

Convergence of virulence and MDR in a single plasmid vector in MDR *Klebsiella pneumoniae* ST15

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Background: MDR and hypervirulence (hv) are typically observed in separate *Klebsiella pneumoniae* populations. However, convergent strains with both properties have been documented and potentially pose a high risk to public health in the form of invasive infections with limited treatment options.

Objectives: Our aim was to characterize the genetic determinants of virulence and antimicrobial resistance (AMR) in two ESBL-producing *K. pneumoniae* isolates belonging to the international MDR clone ST15.

Methods: The complete genome sequences of both isolates, including their plasmids, were resolved using Illumina and Oxford Nanopore sequencing.

Results: Both isolates carried large mosaic plasmids in which AMR and virulence loci have converged within the same vector. These closely related mosaic hv-MDR plasmids include sequences typical of the *K. pneumoniae* virulence plasmid 1 (KpVP-1; including aerobactin synthesis locus *iuc*) fused with sequences typical of IncFII_K conjugative AMR plasmids. One hv-MDR plasmid carried three MDR elements encoding the ESBL gene *bla*_{CTX-M-15} and seven other AMR genes (*bla*_{TEM}, *aac3'-IIa*, *dfrA1*, *satA2*, *bla*_{SHV}, *sul1* and *aadA1*). The other carried remnants of these elements encoding *bla*_{TEM} and *aac3'-IIa*, and *bla*_{CTX-M-15} was located in a second plasmid in this isolate. The two isolates originated from patients hospitalized in Norway but have epidemiological and genomic links to Romania.

Conclusions: The presence of both virulence and AMR determinants on a single vector enables simultaneous transfer in a single event and potentially rapid emergence of hv-MDR *K. pneumoniae* clones. This highlights the importance of monitoring for such convergence events with stringent genomic surveillance.

Introduction

The majority of infections caused by *Klebsiella pneumoniae* (*Kp*) are typically associated with one of two distinct clinical phenomena caused by non-overlapping *Kp* populations: healthcare-associated infections caused by MDR *Kp* strains that also often cause nosocomial outbreaks, and community-acquired, invasive infections caused by hypervirulent (hv) strains.^{1,2} However, convergent strains carrying both MDR and hv genes have been reported.^{3–7} Recently, a high-mortality outbreak of ventilator-associated pneumonia caused by a strain of hv carbapenemase-producing ST11 *Kp* was reported in China, demonstrating that the combination of enhanced virulence potential and difficulties in treatment posed by MDR can be fatal. The Chinese report was particularly notable as ST11 is typically associated with MDR, and

appears to be the most common cause of carbapenemase-producing *Kp* infections reported in China. However, the outbreak strains had additionally acquired a virulence plasmid harbouring *iuc* (aerobactin siderophore) and *rmpA2* (hypermucoidity) loci, which are usually only observed in hv clones, such as ST23.^{2,8,9}

Given that antimicrobial resistance (AMR) and virulence determinants are commonly mobilized on plasmids, their occasional convergence within individual strains is not unexpected. The highly mosaic nature of *Kp* plasmids creates the risk of AMR and virulence determinants converging within a single plasmid. Such hv-AMR vectors could spread amongst *Kp* and confer widespread ability to cause serious infections with very limited treatment options. To our knowledge, only two such plasmids have been reported: pKpST147L, harbouring *iuc*, *rmpA*, *rmpA2* and several AMR determinants (*sul2*, *armA*, *sul1* and *mphA*) in an ST147 carriage isolate

also carrying a *bla*_{NDM-1} carbapenemase and isolated in London;⁴ and pKP70-2, harbouring the typical KpVP-1 virulence plasmid of ST23 (encoding *iuc*, *iro*, *rmpA* and *rmpA2*) with an additional insertion of an MDR transposon including *bla*_{KPC-2} carbapenemase in a K1 ST23 sputum isolate isolated in China.¹⁰

Here we report the complete genome sequences of two Kp ST15 carrying both MDR and virulence determinants, identified during a study of ESBL-producing Kp isolates from Norwegian hospitals.

Materials and methods

Ethics

The isolates presented here were collected and sequenced as part of a larger national study of Kp in Norwegian hospitals between 2001 and 2015 called NOR-KLEB. Ethical approval for NOR-KLEB, including the collection and sequencing of Kp isolates and collection of patient data, was provided by the regional ethics committee: REC west, application ID: 2017/1185.

