

1 **Concurrence of Porin Loss and Modular Amplification of β -Lactamase Encoding Genes**
2 **Drives Carbapenem Resistance in a Cohort of Recurrent *Enterobacterales* Bacteremia**

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29

30 **ABSTRACT**

31 **Background**

32 Carbapenem resistant *Enterobacteriales* (CRE) remain urgent antimicrobial resistance threats.
33 Approximately half of CRE clinical isolates lack carbapenem hydrolyzing enzymes and develop
34 carbapenem resistance through alternative mechanisms. The purpose of this study was to
35 elucidate the development of carbapenem resistance mechanisms from clonal, recurrent
36 extended-spectrum β -lactamase positive *Enterobacteriales* (ESBL-E) bacteremia isolates in a
37 vulnerable patient population.

38 **Methods**

39 This study investigated a historical, retrospective cohort of ESBL-E bacteremia cases in the
40 University of Texas MD Anderson Cancer Center (MDACC) from January 2015 to July 2016.
41 Phylogenetic and comparative genomic analyses were performed to identify clonal, recurrent
42 ESBL-E isolates developing carbapenem resistance. Oxford Nanopore Technology (ONT) long-
43 read and Illumina short-read sequencing data were used to generate consensus assemblies and to
44 identify signatures of mobile genetic element mediated amplification and transposition of
45 antimicrobial resistance genes. Serial passaging experiments were performed on a set of clinical
46 ST131 ESBL-E isolates to recapitulate *in vivo* observations. qPCR and qRT-PCR were used to
47 determine respective copy number and transcript levels of β -lactamase genes.

48 **Results**

49 116 ESBL-E bacteremia cases were identified, 16 of which had documented recurrent infections.
50 Four serial, recurrent isolates displayed a carbapenem resistant phenotype, three without the
51 acquisition of a known carbapenemase. These three isolates had non-carbapenemase-producing
52 CRE (non-CP-CRE) mechanisms driven by IS26- and *ISEcpI*-mediated amplification of

53 respective translocatable units (TU) and transposition units (TPU) harboring both *bla*_{OXA-1} and
54 *bla*_{CTX-M} variants with concomitant outer membrane porin disruption. The TU and TPU
55 structures inserted into the open reading frames of outer membrane porin genes in a subset of
56 non-CP-CRE isolates. Serial passage of an index ST131 ESBL-E isolate under selective
57 carbapenem exposure resulted in chromosomal amplification of modular, TUs harboring β -
58 lactamase genes with concomitant porin inactivation, recapitulating the *in vivo* carbapenem
59 resistance progression. Long-read sequencing of two additional MDACC bacteremia strains
60 identified similar non-CP-CRE mechanisms observed in the serial isolates.

61 **Conclusions**

62 Non-CP-CRE *de novo* mechanisms were the primary driver of CRE development in recurrent
63 bacteremia cases within this vulnerable patient population. The incorporation of long-read ONT
64 data into AMR surveillance platforms is critical to identify high-risk CRE isolates that are
65 difficult to identify with low-resolution phenotypic and molecular characterization methods.

66

67 **Keywords:**

68 Non-carbapenemase-producing carbapenem resistant *Enterobacterales*, Oxford Nanopore
69 Technologies MinION Sequencing, Mobile Genetic Element Amplifications, Mobile Genetic
70 Element Transpositions

71

72 **BACKGROUND**

73 Antimicrobial resistance (AMR) is an emerging global health priority and carbapenem resistant
74 *Enterobacterales* (CRE) are among the most serious AMR threats (1). Carbapenem resistance
75 can develop due to the acquisition of enzymes that hydrolyze carbapenems, known as
76 carbapenemases, as well as through changes in outer membrane permeability and/or drug efflux
77 activity, which decrease intracellular carbapenem concentrations (2). While most CRE research
78 has focused on characterizing carbapenemases (2), recent clinical and molecular epidemiology
79 studies indicate approximately 50% of CRE isolates are not carbapenemase carriers, suggesting a
80 substantial proportion of CRE isolates develop carbapenem resistance through alternative
81 mechanisms (3, 4). Molecular characterization of these alternative mechanisms indicate non
82 carbapenemase-producing carbapenem resistant *Enterobacterales* (non-CP-CRE) generally carry
83 extended-spectrum β -lactamases (ESBL) or AmpC-like enzymes with concomitant mutations
84 that alter porin function (2). Further studies reveal that *Enterobacterales* strains carrying outer
85 membrane porin mutations, but lacking ESBL or AmpC-like enzymes, develop *de novo*
86 carbapenem resistance at lower rates during serial passage under increasing carbapenem
87 concentrations (5, 6). Thus, the presence of ESBL or cephalosporinase genes may be a
88 component cause in non-CP-CRE development. Additionally, van Boxtel et al. demonstrated that
89 serial passaging of an ESBL or AmpC-producing isolate in the presence of a carbapenem can
90 result in amplification of plasmid-borne β -lactamase genes (7). Increased expression of the
91 narrow-spectrum TEM β -lactamases has similarly been reported to result in cefepime (8) and
92 piperacillin-tazobactam resistance (9-11), indicating that β -lactamase gene dosage is a factor in
93 increasing resistance to multiple β -lactam chemotherapies. These findings demonstrate
94 expression level and copy number of β -lactamases without known carbapenemase activity have

95 important effects on carbapenem susceptibility in porin deficient backgrounds (6, 7).
96 Nevertheless, there remains a gap in knowledge regarding how these non-CP-CRE mechanisms
97 evolve *in vivo*, as few existing studies have investigated serially collected isolates from large,
98 patient cohorts (12-16).

99

100 This gap in knowledge is particularly relevant in carbapenem resistant *Escherichia coli*, where
101 the majority of these isolates are non-CP-CRE (3, 4). Additionally, *Klebsiella pneumoniae*
102 isolates with non-CP-CRE phenotypes have also been identified, albeit less frequently (3, 4).
103 Both non-CP-CR *E. coli* and *K. pneumoniae* can cause severe disease, including bacteremia,
104 which have high mortality rates (17). The detection and treatment of infections with these non-
105 CP-CRE isolates remain challenging relative to carbapenemase producing CRE as phenotypic
106 tests may incorrectly identify these isolates as carbapenem susceptible and definitive therapy
107 options may not be readily evident. This highlights the importance to further characterize non-
108 CP-CRE mechanism development, which may provide insights into new surveillance and/or
109 treatment options. Since there remains a lack of studies fully characterizing non-CP-CRE
110 development, we performed a systematic analysis of non-CP-CRE mechanisms using a large
111 cohort of patients with whole genome sequencing (WGS).

112

113 We utilized a cohort of patients with ESBL-E bacteremia from the University of Texas MD
114 Anderson Cancer Center (MDACC). Importantly, analysis of a previous MDACC cohort of non-
115 CP-CRE bacteremia isolates indicated these strains had increased short-read mapping of β -
116 lactamase encoding genes suggestive of β -lactamase encoding gene amplification (18).
117 Nevertheless, the amplification structures and genomic context of these gene amplifications

118 could not be discerned with this short-read sequencing analysis. This is due to inherent
119 limitations with short-read sequencing assemblers, which have difficulties resolving the
120 complex, repetitive mobile genetic elements (MGEs) that carry resistance genes (19). Therefore,
121 we present an analysis utilizing Oxford Nanopore Technologies (ONT) MinION sequencing, a
122 long-read sequencing platform, which overcomes these limitations to elucidate these
123 amplification mechanisms. This study aims to track the *de novo* development of carbapenem
124 resistance mechanisms during recurrent *Enterobacteriales* bacteremia infection by specifically
125 focusing on ESBL positive *E. coli* and *K. pneumoniae* strains and characterizing the non-CP-
126 CRE associated β -lactamase amplification mechanisms within our cohort.

127

128 **METHODS**

129 **Study design and clinical data abstraction**

130 A retrospective review of patients with ESBL-E bacteremia hospitalized from January 2015 to
131 July 2016 was conducted at MDACC in Houston, Texas. All patients with one or more episodes
132 of ESBL-E bacteremia who were 18 years of age or greater were eligible for inclusion. Blood
133 culture isolates are routinely saved at MDACC and stored at -80°C. Clinical and demographic
134 characteristics were manually extracted from electronic medical records and recorded using
135 REDCap software (Vanderbilt University, Nashville, TN) (20).

136

137 **Isolate identification and antimicrobial susceptibility testing**

138 Antibiotic susceptibility testing was performed per routine clinical laboratory practice using an
139 automated system (Vitek2, bioMérieux, Marcy L'Étoile, France) with additional testing
140 performed as needed using individual antibiotic gradient strips (Etest, bioMérieux). ESBL

141 production was assessed per routine laboratory practice on *E. coli*, *K. pneumonia*, and *K. oxytoca*
142 isolates that were resistant to one or more oxyimino-cephalosporins (e.g. cefotaxime,
143 ceftriaxone, or ceftazidime) using either the ESBL Etest (bioMérieux) or the Rapid ESBL Screen
144 kit (ROSCO, Taastrup, Denmark). Carbapenemase production was evaluated in the clinical lab
145 on any *Enterobacterales* isolate resistant to one more of the carbapenems using the Neo-Rapid
146 CARB kit (ROSCO) according to manufacturer's instructions. Carbapenem resistance (CR) was
147 defined as resistance to either ertapenem or meropenem using CLSI criteria (21). Recurrent
148 *Enterobacterales* bacteremia was defined as identification of the same species in blood culture at
149 any point during the follow-up period following at least one negative blood culture and
150 completion of an antibiotic treatment regimen.

151

152 **Illumina short-read sequencing**

153 All available isolates from recurrent bacteremia patients, initially underwent whole genome
154 sequencing (WGS) via Illumina HiSeq as described previously (18). The paired-end short-reads
155 were assessed using the FastQC toolkit (Babraham Institute), and adaptors as well as low-quality
156 reads were trimmed using Trimmomatic v0.33 (22). Genome assembly was performed using
157 SPAdes v3.9.1 (23). Depth of short read mapping to individual genes of interest was quantitated
158 relative to the average read mapping depth for the pubMLST housekeeping gene schema for *E.*
159 *coli* ST10 and ST131 respectively.

160

161 **Oxford Nanopore Technologies (ONT) long-read sequencing**

162 Serial isolates that developed non-CP-CRE, isolates used in the serial passaging experiments,
163 and non-CP-CRE isolates from a previous study (18) underwent Oxford Nanopore Technologies

164 (ONT) long-read sequencing. Library preps were completed using the SQK-RBK004 rapid
165 barcoding kit with ~400 ng of input DNA and run on ONT MinION R9.4.1 flow cells using the
166 ONT GridION X5 (Oxford, UK) per manufacturer’s instructions. ONT fast5 data was generated
167 using ONT MinKNOW software (v3.0.13) with subsequent base-calling using Guppy v3.2.2
168 software (Oxford, UK). qcat-v1.1.0 was used for read demultiplexing, read length filtering
169 (>1000 bp), and barcode removal (nanoporetech qcat GitHub:
170 <https://github.com/nanoporetech/qcat>). A custom python script was used for the generation of
171 polished, consensus assemblies (Shropshire, W flye_hybrid_assembly_pipeline GitHub:
172 https://github.com/wshropshire/flye_hybrid_assembly_pipeline
173). Briefly, the Flye-v2.5 (24) assembler was used for *de novo* assembly, and contigs were
174 circularized using berokka-v0.2 (Seemann, T berokka GitHub:
175 <https://github.com/tseemann/berokka>). The circlator v1.5.5 (25) ‘clean’ command was then used
176 to remove duplicate contigs sharing at least 90% identity. The remaining contigs were polished
177 with Racon-v.1.4.5 (26) using the Oxford Nanopore long reads and then re-oriented with
178 Circlator (25) “fixstart” to standardize the chromosome to the *dnaA* gene. A second long-read
179 polish was then performed with Racon-v1.4.5 and these polished, corrected contigs were used as
180 input for Medaka v0.8.1 long-read polishing (nanoporetech medaka GitHub:
181 <https://github.com/nanoporetech/medaka>) followed by multiple rounds of Illumina short-read
182 polishing using Racon-v1.4.5 (26). ONT assembly metrics are provided in **Table S1**.

