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Citation for final published version:

Paul, Deepjyoti, Babenko, Dmitriy and Toleman, Mark A 2020. Human carriage of cefotaxime-resistant Escherichia coli in North-East India: an analysis of STs and associated resistance mechanisms. Journal of Antimicrobial Chemotherapy 75 (1), pp. 72-76. 10.1093/jac/dkz416

Publishers page: http://dx.doi.org/10.1093/jac/dkz416

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Human carriage of cefotaxime resistant *Escherichia coli* in North-East India: an analysis of

sequence types and associated resistance mechanisms.

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Short running title: E. coli ST in North East India

### 1 **SYNOPSIS**

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2 **Objectives:** To determine the prevalence of *E. coli* sequence types and associated resistance 3 mechanisms carried by the community in North-Eastern India. 4 Methods: E. coli (108) were isolated from sewage collected from 19 sites across the city of 5 Silchar by plating on MacConkey agar with/without selection (50mg/L ceftotaxime). Species identification was confirmed by MALDI-TOF MS for 82 isolates. Common resistance 6 7 mechanisms were determined by WGS of pooled *E. coli* isolates. PFGE combined with specific 8 probes determined the presence of common resistance mechanisms in all isolates. Phylotype, 9 MLST, cgMLST, resistance gene and virulence gene content were determined by in silico 10 analysis of 38 genomes. 11 **Results and conclusions:** Analysis of isolates collected without selection (n=33) indicated that 12 CTX resistance in *E. coli* was 42% (14/33) and estimated meropenem resistance at 9%. The 13 remaining 58% (19/33) were additionally sensitive to ampicillin, trimethoprim, ciprofloxacin 14 and aminoglycosides. The most common ST among the CTX resistant E. coli's was ST167 (29%) followed by ST410 (17%) and ST648 (10%). E. coli ST131 was absent from the collection. 15 16 Sixty-three isolates were resistant to cefotaxime, and harboured blactx-M-15, 54% (34/63) or bla<sub>CMY-42</sub>, 46% (29/63) of which 10% (6/63) harbored both genes. Carbapenem resistance was 17 due to *bla*NDM-5, found in 10/63 CTX resistant isolates and/or *bla*OXA-181 found in 4/63 isolates. 18 19 NDM-5 was encoded by IncX3 and/or IncFII plasmids and CMY-42 was mostly encoded by IncI 20 plasmids. NDM-5 appears to have replaced NDM-1 in this region and CMY-42 appears to be in 21 the process of replacing CTX-M-15. 22

# INTRODUCTION

Escherichia coli is universally carried in the human gut and is one of the most common
bacterial pathogens causing a range of disease manifestations. <sup>1</sup> Importantly, the main source of
infection is the patients' own digestive tract with the main site of infection the urinary tract.
$\it E.~coli$ is the leading cause of urinary tract infections $^2$ and subsequent septicaemia's in many
nations. <sup>3, 4</sup> E. coli has an "open genome" meaning that as a species it easily gains and loses
genetic information. <sup>5</sup> Thus comparison of <i>E. coli</i> genomes has revealed that the core genome
only consists of c. 1000 genes ( $1/5^{th}$ of the genome of each isolate). <sup>6</sup> This diverse genetic make-
up means that any individual isolate may be a serious pathogen or an innocuous commensal
and therefore typing of common community carriage isolates is useful to understand the link
between infection and prevalence of gut carriage of virulent and/or antibiotic resistant clones.
Antibiotic resistance is also an important marker since increasing antibiotic resistance
prevalence is associated with both increasing bacteraemia rates <sup>3</sup> and associated mortality due
to delay in appropriate therapy. Furthermore, rise in the prevalence of gut carriage of a
virulent and antibiotic resistant strain of <i>E. coli</i> has a compounded effect. <sup>3</sup> This is important
since European bacteraemia rates have substantially increased over the last 20 years, partially
due to rising carriage rates of a virulent and often multi-drug resistant <i>E. coli</i> strain, ST131. <sup>3</sup>
We have previously shown that common <i>E. coli</i> sequence types and resistance mechanisms can
vary dramatically in different human communities.8 To further understand this observation we
sought to determine these in the city of Silchar, North East India.