Bacterial isolates

Isolate KP_NORM_BLD_2014_104014 (KP_104014) was cultured from a Romanian male in his eighties admitted to an Oslo hospital in 2014 with cholangiocarcinoma before developing bacteraemia. Isolate KP_NORM_BLD_2015_112126 (KP_112126) was cultured from a female in her seventies admitted to a Western Norway hospital in 2015 to treat a glioblastoma who developed neutropenic fever with pneumonia and bacteraemia. She had been hospitalized in Romania prior to admission in Norway. Antimicrobial susceptibility was determined by disc diffusion and broth microdilution, and hypermucoidy was assessed via the string test.

WGS and analysis

Paired-end reads (of 250 and 150 bp) were generated for *n*=12 ST15 Norwegian Kp isolates on the Illumina MiSeq and HiSeq platforms, respectively, and assembled with Unicycler v0.4.5 with SPAdes v3.11.1. In order to resolve the complete plasmid sequences for strains KP_104014 and

KP_112126, additional long read sequencing on a MinION R9.4 flow cell (Oxford Nanopore Technologies) was performed, and combined with the Illumina short reads to generate hybrid assemblies, using Unicycler as previously described,^{11,12} which were annotated using Prokka v1.11.¹³ Genotyping information including MLST, capsule type, AMR and virulence gene detection was extracted using Kleborate v0.3.0 (<https://github.com/katholt/Kleborate>) and used to curate the annotation of relevant loci in the plasmids.

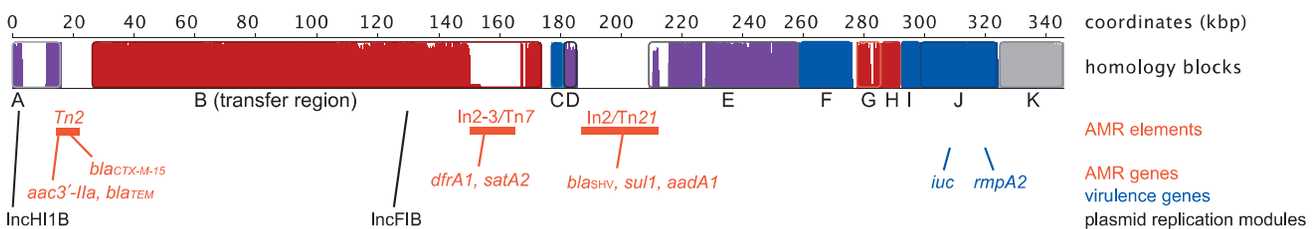
To place the hv-MDR strains in context, we performed comparative genomic analyses (described below) with an additional *n*=10 ST15 strains isolated between 2003 and 2015 from seven hospitals across Norway as part of the NOR-KLEB study (full results to be reported elsewhere), together with publicly available Illumina data identified from papers reporting Kp ST15 genome sequences (genomes and references are listed in Table S1, available as Supplementary data at JAC Online). Illumina read data for Kp genomes collected by the EuSCAPE European survey of carbapenemase-producing Enterobacteriaceae¹⁴ were downloaded and assembled using Unicycler and genotyped using Kleborate to identify ST15 isolates, and the ST15 read sets were included in the comparative analysis.

All read sets were mapped to the genome of KP_104014 using the RedDog v1b 10.2 pipeline (<https://github.com/katholt/RedDog>). An alignment of chromosomal single-nucleotide variants was extracted, recombinant regions were identified and filtered from the alignment using Gubbins v2.0.0¹⁵ and the final alignment was passed to RAXML v8.1.23¹⁶ to infer a core genome maximum likelihood phylogeny. From the mapping data we also extracted the coverage of the pKp104014_1 sequence, the coverage of the hv-MDR plasmid of KP_104014 and the presence of genes annotated in pKp104014_1 (presence defined as ≥95% of the length of the gene covered by five or more reads).

Nucleotide data accessions

Complete annotated sequences for the two novel genomes have been deposited in FigShare (doi: 10.6084/m9.figshare.7222889) and GenBank (BioSamples SAMEA5063299 and SAMEA5063300). The accession numbers for the mosaic plasmids described here are CP034046 (pKp104014_1) and CP034054 (pKp112126_1).

pKp104014_1, 345.8 kbp (ST15 blood isolate, Norway, 2014)



pKp112126_1, 299.2 kbp (ST15 blood isolate, Norway, 2015)

