Complete assembly of Escherichia coli ST131 genomes using long reads demonstrates antibiotic resistance gene variation within diverse plasmid and chromosomal contexts

Running title: Resolving ST131 genomes using GridION sequencing

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- 15 16 **Author contributions:**

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21 22 Abstract

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23 The incidence of infections caused by extraintestinal *Escherichia coli* (ExPEC) is rising globally, 24 which is a major public health concern. ExPEC strains that are resistant to antimicrobials have been 25 associated with excess mortality, prolonged hospital stays and higher healthcare costs. E. coli ST131 is a major ExPEC clonal group worldwide with variable plasmid composition, and has an array of genes 26 27 enabling antimicrobial resistance (AMR). ST131 isolates frequently encode the AMR genes bla_{CTX-M}-28 14/15/27, which are often rearranged, amplified and translocated by mobile genetic elements (MGEs). 29 Short DNA reads do not fully resolve the architecture of repetitive elements on plasmids to allow 30 MGE structures encoding bla_{CTX-M} genes to be fully determined. Here, we performed long read 31 sequencing to decipher the genome structures of six E. coli ST131 isolated from six patients. Most 32 long read assemblies generated entire chromosomes and plasmids as single contigs, contrasting with 33 more fragmented assemblies created with short reads alone. The long read assemblies highlighted 34 diverse accessory genomes with *bla*_{CTX-M-15}, *bla*_{CTX-M-14} and *bla*_{CTX-M-27} genes identified in three, one and one isolates, respectively. One sample had no *bla*_{CTX-M} gene. Two samples had chromosomal 35 36 *bla*_{CTX-M-14} and *bla*_{CTX-M-15} genes, and the latter was at three distinct locations, likely transposed by the 37 adjacent MGEs: ISEcp1, IS903B and Tn2. This study showed that AMR genes exist in multiple 38 different chromosomal and plasmid contexts even between closely-related isolates within a clonal 39 group such as E. coli ST131.

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41 Importance

42 Drug-resistant bacteria are a major cause of illness worldwide and a specific subtype called

43 Escherichia coli ST131 cause a significant amount of these infections. ST131 become resistant to

44 treatment by modifying their DNA and by transferring genes among one another via large packages of

genes called plasmids, like a game of pass-the-parcel. Tackling infections more effectively requires a 45

46 better understanding of what plasmids are being exchanged and their exact contents. To achieve this, we applied new high-resolution DNA sequencing technology to six ST131 samples from infected

47 patients and compared the output to an existing approach. A combination of methods shows that drug-

48 49 resistance genes on plasmids are highly mobile because they can jump into ST131's chromosomes. We

- 50 found that the plasmids are very elastic and undergo extensive rearrangements even in closely related
- 51 samples. This application of DNA sequencing technologies illustrates at a new level the highly

52 dynamic nature of ST131 genomes.

53

54 **Keywords:** Genome assembly, plasmid, MGE, antibiotic resistance, Nanopore, sequencing.

Introduction

Reported cases of bloodstream and urinary tract infections caused by extraintestinal pathogenic *Escherichia coli* (ExPEC) are increasing globally at an alarming rate [1]. As a key source of ExPEC
isolates worldwide, *E. coli* sequence type 131 (ST131) is regarded as a serious threat to public health,
given its high level of antimicrobial resistance (AMR), as well as the broad spectrum of infections it
causes in community and hospital settings [2,3].

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64 E. coli ST131 is virulent [4] and has an expansive range of virulence factors [5,6], especially those 65 linked to uropathogenic E. coli (UPEC) [3,7,8]. AMR and virulence genes allow ST131 to adapt to 66 drug selection pressure and to survive in extraintestinal niches, and are often encoded on mobile genetic elements (MGEs) [9], which means the exact set of virulence and AMR genes in a single 67 ST131 isolate may vary [8,10]. ST131 encodes a range of extended-spectrum β-lactamases (ESBLs) 68 69 that hydrolyse third-line drugs including cephalosporins, the most common of which encode 70 cefotaximase *bla*_{CTX-M-15}. Within ST131, clade C2 has more AMR genes than other clades and is 71 typically *bla*_{CTX-M-15}-positive, differentiating it from clade C1 that can be *bla*_{CTX-M-14} or *bla*_{CTX-M-27}-72 positive [3,8].

