisons. The five functional regions were indicated by arrows on top while the seven sets of long-range PCR targets (locus A to locus F) were indicated by arrows in the bottom. The genes and ORF of interest were indicated in boxes and annotated. Potential deletions were indicated by dotted line. The *rep* genes in the replication and *tra* genes in the transfer region were indicated by white boxes with number and capital letters, respectively. The percentage identity over each indicated block in the transfer region were given above the boxes. The genes or ORFs with sequence variations were indicated in **boldface**. *Orf37* is named according to pHK01. Its counterpart is *orf35* in pHK09 and pHK17 and *orf81* in pKF3-70. The boxes according to shading patterns were: dotted lines (partition-associated genes, *parA/B, ssb, parB-like, psiA/B*), dots (toxin-antitoxin genes, *yacABC*), oblique lines (transportation-related proteins, *malB* and *eitABCD*), gray (insertion sequences, IS903 and ISEcp1) and black (resistance gene, *bla*<sub>CTX-M-14</sub>). The plasmids' origins and their pRFLP profiles were labelled on the two sides..



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