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ORIGINAL RESEARCH

# Comparative genomics of five different resistance plasmids coexisting in a clinical multi-drug resistant *Citrobacter freundii* isolate

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**Background:** Plasmid-mediated multi-drug resistance (MDR) has been widely found in *Citrobacter freundii*. *C. freundii* P10159 was isolated from a human case of postoperative urinary tract infection in a Chinese teaching hospital.

**Methods:** The complete nucleotide sequences of five resistance plasmids pP10159-1, pP10159-2, pP10159-3, pP10159-4 and pP10159-5 from *C. freundii* P10159 were determined through high-throughput genome sequencing, and then compared with related plasmids sequences. Plasmid transfer, CarbaNP test of carbapenemase activity, and bacterial antimicrobial susceptibility test were performed to characterize resistance phenotypes mediated by these plasmids. **Results:** pP10159-1 carrying  $bla_{NDM-1}$  and pP10159-2 harboring  $bla_{IMP-4}$  plus *qnrS1* were almost identical to IncX3 plasmid pNDM-HN380 and IncN1 plasmid pP378-IMP, respectively. The  $bla_{KPC-2}$ -carrying plasmids pP10159-3, pHS062105-3 and pECN49-KPC were highly similar to each other, and constituted a novel group of plasmids belonging to an unknown incomparability group. The MDR plasmids pP10159-4 and pP10159-5 had the backbones highly similar to IncHI4 plasmid pNDM-CIT and type 2 IncC plasmid pR55, respectively, but their accessory resistance regions differed from pNDM-CIT and pR55, respectively. The five plasmids from the P10159 isolate contained a total of 24 different genes or gene loci, which contributed to resistance to 13 distinct antibiotic molecules or toxic compounds.

**Conclusion:** This is the first report of co-occurrence of five different resistance plasmids, with determination of their complete sequences. Data presented here provide a deeper insight into co-selection and maintenance of multiple plasmids and an extremely large number of resistance genes in a single bacterial isolate.

Keywords: Citrobacter freundii, multi-drug resistance, plasmids, mobile elements

## Introduction

*Citrobacter freundii*, a member of the family *Enterobacteriaceae*, is widely found in the environment as well as in the intestinal tract of humans and animals. *C. freundii* is generally considered a low-grade opportunistic pathogen that rarely causes infections, but it has been associated with a wide spectrum of infections of the central nervous system, the respiratory, gastrointestinal, urinary and respiratory tracts, the blood, and many other normally sterile sites in neonates and immunocompromised patients.<sup>1</sup>

Plasmid-mediated multi-drug resistance (MDR) has been widely found in *C. freundii*. This study disclosed the co-occurrence of five resistance plasmids pP10159-1, pP10159-2, pP10159-3, pP10159-4 and pP10159-5, containing a total of 24 different resistance markers, in a single clinical *C. freundii* isolate.

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# Materials and methods Bacterial strain

The use of human specimens and all related experimental protocols were approved by the Committee on Human Research of Southwest Hospital, and carried out in accordance with the approved guidelines. The indicated patient signed a written informed consent.

*C. freundii* P10159 was isolated in 2013 from a midstream urine specimen from an esophageal cancer patient with hospital-acquired postoperative urinary tract infection from a teaching hospital in Chongqing City, China. Bacterial species identification was performed using Bruker MALDI Biotyper (Bruker Daltonics) and 16S rRNA gene sequencing.<sup>2</sup> The sequence type (ST) of P10159 was determined based on the *C. freundii* multilocus sequence typing (MLST) scheme (https://pubmlst.org/cfreundii/). All PCR amplicons were sequenced on an ABI 3730 Sequencer.

# Genomic DNA sequencing and plasmid sequence assembly

Bacterial genomic DNA was isolated using a Qiagen large construct kit and sequenced from a paired-end library with a mate-pair library with average insert size of 5 kb (ranging from 2 to 10 kb) at a mean coverage 108, using a MiSeq sequencer (Illumina, CA, USA). Reads were trimmed to remove poor quality sequences. In order to get the complete plasmid sequences, DNA contigs were assembled based on their contig coverage using Newbler 2.8.<sup>3</sup> Gaps between contigs were filled using a combination of PCR and Sanger sequencing using an ABI 3730 Sequencer.