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74 Most ST131 AMR genes are reported to be encoded on plasmids: circular self-replicating double-

stranded DNA molecules that constitute part of the bacterial accessory genome [11-13]. Plasmids can
 reduce bacterial cell fitness, but a number of post-segregation killing and stable plasmid inheritance

76 mechanisms allow the stable maintenance of IncF plasmids in ST131 [14-16]. The chromosomal

78 integration of plasmid genes is most commonly facilitated by transposons, which can ensure

79 acquisition and conservation of such elements if there is no subsequent local recombination [17-18].

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81 Identifying plasmid conjugation, recombination and transposition could have value in tracking AMR 82 genes associated with disease outbreaks and antibiotic treatment failures. Plasmids may be classified 83 using incompatibility (Inc), relaxase (MOB) and mating pair formation system typing [19], but 84 difficulties in plasmid genetic analysis and reconstruction arise with short read data due to 85 rearrangements driven by recombination, dense arrays of repetitive elements including transposable 86 elements (TEs), changes in gene copy numbers, and high sequence variation. Methods using short

reads alone may fail to detect genomic segments exchanged between plasmids and the chromosome,limiting evaluation of the core and accessory genomes.

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Whole genome sequencing has provided a high resolution of the genomic epidemiology of ST131 and 90 91 plasmid-mediated AMR outbreaks [20]. However, short reads alone are insufficient to resolve 92 plasmids that often have numerous small MGEs of ~1 kb or less in size, e.g. TEs and insertion 93 sequences (ISs) [21]. Complex transposable units (TUs) consisting of multiple TEs or ISs can mobilise AMR genes by transposition, and this can sometimes be followed by recombination within the TU 94 95 between one of the inverted repeats (IRs) flanking the TE and the IR of another local TE or an adjacent 96 homologous sequence, resulting in different TU structures, locations and copy numbers. At present, the exact resolution of complex structural rearrangements of repetitive TUs containing AMR genes 97 98 may be impossible with short reads [22]. Consequently, plasmid assembly is a challenge requiring 99 accurate long reads and sufficient coverage to distinguish between independent plasmids with regions

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102 Long reads, such as those generated using Oxford Nanopore Technologies (ONT) or Pacific

103 Biosciences platforms can provide a solution to this plasmid assembly problem [24-26]. Here, we

sequenced six ST131 using the ONT GridION X5 platform. Using the resulting high-coverage

sequence data, we reconstructed and annotated the plasmids and chromosomal regions carrying bla_{CTX} .

M genes, as well as their genetic context and copy numbers.

of sequence identity [21,23].

Methods

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110 Sample collection

Six ESBL-producing *E. coli* ST131 clinical strains were isolated in June-October 2015 from patients at
Addenbrooke's Hospital, Cambridge, as part of a study on longitudinal surveillance of antibiotic
resistance in the hospital (Supplementary Table 1). Five samples were from faeces, and one was from
blood. These were short-read sequenced in a multiplex run on an Illumina HiSeq 2500 platform and
processed as previously outlined [27].

118 High molecular weight DNA extraction

Frozen stocks of the six isolates were streaked onto LB agar plates and grown overnight at 37°C.
Single colonies were subcultured onto LB agar plates and incubated overnight at 37°C. DNA was
extracted using a Lucigen Masterpure Complete DNA and RNA Purification kit. For each sample, a
swab was used to sweep half a plate of pure colonies, and suspended in 1x phosphate buffer solution
(PBS). Samples were processed according to the manufacturer's instructions, with elution in 70ul of
Nuclease Free water. Pipetting was minimised to reduce shearing of the DNA prior to sequencing.

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127 Oxford Nanopore library preparation and sequencing

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DNA was quantified using a Quant-iT[™] HS (High Sensitivity) kit (Invitrogen). DNA purity was
checked using a Nanodrop (ThermoFisher) and fragment size was confirmed by FEMTO Pulse (Nano
Life Quest). The sequencing libraries were prepared using 1 µg DNA per sample and ligation
sequencing kit 1D SQK-LSK109 with the barcoding extension kit EXP-NPB104 according to ONT
protocols. The samples were combined using equimolar pooling and loaded onto a single 9.4.1 MIN106 flow cell and sequenced on the GridION X5 platform under standard conditions.

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136 Illumina library preparation and sequencing137

The short reads used in this study were created as follows: bacterial genomic DNA was extracted using the QIAxtractor (Qiagen, Valencia, CA, USA) according to the manufacturer's instructions. Library preparation was conducted according to the Illumina protocol and sequenced (96-plex) on an Illumina HiSeq 2500 platform (Illumina, San Diego, CA, USA) using 100 bp paired-end reads.