183

184 **Phylogenetics, genetic variant calling, and clonality analysis**

185 Phylogenetic analysis, *in silico* MLST, and subsequent variant-calling of serial isolates compared
186 against the index strain ONT consensus assembly were used to determine clonality. Initially, the

187 pan-genome pipeline tool Roary-v3.12.0 (27) was used to perform a core genome alignment of
188 the short-read SPAdes assemblies using the probabilistic alignment program PRANK (27, 28). A
189 core genome pairwise SNP distance matrix was generated with these data using a custom Python
190 script (Narechania, A
191 GitHub:https://github.com/narechan/amnh/blob/master/bin/snp_matrixBuilder.pl), which was
192 subsequently used to build a maximum likelihood (ML) phylogenetic tree using RAxML-v8.2.12
193 (29). *In silico* multi-locus sequencing typing (MLST) was performed on the short-read SPAdes
194 assemblies using mlst-v2.15.1 (Seemann, T mlst Github: <https://github.com/tseemann/mlst>; (30).
195 Recurrent isolates that became non-CP-CRE were checked for clonality using the variant calling
196 pipeline tool Snippy-v4.3.6 (Seemann, T snippy GitHub: <https://github.com/tseemann/snippy>)
197 with INDELS removed.

198

199 **Genome annotation, AMR gene, and MGE identification**

200 Gene calling and functional annotation was performed using Prokka-v1.14.0 (31). Annotated
201 consensus genomes, as well as individual contigs, were parsed with ABRicate (Seemann, T
202 ABRicate Github: <https://github.com/tseemann/abricate>) using the Comprehensive Antibiotic
203 Resistance Database (CARD) (32) and PlasmidFinder (33) to search for AMR determinants and
204 plasmid signatures respectively. Additionally, annotated Prokka gbk files were used to
205 characterize ORFs of interests as well as confirm the results found with ABRicate. CARD,
206 PlasmidFinder, ISFinder (34) and BLAST webtools were utilized during manual inspection of
207 the assemblies to ensure correct context of annotated features, identify inverted repeat regions
208 and target site duplications of insertion sequences, and confirm likely AMR gene mutations if
209 present.

210

211 In cases where assemblies were not completely resolved due to putative large repeat regions that
212 could not be fully captured on ONT long reads, we used a newly developed tool SVAnts-v0.1
213 (35) (Hanson, Blake SVAnts Github: <https://github.com/EpiBlake/SVAnts>) to investigate subsets
214 of ONT long reads. This tool enabled us to find individual ONT long reads containing MGEs of
215 interest and align the DNA bordering these MGEs to our index assemblies to identify the regions
216 in which the MGE was inserted within each bacterial genome.

217

218 **B-lactamase encoding gene and gene transcript level analysis**

219 Quantitative PCR (qPCR) and quantitative real-time PCR (qRT-PCR) was used to assess both
220 DNA copy number and RNA transcript levels respectively. Strains were grown in triplicate on
221 two separate days (six biologic replicates) to mid-exponential phase ($OD_{600} \sim 0.5$) in Luria-
222 Bertani (LB) broth (ThermoFisher) at 37° C shaking at 220 rpm. DNA isolation was performed
223 using the DNEasy kit (Qiagen) and qPCR was performed using TaqMan reagents on the StepOne
224 Plus Real Time PCR platform (Applied Biosystems). The DNA levels of *bla_{OXA-1}* and *bla_{CTX-M}*
225 were determined relative to the *rpsL* control gene using the ΔC_t method (6).

226

227 For RNA transcript level analysis, cells were mixed 1:2 with RNAProtect (Qiagen) and
228 harvested via centrifugation. RNA was isolated from cell pellets using the RNEasy kit (Qiagen)
229 and converted to cDNA using the High Capacity cDNA Reverse Transcription Kit (Applied
230 Biosystems). Relative transcript levels of the β -lactamase encoding genes (*bla_{OXA-1}* and *bla_{CTX-M}*)
231 were assayed using TaqMan reagents on the StepOne Plus Real Time PCR machine (Applied
232 Biosystems). The transcript level of *bla_{OXA-1}* and *bla_{CTX-M}* were determined relative to the

233 endogenous control gene *rpsL* (36) using the Δ Ct method. qPCR and qRT-PCR primers and
234 probes are provided in **Table S2**.

235

236 **Serial passaging experiments on clinical *E. coli* ST131 index bacteremia isolate**

237 p4A passaging experiments with antibiotic selection were performed in LB broth under shaking

238 conditions at 37°C. A single colony from the index strain of patient 4 (p4A) was grown

239 overnight, then diluted 1:100 into fresh LB containing ertapenem (ETP) (Sigma-Aldrich) at 0.5

240 MIC ETP. The process was repeated with increasing concentrations of ETP until growth was

241 observed at an ETP concentration of 32 μ g/mL. Passaging experiments were performed twice

242 with growth occurring at an ETP MIC of \geq 32 μ g/mL within 3 passages (i.e. 72 hours) on each

243 occasion. For the first passaging experiment, cells were collected on four consecutive days

244 (strains p4A_1 to p4A_4) while continuing to passage at an ETP concentration of 32 μ g/mL.

245 This was done in order to determine whether progressive β -lactamase encoding gene

246 amplification would be observed. On the second set of passaging experiments, cells were

247 collected on the first day that the ETP MIC reached \geq 32 μ g/mL, serially diluted onto agar plates,

248 and two different isolates were studied to assess for heterogeneity (strains p4A_H1 and

249 p4A_H2). Additional serial passaging experiments were conducted using p4C and p4D isolates

250 in the absence of antibiotic selection to determine the stability of the amplified units with

251 protocol adapted from previously published methods (37).

252

253 **β -lactamase cloning and expression analysis**

254 The open reading frames of β -lactamases were amplified from genomic DNA of the *K.*

255 *pneumoniae* strain MB101 using Q5 polymerase and the primers listed in **Table S2**. Cloned

256 ORFs were inserted into the arabinose inducible vector pBAD33 by Gibson assembly (38) of
257 purified products. pBAD33 $bla_{CTX-M-15}$ and pBAD33 bla_{OXA-1} were transformed into DH5 α *E. coli*
258 and were maintained with 50 μ g/mL chloramphenicol in cation adjusted Mueller Hinton (MHII)
259 media. MIC assays were performed with ceftriaxone (Sandoz GmbH), or piperacillin-tazobactam
260 (Fresenius Kabi USA) as follows. DH5 α strains carrying pBAD33 $bla_{CTX-M-15}$, pBAD33 bla_{OXA-1} ,
261 or control vector were grown for 18 hours at 220 rpm at 37°C in MHII with 50 μ g/mL
262 chloramphenicol. These cultures were diluted to approximately 5×10^5 CFU/mL in MHII with
263 0.2% L-Arabinose or vehicle control, without chloramphenicol. Each strain was exposed to serial
264 dilutions of the above drugs in microtiter plates sealed with gas-permeable membranes (Midsco).
265 Microtiter plates were incubated for 18 hours at 220 rpm at 37°C, followed by OD₆₀₀
266 measurement in a Biotek Synergy HT plate reader. The lowest tested antibiotic concentration
267 yielding OD₆₀₀ measurement of 0.06 or less in at least two of three replicate wells was
268 considered to be the MIC.

269

270 **Statistical analyses**

271 All statistical analyses were performed using Stata v13.1 (StataCorp LP, College Station, TX).
272 Bivariate comparisons between patients with recurrent bacteremia and patients with a single
273 bacteremia episode were made with the Wilcoxon Rank-sum test and Fisher's exact test as
274 appropriate based on covariate distributions. Comparisons of DNA and RNA levels among
275 strains was performed using the Kruskal-Wallis test when more than two strains were analyzed
276 or the Wilcoxon Rank-sum test when two strains were analyzed. MIC comparisons were
277 performed using ANOVA with Dunnett's test of multiple comparisons. Statistical significance
278 was assigned as a two-sided *P* value < 0.05.

279

280 **Data availability**

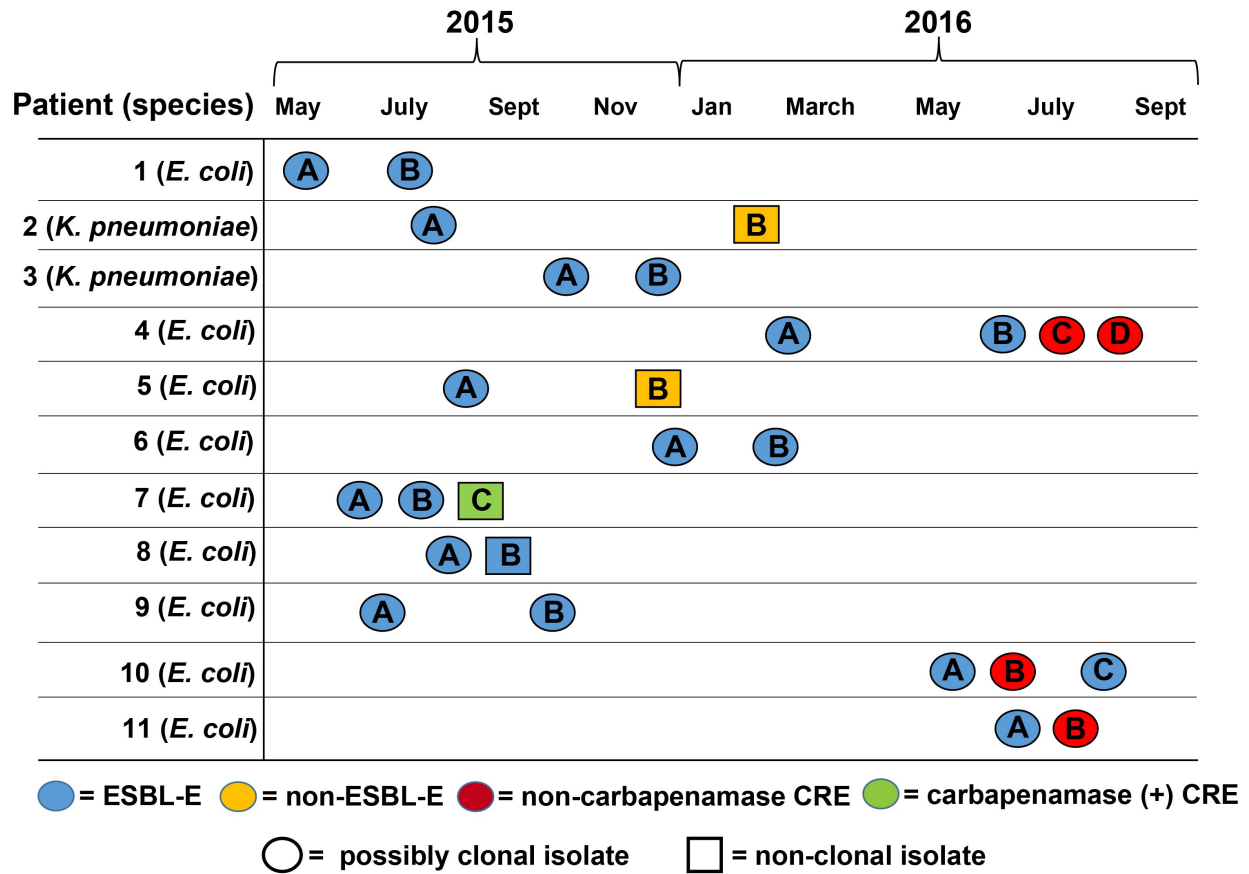
281 The index isolate assemblies from recurrent bacteremia patients that developed non-CP-CRE as
282 well as the long-read and short-read data for all isolates have been deposited in the National
283 Center for Biotechnology (NCBI) BioProject database PRJNA603908. ONT-sequencing data of
284 non-CP-CRE isolates from a previous study (18) were deposited in NCBI BioProject database
285 PRJNA388450. All other data analyzed during this study are available in the supplemental
286 materials and/or available upon request from the corresponding author.