# Sequence annotation and genome comparison

Open reading frames and pseudogenes were predicted using RAST 2.0<sup>4</sup> combined with BLASTP/BLASTN searches against the UniProtKB/Swiss-Prot<sup>5</sup> and RefSeq<sup>6</sup> databases. Annotation of resistance genes, mobile elements and other features was carried out using CARD,<sup>7</sup> ResFinder,<sup>8</sup> ISfinder<sup>9</sup> and INTEGRALL.<sup>10</sup> Multiple and pairwise sequence comparisons were performed using MUSCLE 3.8.31<sup>11</sup> and BLASTN, respectively. Gene organization diagrams were drawn in Inkscape 0.48.1 (https://inkscape.org/en/).

# Plasmid conjugal transfer

Plasmid conjugal transfer experiments were carried out with the rifampin-resistant *Escherichia coli* EC600 being used as recipient and the P10159 strain as donor. Three milliliters of overnight cultures of each of donor and recipient bacteria were mixed together, harvested and resuspended in 80  $\mu$ L of Brain Heart Infusion (BHI) broth (BD Biosciences). The mixture was spotted on a 1 cm<sup>2</sup> hydrophilic nylon membrane filter with a 0.45  $\mu$ m pore size (Millipore) that was placed on a BHI agar (BD Biosciences) plate and then incubated for mating at 37°C for 12–18 h. Bacteria were washed from the filter membrane and spotted on Muller-Hinton (MH) agar (BD Biosciences) plates containing 1,000  $\mu$ g/mL rifampin together with indicated additional antibiotics for selecting an *E. coli* transconjugant carrying one of the following resistance markers: 4  $\mu$ g/mL meropenem for *bla*<sub>NDM-1</sub> (pP10159-1), *bla*<sub>IMP-4</sub> (pP10159-2) and *bla*<sub>KPC-2</sub> (pP10159-3); 10  $\mu$ g/mL chloramphenicol for *catB3* (pP10159-4); and 10  $\mu$ g/mL azithromycin for *mph*(A) (pP10159-5).

# Plasmid electroporation

To prepare competent cells for plasmid electroporation, 200 mL of overnight culture of *E. coli* TOP10 in Super Optimal Broth (SOB) at an optical density (OD<sub>600</sub>) of 0.4 to 0.6 was washed three times with electroporation buffer (0.5 M mannitol and 10% glycerol) and concentrated into a final volume of 2 mL. One microgram of plasmid DNA, which was isolated from the P10159 strain using a Qiagen Plasmid Midi Kit, was mixed with 100  $\mu$ L of competent cells for electroporation at 25  $\mu$ F, 200  $\Omega$  and 2.5 Kv. The resulting cells were suspended in 500  $\mu$ L of SOB and an appropriate aliquot was spotted on SOB agar plates containing the above indicated antibiotics for selecting of an *E. coli* electroporant carrying one of *bla*<sub>NDM-1</sub>, *bla*<sub>IMP-4</sub>, *bla*<sub>KPC-2</sub>, *catB3* and *mph*(A).

# Detection of carbapenemase activity

Activity of class A/B/D carbapenemases in bacterial cell extracts was determined via a modified CarbaNP test.12 Overnight bacterial cell culture in MH broth was diluted 1:100 into 3 mL of fresh MH broth and bacteria were allowed to grow at 37°C with shaking at 200 rpm to reach an OD<sub>600</sub> of 1.0 to 1.4. If required, ampicillin was used at 200  $\mu$ g/ mL. Bacterial cells were harvested from 2 mL of the above culture and washed twice with 20 mM Tris-HCl (pH 7.8). Cell pellets were resuspended in 500 µL of 20 mM Tris-HCl (pH 7.8) and lysed by sonication, followed by centrifugation at  $10,000 \times g$  at 4°C for 5 minutes. Fifty microliters of the supernatant (the enzymatic bacterial suspension) were mixed with 50 µL of substrate I to V, respectively, followed by incubation at 37°C for 2 hours. Substrate I: 0.054% phenol red plus 0.1 mM ZnSO<sub>4</sub> (pH7.8). Substrate II: 0.054% phenol red plus 0.1 mM ZnSO<sub>4</sub> (pH7.8) and 0.6 mg/µL imipenem. Substrate III: 0.054% phenol red plus 0.1 mM