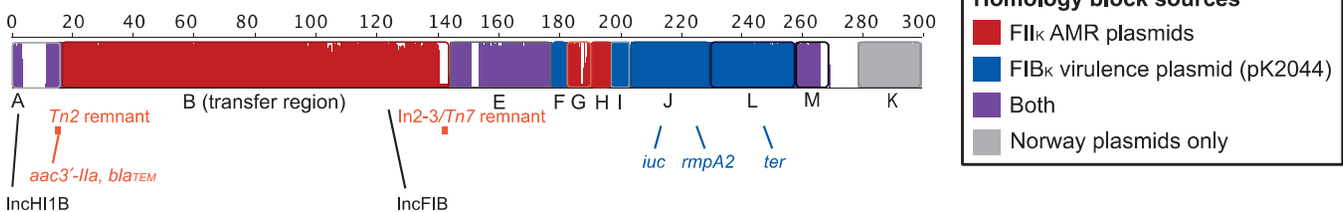


Figure 1. Map of novel mosaic hv-MDR plasmids, showing regions of homology with closely related AMR (pKp_Goe_579-1) and virulence (pK2044) plasmids, generated using Mauve. The location of known virulence genes (blue), as well as AMR genes and their associated mobile elements (red), and the plasmid replication modules are also indicated.