143 Oxford Nanopore base-calling and adapter trimming

144 The resulting fast5 read files were transferred to a separate Linux server 4.4.0 (Ubuntu 16.04.4) for 145 analysis. Basecalling was performed during the GridION run using ONT's Guppy v0.5.1 and the 146 147 resulting fast5 from the initial run was converted to fastq format with Albacore v2.0 (ONT). The 148 statistical data of the sequencing run was processed with MinIONQC v1.3.5 [28] based on the default 149 Q score cut-off of seven. Adapters and chimeric reads were removed from fastq files using Porechop 150 v0.2.4 [29] with demultiplex settings (Supplementary Figure 1). Standard outputs were saved as log 151 files and were then parsed. The quality of the final fastq files was assessed using FastQC v0.11.8 152 (https://www.bioinformatics.babraham.ac.uk/projects/fastqc/) and MultiOC v1.4 [30].

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154 Genome assembly and improvement

We assembled the genomes using the conservative, normal and bold modes of the long read-only

assembly pipeline in Unicycler v4.6. Previous work has suggested that Unicycler outperforms

alternatives [21] that struggle to resolve plasmids [31]. This workflow included the assembly polisher,

Racon, which ran iteratively to minimise error rates of called bases [29]. For comparison, short read-

only and hybrid assemblies were also created using Unicycler v4.6. Briefly, during short read-only
 assembly, Unicycler v4.6 employed SPAdes v3.12 to assemble short reads then used Pilon to polish

162 the assembly. In hybrid assemblies, Unicycler v4.6 used Miniasm to piece long reads together first and

applied SPAdes v3.12 to incorporate short reads and bridge gaps. Pilon was run several times toachieve the most contiguous and completed genome assemblies.

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166 Genome assembly assessment and error rate quantification

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168 The quality of resulting assemblies was assessed using Quast 3.0 [32] according to the total assembly 169 length, number of contigs, N50, GC content and degree of replicon circularization. Assembly graphs 170 were visualized with Bandage [33]. The resulting contigs in each assembly were classified as 171 chromosomal or plasmid using machine learning algorithms implemented in mlplasmids [22].

173 **Read depth estimation**

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175 The read depth of each replicon was estimated by aligning the short Illumina and long Oxford

Nanopore reads to the completed genomes using Smalt v0.7.6 and BWA-MEM v0.7.17 (with the flag
-x ont2d for ONT reads), respectively. SAMtools v1.7 was used to process the SAM files to BAM
format, remove duplicates, and identify the coverage at each base of each assembly. The median value
for each replicon was noted and was normalized using the median chromosomal depth of the same
assembly.

182 Genome annotation

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The genomes were annotated using Prokka v1.13.3 [34]. *Bla*_{CTX-M} alleles and their contexts were
detected using the Multiple Antibiotic Resistance Annotator (MARA) [35] and by aligning the
assemblies against the Comprehensive Antibiotic Resistance Database (CARD v3.0). Information on
the detected AMR features and MGEs are retrieved from Galileo AMR

(https://galileoamr.arcbio.com/mara/feature/list). Plasmid identification and typing was carried out
 using PlasmidFinder v2.0 [36]. The plasmid-derived contigs from the assembled genomes were
 compared using BLAST v2.6.0 and their homology and annotation was visualised using EasyFig
 v2.2.2 [37].

193 Phylogenetic analysis

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195 To provide a phylogenetic context for these six isolates, the short Illumina reads of 63 from [8] and 56 196 from [38] published ST131 short read libraries were cleaned and trimmed using Fastp v0.12.3 [39], as were the six isolates' short read libraries from this study. These 125 libraries were de novo assembled 197 with Unicycler v4.6 using NCTC13441 as a reference and annotated using Prokka. The 126 genomes 198 199 were processed using Roary v3.11.2 [40] with a 95% BLAST v2.6.0 identity threshold to create a core 200 genome alignment containing 4,457 SNPs using MAFFT v7.310 [41] spanning 3,250,343 bases and 3,350 genes of the NCTC13441 chromosome (a length similar to [20]). This core genome was used to 201 202 construct a maximum likelihood phylogeny using RAxML v8.2.11 with the GTR model with gamma 203 rate heterogeneity [42]. Clade classification of the six isolates was based on published ST131 204 phylogenetic analysis [8] with associated classification and bla_{CTX-M} allele data from [8] and [38]. 205

Results

209 Oxford Nanopore long read quality control and filtering

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High molecular weight DNA from six *E. coli* ST131 isolates was sequenced using long Oxford