ZnSO<sub>4</sub> (pH7.8), 0.6 mg/ $\mu$ L mg imipenem and 0.8 mg/ $\mu$ L tazobactam. Substrate IV: 0.054% phenol red plus 0.1 mM  $ZnSO_4$  (pH7.8), 0.6 mg/µL mg imipenem and 3 mM EDTA (pH7.8). Substrate V: 0.054% phenol red plus 0.1 mM ZnSO<sub>4</sub> (pH7.8), 0.6 mg/µL mg imipenem, 0.8 mg/µL tazobactam and 3 mM EDTA (pH7.8).

## Bacterial antimicrobial susceptibility test

Bacterial antimicrobial susceptibility was tested by the broth dilution method and interpreted as per Clinical and Laboratory Standards Institute (CLSI) guidelines.13

## Nucleotide sequence accession numb

The pP10159-1, pP10159-2, pP10159-3, pP10159pP10159-5 sequences were submitted to GenBank accession numbers MF072961 to MF072965, respect

### Results

### C. freundii co-harboring five resistance plasmids C. freundii

P10159 belonged to a novel ST252, with an allelic 108-62-71-7-1-1-57 corresponding to the seven h keeping genes aspC, clpX, fadD, mdh, arcA, dnaG, and High-throughput sequencing with the genomic DNA P10159 strain revealed the circularly closed sequences five plasmids pP10159-1, pP10159-2, pP10159-3, pP10 and pP10159-5, 42.8 kb to 228.7 kb in length with 53 predicted open reading frames (ORFs) (Figure S1 and Table 1). The modular structure of each plasmid was divided into the backbone regions and separate accessory modules, which were defined as acquired DNA regions associated with and bordered by mobile elements (Figure S1 and Table 1). Some of these accessory modules harbored drug resistance genes (Table 2).

pP10159-1, pP10159-2 and pP10159-3 could be transferred into E. coli through conjugation or electroporation, which generated the corresponding transconjugants NDM-EC600, IMP-EC600 and KPC-EC600 plus the respective electroporants NDM-TOP10, IMP-TOP10 and KPC-TOP10 (Table 3). This assay indicated that these three plasmids were conjugative. Repeated attempts failed to transfer pP10159-4 or pP10159-5 into E. coli through conjugation and electroporation.

P10159 and all the transconjugants and transformants were resistant to ampicillin, ceftazidime, meropenem and cefoxitin (Table 3). P10159 had class A + B carbapenemase activity, while the transconjugants and transformants harboring pP10159-1 or pP10159-2 had B activity and those harboring pP10159-3 had A activity (Table S1). These resistance phenotypes were consistent with the production

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Table I Major features of plasmids analyzed

Category	Plasmid										
	pP10159-1	I pNDM- HN380	рР10159-1 рИDM- рР10159-2 НИ380	рР378- IMP	pP10159-3	PP378- pP10159-3 pHS062105-3 pECN49- pP10159-4 IMP KPC	pECN49- KPC	pP10159-4	pNDM-CIT	pP10159-5	pR55
Incompatibility group Total length (bp) Total number of ORFs Mean G + C content, %	IncX3 54,034 62 49.0	54,035 62 49.0	IncNI 51,104 50.4 50.4	51,207 64 50.5		42,848 53 49.7	41,317 52 49.1	IncH14 228,709 270 47.0	288,920 314 47.7	Type 2 IncC 152,132 190 51.1	170,810 203 53.0
Lengun of the backbone (bp) Accessory modules	э4,/э1 54,/э. The bla <sub>NDM-1</sub> region <sup>#</sup> and ISKox3	J4,/J2	57,474 57,545 27,082 In823b <sup>#</sup> , △Tn6292 <sup>#</sup> and △Tn6296 <sup>#</sup> the IS <i>I</i> remnant	5,52 6292 <sup>#</sup> and ant	∠ <del>3</del> ,082 ∆Tn6296#	5 to	24,045	170,385 The MDR region", the bla <sub>CTKAN3</sub> region", ISLad2, ISKpn26-ISI <i>R</i> , ISCfi44-∆ISCfi9, IS903B and ISCfi6:ISEc21	206,301 The MDR region <sup>#</sup> , ⊕CP4- 6/57 <sup>#</sup> , ISKpn26, ISKpn34, ∆ISCfr9 and ISSen3	67,436 The MDR region#, ∆Tn6292-3' and two separate copies of ISEc52	The floR-ISCR2 region <sup>#</sup> and Tn6/87#
Notes: pPI0159-1, pP10159-2, pP10159-3, pP10159-4 and pP10159-5 from C freundii P10159 were sequenced in this work, while the other plasmids were derived fr of these five sequenced plasmids with their reference plasmid(s) was interpreted in the text. "These accessory modules contained resistance genes as listed in Table 2.	1-2, pP10159-3, nids with their 1	pP10159-4 an "eference plası	d pP10159-5 from mid(s) was interpr	C. freundii P eted in the t	10159 were sequext. #These acces	Jenced in this work, v ssory modules contai	while the other ned resistance	Notes: pPI0159-1, pPI0159-3, pPI0159-4 and pPI0159-5 from C. freundii PI0159 were sequenced in this work, while the other plasmids were derived from GenBank and used as references. Sequence comparison of each of these five sequenced plasmids with their reference plasmid(s) was interpreted in the text. "These accessory modules contained resistance genes as listed in Table 2.	GenBank and used a	s references. Sequence	comparison of each