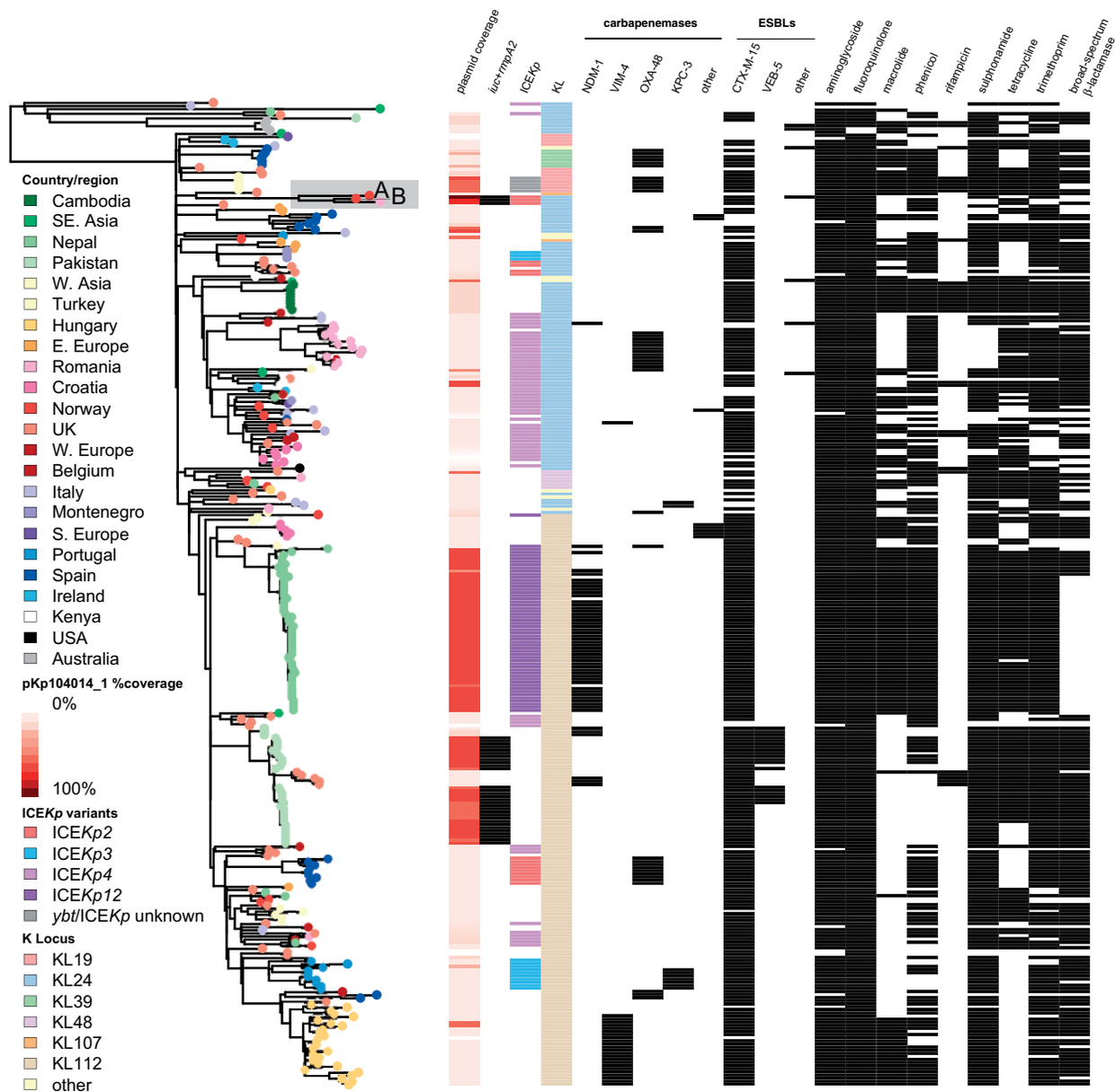


Figure 2. Recombination-free maximum likelihood phylogeny showing virulence and AMR properties of 318 ST15 isolates. Tips are coloured by the country of isolation (for $n \geq 4$ genomes) or geographical region as indicated; Norwegian convergent hv-MDR strains KP_NORM_BLD_2014_104014 and KP_NORM_BLD_2015_112126 are labelled as A and B, respectively, and highlighted in a grey box along with the closely related Romanian isolate. Columns are as follows: (i) percentage coverage of the *bla*_{CTX-M-15}/*iuc* plasmid pKp104014_1, determined by read mapping; (ii) presence of the aerobactin synthesis locus *iuc* and the hypermucooidy *rmpA2* gene (coloured in black); (iii) yersiniabactin ICEKp variants detected; and (iv) capsule K-locus (KL), followed by AMR determinants as labelled (coloured in black for presence).

Results and discussion

Isolate KP_104014 displayed resistance to cefotaxime, ceftazidime, ciprofloxacin, gentamicin, piperacillin/tazobactam and co-trimoxazole, and susceptibility to meropenem, colistin and tigecycline. The complete genome sequence resolved seven plasmids (Table S2), including a novel 346 kbp mosaic hv-MDR plasmid pKp104014_1, which harboured *bla*_{CTX-M-15} and seven additional AMR genes (Figure 1). Plasmid pKp104014_1 shares regions of homology with typical KpVP-1-type Kp IncFIB_K virulence plasmids,

such as pK2044 (40% coverage, including *iuc* and *rmpA2*), in addition to regions of homology to IncFII_K conjugative AMR plasmids (closest match: 246 kbp plasmid pKp_Goe_579-1, accession CP018313.1, from an ST147 Kp isolated in Germany, 59% coverage). This region (labelled as block B in Figure 1) is flanked by blocks present in both the virulence and AMR plasmids, probably facilitating homologous recombination forming the resulting mosaic plasmids. The IncFII_K regions include genes for conjugative transfer, suggesting that the plasmid may be self-transmissible. The plasmid harboured eight AMR genes mobilized via various elements

including: *bla*_{CTX-M-15} (mobilized by *ISEcp1*, flanked by 5 bp DRs); *bla*_{TEM-1} and *aac3*^{-IIa} (Tn2); *dfrA1* and *sat2* (In2-3/Tn7); *bla*_{SHV-5} (IS26); and *sul1* and *aadA1* (In2/Tn21) (Figure 1). A second copy of *bla*_{CTX-M-15} was also inserted into the chromosomal gene *phoE* (via *ISEcp1*, flanked by 5 bp DRs), and additional AMR genes (*aacA4*, *bla*_{OXA-1}, *bla*_{TEM-1} and *cat*) were carried on a 76 kbp IncFII plasmid, pKp104014_3 (Table S2).

Isolate KP_112126 displayed resistance to cefotaxime, ceftazidime, ciprofloxacin and gentamicin, intermediate susceptibility to piperacillin/tazobactam and tigecycline, and susceptibility to meropenem, colistin and co-trimoxazole. The complete genome sequence resolved four plasmids, including a mosaic 299 kbp hv-MDR plasmid pKp112126_1, with similarity to pKp104014_1 (99.99% nucleotide identity) including *iuc*, *rmpA2* and the IncFII_K transfer region. This plasmid lacks most of the AMR genes, although remnants of two AMR regions of pKp104014_1, namely one end of Tn2 (encoding *bla*_{TEM}, *aac3*^{-IIa}) and one end of In2/Tn7 (integrase only), were present (Figure 1). Plasmid pKp112126_1 also carried an additional region with homology to KpVP-1 virulence plasmids, including the *ter* locus encoding tellurite resistance (block L in Figure 1). *bla*_{CTX-M-15} was present in a distinct 90 kbp IncFII plasmid, pKp112126_3, which displayed homology with pKp104014_3 and *Shigella flexneri* plasmid R100 (accession AP000342.1). Carriage of multiple IncFII plasmids, which was detected in both strains, is unusual but has been reported previously.¹⁷