212 Nanopore reads and short Illumina reads to assemble their genomes allowing for plasmid

213 reconstruction and resolution of AMR genes, MGEs and associated rearrangements. The ONT

214 GridION X5 sequencing generated 8.9 Gbases in total across 1,406,087 reads (mean length of 6.3 Kb,

Table 1). The number of reads generated per hour, total yield of bases over time, read length

distribution, and read Q score distribution were examined (Supplementary Figure 2). Half of the bases

with Q > 7 were on reads of 18 Kb or longer (Supplementary Figure 3). These metrics indicated sufficient GridION data in terms of quantity and quality. Initial screening removed reads with Q < 7,

leaving 1,142,067 reads with 8.2 Gbases with a mean Q score of 10.2 and a mean length of 7.2 Kb 219 220 (Table 1) for analysis. This included 81 reads longer than 100 Kb, including one of 155,312 bases.

This corresponded to 257-fold theoretical coverage for six 5.3 Mb genomes.

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Parameter	All reads	Reads with 2247
Total bases	8,908,946	8,19 3,95 1
Total reads	1,406,087	114 2,9 <u>6</u> 7
Mean length (bp)	6,336	7,175
Median length (bp)	2,273	2,897
Mean Q score	9.1	2202
Median Q score	10.0	21015
Reads >100 Kb	85	2321

Table 1. Quality parameters indicated highquality read libraries for the six ST131 samples from GridION X5 sequence data. A total of 264.020 of low-quality reads (with O<7) totalling 715,024,800 bases were excluded.

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235 The initial number of reads per library ranged from 127,118 to 510,253 and these were filtered using a series of steps to ensure that the reads used for each of the six assemblies had high quality. Bases were 236 successfully called at an average of 97.9% of reads (Table 2). Identifying the consensus demultiplexed, 237 238 duplicate-free and adapter-free reads from Porechop v0.2.4 eliminated a further 2.9% of the basecalled 239 reads, yielding 120,123 to 487,482 reads per library (Table 2).

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Strain	Initial reads (fast5)	Basecalled (fastq)	Adapter-free (fastq)	Average length (bp)
VRES1160	358,829	351,636	345,033	7,037
VREC0693	208,478	204,904	194,413	8,982
VRES0739	163,349	160,693	155,900	9,171
VREC1013	510,253	497,646	487,482	6,657
VREC1073	313,627	304,218	298,658	7,256
VREC1428	127,118	124,539	120,123	9,301

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Table 2. Number of reads generated from GridION X5 sequencing data per library that passed filtering
 243 during basecalling with Albacore v2.0 and those that were adapter-free (using Porechop v0.2.4). The 244 latter totalling 1,601,609 reads were used for downstream analyses. 80,045 reads were excluded during 245 basecalling or adapter-trimming.

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Long read genome assembly illuminates highly diverse accessory genomes 247

248 249 All six genome assemblies produced chromosomes of 4.81-5.38 Mb with differing numbers of plasmids with lengths spanning 4-156 Kb (Supplementary Figure 4; Table 3). The numbers of contigs 250 produced by long read assemblies of three samples (VREC0693, VRES0739, VREC1073) 251 252 corresponded exactly to the chromosome and plasmids. The other three isolates had either one 253 (VRES1160 and VREC1013) or two (VREC1428) additional chromosomal contigs (Supplementary 254 Table 2).

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Contigs were classified as chromosomal or plasmid-derived using mlplasmids given a probability 256 257 threshold of 60% [22], with further screening for plasmid-related gene content using MARA, CARD and PlasmidFinder (Supplementary Table 2). The largest plasmid was a 156.3 Kb IncFIA one in 258 259 VREC1073, its sole plasmid. VREC1428 and VRES1160 had 92.8 and 61.9 Kb IncFIA plasmids, respectively, along with three small Col plasmids each (Table 3). VREC0693 had a 132.0 Kb IncFIB 260 plasmid and an 88.8 Kb IncB plasmid - IncB plasmids have the same Rep domains as IncFII plasmids 261 262 [42]. VREC3013 had one 89.9 Kb IncFII plasmid. VRES0739 alone had no large plasmid, which was 263 verified with the short read data.