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#### Table 2 Drug resistance genes in plasmids analyzed

Plasmid	Resistance	Resistance phenotype	Nucleotide position	Accessory or backbone
	gene		in plasmid	region located
pP10159-1	bla <sub>sHV-12</sub>	Beta-lactam resistance	9,324–10,184	The <i>bla</i> <sub>NDM-1</sub>
	ble	Bleomycin resistance	17,458–17,823	region
	bla <sub>NDM-1</sub>	Beta-lactam resistance	17,827–18,639	
oP10159-2	bla <sub>IMP-4</sub>	Beta-lactam resistance	4,295–5,035	In823b
	qnrS I	Quinolone resistance	27,662–28,318	∆ <b>T</b> n6292
P10159-3	Ыа <sub>крс-2</sub>	Beta-lactam resistance	7,737–8,618	∆ <b>T</b> n6296
pP10159-4	aacA4cr	Quinolone resistance	85,056–85,655	The MDR region
	bla <sub>oxA-1</sub>	Beta-lactam resistance	85,786–86,616	
	catB3	Phenicol resistance	86,754–87,386	
	arr3	Rifampicin resistance	87,471–87,923	
	qacE∆I	Quaternary ammonium compound resistance	88,146-88,493	
	sull	Sulfonamide resistance	88,487-89,326	
	bla <sub>CTX-M-3</sub>	Beta-lactam resistance	114,790–115,665	The <i>bla<sub>ctx-M-3</sub></i> region
	The ter locus	Tellurium resistance	116,056-136,305	The plasmid backbone
pP10159-5	bla <sub>sFO-1</sub>	Beta-lactam resistance	87,807–88,694	The MDR region
	chrA	Chromate resistance	98,473–99,678	-
	mph(A)	Macrolide resistance	103,297-104,202	
	aacC2	Aminoglycoside resistance	105,261-106,121	
	tmrB	Tunicamycin resistance	106,134–106,676	
	Ыа <sub>тем-1</sub>	Beta-lactam resistance	118,230-119,090	
	The mer locus	Mercuric resistance	120,698-124,660	
	qacE∆I	Quaternary ammonium compound resistance	128,336-128,683	
	sull	Sulfonamide resistance	128,677-129,516	
	armA	Aminoglycoside resistance	132,848-133,621	
	msr(E)	Macrolide resistance	135,920-137,395	
	mph(E)	Macrolide resistance	37,45 - 38,335	

Abbreviation: MDR, multi-drug resistant.