287

288 **RESULTS**

289 **116 patients with ESBL-E bacteremia were identified from the University of Texas MD** 290 **Anderson Cancer Center (MDACC) from January 2015 to July 2016**

291 Clinical and demographic features are presented on **Table S3**. *E. coli* was the most common
292 organism isolated (100/116; 86.2%), followed by *K. pneumoniae* (14/116; 12.1%), and *K.*
293 *oxytoca* (2/116; 1.7%). Carbapenems were used as primary treatment in 92% of index cases.
294 Recurrent bacteremia was identified in 16/116 (13.8%) patients and primarily occurred either in
295 patients with leukemia or recipients of hematopoietic stem cell transplants (14/16 cases, **Table**
296 **S3**). The majority (14/16) of recurrent bacteremia cases had *E. coli* isolated, with the remaining
297 two cases being due to *K. pneumoniae* infections. Carbapenem-resistant isolates were present in
298 4/16 (25%) recurrent bacteremia patients (**Fig. 1**). All four recurrent isolates that developed
299 carbapenem resistance were *E. coli* isolates from leukemia patients. The full set of serial isolates
300 was available for 11/16 (68.8%) patients, including all strains from the four patients that had at
301 least one recurrent isolate that developed a carbapenem resistant phenotype.



302

303 **FIG 1** Overview of strains using Illumina short-read data. Timeline showing date of serial
 304 isolation from blood cultures. Patient numbers are in the first column. The shape and color of the
 305 isolates as labelled in the legend indicate clonality and antimicrobial resistance respectively.
 306 Strains were considered possibly clonal if they were the same sequence type and clustered on the
 307 phylogenetic tree. Patient subgroups (e.g. Patient 1; isolate A and B) refers to the order of
 308 isolation. Abbreviations are as follows: ESBL-E = extended spectrum β -lactamase producing
 309 *Enterobacteriales*, CRE = carbapenem resistant *Enterobacteriales*.

310

311

312 **Detection of non-CP-CRE emergence from three sets of clonal, ESBL-E recurrent**
313 **bacteremia isolates**

314 The 11 sets of patient serial isolates included nine *E. coli* serial isolates and two *K. pneumoniae*
315 serial isolates. We performed Illumina short-read sequencing on the 26 strains identified from the
316 11 sets of serial isolates. Strain details are provided in **Table S4**. Serial strain relatedness was
317 assessed using phylogenetic analysis with strains considered possibly clonal if they were the
318 same sequence type, had the same Bayesian population structure, and clustered together on the
319 phylogenetic tree (**Fig. S1**). The temporal collection, antimicrobial resistance, and strain
320 relatedness are depicted in **Fig. 1**. Only a single strain, the 3rd isolate from patient 7, which we
321 will abbreviate as p7C, had a carbapenemase based on a positive Neo-Rapid CARB Kit Test
322 result (**Fig. 1**). Whole genome, short-read sequencing analysis confirmed the presence of the
323 class D carbapenemase, *bla*_{OXA-181} in p7C. Furthermore, p7C was a different sequence type from
324 the patient 7 index strain (p7A) indicating new strain acquisition.

325
326 We focused our subsequent analyses on the isolates from patient 4 (p4), patient 10 (p10), and
327 patient 11 (p11), as each had at least one carbapenem resistant recurring non-CP-CRE isolate.
328 Our HiSeq WGS data indicated that all three patients had recurrent *E. coli* isolates that
329 respectively clustered together within a core genome phylogenetic tree, had the same Bayesian
330 hierarchical population structure, and belonged to the same sequence type (**Fig. S1**). We
331 confirmed clonality by measuring pairwise SNP distances between each recurrent isolate and
332 their respective index strain using their highly resolved, ONT consensus assemblies. Our analysis
333 indicated that there were less than 20 SNPs for all respective recurrent strains relative to their

334 index strain suggesting that these three patients had clonal, re-infecting strains that had
335 developed carbapenem resistance through a non-carbapenemase mechanism (**Table 1**).
336

Table 1. Clonality, Antibiotic Minimum Inhibitory Concentration (MIC), Outer Membrane Protein, and β -lactamase Characterization of *Enterobacteriales* Isolates

Patient	Isolate	ST	Species	Pairwise SNP Distance	CAZ MIC _a	CEP MIC _a	TZP MIC _a	ETP MIC _a	MEM MIC _a	<i>ompC</i> ^{b,d}	<i>ompF</i> ^{c,d}	ESBL Amplification
patient 4	p4A	131	<i>E. coli</i>	Ref	16	≥64	8	≤0.5	≤0.25	WT	WT	None
	p4B	131	<i>E. coli</i>	2	16	8	≥128	≤0.5	≤0.25	WT	WT	<i>bla</i> _{OXA-1}
	p4C	131	<i>E. coli</i>	14	16	64	≥128	≥32	4	c.504_505ins ^e	c.548delA	<i>bla</i> _{OXA-1}
	p4D	131	<i>E. coli</i>	15	≥64	≥64	≥128	≥32	4	c.504_505ins ^e	c.548delA	<i>bla</i> _{OXA-1}
patient 10	p10A	10	<i>E. coli</i>	Ref	16	2	8	≤0.5	≤0.25	WT	WT	None
	p10B	10	<i>E. coli</i>	1	≥64	≥64	≥128	≥32	8	c.634_649del	c.77_87del	<i>bla</i> _{CTX-M-55}
	p10C	10	<i>E. coli</i>	0	≥64	≥64	16	≤0.5	≤0.25	WT	WT	<i>bla</i> _{CTX-M-55}
patient 11	p11A	131	<i>E. coli</i>	Ref	≥64	≥64	64	≤0.5	≤0.5	c.508_521delins ^f	c.305_312del	<i>bla</i> _{OXA-1} , <i>bla</i> _{CTX-M-15}
	p11B	131	<i>E. coli</i>	0	≥64	≥64	≥128	4	1	c.508_518del	c.305_312del	<i>bla</i> _{OXA-1} , <i>bla</i> _{CTX-M-15}
NA	MB101	37	<i>K. pneumoniae</i>	NA	≥64	≥64	≥128	≥32	8	c.607_626ins ^e	NA	<i>bla</i> _{OXA-1} , <i>bla</i> _{CTX-M-15}
NA	MB746	405	<i>E. coli</i>	NA	64	≥64	≥128	≥32	4	c.131insA	c.760_763ins	<i>bla</i> _{OXA-1} , <i>bla</i> _{CTX-M-15}

338 ^aAll minimum inhibitory concentrations (MICs) reported in μ g/ml; abbreviations for each antibiotic are as follows: CAZ

339 (ceftazidime), CEP (cefepime), TZP (piperacillin-tazobactam), ETP (ertapenem), MEM (meropenem)

340 ^bUniprot reference entry names for *ompC* are A0A192C9D6_ECOLX, OMPC_ECOLI, and U9Y7F2_ECOLX for ST131, ST10, and

341 ST405 *E. coli* isolates respectively; MB101 porin is *ompK36* with Uniprot reference entry name A0A0H3H0Y2_KLEPH

342 ^cUniprot reference entry names for *ompF* are A0A192CJUO_ECOLX, OMPF_ECOLI, and S0Z171_ECOLX for ST131, ST10, and

343 ST405 *E. coli* isolates respectively

344

345 ^dWT indicates ‘wild type’ coding DNA sequence, ‘ins’ indicates insertion, ‘del’ indicates deletion, and ‘delins’ indicates

346 insertion/deletion; all indel events are frame-shift mutations that leave premature, truncated coding DNA sequences unless otherwise

347 noted.

348 ^eInsertion of MB1860TU_A with variable number of repeating units

349 ^fINDEL, Y170_N174delinsKR, creates an in-frame OmpC protein

350 ^eInsertion of 1X copy of MB101TPU

351

352

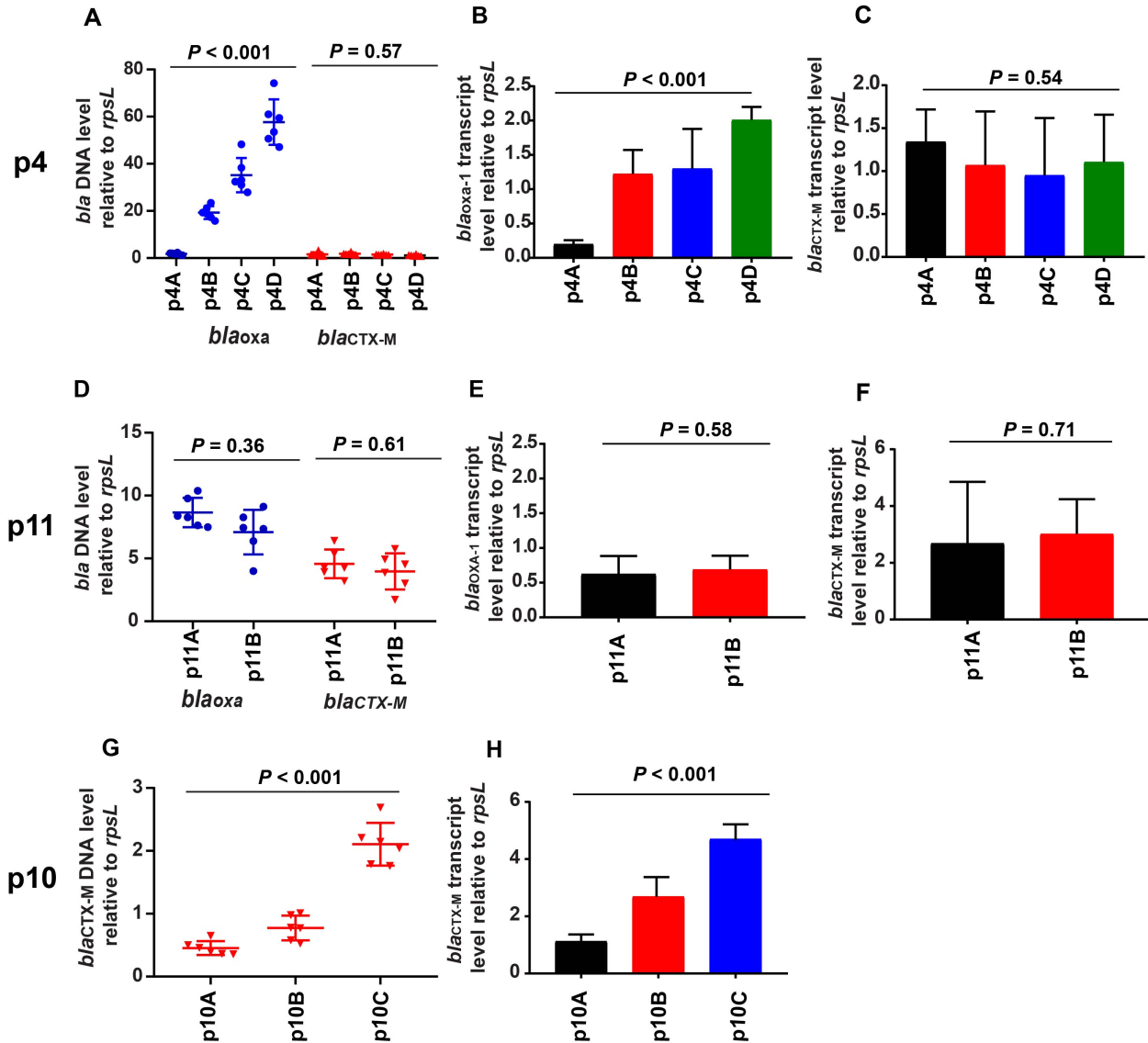
353 **Gene amplification and porin loss associated with emergence of non-CP-CRE**