51	Materials and Methods:		
52	Collection of samples		
53	Human sewage samples (30mL) were collected in January 2018 from 19 sites across the city of		
54	Silchar, Assam, India.		
55	Bacterial Identification		
56	Bacteria were pelleted by centrifugation and resuspended in 500μL LB broth. 50μL of several		
57	serial dilutions were then spread on MacConkey agar plates with and without 50mg/L		
58	cefotaxime. Up to 10 colonies with typical <i>E. coli</i> morphology were randomly collected from		
59	each site, 5 with and 5 without selection (55 and 53 with and without selection, respectively)		
60	of which 49 and 33 were confirmed as <i>E. coli</i> by MALDI-TOF MS.		
61	Genomic DNA extraction		
62	Genomic DNA was extracted using the Qiagen genomic DNA kit.		
63	MiSeq sequencing		
64	DNA libraries were prepared using the Nextera XT sample kit and sequenced at 30X coverage		
65	with a standard 2X 100 base protocol on a MiSeq instrument (illumina, San Diego, CA, USA).		
66	Initial WGS analysis		
67	Isolates were pooled into ten pools consisting of 8 isolates per pool and sequenced at 30X		
68	coverage by illumina Miseq to give an indication of the range of resistance mechanisms in the		
69	samples.		
70	PFGE and specific probing		
71	PFGE was performed as described previously. <sup>9</sup> Gels were probed directly using radio-labeled		
72	probes for $bla_{\text{NDM}}$ , $bla_{\text{CTX-M-15}}$ , $bla_{\text{CMY-42}}$ , $qnrS1$ .		
73	ST and virulence gene detection		
74	The MLST of sequenced strains was determined with StringMLST using short read data in		
75	FASTQ format and Ridom Seqsphere + (version 3.5.0) using assembled data in FASTA format.		

CH typing (fumC/fimH) was used to indicate the MLST group of non-sequenced strains. E. coli
strains were clustered based on core genome MLST (cgMLST). Antimicrobial resistance genes
were detected using CLC Biogenomic workbench. CH types, plasmid and virulence genes were
determined online using CH typer, Plasmid finder and Virulence finder

(<a href="http://www.genomicepidemiology.org/">http://www.genomicepidemiology.org/</a>).

## Phylogroup analysis

The *E. coli* phylotypes were determined with in-silico searches for *chuA*, *yjaA*, *tspE4C2*, *arpAgpE* and *tnpAgpC* using geneious software based on the Clermont method.<sup>11, 12</sup>

## **Results and discussion**

We isolated 82 *E. coli* collected from 19 sites across the city of Silchar to determine the *E. coli* ST and associated resistance mechanisms carried by the local population. The 82 *E. coli* included 33 isolates collected without selection and 49 by selection on 50mg/L cefotaxime. Fifty-eight percent of isolates collected without selection (19/33) were cefotaxime susceptible and also susceptible to all tested antimicrobials including ciprofloxacin, trimethoprim and gentamicin. Sixty-three isolates were cefotaxime resistant including 14/33 collected without selection, thus 42% of carriage *E. coli* in Silchar are cefotaxime resistant. Twenty-percent (10/49) of cefotaxime resistant isolates were also resistant to carbapenems giving an estimate of carriage of carbapenem resistance in *E. coli* of 9% (42/100 X 10/49). Cefotaxime and carbapenem resistant isolates were found at 100% and 47% (9/19) of sample sites, respectively (Figure 1, Table S1). The total complement of resistance mechanisms was initially determined by MiSeq sequencing of ten pools of eight isolates at 30X coverage. This indicated a high prevalence of resistance genes *blactx.m.-15, blacmy-42, blandm-5* and *qnrS1* and allowed us to target their presence by PCR/sequencing and genomic location by in-gel radio-labeled probing of PFGE gels (Table S1). The most common gene conferring cefotaxime resistance was *blactx.m.* 