#### Table 3 Antimicrobial drug susceptibility profiles

Antibiotics	MIC (mg/L)/antimicrobial susceptibility														
	P10159	NDM-	NDM-	IMP-	IMP-	KPC-	KPC-	TOP10	EC600						
		EC600	TOPIO	EC600	TOP10	EC600	TOP10								
Ampicillin	>1,024/R	>1,024/R	>1,024/R	512/R	256/R	>1,024/R	>1,024/R	<4/S	<4/S						
Ceftazidime	>512/R	>512/R	>512/R	>512/R	512/R	64/R	16/R	<4/S	<4/S						
Imipenem	64/R	64/R	64/R	64/R	32/R	64/R	64/R	<1/S	<1/S						
Meropenem	64/R	16/R	8/R	4/R	4/R	32/R	4/R	<1/S	<1/S						
Cefoxitin	>1,024/R	I,024/R	512/R	>1,024/R	256/R	128/R	32/R	<8/S	<8/S						
Ciprofloxacin	64/R	<1/S	<1/S	2/I	2/I	<1/S	<1/S	<1/S	<1/S						
Amikacin	>1,024/R	<8/S	<8/S	<8/S	<8/S	<8/S	<8/S	<8/S	<8/S						
Azithromycin	>512/R	<4/S	<4/S	<4/S	<4/S	8/S	<4/S	4/S	4/S						
Trimethoprime	>32R	<0.25/S	<0.25/S	<0.25/S	<0.25/S	<1/S	<0.25/S	<0.25/S	<0.25/S						
Sulfamethoxazole	>608/R	<4.75/S	<4.75/S	<4.75/S	<4.75/S	19/S	<4.75/S	<4.75/S	<4.75/S						
Chloramphenicol	32/R	<8/S	<8/S	<8/S	<8/S	<8/S	<8/S	<8/S	<8/S						
Nitrofurantoin	16/S	16/S	<4/S	16/S	<4/S	32/S	<4/S	<4/S	8/S						
Minocycline	2/S	<1/S	4/S	<1/S	4/S	<1/S	4/S	4/S	<1/S						
Fosfomycin	<64/S	<64/S	<64/S	<64/S	<64/S	<64/S	<64/S	<64/S	<64/S						
Tigecycline	<1/S	<1/S	<1/S	<1/S	<1/S	<1/S	<1/S	<1/S	<1/S						
Colistin	<1/S	<1/S	<1/S	<1/S	<1/S	<1/S	<1/S	<1/S	<1/S						

Abbreviations: MIC, minimum inhibitory concentration; S, sensitive; R, resistant; I, intermediately resistant.

of one or all of NDM, IMP and KPC enzymes in the corresponding strains.

# Comparison of pP10159-1 with pNDM-HN380

pP10159-1 showed 99% nucleotide identity (with 100% query coverage) to nine  $bla_{\rm NDM}$ -carrying IncX3 plasmids, including the first fully sequenced pNDM-HN380<sup>14</sup> (Figure 1A). pP10159-1/pNDM-HN380 contained three resistance genes  $bla_{\rm NDM-1}$ , ble and  $bla_{\rm SHV-12}$ , which were all located in the  $bla_{\rm NDM-1}$  regions (Figure 1B). The  $bla_{\rm NDM-1}$  regions of pP10159-1/pNDM-HN380 were organized in order of a 4.1-kb truncated version of the composite transposon-like IS26- $bla_{\rm SHV-12}$ -IS26 unit,<sup>15</sup> a  $bla_{\rm NDM-1}$ -containing $\Delta$ Tn125 element derived from the ISAba125-flanked composite transposon Tn125,<sup>16</sup> IS3000 and a 583-bp Tn3 remnant (Figure 1B). An inversion of the truncated IS26- $bla_{\rm SHV-12}$ -IS26 unit in the  $bla_{\rm NDM-1}$  regions represented the only modular difference between pP10159-1 and pNDM-HN380 (Figure 1B).

### Comparison of pP10159-2 with pP378-IMP

pP10159-2 displayed >99% nucleotide identity (with >97% query coverage) to 16  $bla_{IMP}$ -carrying IncN1 plasmids, including pIMP-HZ1<sup>17</sup> and pP378-IMP<sup>18</sup> (Figure 2A). Although pIMP-HZ1 was the first fully sequenced  $bla_{IMP}$ -carrying IncN1 plasmid, pP378-IMP<sup>18</sup> was more appropriate as the reference for genomic comparison because it contained two relatively complete mobile elements, namely a class 1 integron In823 and a truncated Tn3-family unit transposon  $\Delta Tn6292$ ,<sup>19</sup> harboring  $bla_{IMP-4}$  and *qnrS1*, respectively. pP10159-2 and pP378-IMP shared the same In823 and also two resistance genes  $bla_{IMP-4}$  and *qnrS1*.