354 Each of the serial isolates that developed a non-CP-CRE phenotype (p4A-D, p10A-C, and p11A-
355 B) had increased Illumina short-read and ONT long-read coverage depth for β -lactamase
356 encoding genes indicating amplification (**Fig. 2, Table S5 and Table S6**). Specifically, we noted
357 increased coverage depth of *bla*_{OXA-1} in all p4 recurrent isolates, *bla*_{CTX-M-55} in p10 recurrent
358 isolates, and *bla*_{OXA-1} and *bla*_{CTX-M-15} in both the index and recurrent isolate of p11 (**Table 1**). We
359 used qPCR to confirm the increased short-read and long-read coverage depth of our WGS
360 analyses and found that the increased coverage depth corresponded to increased transcript levels
361 of the amplified β -lactamase encoding genes (**Fig. 2**).

362

363 We further characterized outer membrane porin genes with our sequencing data as disruptions of
364 these genes are often correlated with carbapenem resistance. The only consistently identified
365 variation in the non-CP-CRE strains relative to their ESBL-E index strains were mutations that
366 disrupted the ORFs of the porin proteins OmpC and OmpF (**Table 1**). The initial sequencing
367 results indicated that an insertion sequence (IS) as well as nucleotide deletions that resulted in
368 frame-shifts mediated *ompC* gene disruption in isolates that developed non-CP-CRE.

369 Conversely, we consistently found frame-shift inducing deletions in *ompF* genes for both the
370 carbapenem susceptible and resistant isolates (**Table 1**). For p10 isolates, interruption of OmpC
371 and OmpF in the second, non-CP-CRE isolate (p10B) was followed by reversion to the WT
372 OmpC and OmpF genes in the third, carbapenem-susceptible isolate (p10C).



373

374 **FIG 2** PCR analysis of β -lactamase encoding gene levels and transcript levels. Rows from top to

375 bottom are data from patient 4 (p4), patient 11 (p11), and patient 10 (p10). (A, D, G) Taq-Man

376 qPCR of genomic DNA results collected in triplicate on two separate days ($n = 6$) for either

377 *bla*_{OXA-1} (blue) or *bla*_{CTX-M} (red) relative to the endogenous control gene *rpsL*. Data shown are

378 individual data points with mean \pm SD superimposed. (B, E, H) *bla*_{OXA-1} transcript level relative

379 to endogenous control gene *rpsL*. RNA was collected from mid-exponential phase in triplicate on

380 two separate days ($n = 6$). Data shown are mean \pm SD. (C, F) similar analysis to (B, E, H) except

381 that data are for *bla*_{CTX-M}. Note that the p10 strains do not contain *bla*_{OXA-1}. P values refer to

382 measurements in the serial isolates relative to initial isolate using the Kruskal-Wallis (p4 and
383 p10) or Wilcoxon rank-sum tests (p11).

384

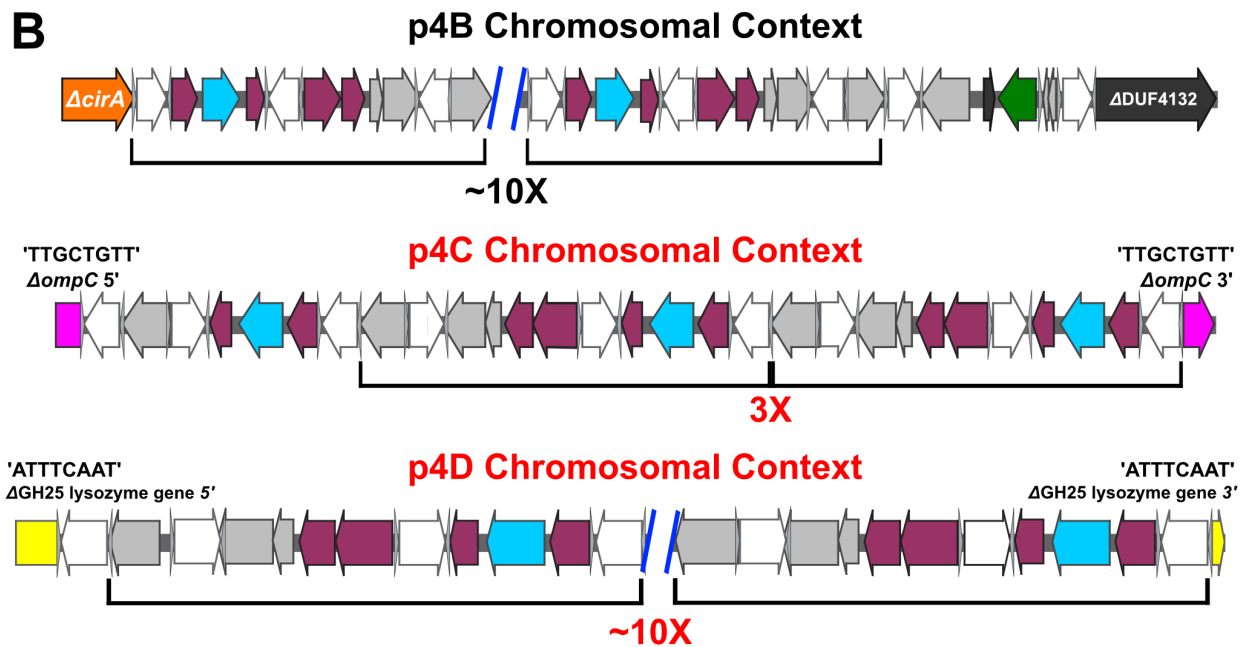
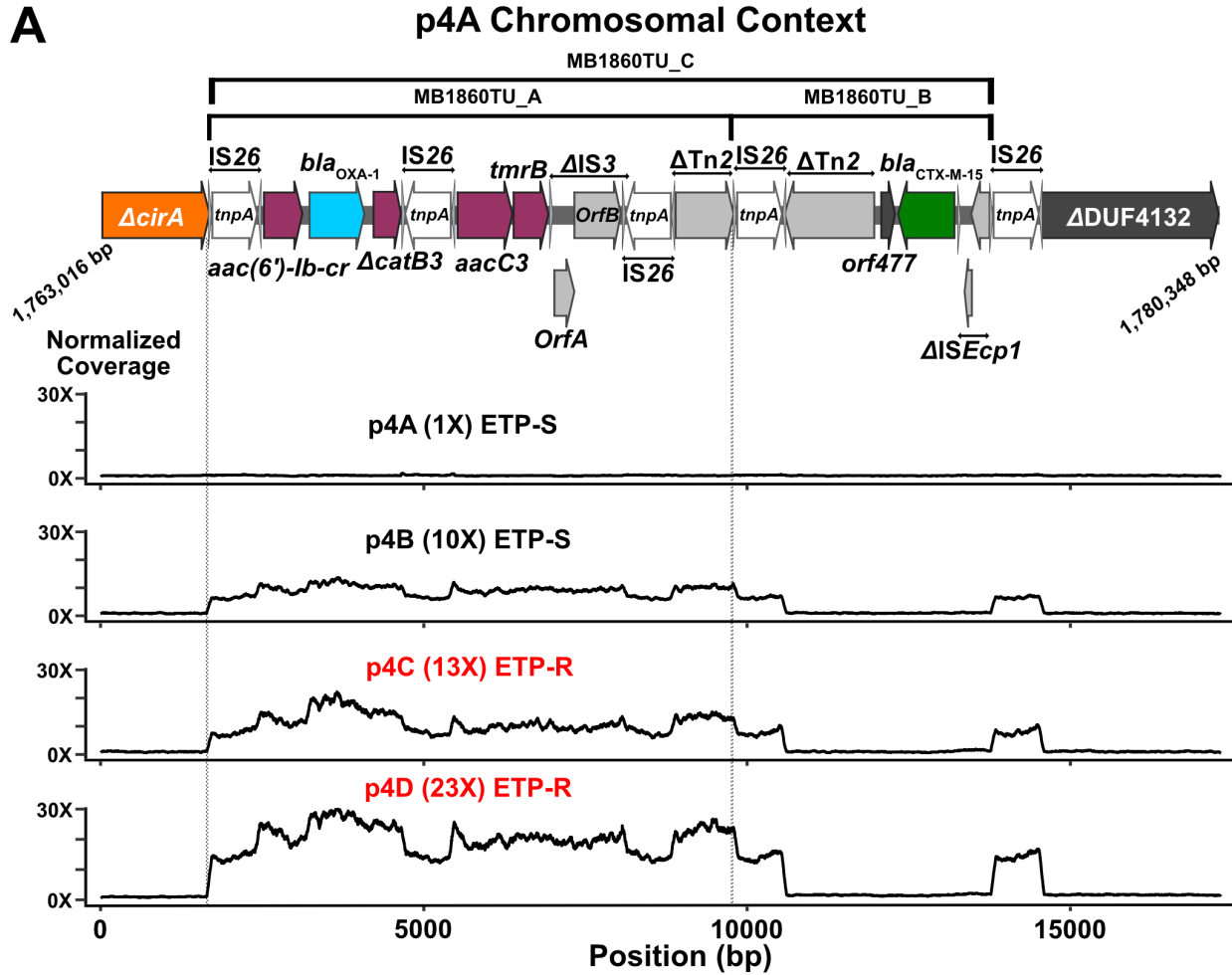
385 **Class 1 transposon TnMB1860 found in both ST131 p4 and p11 serial isolates.**

386 The consensus ONT assembly index strains for patient 4 (p4A) and patient 11 (p11A) both
387 contain an 8,147 bp, IS26-mediated translocatable unit (TU), designated as MB1860TU_A,
388 which carries *bla_{OXA-1}*, and inserts upstream adjacent to another 3,985 bp TU, designated
389 MB1860TU_B, that carries *bla_{CTX-M-15}* (**Fig. 3A**). MB1860TU_C is the combination of both
390 respective TUs that is 12,132 bp in length. The entire putative class 1 transposon structure
391 designated TnMB1860 is 12,952 bp (**Fig. 3A**).