Both of the Norwegian hv-MDR isolates belonged to ST15 and carried the siderophore yersiniabactin (in genomic island ICEKp2) and the KL24 locus encoding capsular serotype K24. ST15 is a well-documented international ESBL-producing clone associated with nosocomial outbreaks worldwide, which frequently carries *bla*_{CTX-M-15}-encoding IncFII plasmids.^{18–21} To explore the relatedness of the Norwegian isolates to one another and to the wider ST15 *Kp* population, we constructed a recombination-filtered, core-genome maximum likelihood phylogeny including KP_104014, KP_112126, 10 additional ST15 isolates from Norway and 306 publicly available ST15 genomes from 29 other countries (Figure 2 and Table S1). The tree showed that the two Norwegian hv-MDR isolates were closely related to one another (77 SNPs, 0.001% nucleotide divergence) and to a urine isolate from Romania collected in 2013 (110 SNPs), but quite distant (>0.003% divergent) from the other Norwegian and global isolates.

Interestingly, both of the Norwegian hv-MDR plasmids were isolated from patients in Norway with epidemiological links to Romania (one of Romanian descent and one with recent travel history to Romania), suggesting that the convergence of AMR and virulence plasmids may have occurred in that country prior to importation and detection in Norway. The closely related Romanian isolate genome (ENA accession ERR1415588) carried *iuc* and *rmpA2*, and its reads covered 98% of the pKp112126_1 sequence and only 54% of the typical virulence plasmid pK2044 sequence. This is consistent with the presence of a mosaic plasmid in this isolate, although the available Illumina reads were not sufficient to resolve the full sequence of the Romanian plasmid containing *iuc*.

*bla*_{CTX-M-15} was present in most (87%) of the ST15 genomes, along with other AMR genes (see Figure 2 and Table S1). There were also multiple independent acquisitions of the ICEKp genomic island encoding yersiniabactin, affecting 48% of all ST15 isolates

including 50% of ESBL isolates (Figure 2). The only non-Norwegian ST15 isolates harbouring *iuc* were 30 isolates from Pakistan and the closely related Romanian isolate, all of which carried *iuc* and *rmpA2* loci in addition to *bla*_{CTX-M-15} and multiple other AMR genes. The convergence of AMR and virulence was noted in the original study reporting these genomes from Pakistan,²² however, it is not possible to determine from the draft genomes whether *iuc* is co-localized on the same plasmid as AMR genes. Mapping of all ST15 read sets to pKp104014_1 showed that *iuc*+ isolates from Pakistan and *iuc*- isolates from Nepal (alongside a small number of *iuc*- isolates from other countries) share many genes with the mosaic plasmid pKp104014_1 (55.7%–68.5% coverage for Pakistan isolates and 52.2%–0.2% coverage for Nepal isolates) (Figure 2 and Figure S1). This confirms that IncFII_K and IncFIB_K AMR and virulence plasmids circulate in South Asian *Kp* ST15 populations and could potentially fuse to form hybrid hv-MDR plasmids.

Concerningly, our findings reveal mosaic plasmids carrying both virulence determinants (*iuc* and *rmpA2*) and AMR determinants in ESBL-producing isolates of a well-established MDR *Kp* clone that has been associated with nosocomial infections and outbreaks worldwide. Although the plasmids uncovered here date back to 2014–15, it is not yet known whether they and/or other convergent plasmids are already widespread, since genomic surveillance data on *Kp* remain limited beyond CP outbreaks, and available studies rarely address virulence or utilize long reads to investigate linkage between AMR and virulence loci. The co-presence of these loci in a single plasmid vector, as in the previously reported ST147 and ST23 lineages,^{4,10} poses a substantial public health threat with the ability to spread AMR and virulence simultaneously, and highlights the need for surveillance of virulence alongside AMR before such strains become widespread.

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Transparency declarations

None to declare.

Author contributions

M. M. C. L., K. E. H. and I. H. L. conceived the study, performed data analyses and wrote the manuscript. K. L. W., R. R. W., A. F., K. E. H. and I. H. L. contributed additional data analysis and interpretation. A. F. and I. H. L. provided isolates. L. M. J. performed DNA extractions and Nanopore sequencing. All authors edited and approved the manuscript.

Supplementary data

Tables S1 and S2 and Figure S1 are available as Supplementary data at JAC Online.

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