There were two major modular differences between pP10159-2 and pP378-IMP. First, the CUP (conserved upstream repeat)-controlled regulon<sup>18</sup> was composed of seven sequentially arranged operons CUPA, CUPB, CUP5, CUP4, CUP3, CUP2 and CUP1 in the backbone of pP10159-2, while the translocation of the CUP2 operon occurred in pP378-IMP (Figure 2B).

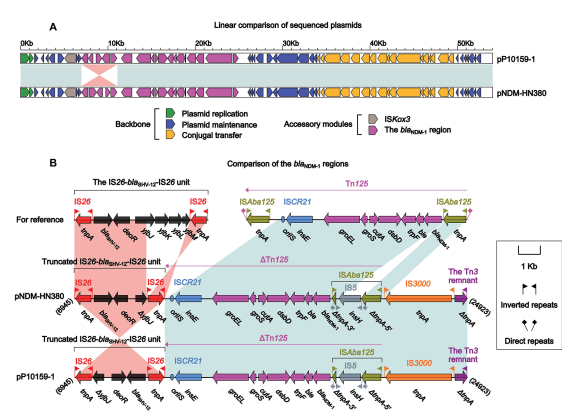


Figure I Comparison of pP10159-1 with pNDM-HN380.

Notes: Shown are linear comparison of the two sequenced plasmids pP10159-1 and pNDM-HN380 (**A**), and that of the *bla*<sub>NDM-1</sub> regions of these two plasmids (**B**). Genes are denoted by arrows. Genes, mobile elements and other features are colored based on function classification. Shading denotes regions of homology (>95% nucleotide identity). Numbers in brackets indicate the nucleotide positions within the corresponding plasmids.

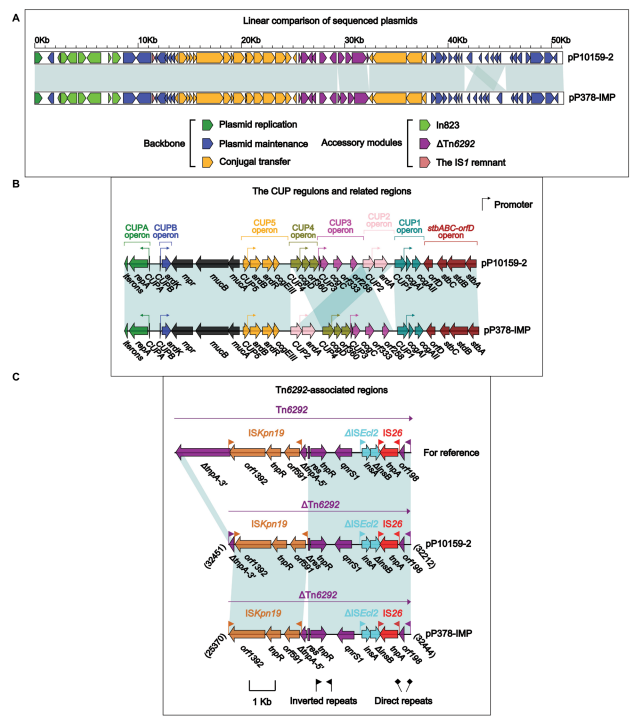


Figure 2 Comparison of pP10159-2 with pP378-IMP.

Notes: Shown are linear comparison of the two sequenced plasmids pP10159-2 and pP378-IMP (**A**), that of the CUP regions of these two plasmids (**B**), and that of the Tn6292-associated regions of these two plasmids (**C**). Genes are denoted by arrows. Genes, mobile elements and other features are colored based on function classification. Shading denotes regions of homology (>95% nucleotide identity). Numbers in brackets indicate the nucleotide positions within the corresponding plasmids.

Second, the core transposition module *tnpA* (transposase)-*res* (resolution site)-*tnpR* (resolvase) of Tn6292 was interrupted by IS*Kpn19*, breaking *tnpA* into separate  $\Delta$ *tnpA-3'* and  $\Delta$ *tnpA-5'*,<sup>18</sup> whereas distinct additional deletion events occurred within the *tnpA-res-tnpR* modules of the  $\Delta$ Tn6292 elements from P10159-2 and pP378-IMP (Figure 2C).