392

393 When moving from the 5' to 3' end of TnMB1860 using the p4A chromosome (**Fig. 3A**;
394 GenBank Accession #: CP049085) as the reference, the first resistance island contains two
395 flanking IS26 *tnpA* genes in opposite orientation with the aminoglycoside N6'-acetyltransferase
396 variant gene (*aac(6')-Ib-cr*), the oxacillinase gene (*bla_{OXA-1}*), and a truncated chloramphenicol
397 resistance determinant (*ΔcatB3*). Following the second IS26 element in TnMB1860, there is an
398 aminoglycoside N3'-acetyltransferase III variant gene, *aacC3*, and the tunicamycin resistance
399 gene, *tmrB*. Downstream of the *tmrB* gene is an IS3 element, which has a modified left inverted
400 repeat (IR_L) and a frame-shifted transposase that has been truncated by a third IS26 element at
401 the 3' end of the IS3 transposase. Immediately downstream of this IS26 element is a truncated
402 Tn2-like transposase which marks the 3' boundary of MB1860TU_A. MB1860TU_A is inserted
403 adjacent to the smaller translocatable unit, MB1860TU_B, which includes another fragment of a
404 Tn2-like transposase, as well as *orf477*, *bla_{CTX-M-15}*, and an *ISEcpI* truncated by an intact IS26,

405 which marks the 3' boundary of both MB1860TU_B, as well as serves as the 3' flanking IS26
406 element for the full length TnMB1860. Similar IS26/Tn2 family transposable elements that have
407 putatively formed via homologous recombination events have previously been reported in
408 association with IS26-mediated AMR transfer in *Enterobacteriales* (6, 39, 40).
409
410 TnMB1860 is located on the p4 chromosome (1,764,660 – 1,777,611 bp; GenBank Accession #:
411 CP049085) and the p11 chromosome (1,812,524 – 1,825,475 bp; GenBank Accession #:
412 CP049077). One of the signatures of transposition are variable sized direct repeats called target
413 site duplications (TSDs) that flank insertion sequences and are created during the transposition
414 process (41-43). The TnMB1860 composite transposon on the p11A chromosome has 7-bp TSDs
415 that indicate a transposition within a 3,762 bp ORF that putatively is involved in molybdopterin
416 cofactor biosynthesis (**Fig. S2**). Interestingly, p4A differs in chromosomal context relative to
417 p11A due to an intramolecular transposition event that occurred in reverse orientation (**Fig. S3**).
418 This is evidenced by the fact that p4A has an approximately 61 kbp region that is inverted with a
419 downstream IS26 in inverse orientation that has an 8-bp TSD within the colicin I receptor gene,
420 *cirA*, in reverse complement orientation (**Fig. S3**). An alignment of p4A and p11A with two
421 other *E. coli* ST131 chromosomes, TO217 (GenBank Accession #: LS992192.1) and
422 Ecol_AZ146 (GenBank Accession #: CP018991.1) indicate similar chromosomal carriage of
423 TnMB1860 with the noted inversion event that has occurred in the p4A isolate (**Fig. S3**).
424



426 **FIG 3** Characterization of IS26-flanked composite transposon with amplification and
427 transposition of modular translocatable units in patient 4 serial isolates (i.e., p4A – p4D).
428 Terminal left and right inverted repeats (IR_L and IR_R respectively) of insertion sequences (ISs)
429 are specified by grey triangles that bracket respective complete and incomplete *tnpA* genes.
430 ORFs are colored as follows: non-β-lactam AMR encoding genes (maroon), *bla*_{OXA-1} (blue),
431 *bla*_{CTX-M-15} (green), IS26 *tnpA* (white), and other IS/Tn elements (gray). (A) Schematic indicates
432 chromosomal context (1,763,016 – 1,780,348 bp; GenBank Accession #: CP049085) of
433 TnMB1860 locus flanked by directly oriented IS26 transposases found in p4A isolate.
434 Immediately below schematic are normalized, short-read coverage depth line graphs for the four
435 p4 serial isolates with MB1860TU_A bracketed by dotted lines. ‘ETP-S’ = ertapenem
436 susceptible; ‘ETP-R’ = ertapenem resistant. Font color for each serial isolate labelled in
437 normalized coverage graph representing carbapenem susceptibility (black) or carbapenem
438 resistance (red). (B) Characterization of amplification, transposition, and ORF disruption events
439 in each of the respective patient 4 recurrent episode isolates. Black brackets beneath ORFs
440 indicate MB168TU_A. The p4B chromosomal context shows a ~10X MB1860TU_A
441 amplification event in the original p4A locus. The p4C chromosomal context additionally
442 contains transposition and disruption of the *ompC* porin gene (pink) ~63 kbp upstream of
443 original p4A TnMB1860 locus with subsequent amplification. p4D isolate has previous
444 amplification and transposition events found in p4B and p4C as well as another MB1860TU_A
445 transposition and disruption of a putative glycoside hydrolase gene (i.e. GH25; yellow) ~71 kbp
446 downstream of TnMB1860 locus with an amplification. The target site duplications (TSDs)
447 created by the transposition of the TU are indicated above each respective junction site that flank
448 the insertion and disruption of the *OmpC* and GH25 genes respectively.
449

450 **Delineation of MB1860TU_A amplification and transposition in ST131 p4 and p11 serial**
451 **isolates**

452 Patient 11 isolates p11A and p11B had relatively the same increase in mapping coverage for both
453 *bla*_{OXA-1} and *bla*_{CTX-M-15} (**Table S3**). The consensus assemblies of both p11 isolates revealed that
454 this increase in relative short-read coverage was due to two copies of *bla*_{OXA-1} and *bla*_{CTX-M-15}
455 being present on a chromosomally located Tn*MB1860* as well as on IS26-mediated genomic
456 resistant modules present on a 180,963bp multireplicon, F-type plasmid, p11A_p2 (**Fig. S2, Fig.**
457 **S3**; GenBank Accession #: CP049079). The CRE phenotype of p11B relative to p11A appears to
458 be driven by additional inactivation of the *ompC* gene given that the *ompF* gene is truncated in
459 both strains (**Table 1**).

460

461 In contrast to the p11 isolates, the amplification and transposition of the modular, translocatable
462 units that compose Tn*MB1860* in p4 was completely in a chromosomal context (**Fig. 3**). There
463 was a consistent increase in short-read coverage depth (**Fig. 3A**) of the entire MB1860TU_A
464 structure up to approximately 23-fold in p4D relative to the seven housekeeping pubMLST genes
465 for ST131. We first analyzed strain p4B, which had developed resistance to piperacillin-
466 tazobactam (TZP) but remained carbapenem susceptible (**Table 1**). In p4B, MB1860TU_A
467 generated a ~10X tandem array *in situ* most likely through a conservative, IS26-mediated
468 transposition mechanism or homologous recombination (**Fig. 3B**) (44, 45). We were unable to
469 assemble the full-length tandem array due to limitations in read length size. However, we were
470 able to use the SVAnts tool to identify thirty individual reads with two or greater number of
471 tandem arrays of MB1860TU_A with 7 individual reads having 4X copies of *bla*_{OXA-1}. The
472 occurrence of the 10X TU amplification at the original Tn*MB1860* locus was confirmed by

473 aligning the p4B long-reads to the reference chromosome p4A using SVAnts in conjunction with
474 a short-read pileup analysis. Additionally, all outer membrane protein genes were WT and
475 remained intact with p4B. Given that p4B had developed resistance to TZP relative to p4A
476 (**Table 1**), we sought to determine whether overexpression of *bla*_{OXA-1} could drive TZP
477 resistance. Inducing *bla*_{OXA-1} expression through cloning under an arabinose responsive promoter
478 increased TZP MIC 6.8-fold relative to uninduced cells (**Fig. S4**).

479
480 Next, we examined the non-CP-CRE serial strains p4C and p4D. Interestingly, in both p4C and
481 p4D a transposition and insertion of MB1860TU_A into *ompC* was present approximately 63
482 kbp upstream of the original *TnMB1860* chromosomal locus (**Fig. 3B**). This *ompC* gene
483 disruption was confirmed through the identification of multiple p4C long-reads > 30 kbp that
484 covered the full transposition site as well as identifying this insertion on the p4C and p4D
485 incomplete chromosomal assemblies. We found two individual long reads that were able to span
486 the full length of the MB1860TU_A array (3X copies) which disrupts *ompC* for p4C and
487 confirmed the exact MB1860TU insertion location within *ompC* (c.504_505ins; **Table 1**). We
488 also were able to identify 8-bp TSDs ('5-TTGCTGTT-3') at each end of the *ompC* insertion
489 sites, which indicates MB1860TU_A replicative transposition. The p4D assembly and individual
490 long-reads indicated a second MB1860TU_A transposition and insertion ~67 kbp downstream of
491 the *TnMB1860* within a GH25 lysozyme gene (**Fig. 3B**). This transposition event could be
492 confirmed by 8-bp TSDs (5'-TTGCTGTT-3') flanking the full insertion site (**Fig 3B**). A
493 progressive increase in both DNA and RNA levels for *bla*_{OXA-1} but not *bla*_{CTX-M-15} in the p4 serial
494 isolates was confirmed using qPCR and qRT-PCR, respectively (**Fig. 2A-C**).

495

496 **Serial passage of patient 4 index strain with ETP elicits *bla*_{OXA-1} and *bla*_{CTX-M-15}**
 497 **amplification through unique translocatable units relative to *in vivo* recurrent strains**
 498 We passaged strain p4A in increasing concentration of ertapenem (ETP) to determine whether
 499 antimicrobial exposure was driving the genetic changes observed in our serial clinical isolates.
 500 Resistance to ETP developed within three passages corresponding to three days, upon which we
 501 collected strains for the next four days (p4A_1- p4A_4). Short-read and qPCR analyses
 502 demonstrated all four isolates had amplification of both *bla*_{OXA-1} and *bla*_{CTX-M-15} relative to the
 503 p4A index strain prior to ETP exposure (**Fig 4**). ONT sequencing on all four isolates indicated
 504 that similar to p4B, there was *in situ* amplification occurring at the original TnMB1680
 505 chromosomal locus. However, the full-length TU, MB1860TU_C, which consists of
 506 MB1860TU_A and MB1860TU_B (**Fig 4A**) that harbor *bla*_{OXA-1} and *bla*_{CTX-M-15} respectively,
 507 was the amplifying structure for the passaged isolates in contrast to what we saw in the *in vivo*
 508 isolates where MB1860TU_A was the sole amplifying structure (**Table 2**).

Table 2. p4A Serial Passaging Characterization

Sample	ETP MIC (µg/ml)	<i>ompC</i>	<i>ompF</i>	<i>bla</i> _{OXA-1} copy # ^a	<i>bla</i> _{CTX-M-15} copy # ^a
p4A_1	≥ 32	c.208T>C; p.Q70X	c.17_18insIS1A	6	7
P4A_2	≥ 32	c.208T>C; p.Q70X	WT	8	9
P4A_3	≥ 32	c.208T>C; p.Q70X	c.62_63insIS1A	6	7
P4A_4	≥ 32	c.208T>C; p.Q70X	c.517_518insIS1A	6	7
p4A_H1	≥ 32	IS1A variant mediated insertion 96 bp upstream of +1 <i>ompC</i> start site	WT	1	9
p4A_H2	≥ 32	IS1A variant mediated insertion 96 bp upstream of +1 <i>ompC</i> start site	WT	1	17