101 15 found in 34/63 cefotaxime resistant isolates followed by blacmy-42 (29 isolates) and blandm-5 102 (10 isolates). One isolate contained all three ß-lactamases and six isolates contained both 103 blactx-M-15 and blacmy-42 genes, several isolates contained multiple ß-lactamase genes (Figure 1, 104 Table S1). The *bla*<sub>NDM-5</sub> and *bla*<sub>CMY-42</sub> genes were found on plasmids ranging in size from 85-105 140kb and 25-130kb (Table S1), respectively. However, blactx-M-15 and gnrS1 genes were found 106 mostly on the chromosome 66% (22/33) or on plasmids of 40-175kb. 38 individual isolates 107 were further chosen for whole genome sequencing to determine the MLST types and resistance 108 and virulence gene complements (Figure 1, Figure S1, Table S1). In addition, a further 10 109 isolates were typed by the Weissman 2 locus scheme. MLST was determined for 48 isolates 110 (Figure 1, Figure S1, Table S1), the most prevalent being ST167, 29% (14/48), followed by 111 ST410 17% (8/48), ST648 10% (5/48), ST224 (2/48), ST609 (2/48), ST973 (2/48), ST2083 112 (2/48) and single isolates of ST46, ST84, ST101, ST156, ST215, ST315, ST361, ST617, ST405, 113 ST2521, ST3268 and ST4450 and one new ST (Figure 1, Table S1). ST167 was found at 9/19 114 sites and ST410 was found at 6/19 sites. Over half of all of isolates (56%) belonged to ST167, 115 ST410 and ST648 (Figure 1). In a recent UK nationwide study of cefotaxime resistant *E. coli* 116 isolated from human faeces (360 isolates), sewage (65 isolates) and bacteraemia's (293) 13, 14 117 the most prevalent *E. coli* were ST131 and ST38 (44%, 31% and 70% of all isolates from faeces, 118 sewage and bacteraemia's, respectively). Interestingly, in this study both ST38 and ST131 were 119 absent. Thus, the prevalence of *E. coli* ST carriage varies greatly by geographic location. In 120 Silchar, the common sequence type ST167 was closely associated with  $bla_{NDM-5}$ , 50% (7/14) 121 and both ST167 and ST410 with *bla*<sub>CMY-42</sub>, 65% (9/14) and 100% (8/8), respectively (Figure 1). 122 This high prevalence of *bla*<sub>CMY-42</sub> has not been documented before in India or elsewhere and 123 appears to be in the process of replacing  $bla_{CTX-M-15}$  at this location. This may be related to the 124 wider spectrum of ß-lactam hydrolysis of CMY-42 as compared to CTX-M-15. Similarly blandm-5 125 appears to have replaced *bla*NDM-1 in Silchar as no isolates were detected with *bla*NDM-1. Whilst

this may be a local phenomenon, it is interesting that many nations have reported increased detection of *bla*<sub>NDM-5</sub> over the last few years. <sup>15, 16, 17, 18</sup> Many isolates harboured other resistance mechanisms (Figure S1, Table S1) and chromosomal mutations conferring high level fluoroquinolone resistance making many strains MDR (Figure S1, Table S1). However, fosfomycin and chloramphenicol resistance was low (only 3 strains produced a full-length chloramphenicol resistance gene) suggesting that these antibiotics could be useful treatment options in this locale. Phylotype and virulence gene analysis indicated that most isolates belonged to non-pathogenic phylogroups A and B1, which included the majority of isolates carrying carbapenemase genes (ST167 and ST410). Six isolates belonged to the pathogenic phylotype D (ST268, ST315, STNEW, ST405, 2X ST 973, Figure S1) yet none belonged to the B2 group. Many of the isolates (20/48) including 2x ST648 and the majority of ST167 isolates were missing the type 1 fimbrial adhesion *fimH* which has been shown to be essential for colonizing the urinary tract. The cgMLST (Figure S1, Figure 2) agreed with the phylogroup analysis and demonstrated that all sequenced strains were unique confirming the random nature of the selection process. The within ST SNP analysis further demonstrated this and highlighted that differences within ST648 and ST167 were more numerous than within ST410, perhaps suggesting that the ST410 have expanded within the Silchar population in more recent history. In conclusion, our survey has revealed that the majority of *E. coli* strains in this area are fully sensitive to antibiotics. However, 42% and 9% of isolates are cefotaxime and carbapenem resistant, respectively. We also found that the majority of resistant strains belonged to just three prevalent ST, which are different to the prevalent E. coli ST in Europe. Notably, E. coli ST131, the dominant cefotaxime resistant strain found throughout Europe and North America was absent from our study. Since UTI infections typically originate in the community and from