# Comparison of pP10159-3 with pHS062105-3 and pECN49-KPC

pP10159-3, pHS062105-3 and pECN49-KPC constituted a novel group of plasmids with almost identical backbones (>99% query coverage and >99% nucleotide identity). Their key backbone gene loci included *repA* for plasmid replication initiation, *parA* for plasmid partition, a toxin-antitoxin system *relEB* for post-segregational killing and a P-type IV secretion system for plasmid conjugal transfer (Figures S1C and 3A). Their RepA proteins belonging to the Rep\_3 superfamily (pfam10134) and matched the RepA proteins from *Klebsiella pneumoniae* and *Pantoea stewartii* with >96% query coverage and >99% amino acid identity; all these RepA proteins could not be assigned into any known incompatibility groups.

The insertion of the core  $bla_{\rm KPC}$  platform (ie, the Tn6376 to  $\Delta repB$  region) into the cryptic Tn3-family transposon Tn1722, truncating and splitting it into  $\Delta Tn1722-5'$  and  $\Delta Tn 1722-3'$ , generated Tn 6296 (Figure 3B) as initially observed in pKP048.<sup>20</sup> Each of pP10159-3, pHS062105-3 and pECN49-KPC carried a single accessory region  $\Delta Tn6296$ , which carried the  $bla_{KDC,2}$  gene serving as the sole resistance determinant of these plasmids (Figure 3B). The three  $\Delta Tn6296$  elements of pP10159-3, pHS062105-3 and pECN49-KPC had undergone deletions and insertions relative to the prototype Tn6296: 1)  $\Delta$ Tn1722-3' was lost from all the three  $\Delta Tn6296$  elements; 2) a 123-bp deletion at the 3'-terminal region of Tn6376 was found in pP10159-3; and 3) a 1,754-bp deletion within  $\Delta Tn 1722-5'$  as well as a 73-bp insertion within the variable number tandem repeat (VNTR) region of orf396 was identified in pECN49-KPC. Notably, the above variations within  $\Delta Tn6296$  accounted for the only modular difference between pHS062105-3, pP10159-3 and pECN49-KPC.

# Comparison of pP10159-4 with pNDM-CIT

The pP10159-4 backbone was highly similar (98% query coverage and 98% nucleotide identity) to the reference IncHI4 plasmid pNDM-CIT.<sup>19,21</sup> pP10159-4 and pNDM-CIT shared the core IncHI4 backbone gene loci,<sup>19</sup> including repHI4A and repHI4B for replication initiation, parAB and parMR for partition, and the tra1 and tra2 regions for conjugal transfer. There were two major modular differences between these two backbones (Figure S2): 1) the deletion of an 11-gene region [downstream of ISKpn26; containing the arsenic resistance (ars) locus] and a 4-gene region (downstream of ISKpn34) from pP10159-4 relative to pNDM-CIT, and 2) the deletion of a distinct 4-gene region (downstream of ISLad2) from pNDM-CIT compared with pP10159-4. These large-fragment deletions might result from the insertion of the corresponding IS elements ISKpn26, ISKpn34 and ISLad2. Of the accessory regions of pP10159-4, only the 25.3-kb MDR region and the 2.9-kb bla<sub>CTX-M-3</sub> region contained the resistance genes (Tables 1 and 2).