509 ^aSee Materials and Methods for Calculation

510

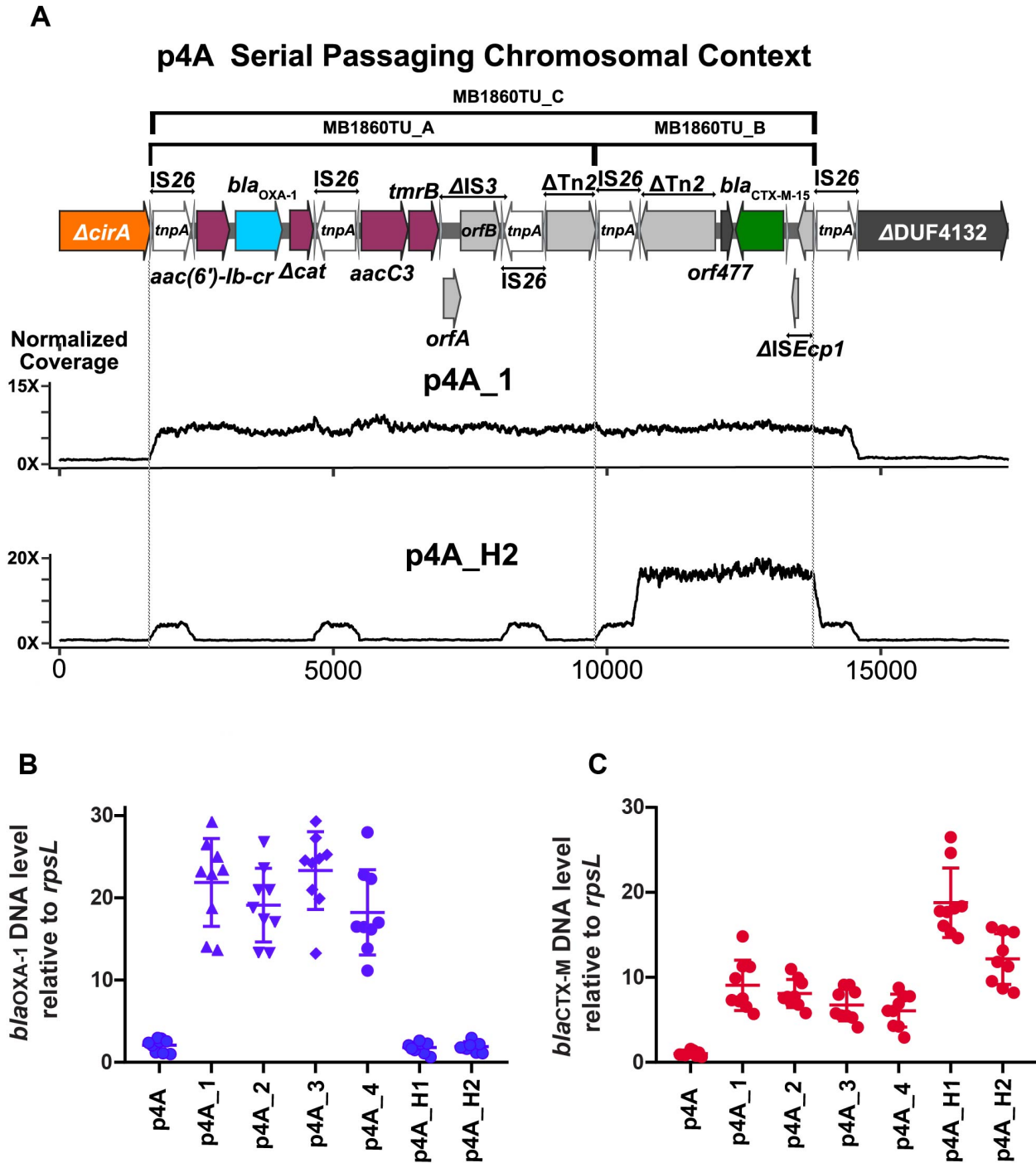
511 We repeated the experiment and again found that ETP resistance developed within three
 512 passages from the p4A isolate. ONT sequencing of the 2nd round of passaged isolates (p4A_H1
 513 and p4A_H2) revealed amplification of MB1860TU_B that harbors *bla*_{CTX-M-15} solely, which we

514 verified using qPCR (**Fig. 4**). Similar to what we observed *in vivo*, the serially passaged ETP
515 resistant isolates contained inactivating mutations in *ompC* although we did not observe any
516 MB1860TU mediated interruptions (**Table 2**). INDELS inactivating the *ompF* gene were
517 observed in a fraction of the serial isolates suggesting *ompF* inactivation may not be necessary
518 for the development of non-CP-CRE (**Table 2**). We found that p4A responded *in vitro* to ETP
519 similarly to what was observed in our serial clinical isolates by amplification of modular
520 MB1860TU elements with concomitant porin disruption. Nevertheless, there was differential
521 amplification of IS26 translocatable units demonstrating the modularity of these MGEs.

522

523 In order to determine the chromosomal stability of the MB1860TU tandem arrays in the absence
524 of antibiotic selective pressure, both ertapenem resistant (ETP-R) strains p4C and p4D were
525 passaged for 10 days (~60 generations) without supplemented ertapenem. Both ETP-R recurrent
526 strains consistently maintained carbapenem resistance through 60 generations of growth (**Fig.**
527 **S5**). The p4C and p4D strains had relative copy number decreases of *bla*_{OXA-1} from 33 to 18X
528 (45% decrease) and 53 to 42X (21% decrease) respectively (**Fig. S5**). These results indicate that
529 both adapted strains can have persisting carbapenem resistance with associated tandem arrays of
530 amplified *bla*_{OXA-1} maintained in the absence of antibiotic exposure.

531



532

533 **FIG 4** Identification and characterization of β -lactamase gene amplification following p4A serial
 534 passaging under ertapenem (ETP) exposure. Strain p4A was grown in ETP with isolates p4A_1-
 535 4 collected during the 1st round of passaging and p4A_H1 and p4A_H2 collected during the 2nd
 536 round. (A) Schematic of TnMB1860 locus from p4A as detailed in FIG 2. Immediately below

537 the schematic are normalized, short-read coverage depth line graphs for p4A_1 and p4A_H2
538 aligned to p4A with location of MB1860TU_A and MB1860TU_B bracketed by dotted lines.
539 Note amplification of MB1860 TU_C in strain p4A_1 whereas only MB1860TU_B amplified in
540 strain p4A_H2. (B) and (C) are Taq-Man qPCR of genomic DNA collected in triplicate on two
541 separate days (n = 6) for either *bla*_{OXA-1} (B) or *bla*_{CTX-M} (C) relative to the endogenous control
542 gene *rpsL*. Data shown are individual data points with mean ± SD superimposed.

543

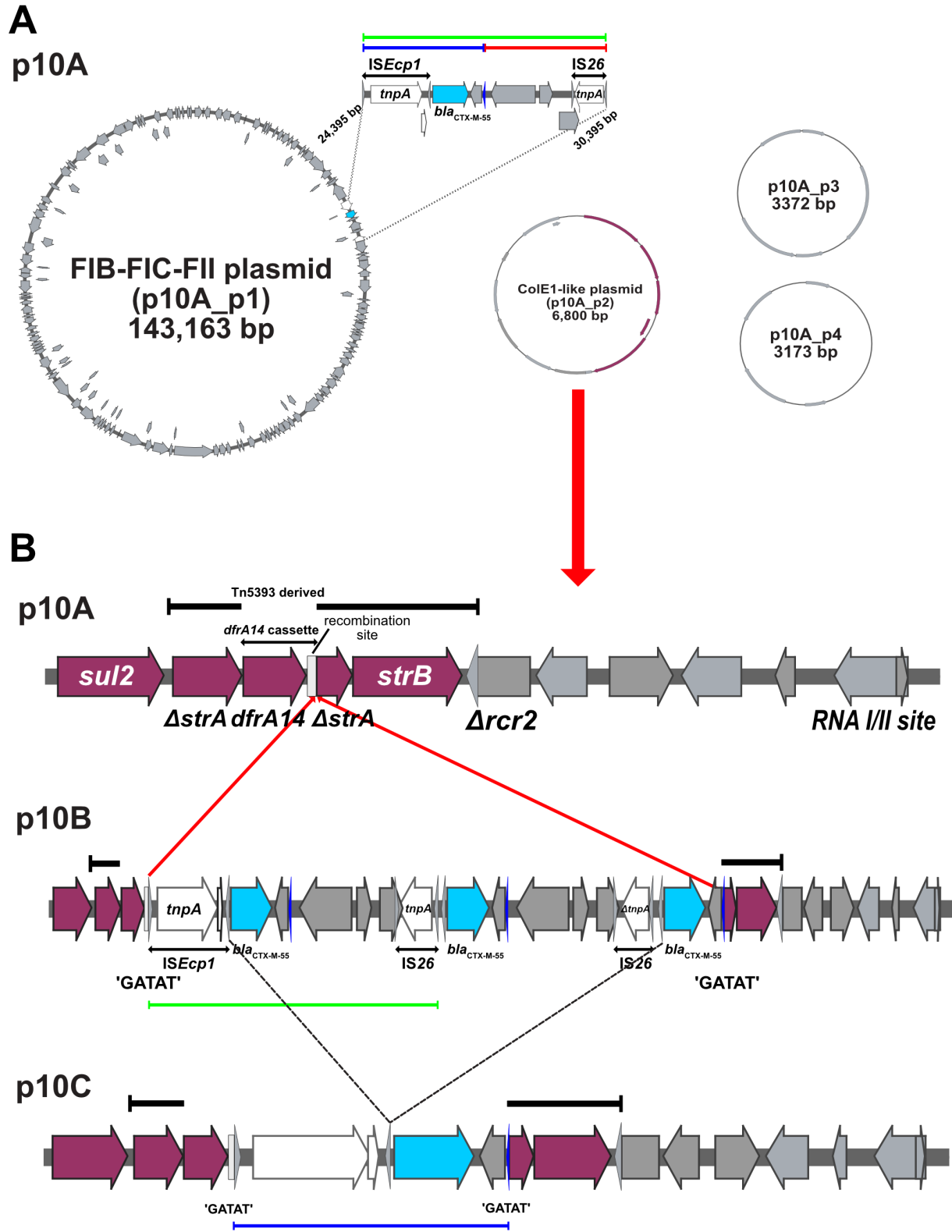
544 **Unique *ISEcpI*-mediated plasmid transposition of *bla*_{CTX-M-55} in patient 10 serial ST10 *E.***

545 ***coli* isolates** Compared to the p4 and p11 strains, the serial isolates from patient 10 (p10A –
546 p10C) contain a different MGE that putatively drives the amplification of the ESBL encoding
547 gene, *bla*_{CTX-M-55}. Index strain p10A harbors *bla*_{CTX-M-55} located on an *ISEcpI* transposition unit
548 designated EC215TPU that resides on a multireplicon FIB-FIC-FII plasmid (**Fig. 5; GenBank**
549 **Accession #: CP049082**). ‘Transposition unit’ (TPU) is used here to distinguish from
550 translocatable unit (TU) as they have different transposition mechanisms that are mediated by
551 *ISEcpI* and IS26 respectively. There are three additional small plasmids present in p10A, one of
552 which is a Cole1-like 6.8 kbp plasmid designated as p10A_p2 (GenBank Accession #:
553 CP049083), that is highly comparable to pCERC1 (GenBank Accession #: JN012467) (46).