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150 the human gut, the survey of resistant human carriage isolates is a useful surveillance 151 approach to quickly identify common ST and resistance mechanisms and guide local therapy. 152 **Funding** 153 This work was funded by an International Society for Infectious Diseases/European Society of 154 Clinical Microbiology and Infectious Diseases joint fellowship awarded to Dr Deepjyoti Paul 155 2017 and was presented at the European Congress of Clinical Microbiology and Infectious 156 Diseases April 13-16, 2019. **Transparancy declaration** 157 158 All authors have none to declare. 159 160 References 161 1. Russo TA, Johnson JR. Medical and economic impact of extraintestinal infections due to 162 Escherichia coli: focus on an increasingly important endemic problem. Microbes Infect 163 2003; **5**: 449-56. 164 Schito GC, Naber KG, Botto H et al. The ARESC study: an international survey on the 2. 165 antimicrobial resistance of pathogens involved in uncomplicated urinary tract 166 infections. *Int J Antimicrob Agents* 2009; **34**: 407-13. 167 3. de Kraker ME, Jarlier V, Monen JC et al. The changing epidemiology of bacteraemias in 168 Europe: trends from the European Antimicrobial Resistance Surveillance System. *Clin* 169 Microbiol Infect 2013; **19**: 860-8. 170 Laupland KB. Incidence of bloodstream infection: a review of population-based studies. 4. Clin Microbiol Infect 2013; **19**: 492-500. 171 172 5. Lukjancenko O, Wassenaar TM, Ussery DW. Comparison of 61 sequenced Escherichia 173 coli genomes. Microb Ecol 2010; 60: 708-20.

- Land M, Hauser L, Jun SR *et al.* Insights from 20 years of bacterial genome sequencing.
   Funct Integr Genomics 2015; 15: 141-61.
- 7. Schwaber MJ, Carmeli Y. Mortality and delay in effective therapy associated with
  extended-spectrum beta-lactamase production in Enterobacteriaceae bacteraemia: a
  systematic review and meta-analysis. *J Antimicrob Chemother* 2007; **60**: 913-20.
- Zahra R, Javeed S, Malala B *et al.* Analysis of *Escherichia coli* STs and resistance
   mechanisms in sewage from Islamabad, Pakistan indicates a difference in E. coli
   carriage types between South Asia and Europe. *J Antimicrob Chemother* 2018.
- Toleman MA. Direct in Gel Genomic Detection of Antibiotic Resistance Genes in S1
   Pulsed Field Electrophoresis Gels. *Methods Mol Biol* 2018; **1736**: 129-36.
- 184 10. Weissman SJ, Johnson JR, Tchesnokova V *et al.* High-resolution two-locus clonal typing
   185 of extraintestinal pathogenic *Escherichia coli. Appl Environ Microbiol* 2012; **78**: 1353-60.
- 11. Clermont O, Christenson JK, Denamur E *et al.* The Clermont *Escherichia coli* phylo-typing method revisited: improvement of specificity and detection of new phylo-groups.
- 188 *Environ Microbiol Rep* 2013; **5**: 58-65.

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4.

- 189 12. Beghain J, Bridier-Nahmias A, Le Nagard H *et al.* ClermonTyping: an easy-to-use and accurate in silico method for *Escherichia* genus strain phylotyping. *Microb Genom* 2018;
- 13. Livermore DM, Day M, Cleary P et al. OXA-1 beta-lactamase and non-susceptibility to
   penicillin/beta-lactamase inhibitor combinations among ESBL-producing Escherichia
- 194 *coli. J Antimicrob Chemother* 2019; **74**: 326-33.
- Sattar H, Toleman M, Nahid F et al. Co-existence of bla(NDM-1) and bla(KPC-2) in
   clinical isolates of Klebsiella pneumoniae from Pakistan. Journal of Chemotherapy 2016;
   28: 346-9.

