Analysis of *Escherichia coli* sequence types and resistance mechanisms in sewage from Islamabad, Pakistan indicates difference in *E. coli* carriage types between S. Asia and Europe

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Running title: Carriage *E. coli* sequence types in Islamabad, Pakistan.
Abstract:
Objectives: To discover the *E. coli* carriage sequence types and associated resistance mechanisms in the community in Islamabad, Pakistan by analysis of *E. coli* isolates in sewage.

Methods: 110 *E. coli* were isolated from sewage across the city of Islamabad without antibiotic bias and confirmed as *E. coli* by MALDI-TOF. Isolates were characterized by CH typing and cgMLST. Resistance mechanisms, virulence genes, phylotypes and plasmid incompatibility types were determined in a subset of isolates by in-silico analysis. The genomic position of *bla*\textsubscript{CTX-M-15} was determined using S1 PFGE, probing and Nanopore minion sequencing.

Results and conclusions: The most prevalent sequence types were ST394, ST10 and ST648 accounting for 39% of all isolates collected and were found at many sites across Islamabad. Carbapenemase genes were absent and only a single isolate of ST131 was found. *qnr*S1 and *bla*\textsubscript{CTX-M-15} were the most prevalent resistance mechanisms with *bla*\textsubscript{CTX-M-15} penetrating many sequence types and 31% of all collected isolates. However, the majority of the successful sequence types were *bla*\textsubscript{CTX-M-15} negative indicating that resistance is not the main driver of prevalence. 23% of *bla*\textsubscript{CTX-M-15} genes were chromosomally encoded and large ISEcP1 mediated insertions included *qnr*S1 and several plasmid genes. In all chromosomally encoded isolates no plasmid copies of *bla*\textsubscript{CTX-M-15} were found. The most prevalent ST (ST394) contained many Enteroaggregative *E. coli* (EAEC) virulence genes and the *fimH*30 variant allele previously linked to the success of ST131.
Introduction:

Antibiotic resistance in *Escherichia coli* is a major concern as *E. coli* is both a leading cause of human infection and an ever-present gut colonist. Carriage of ESBL *E. coli* can lead to difficult to treat urinary tract infections (UTIs), delay appropriate therapy and lead to poor outcomes.\(^1\)

The *E. coli* species consists of many different sequence types (>7,000 http://pubmlst.org/databases.shtml) and strains.\(^2\) Individual strains vary in ability to cause disease and to resist antibiotics. Several molecular methodologies have enabled comparisons of strains between diverse geographic locations including: MLST; *fimH/fumC* (CH typing); \(^3\) Phylo-typing \(^4\) and WGS. Comparisons have revealed that resistance is associated with a small number of successful strains and sequence types, for example ST131 *E. coli* belonging to phylogroup B2 \(^5\) and *E. coli* ST101 belonging to the phylogroup B1.\(^6\) We have also seen expansion of ST131 ESBL *E. coli* causing severe infections.\(^7\) A recent study investigating ESBL *E. coli* in the UK found ST131 was the primary cause of cephalosporin resistant bacteraemia’s and the most prevalent ST in faeces and sewage (H. Cadden, P. Cleary, M. Day, M. Doumith, M. Ellington, N. Elviss, J. Findlay, K. Hopkins, B. Jones, D. Livermore, L. Randall, C. Teale, M. Toleman, D. Wareham, C. Wiuff, N. Woodford unpublished results). This suggests that analysis of *E. coli* in sewage is a good proxy for gut carriage in the community.

In this study, we sought to determine the carriage *E. coli* sequence types/resistance mechanisms by analysis of *E. coli* collected without antibiotic selection from 18 sewage outfall sites across the city of Islamabad.
Materials and Methods:

Collection of samples: Samples (30 mL) were collected from 18 sites in 9 sectors of Islamabad (Figure S1).

Bacterial isolation and identification: Samples were centrifuged, pelleted and re-suspended in 0.5mL LB broth. 50 µL was spread on MacConkey plates without antibiotic and grown at 37°C overnight. Ten colonies with typical E. coli morphology were randomly collected from each site with a total of 110 colonies confirmed as E. coli by MALDI-TOF.

Detection of bla_{CTX-M} and bla_{NDM-1} genes: PCR utilized custom primers and ReddyMix Extensor PCR Master Mix 1 (Thermo Scientific) with appropriate controls.

Two Locus CH typing: fimH and fumC genes were amplified by PCR as described by Weissman.³ fumC and fimH alleles were assigned using (http://mlst.warwick.ac.uk/mlst/dbs/Ecoli) and (https://cge.cbs.dtu.dk/services/FimTyper/) websites, respectively.

bla_{CTX-M-15} genomic location: S1 PFGE was performed as described.⁸ The probe was amplified using primers CTXMF/R (GTTCACGCTGATGGCGACGGC, ACGCTAATACATCGCGACGACGC) and radio-labeled using P³² dCTP as described previously.⁸

Genomic DNA extraction: Genomic DNA was extracted using the Qiagen genomic DNA kit.