554

555 EC215TPU translocates from the multireplicon, type F plasmid in p10A, designated p10A_p1, to
556 p10A_p2 in the second serial isolate p10B (**Fig. 5B**). The *ISEcpI*-mediated transposition can be
557 identified and confirmed by the 5-bp TSDs that are immediately upstream of the *ISEcpI* 14 bp
558 IRL and downstream of another alternative, 14 bp right inverted repeat (IRRalt) respectively (41,
559 42, 47). There are TSDs flanking EC215TPU at the p10A_p2 recombination site with both the

560 IR_L and IR_{Ralt} represented as a blue triangle (5'-CCTCACACCTTCGA-3') on **Fig. 5B**.
561 Notably, as the entire insertion region with three *bla*_{CTX-M-55} copies on p10B has 5 bp TSDs (5'-
562 GATAT-3') that flank IR_L and IR_{Ralt} respectively, one can postulate that an amplification event
563 occurred on p10A_p1, which then subsequently inserted into p10A_p2 (**Fig. 5B**). p10B is also
564 the only carbapenem-resistant p10 serial isolate. This likely is due in part to deletions in *ompC*
565 and *ompF* that create frame-shift truncations of each respective porin in contrast to the WT
566 genotypes of each respective porin found in p10A and p10C. When analyzing the third isolate
567 p10C, there only is one copy of *bla*_{CTX-M-55} identified in the assembly and short-reads on
568 p10A_p2. However, the short-read and long-read coverage depth analysis suggests an increase in
569 copy number (**Table S6**). The subsequent increase in initiation RNA genes, a marker for ColE1-
570 like plasmids with *bla*_{CTX-M-55} suggests that p10C amplification occurs by an increase in the
571 overall copy number of p10A_p2. These data are consistent with our qPCR analysis which
572 demonstrate progressively higher DNA levels of *bla*_{CTX-M-55} for each serial isolate (**Fig. 2**).
573



575 **FIG 5** Genomic analysis of *ISEcpI*-mediated mobilization of *bla*_{CTX-M-55} from a multireplicon F
576 type plasmid (GenBank Accession #: CP049082) to a ColE1-like, high copy number plasmid
577 (GenBank Accession #: CP049083) in p10 isolates. (A) Non-chromosomal genomic context for
578 patient 10 index strain (p10A) harboring four circular plasmid structures. The *bla*_{CTX-M-55} gene is
579 located on a 143,163 bp FIB-FIC-FII plasmid (p10A_p1) on an *ISEcpI*-*bla*_{CTX-M-55}-IS26 5945
580 bp transposition unit (TPU) indicated by a green bracket. The blue and red regions are the union
581 of this green bracketed region and represent regions highlighted in the following section. (B)
582 Progression of TPU insertion into ColE1-like plasmid (p10A_p2). p10A illustrates the full
583 genome context of p10A_p2 including the *sul2-ΔstrA-dfrA14-ΔstrA-strB* resistance island. The
584 top black, bracketed region indicates genome that is derived from the Tn5393 transposon. The
585 region indicated by double-ended arrows indicates where the *dfrA14* cassette inserted itself into
586 Tn5393. The box downstream of *dfrA14* indicates the *attC* recombination site. p10B shows the
587 amplified TPU (11,939 bp) insertion into the p10A_p2 recombination site (indicated by red
588 arrows). The region is bracketed by 5-bp TSDs (5'-GATAT-3') indicative of *ISEcpI* mediated
589 transposition. An alternative, right inverted repeat (IRRalt) is indicated by blue triangle. p10C
590 contains an 8,998 bp deletion (black dotted lines) where two copies of *bla*_{CTX-M-55} are dropped.
591 This creates a TPU flanked by the IRL and IRRalt of *ISEcpI* (blue bracket).

592

593 **Detection of β-lactamase gene amplification and porin disruption by AMR elements in non-** 594 **serial *Enterobacteriales* strains**

595 There were two non-CP-CRE strains, MB101 (*K. pneumoniae*) and MB746 (*E. coli*), identified
596 in a previous study examining the role of short-read WGS in predicting β-lactam resistance
597 (**Table 1**) (18). We performed ONT sequencing on the two isolates to determine whether

598 mechanisms of gene amplification and porin disruption were similar to what was observed in the
599 cohort of non-CP-CRE bacteremia serial isolates. Both isolates contain IS26-mediated TUs
600 carrying *bla*_{OXA-1} as previously characterized in TnMB1860 (**Fig. 6**).

601
602 For strain MB101 (**Fig. 6A-B**), a ST37 *K. pneumoniae* isolate, the ~9310 bp IS26-mediated TU,
603 designated MB101TU harbors *aac*-(6')-*Ib-cr*, *bla*_{OXA-1}, Δ *catB3*, *aacC3*, and *tmrB* and is nearly
604 identical (100% coverage; 99.9% ID) to MB1860TU_A found in serial *E. coli* isolates from
605 patients 4 and 11 (**Fig. 6A**). An ISEcpI-mediated TPU harboring *bla*_{CTX-M-15}, designated
606 MB101TPU, was also present on the MB101 chromosome as well as an FIB_K type plasmid, with
607 at least five copies present on the chromosome based on the Flye assembly and individual long
608 reads. The assembly and long reads indicate the transposition of MB101TPU into a putative
609 glycoporin encoding gene which was followed by the transposition of MB101TU to an existing
610 IS26 element downstream of MB101TPU (**Fig. 6A**). We identified an amplification of
611 MB101TU at this locus that created a tandem array of at least 8X copies based on the
612 identification of multiple long-reads with multiple MB101TU copies. Interestingly, there was
613 also a transposition event of MB101TPU into the *ompK36* encoding gene, a homolog of *ompC* in
614 *E. coli* (**Fig. 6B**). Thus, MB101 has amplification of both *bla*_{OXA-1} and *bla*_{CTX-M-15} via distinct
615 mechanisms along with TPU mediated porin disruption.

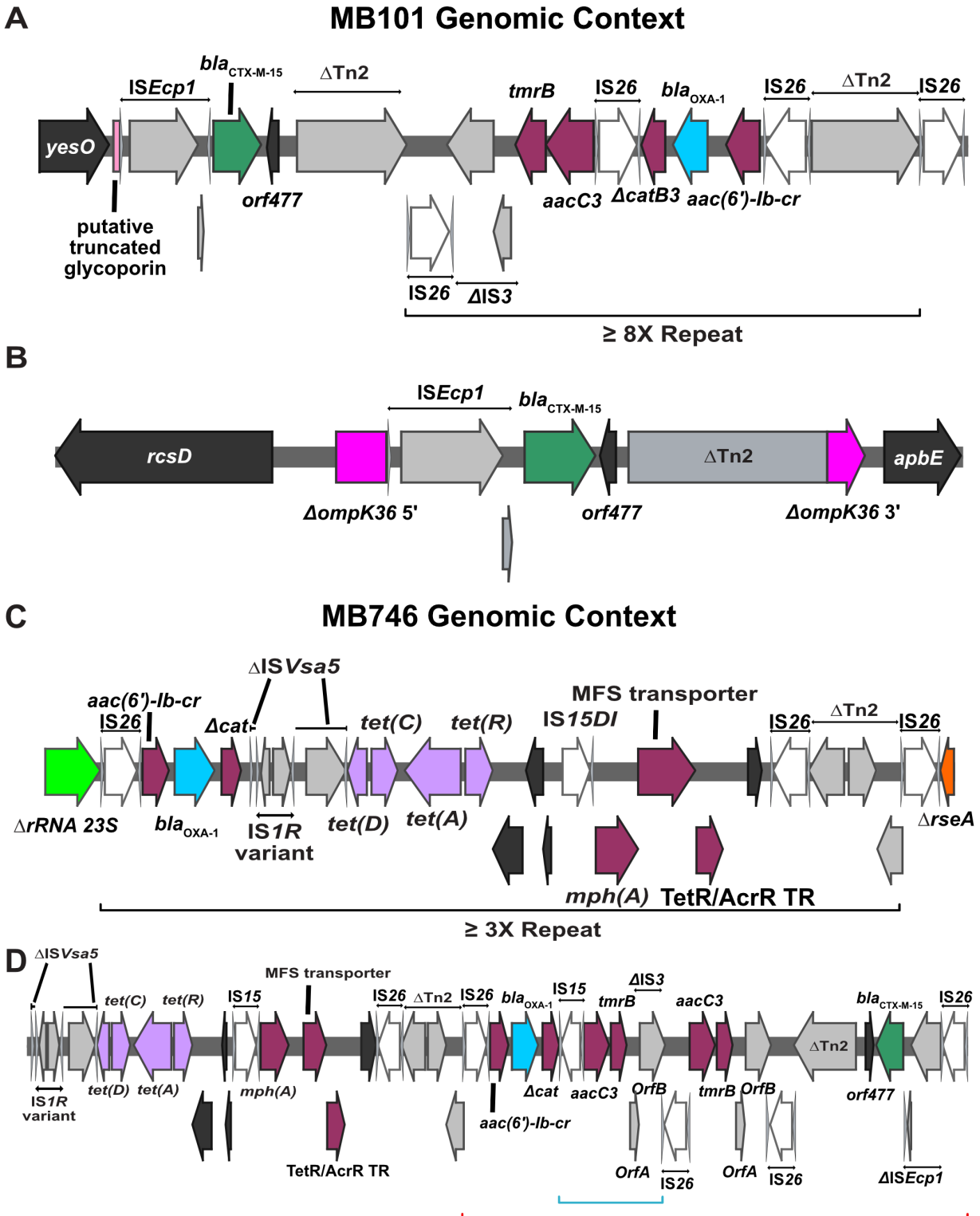
616
617 The ST405 *E. coli* isolate, MB746, has an IS26-mediated translocatable unit, designated
618 MB746TU, which also includes a genomic resistance module that carries *aac*-(6')-*Ib-cr*, *bla*_{OXA-1},
619 and a truncated chloramphenicol resistance determinant (**Fig. 6C-D**) similar to what is seen on
620 MB1860TU_A. MB746TU carries an additional tetracycline resistance operon and macrolide

621 resistance operon, not present on TnMB1860. We found four individual long-reads that carried at
622 least 3X copies of MB746TU present in the same MB746 chromosomal location (**Fig. 6C**).

623 Additionally, we identified a FIB plasmid that harbored both *bla*_{OXA-1} and *bla*_{CTX-M-15} which has a
624 similar configuration and orientation to TnMB1860 (Red bracket; **Fig. 6D**). The IS26-mediated
625 TU carrying *bla*_{CTX-M-15} has 99.7% BLAST identity with MB1860_B with the only substantial
626 difference being a size of 5558 bp vs the 3985 bp MB1860TU_B (**Fig. 6D**). Notably, MB746
627 also had *ompC* and *ompF* gene disruptions (**Table 1**) comparable to what was observed in serial
628 isolates that developed non-CP-CRE phenotypes with concomitant β -lactamase amplifications.

629 These data show that there are multiple, non-CP-CRE clinical isolates within our region that
630 have amplification and transposition of similar IS26 and *ISEcpI*-mediated elements that harbor
631 β -lactamase genes encoding genes in conjunction with porin disruption.

632



633

634 **FIG 6** Genomic context of end-stage MB101 and MB746 isolates respectively. Terminal left and

635 right inverted repeats (IR_L and IR_R respectively) of insertion sequences (ISs) are specified by

636 grey triangles that bracket respective complete and incomplete *tnpA* genes. ORFs are colored as
637 follows: AMR genes (maroon), *bla*_{OXA-1} (blue), *bla*_{CTX-M-15} (green), IS26 *tnpA* (white), and other
638 IS/Tn elements (gray). Delta (Δ) next to annotated genetic region indicates a truncation or
639 disruption (A-B) Chromosomal locations of MB101TU (Fig. 6A) and MB101TPU (Fig. 6B)
640 indicating respective amplification and transposition of each element. ~8X MB101TU repeat
641 indicated by black bracket. Truncated *ompK36* gene labelled in pink (Fig. 6B). (C-D) Genomic
642 context of MB746TU and respective genomic resistance modules carrying AMR genes. Black
643 brackets beneath Fig. 6C schematic indicate repeating MB101TU unit. Fig. 6D indicates FIB
644 plasmid carriage of genomic resistance modules. Red bracket indicates IS26 transposon structure
645 that shares 100% coverage; 99% BLAST similarity with TnMB1860. Blue bracket indicates
646 small, 2X repeat structure.