198	15.	Li X, Fu Y, Shen M <i>et al.</i> Dissemination of blaNDM-5 gene via an IncX3-type plasmid	
199		among non-clonal <i>Escherichia coli</i> in China. <i>Antimicrob Resist Infect Control</i> 2018; <b>7</b> : 59.	
200	16.	Huang Y, Yu X, Xie M et al. Widespread Dissemination of Carbapenem-Resistant	
201		Escherichia coli Sequence Type 167 Strains Harboring blaNDM-5 in Clinical Settings in	
202		China. Antimicrob Agents Chemother 2016; <b>60</b> : 4364-8.	
203	17.	Howard JC, Creighton J, Heffernan H et al. Evidence of transmission of an NDM-5-	
204		producing Klebsiella pneumoniae in a healthcare facility in New Zealand. J Antimicrob	
205		Chemother 2017; <b>72</b> : 949-51.	
206	18.	Reynolds ME, Phan HTT, George S et al. Occurrence and characterization of Escherichia	
207		coli ST410 co-harbouring blaNDM-5, blaCMY-42 and blaTEM-190 in a dog from the UK. J	
208		Antimicrob Chemother 2019.	
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210	Legend Figure 1		
211	This figure appears in colour in the online version of JAC and in Black and hite in the printed		
212	version of JAC.		
213	The association of ß-lactamase resistance mechanism with <i>E. coli</i> ST for 48 carriage isolates.		
214	The Inner ring corresponds to the percentage of cefotaxime resistant <i>E. coli</i> belonging to each		
215	sequence type. MLST was determined in silico for 38 isolates that had been whole genome		
216	sequenced (Strain ID and isolation site in bold) and a further ten isolates by CH typing (normal		
217	font). The outer ring indicates the ß-lactamase resistance mechanisms associated with each		
218	isolate. Isolate ID and isolation site is indicated outside the second ring. Isolates coloured		
219	yellov	w and green belong to commensal phylotype A. Isolates coloured white-blue belong to	
220	phylo	type B1 and isolates coloured orange to red belong to pathogenic phylogroups F and D.	

Legend figure S1

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222 This figure appears in colour in the online version of JAC and in Black and hite in the printed 223 version of JAC. 224 Core genome analysis of sequenced isolates (38) with detail of non-ß-lactam resistance 225 mechanisms. Chromosomal mutations in gyrase A and parC genes rthat result in amino acid 226 substitutions known to confer ciprofloxacin resistance in *E. coli* are given in the quinolone 227 column along with acquired genes that also confer quinolone resistance. Virulence genes and 228 fimH types as well as phylogroup are given for each strain. Isolates coloured yellow and green 229 belong to commensal phylotype A. Isolates coloured white-blue belong to phylotype B1 and 230 isolates coloured orange to red belong to pathogenic phylogroups F and D. 231 Legend figure 2 232 This figure appears in colour in the online version of JAC and in Black and hite in the printed 233 version of JAC. 234 SNP variation found among and between sequence types. Isolates belonging to the same 235 sequence type are highlighted as clusters 1-6. The number of SNP's found between members of 236 the same sequence types and between sequence types are given in bold adjacent to each isolate 237 pair. Cluster 2 ST410 isolates shared the least within ST SNP variation. Colours represent 238 different ST groups as in other figures. 239 Table S1 240 Table gives ST and CH data of all 63 cefotaxime resistant *E. coli* isolates collected in this study 241 together with entire resistance gene complements and plasmid size, incompatibility group and 242 genomic location of the most prevalent resistance genes. Whole genome sequenced isolates are 243 highlighted in bold. Isolates with  $bla_{CMY}$  genes other than  $bla_{CMY-42}$  are highlighted with an 244 asterix in the table ie DJ-95 produces blacmy-145 and isolate DJ58 produces blacmy-4. The genomic 245 location of resistance genes is highlighted by plasmid size and chromosomal location of CTX-M-

- 246 15 genes by c. Selection or non selection is indicated by CTX (50mg/L cefotaxime) or NS,
- 247 respectively.
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