Miseq Sequencing: DNA libraries were prepared using the NexteraXT sample kit and sequenced (20-30× coverage) with a standard 2x100 base protocol on a MiSeq Instrument (Illumina, San Diego, CA, USA) in house at Cardiff.
**Sequence type and resistance/virulence gene detection:** MLST was determined with StringMLST using short read data in fastq format and Ridom SeqSphere+ (version 3.5.0) using assembled data in fasta format. *E. coli* strains were clustered based on cgMLST typing with SeqSphere+. A core genome MLST (cgMLST) scheme was based on *E. coli* ATCC25922 with 1907 targets. Antimicrobial resistance genes were detected using CLC-Biogenomic workbench. Plasmid and virulence genes were detected using Plasmid finder (https://cge.cbs.dtu.dk/services/PlasmidFinder/); and Virulence finder (https://cge.cbs.dtu.dk/services/VirulenceFinder/).

**Minion sequencing:** Isolates with chromosomally encoded *bla*<sub>CTX-M-15</sub> were sequenced using the nanopore minion RAD-002 rapid sequencing kit following isolation of high molecular weight DNA by the CTAB (hexadecyltrimethylammonium bromide) method. Single reads were used as scaffold to assemble miseq data and annotated using geneious.

**Phylo-group analysis:** The *E. coli* phylogroups were determined with in-silico searches for *chuA*, *yjaA*, *tspE4.C2*, *arpAgpE*, *tnpAgpC* using geneious software based on the Clermont method.
Results and discussion

This study was designed to discover *E. coli* carriage ST and resistance mechanisms in Islamabad. We collected *E. coli* at sewage outfalls across the city and isolated 110 *E. coli* strains without antibiotic selection. PCR indicated 34 isolates carried *bla*$_{CTX-M-15}$ and a single isolate carried *bla*$_{CTX-M-27}$. All isolates were *bla*$_{NDM-1}$ negative and carbapenem susceptible indicating low carbapenemase carriage rates in Islamabad as compared to other S. Asian sites. 10 11 12 13

CH clonotyping revealed 24 known MLST in the collection of 110 *E. coli*’s. These were *fumC- fimH*: 35-30 (ST394); 11-54 (ST10); 4-0 (ST648); 65-32/ 54-0 (ST1431) and 7-54 (ST45) which accounted for 15%, 13%, 7%, 3.6% and 2.7% of isolates respectively. Two isolates of clonotypes, *fumC- fimH*: 4-27 (ST88); 29-38 (ST156); 6-0 (ST688); 7-54 (ST2325); 636-34 (ST5176) and 11-24 (ST43) which equates to 1% prevalence each and single isolates of each clonotype including: 11-27 (ST93); 40-30 (ST131); 6-31 (ST154); 4-32 (ST155); 23-54 (ST205); 11-69 (ST216); 4-61 (ST224); 7-25 (ST398); 136-331 (ST543); 65-32 (ST1128) and 27-0 (ST4121). Details of location, numbers of ST and respective clonotypes are shown in Figure S1. Of 110 *E. coli* isolates, 33 were whole genome sequenced (WGS) including examples of all MLST sequence types and multiple examples of common ones (Table S1). The in silico MLST agreed 100% with MLST derived from CH clonotyping. Most frequently found sequence types were ST394, ST10, ST648 and ST1431 found at 44%, 44%, 28% and 11% of the sampling sites, respectively (Figure S1). Interestingly, ST131 was particularly rare (a single isolate <1%). Isolation of *E. coli* without antibiotic selection enabled us to measure penetration of resistance mechanisms through this species. *bla*$_{CTX-M-15}$ is highly prevalent worldwide 14 and colonized 23% of *E. coli*
isolates in Islamabad including 63% of MLST. However, it was found only in 14%, 24% and 50% of individual isolates of the most prevalent types: ST648; ST394 and ST10, respectively. This indicates that $\text{bla}_{\text{CTX-M-15}}$ E. coli are overall less fit than parental isolates and that the prevalence of individual E. coli ST is not directly related to cephalosporin resistance.

WGS revealed all resistance and virulence mechanisms in 33 isolates including 24 positive and 9 negative CTX-M isolates (Table S1). The $\text{qnrS1}$ gene was most common, found in 80 isolates by PCR. This was often associated with $\text{bla}_{\text{CTX-M-15}}$ (91%). Other prevalent mechanisms were: $\text{dhfr}$ genes found in 60%, $\text{bla}_{\text{TEM-1B}}$ in 33%, $\text{tetA}$ in 42%, $\text{su}l2$ in 33%, $\text{su}l1$ 21% and $\text{tetB}$ 9% of WGS isolates. Other mechanisms were rarely found: $\text{bla}_{\text{OXA-1}}$ in 3; $\text{bla}_{\text{TEM-199}}$ in 2; and $\text{bla}_{\text{TEM-158}}, \text{bla}_{\text{CMY-42}}$, and $\text{bla}_{\text{CMY-44}}$ and in individual isolates Table S1.