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265 By mapping the long reads to the optimal assemblies, the read coverage of each chromosome and 266 plasmid was estimated (Supplementary Table 2). Each chromosome had between 126- and 310-fold

median coverage, and the median coverage levels of large plasmids ranged from 85- to 282-fold, 267 268 except for VREC1013's IncFII plasmid that had 1,015-fold coverage and a normalized depth of 3.3fold. The normalised depth of plasmids compared to chromosomes suggested some cells in VREC1428 269 270 and VREC1073 may have lost their IncFIA plasmid, and the same for VREC0693 and its IncFIB 271 plasmid. However, the IncFIA plasmid in VRES1160 and the IncB plasmid in VREC0693 had higher than expected copy numbers (by 9% after normalisation), potentially indicating stable plasmid 272 retention.

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	Genome	Number of contigs				Number	
Strain	length (bp)	Assembled	Minimum possible	N50	Chromosome size (Mb)	of plasmids	Plasmid sizes (Kb)
VRES1160	5,326,801	6	5	5,126,679	5.23	4	62, 16, 5, 4
VREC0693	5,260,741	3	3	5,039,909	5.04	2	132, 89
VRES0739	4,806,912	3	3	4,797,749	4.81	2	5, 4
VREC1013	5,223,433	3	2	3,699,451	5.14	1	90
VREC1073	5,539,158	2	2	5,286,804	5.38	1	156
VREC1428	5,236,419	7	5	4,924,536	5.13	4	92, 5, 5, 4

Table 3. Total size of assemblies, chromosomes and plasmids found in each strain based on their 276 optimal whole genome assemblies using the GridION X5 long reads. Each assembly had seven or less 277 contigs, and in three cases no fewer contigs were possible, consistent with full genome assembly (for 278 VREC0693, VRES0739 and VREC1073). The optimal assembly with Unicycler used long reads alone 279

(in bold mode), with exception of VREC1013, where a hybrid combining short Illumina reads with 280

281 long Oxford Nanopore reads was best, with minor manual screening (Supplementary Results).

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284 Across five assemblies in the Unicycler normal mode, the median indel error rates for short reads and hybrid assemblies were similar (0.21 and 0.28 per 100 Kb, respectively), but was much higher for long 285 286 read assemblies (265.0 per 100 Kb, Supplementary Table 3). Likewise, the median mismatch error 287 rates for short reads and hybrid assemblies were comparable (4.25 and 2.28 per 100 Kb, respectively), 288 but was much higher for long read assemblies (332.8 per 100 Kb, Supplementary Table 3). These rates 289 excluded VREC1073, for which some Quast metrics were zero values. 290

The dynamic locations and genomic contexts of *bla*_{CTX-M} genes in long read assemblies 291 292

293 The optimised assemblies provided an improved view of the genomic context of each *bla*_{CTX-M} allele, 294 whose effectiveness as a marker for ST131 clade classification and origin [8] we explored here. The 295 deeper resolution of genome architecture revealed surprising differences in *bla*_{CTX-M} gene context 296 (Figure 1; Supplementary Table 2), including the discovery of chromosomal *bla*_{CTX-M} genes in 297 VREC0693 (three copies of *bla*_{CTX-M-15}) and VREC1073 (one copy of *bla*_{CTX-M-14}). All *bla*_{CTX-M} genes 298 were complete (876 bp) with adjacent ISEcp1 (1,658 bp with flanking IRs of 14-16 bp) and Tn2 (5.8 299 Kb) elements: ISEcp1 and Tn2 can transpose bla_{CTX-M} and other ESBL genes [44-45]. The VRES0739 300 genome did not contain any region homologous to *bla*_{CTX-M}, most likely because it had lost an IncF 301 plasmid, unlike the other isolates.

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303 VRES1160, VREC0693 and VREC1013 all had *bla*_{CTX-M-15} genes linked to isoforms of ISEcp1, IS26 and Tn2, implicating them in driving transposition of the TU (Supplementary Figure 5). Each was 304 similar to the ST131 clade C2 *ISEcp1-bla_{CTX-M-15}*-orf477 Δ TU [8,46] but with distinct structural 305 differences. VRES1160's single *bla*_{CTX-M-15} gene was at 2,296 bp on its IncFIA plasmid and was 306 flanked by ISEcp1 to its 5' and Tn2 followed by IS26 at its 3' end, with another Tn2 5' of ISEcp1. 307 VREC0693's three chromosomal bla_{CTX-M-15} genes were not tandem repeats (chromosomal locations 308 309 2,781,074, 3,696,068 and 3,970,927), but each of these TUs were identical: all had ISEcp1 at the 5' 310 ends and truncated Tn2s at the 3' ends. VREC1013's sole *bla*_{CTX-M-15} gene was located at 13,226 bp on

its IncFII plasmid and was flanked by a truncated ISEcp1 at its 5' end and Tn2 at its 3' end, with IS26 311 312 copies 5' and 3' of these segments.