647

648 **DISCUSSION**

649 While recent systematic surveillance studies indicate that the prevalence of non-CP-CRE
650 remains high (3, 4), the mechanisms contributing to non-CP-CRE emergence within clinical
651 settings are largely unknown. This public health issue is particularly troublesome given the
652 difficulties in properly diagnosing non-CP-CRE infections and prescribing effective
653 antimicrobials due to the transient nature of their carbapenem resistance phenotypes as well as
654 their lack of a carbapenemase. This analysis demonstrates that multiple TU- and TPU-mediated
655 modular amplifications of β -lactamase encoding genes in conjunction with porin inactivation are
656 driving non-CP-CRE emergence in our cohort of cancer patients with recurrent bacteremia . It is
657 well established that “copy-in” replicative transposition is a common mechanism by which IS26-
658 mediated TUs mobilize to regions lacking a pre-existing IS26 element (43, 45, 48). Recent

659 studies have shown that TUs can create tandem resistance gene arrays by targeting genomes with
660 an existing copy of IS26 through an intermolecular, conservative transposition mechanism and,
661 less frequently, through RecA-dependent homologous recombination (43-45, 49). We were able
662 to determine that the initial incorporation of the composite transposon TnMB1860 as well as
663 MB1860TU_A transposition into *ompC* and the predicted glycoside hydrolase gene occurred
664 through a replicative transposition mechanism based on characteristic TSDs at insertion sites.
665 The amplification of MB1860TU_A, MB1860TU_B, and MB1860TU_C that created tandem
666 arrays of resistance genes observed in our study likely occurred through a conservative
667 transposition mechanism although RecA-dependent homologous recombination could also be
668 contributing to the generation of these tandem arrays. The ability to detect genetic structures
669 capable of amplifying β -lactamase encoding genes that synergistically insert into porin encoding
670 genes will be essential for diagnostic and surveillance purposes in order to ensure we are
671 properly identifying clinical isolates that have the ability to develop CRE.

672
673 Amplification of *bla*_{OXA-1} in both serial and non-serial isolates consistently included *aac*-(6')-Ib-
674 *cr* and Δ *catB3* in association with other modular, IS26-mediated translocatable units carrying
675 resistance genes. Livermore et al. recently observed the co-occurrence of *aac*-(6')-Ib' and
676 *bla*_{OXA-1} in 147/149 *E. coli* strains, primarily ST131 isolates, suggesting that many *E. coli*
677 isolates carrying *bla*_{OXA-1} may be capable of this amplification (40, 50). The finding that *bla*_{OXA-1}
678 amplification was consistently associated with progressive development of β -lactam resistance
679 was somewhat surprising given that this enzyme is typically considered a narrow spectrum β -
680 lactamase (51). However, overexpression of *bla*_{OXA-1} using an arabinose inducing promoter was
681 shown to generate resistance to ertapenem in a porin deficient strain (6). This study, along with

682 our own results, indicates that *bla*_{OXA-1} likely contains sufficient carbapenem hydrolysis activity
683 to generate resistance with the combination of augmented gene copy number and decreased
684 carbapenem concentration due to porin inactivation. Additionally, we demonstrate that
685 overexpression of *bla*_{OXA-1} without porin inactivation produced TZP resistance, a finding that
686 could help resolve previously noted discrepancies between β -lactamase gene content and TZP
687 susceptibility (18, 52). Similar amplifications of the narrow-spectrum, TEM β -lactamases have
688 also been shown to be associated with TZP resistance, suggesting amplification of narrow-
689 spectrum β -lactamases may be a substantial mechanism contributing to TZP resistance (9-11).

690
691 The other amplified β -lactamase in our cohort was *bla*_{CTX-M}. CTX-M is the most commonly
692 identified ESBL enzyme in *Enterobacterales* (53) and high level CTX-M production has
693 previously been shown to confer carbapenem resistance in porin-deficient *E. coli* (6) as well as
694 *K. pneumoniae* (54). Amplification of MB1860TU_B harboring *bla*_{CTX-M-15} with concomitant
695 porin disruption was associated with development of ETP resistance in one of our two passaging
696 experiments, demonstrating the versatility of various modules of TnMB1860 in responding to
697 carbapenem exposure. Unlike *bla*_{OXA-1}, we did not identify *bla*_{CTX-M} genes in association with
698 other AMR encoding elements, but rather *bla*_{CTX-M} genes were consistently present with an
699 *ISEcpI* element, which has been previously described for *E. coli* (55, 56). The array of
700 mechanisms by which *bla*_{CTX-M} copy numbers could increase included (1) *in situ* IS26-mediated
701 amplification of MB1860TU_B (strain P4A_H1) and MB1860TU_C (strain p4A1 – p4A4); (2)
702 *ISEcpI*-mediated transposition of EC215TPU from a low-copy, F-type plasmid to a multi-copy,
703 ColE1-like plasmid (strain p10B); (3) and multiple chromosomal *ISEcpI*-mediated transposition
704 unit insertions including into the *ompK36* porin encoding gene (strain MB101). Thus, the various

705 *bla*_{CTX-M} amplification mechanisms suggest that augmentation of *bla*_{CTX-M} copy number may be a
706 major driver of non-CP-CRE development.

707

708 Along with permitting direct visualization of genomic resistance module amplifications
709 containing various β -lactamase genes, ONT sequencing also allowed for identifying porin-
710 mediated disruption by TU and TPU structures. Laboratory studies that generated CRE strains
711 through serial passaging have consistently found that OmpC and OmpF porin production is
712 reduced in carbapenem resistant isolates, but mechanisms have generally involved alterations in
713 the porin regulatory protein OmpR or in OmpR binding sites (6, 7). Similarly, a recent study of a
714 single patient with recurrent *E. coli* infection identified a single amino acid change in OmpR as
715 leading to loss of OmpC and OmpF expression with development of ertapenem resistance (15).
716 Conversely, we observed direct inactivation of OmpC and OmpF either through TU/TPU
717 transposition into their respective ORFs or through INDELS that resulted in frame-shift
718 mutations whereas no alterations in OmpR were identified in our cohort. To our knowledge,
719 there have been only two other reports of an *ISEcpI*-mediated TPU harboring a β -lactamase
720 encoding gene causing the disruption of a porin, in both cases OmpK35 from *K. pneumoniae* (57,
721 58). While there have been a number of studies that have implicated IS-mediated porin
722 disruptions, including IS26 mediated disruptions (54, 59-61), to our knowledge, this is the first
723 documented IS26-mediated translocatable unit disruption of an outer membrane porin followed
724 by amplification of a β -lactamase gene. Given the length of the TUs, as well as the fact that they
725 have the ability to amplify and create tandem arrays once inserted into the porin encoding genes,
726 it is unlikely that targeted PCR based strategies or commonly used short-read approaches alone
727 would have been able to identify the full extent of the TUs observed in our study. Thus,

728 performing long read sequencing on larger cohorts of *Enterobacterales* with porin deficient
729 backgrounds should reveal whether TU/TPUs harboring AMR genes are a frequent mediator of
730 porin gene disruption, but have not been previously identified due to the long and repetitive
731 nature of the involved DNA structures.

732

733 The clinical impact of β -lactamase gene amplification and porin loss driving carbapenem
734 resistance has been postulated to be mitigated by the fitness costs imposed on the organism by
735 such genetic changes (37, 62-64). However, such AMR mechanisms are increasingly being
736 recognized as commonly occurring in clinical isolates, including in the serious infections
737 described in our cohort (37, 62-64). This would suggest that organisms with the capability to
738 amplify AMR genes are widespread and capable of causing significant human infections
739 especially under antibiotic selective pressure (7, 9, 10, 65). Moreover, the most recent systematic
740 data on CRE in the U.S. found no difference in outcomes between patients with CRE and non-
741 CP-CRE suggesting that circulating non-CP-CRE organisms may not have significant fitness
742 defects (4). Furthermore, our own results from serial passaging p4C and p4D without ETP
743 exposure suggests these amplified structures harboring AMR genes associated with increased
744 MICs to ertapenem and meropenem may be fairly stable in the absence of antibiotic selective
745 pressure. The chromosomal context of *TnMB1860* may provide insights into the stability of this
746 structure as a previous study had noted stable transposon carriage when mobilized from plasmid
747 to chromosome at very low levels of antibiotic exposure (66).

748

749 In addition to possibly imposing a fitness cost on the organism, AMR gene amplifications have
750 also been associated with the presence of antimicrobial heteroresistance (65, 67). The interplay

751 between AMR gene amplification, heteroresistance and fitness is likely reflected in the genotype
752 changes of patient 10 isolates given the development and subsequent reversion of porin
753 mutations observed in those strains (Table 1). Recent systematic surveys have indicated
754 approximately 25% of strains reported as carbapenem resistant tested as susceptible at a central
755 laboratory, likely reflecting the transient phenomenon of AMR heteroresistance (4, 64). A
756 limitation of our study is that we only sequenced a single colony for each isolate, which almost
757 certainly underestimates the genetic complexity of a bacterial population responding to
758 antimicrobial therapy. Nevertheless, a challenge presents itself in capturing population
759 heterogeneity with a single genome assembly as multiple, heterogeneous sequencing reads will
760 break current assembler algorithms.

761

762 **CONCLUSIONS**

763 We have used serial, clinical isolates subjected to complementary WGS approaches to identify
764 that a combination of porin inactivation and IS-mediated amplification of TU and TPU elements
765 harboring β -lactamase encoding genes underlies the emergence of non-CP-CRE from ESBL-E
766 parental isolates within our study population. We predict that more widespread application of
767 long-read sequencing technologies will facilitate appreciation of the mechanisms and impact of
768 TU- and TPU-mediated transpositions, porin disruptions, and gene amplifications on a diverse
769 array of AMR pathogens in the clinical setting.

770

771 **LIST OF ABBREVIATIONS**

772 non-carbapenemase-producing carbapenem resistant *Enterobacterales* = non-CP-CRE;
773 translocatable unit = TU; transposition unit = TPU

774

775 **DECLARATIONS**

776 **Ethics approval and consent to participate:** A waiver of informed consent to collect clinical
777 data from electronic medical records and analyze the isolates was provided by the MDACC IRB
778 (PA15-0799). A waiver of consent from UTHealth Science Center at Houston (IRB #: HSC-
779 SPH-20-0032) was obtained to perform bacterial genomics sequencing and analysis.

780

781 **Consent for publication:** Not applicable

782

783 **Availability of data and materials:** The index isolate assemblies from recurrent bacteremia
784 patients that developed non-CP-CRE as well as the long-read and short-read data for all isolates
785 have been deposited in the National Center for Biotechnology (NCBI) BioProject database
786 PRJNA603908. ONT-sequencing data of non-CP-CRE isolates from a previous study (18) were
787 deposited in NCBI BioProject database PRJNA388450. All other data analyzed during this study
788 are available in the supplemental materials and/or available upon request from the corresponding
789 author.

790

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792

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802

803 **Author contributions:** WCS developed the database, generated and analyzed the data, and was
804 a major contributor to the writing of the manuscript. SLA designed the study, collected and
805 analyzed the data, and significant contributor to writing the manuscript. RP helped designed the
806 study, performed the experiments, and analyzed the data. JK performed experiments and
807 analyzed the data. MB performed the experiments and contributed to writing the paper. XL
808 helped perform phylogenomic analyses. AK helped perform phylogenomic analyses and
809 contributed to writing the paper. JGP contributed to the design of the study, analyzed the isolates,
810 and writing of the paper. PS collected and curated the isolates. CAA helped design the study,
811 analyze the data, and writing of the paper. DEG analyzed the data and contributed to the writing
812 of the paper. BMH wrote scripts to analyze the data, analyzed the data, and contributed to
813 writing the paper. SAS designed the study, analyzed the data, and major contributor to the
814 writing of the manuscript. All authors read and approved the final manuscript.

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819

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