WGS detected 19 different virulence genes (Table S1). Overall the majority of isolates were commensals with few virulence factors belonging to phylogenotypes A, C and B1. The ST394 isolates belonged to phylogroup E and harboured entero-aggregative virulence factors typical of the EAEC group of E. coli. ST394 is associated with diarrheal disease but also commonly recovered from healthy people and has been implicated in acute and persistent sporadic diarrhea, and outbreaks, in both industrialized and developing countries. ST131 and ST648 were the only EXPEC pathogens recovered belonging to phylogenotypes B2 and D, respectively. They are known to cause UTI’s and sepsis, though they do not carry the classical EXPEC virulence gene repertoire.

S1 PFGE analysis revealed 1-4 plasmids present in each strain. Plasmids were of a range of incompatibility groups (Table S1). The $\text{bla}_{\text{CTX-M-15}}$ plasmids ranged in size from 50-150 kb. We found $\text{bla}_{\text{CTX-M-15}}$ genes on the chromosome in 8 isolates
and 6 ST. We determined the insertion sites in 4 isolates including identical sites in ST394 and a unique insertion in ST4121 (Figure S2). In all cases movement to the chromosome was \textit{ISEcP1} mediated and confirmed by the presence of direct repeats generated at the site of insertion (Figure S2). Notably these movement events not only transposed \textit{bla} \textit{CTX-M-15} to the chromosome but also sections of a common plasmid containing \textit{qnrS1}. Movement to the chromosome is typically associated with subsequent loss of the \textit{bla} \textit{CTX-M-15} plasmid as evidenced by lack of \textit{bla} \textit{CTX-M-15} harboring plasmids in these strains. This process likely enhances the fitness of the strain overall.\cite{17}

We used cgMLST to relate the various \textit{E. coli} to each other (Figure 1). ST were correctly grouped together. The Clermont PCR phylogroup analysis gave good agreement in general with cgMLST. Interestingly, cgMLST aligned the ST648 isolates as group F instead of D, confirming the recent observation of Johnson\cite{18} and highlighting the superior discriminatory power of cgMLST.

Analysis of the \textit{fimH} data revealed that several strains were \textit{fimH} null. This included all ST648 isolates and is typical of this ST.\cite{6} Its prevalence as an EXPEC pathogen suggests that another adhesion is substituting for \textit{fimH}. The worldwide success of the \textit{bla} \textit{CTX-M-15} gene in ST131 \textit{E. coli} has been closely linked to a highly adherent \textit{fimH} variant allele, \textit{fimH30}.\cite{5} In this study despite ST131 being particularly rare, \textit{fimH30} was the second most prevalent variant of \textit{fimH} and was present in all ST394 isolates. The possession of \textit{fimH30} by ST394 is likely advantageous. However, the fact that ST131 \textit{fimH30} \textit{E. coli} is particularly rare suggests that other as yet unknown factors are responsible for the success of ST394 in Islamabad.
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Figure and Table legends:

**Figure 1. cgMLST of Islamabad sewage isolates.**

cgMLST was generated using a core genome of 1907 target genes identified in each sequenced isolate using Ridom SeqSphere+ software. Phylogroup analysis was based on in silico PCR using targets identified by Clermont et al 2013.

**Figure 2. Chromosomal insertion sites of bla_{CTX-M-15}.**

Chromosomal insertion sites were identified using a combination of miseq and Nanopore Minion single reads sequences. (A) represents the insertion site found in the ST4121 isolate 8 and (B) represents the insertion site found in ST394 isolates 2, 12 and 15. The 11,389bp insertion found in isolates 2, 12 and 15 included bla_{CTX-M-15} and qnrS1 and was identical to the first 11,389bp of the 23,174bp insertion in ST4121 isolate 8. The larger insertion in ST4121 isolate 8 included additional tetA and tetR resistance genes as well as several other genes of plasmid origin. The chromosomal insertion in the ST394 isolates was in a low GC % intergenic region between ydcs and ydcR genes encoding a putative DNA binding transcriptional regulator and a putative spermidine putrescine transporter subunit, respectively. The insertion in the ST4121 isolate was in a hydrolase gene. Each insertion had the mobile element ISEcp1 at the LH terminus and was flanked by target site direct repeats of TATGA for isolate 8 and TTAAA for isolates 2, 12 and 15. This figure appears in colour in the online version of JAC and in black and white in the printed version of JAC.
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S2 legend

(Figure 3). This insertion was 11,398 bp and was ISEcP1 mediated with a typical 
5bp target site duplication of host DNA adjacent to the ISEcP1 left hand terminus 
(TATGA) and a surrogate ISEcP1 right hand terminus at the insertion site 
boundaries. The insertion included blac\text{CTX-M-15} and qnrS1 resistance genes as well 
as several partial or complete transposon genes and was similar to blac\text{CTX-M-15} 
containing sections of several plasmids suggesting a plasmid origin. The 
chromosomal insertion in ST4121 included an identical section of DNA adjacent 
to ISEcP1 but was considerably longer including 23,174 bp and included tetA and 
tetR genes in addition to blac\text{CTX-M-15} and qnrS1 genes as well as several other 
genes of plasmid origin. This insertion was also ISEcP1 mediated with a typical 
direct repeat of target DNA at the insertion site (TTAAAA), which was at position 
112bp of a hydrolase gene (Figure 3).