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314 VREC1428's single *bla*_{CTX-M-27} gene was on its IncFIA plasmid at position 6,018, and VREC1073's 315 single chromosomal *bla*_{CTX-M-14} gene was at contig position 19,746 (Supplementary Figure 5). Both their *bla*_{CTX-M} genes were flanked by a truncated IS*Ecp1* at the 5' ends and a shortened IS*903B* at the 316 3' ends suggesting that ISEcp1 and IS903B may have facilitated the transposition of the TU from the 317 plasmid. Similar bla_{CTX-M} gene transposition events have been observed in ST131 clade C1 [8].

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319 320 Alignment of the plasmid-derived contigs of VRES1160 (IncFIA) to VREC1013 (IncFIB) showed that 321 the *bla*_{CTX-M-15}-positive plasmids were much more similar (>83% identity) relative to VREC1428's *bla*_{CTX-M-27}-positive IncFIA plasmid, which was more distinct (Figure 2). This suggested that the 322

323 plasmid homology corresponded well with VRES1160 and VREC1013 blacTX-M gene and subclade

324 classification, but not IncF replicon type. 325

326 Phylogenetic context of analysed isolates

327 Comparison of these six samples with 119 published ST131 [8,38] as short read assemblies scaffolded 328 using reference genome NCTC13441 showed that all clustered in ST131 clade C (Supplementary 329 330 Figure 6). There was sufficient resolution across 4,457 core genome SNPs to confidently assign them 331 to subclades C1 (n=1) or C2 (n=5) (Figure 3). VRES1160, VREC0693, VREC1013, VRES0739 and 332 VREC1073 clustered with C2, whereas the *bla*_{CTX-M-27}-positive VREC1428 was in C1. VRES1160, VREC0693 and VREC1013 all had IncF plasmids (IncFIA, IncFIB, IncFII) and *bla*_{CTX-M-15} genes, 333 334 consistent with C2 are typically *bla*_{CTX-M-15}-positive, which was observed for 77% of C2 isolates here 335 (48 out of 62). However, VREC1073 was in C2 but had an IncFIA plasmid with a *bla*_{CTX-M-14} gene, 336 contradicting this pattern and was the sole *bla*_{CTX-M-14}-positive C2 isolate found here. The core 337 genomes of VRES0739 and VREC0693 were identical, implying that VRES0739 has very recently 338 lost its (*bla*_{CTX-M}-positive IncF) plasmid. The sole isolate clustering with C1 was VREC1428, which 339 had an IncFIA plasmid with a *bla*_{CTX-M-27} gene, and so may belong to the emerging subclade C1-M27 340 as evidenced by the presence of prophage-like regions like M27PP1/2 [38]. 341

Discussion

344 345 Our study resolved the plasmid architecture of several recent E. coli ST131 isolates, allowing investigation of AMR gene location, copy number and potential transposon-driven rearrangements. 346 347 This advance was facilitated by the careful DNA handling during extraction to produce large volumes 348 of high molecular weight DNA that was pure and free from contamination, which was avoided by 349 performing separate extraction steps to obtain small plasmids [47] overcoming a limitation for 350 MinION sequencing [21]. 351

The long read genome assemblies illuminated significant variation in plasmids, MGEs and *bla*_{CTX-M} 352 353 gene composition that was not captured by short reads. ST131 is a globally pandemic E. coli clonal 354 group [15] with diverse sources of transmission [25]. Phylogenetic comparison with published 355 genomes [8,42] showed that five out of six isolates were from subclade C2 with one from C1. The 356 emergence of clade C has been associated with IncF plasmids, and clade C2 with ISEcp1 and Tn2 357 elements flanking *bla*_{CTX-M-15} genes [47-48]. Our long read assemblies showed the excision of the entire TU from the IncFIB plasmid and chromosomal integration at three distinct locations for 358 359 VREC0693, and similarly chromosomal translocation of the *bla*_{CTX-M-14} gene from an IncFIA plasmid 360 for VREC1073, mediated by ISEcp1 and IS903B based on previous work [8]. These transposition 361 events were likely driven by recombination at adjacent transposable elements. This highlights the 362 value of long read sequencing to resolve the location of *bla*_{CTX-M} genes and that chromosomal 363 translocations are not rare in ST131.

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A high resolution of the AMR gene structure, context and copy number is highly predictive of AMR 365 phenotypes [45], and could lead to new insights into AMR mechanisms. However, the high indel and 366

- 367 mismatch errors in long Oxford Nanopore reads [31,47,51-52] limits power to identify AMR isoforms
- that could permit genome-based antimicrobial susceptibility testing [43,53-54]. Here, the five ONT
- assemblies together had an average of 447-fold higher indel and 48-fold higher mismatch error rates
- than those for the corresponding Illumina reads, similar to previous work with MinION reads [23].
- 371 Consequently, short reads and assembly polishing methods remain important for SNP identification
- and error detection until long read error rates can be reduced [55].
- 373
- Our findings illustrate the diversity of AMR gene context even within recently emerged clones such as
 ExPEC ST131. Further studies are needed with larger sample sizes to identify the rates and
- 376 mechanisms of these dynamic changes.
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- 378

379 Data Summary

- 1. Illumina reads accession numbers: ERR2138475, ERR2138200, ERR2138591, ERR1878196,
 ERR2137889 and ERR1878359 in the European Nucleotide Archive (ENA) under BioProjects
 PRJEB21499 and PRJEB19918.
- 2. ONT reads ENA accession numbers: www.ebi.ac.uk/ena/data/view/PRJXXXXX, Figshare
 https://doi.org/10.6084/m9.figshare.7554293.v1
- 385 3. Unicycler assemblies, Figshare https://doi.org/10.6084/m9.figshare.7560458.v2

386387 Ethical approval

The study protocol was approved by the National Research Ethics Service (ref:14/EE/1123), and the Cambridge University Hospitals NHS Foundation Trust Research and Development Department (ref: A093285).

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- 397

398 Conflicts of interest

399 JP is a consultant to Next Gen Diagnostics Llc.400

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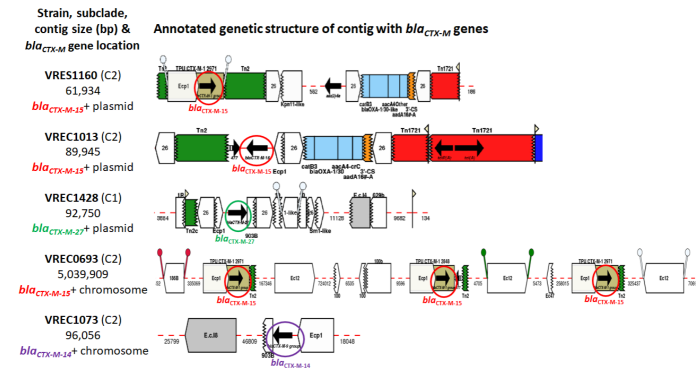
410 Figure legends

411

Figure 1. Two of the ST131's bla_{CTX-M} genes were on chromosomal contigs (VREC0693 and VREC1073). VRES1160 and VREC1013 had IncFIA and IncFII plasmids, respectively, both of which had $bla_{CTX-M-15}$ genes. VREC1428 had an IncFIA plasmid with $bla_{CTX-M-14}$ gene. VRES0739 is not shown because it was bla_{CTX-M} -negative and had no large plasmid. The contigs were classified as chromosomal or plasmid-derived by mlplasmids so that the bla_{CTX-M} genes and their genetic flanking context could be examined. Annotation was derived from GalileoTM AMR based on the Multiple Antibiotic Resistance Annotator (MARA) and database. The bla_{CTX-M} variants are labelled and

419 encircled in red ($bla_{CTX-M-15}$), purple ($bla_{CTX-M-14}$) or green ($bla_{CTX-M-27}$).

420



422 Figure 2. Pairwise comparisons of the three *bla*_{CTX-M}-positive plasmid-associated contigs showed high 423 homology for the two from subclade C2 (VREC1013 and VRES1160) relative to one from C1 424 (VREC1428, top). BLAST alignment homology was visualised with EasyFig v2.2.2 such that the 425 middle blocks connecting regions of the contigs represent nucleotide homology: blue for homologous 426 regions in the same direction, and yellow for inversions. Gaps or white spaces denote unique loci or regions present in a contig but not in the other. Gene models are in green with the direction of 427 transcription shown by arrows. Genes of interest are labelled above each arrow. The *bla*_{CTX-M-27} grey 428 (top) is in mauve and the two *bla*_{CTX-M-15} genes (middle, bottom) are in red. The table below shows the 429 430 contig size, plasmid type and the number of genes per strain. Products of the annotated genes are in 431 Supplementary Table 4.

432

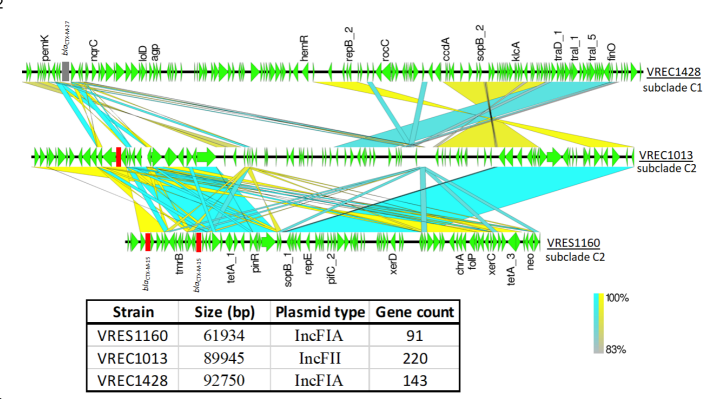
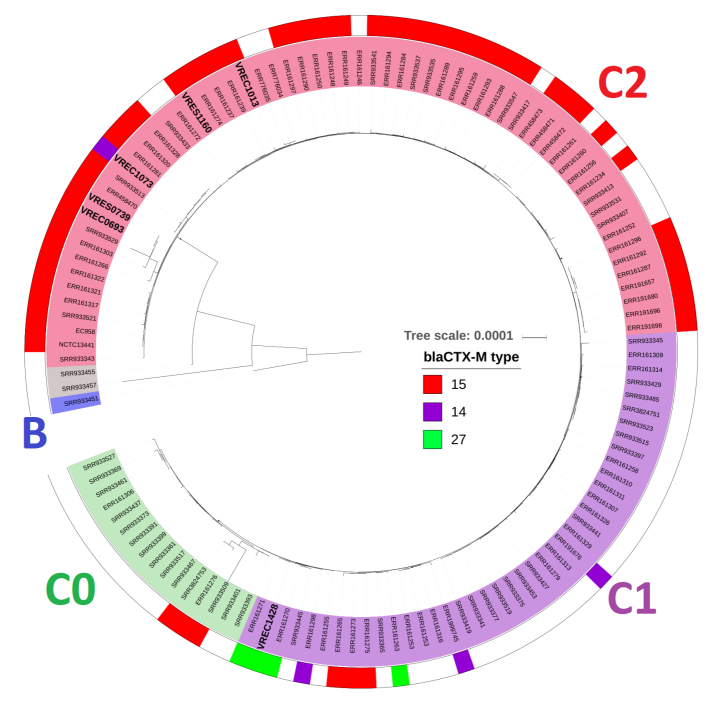


Figure 3. The phylogenetic context of the six ST131 genomes (names are in large bold font) showed 434 435 that all except VREC1428 were in ST131 subclade C2 (red inner ring: VRES1160, VREC1073, VRES0739, VREC0693 and VREC1013). VREC1428 clustered in subclade C1 (purple inner ring). No 436 new isolate clustered in C0 (green inner ring), B (blue inner ring) or an intermediate cluster (grey inner 437 438 ring). Clade classification was based on phylogenetic analysis by [8] by including the reference 439 NCTC13441, n=63 isolates from [8] and n=56 from [38] with associated classification and *bla*_{CTX-M} allele data. VREC1073, and VREC0693 had chromosomal *bla*_{CTX-M} genes. The outer ring shows 440 441 *bla*_{CTX-M-15} (red), *bla*_{CTX-M-14} (purple) and *bla*_{CTX-M-27} alleles (green). The phylogeny was built with RAxML v8.2.11 using 4,457 SNPs from a core genome alignment generated with Roary v3.11.2 and 442 was visualised with iTOL v4.3. Branch support was performed by 100 bootstrap replicates, and the 443 444 scale bar indicates the number of substitutions per site. This mid-pointed rooted phylogeny includes reference genome isolates EC958 and NCTC13441 (both in C2). 445 446



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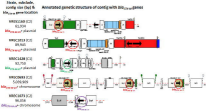
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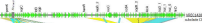
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times	Price P	100E	Que con	A S C S	subclade C2
2 6	2	8 a.		2	

Strain	Size (bp)	Plasmid type	Gene count	
VRES1160	61934	IncFIA	91	
VREC1013	89945	IncFII	220	
VREC1428	92750	Incl ⁻ IA	143	