The MDR region (Figure 4) of pP10159-4 was organized as IS1R, a 14.1-kb 11-gene region encoding unknown functions, ΔIS1R, ISKpn19, ΔISPa38, orf402 (hypothetical protein), orf291 (DNA polymerase),  $\Delta Tn6308$  and ISKpn19. The unit transposon Tn6308 belonged to the Tn21 subgroup of the Tn3 family and was initially identified in plasmid pP10164-3.22 Tn6308 (Figure 4) had a hybrid backbone, which was composed of the core transposition module tnpAR-res of Tn1696 and the mercury resistance (mer) region of Tn21 and bordered by an intact 38-bp IRL and an IS5075-disrupted IRR (inverted repeat right); the class 1 integron In37 was inserted into the res site of Tn6308, truncating it from an original 120-bp fragment into an 83-bp remnant. Compared with the prototype class 1 integron, In37 in Tn6308 had a very complex mosaic structure and had undergone two major events: 1) an IS26-flanked composite transposon Tn6309, carrying the class C tetracycline resistance module tetA(C)-tetR(C), was inserted into intI1; and 2) an 8.1-kb region (8KR) was inserted at a site between 3'-conserved segment (3'-CS), and the Tn402 tni module, leading to truncation of 3'-CS and tni. 8KR included the chromate-resistance unit IRL<sub>chrd</sub>-chrAorf98 and the macrolide-resistance unit IS26-mph(A)-mrxmphR(A)-IS6100;<sup>23</sup> IRL<sub>chrA</sub> was further disrupted by IS5075 in 8KR. Tn6309, a 217-bp region containing attI1, and a 15.5-kb region (8KR-Δ*tniA-mer*:IS5075-IRR<sub>Tn6308</sub>) were not found in  $\Delta$ Tn6308 of pP10159-4 relative to Tn6308.

ISEcp1 was able to capture and arrange  $bla_{CTX-M-3}$ - $\Delta orf477$  at its downstream end while it moved,<sup>24</sup> while the  $bla_{CTX-M-3}$  region of pP10159-4 was a close derivative of this ISEcp1-base unit with truncation at its 3'-end region.

### Comparison of pP10159-5 with pR55

pP10159-5 shared 94% of its backbone with the reference type 2 IncC plasmid pR55,<sup>25</sup> with 99% nucleotide identity. All the type 2 IncC signature sequences, including two small insertions i1 and i2 as well as two genes *orf1847* and *rhs2*,<sup>26</sup> were found in pR55 and pP10159-5, with the only exception that a 246-bp deletion occurred at the 3'-terminus of *rhs2* in pP10159-5 (Figure S1E). pP10159-5 and pR55 possessed the core IncC backbone gene loci including *repA* for replication initiation, *parAB* and *parM* for partition, and the *tra1* and *tra2* regions for conjugal transfer (Figure S1E).

Linear sequence comparison of pP10159-5 and pR55 revealed five different regions (DIERs), namely DIER-1 to DIER-5 (Figure S3). DIER-1 was located between *orf564* and *orf312*, and manifested as the *traE* and *orf225* region composed of a 3'-terminal Tn6292<sup>18</sup> remnant (designated  $\Delta$ Tn6292; harboring no resistance genes) and eight con-

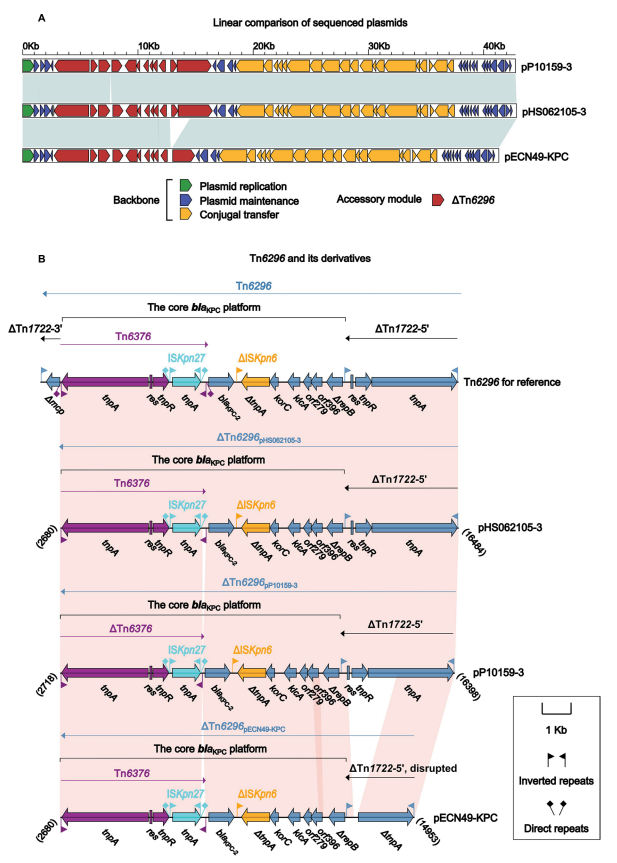


Figure 3 Comparison of pP10159-3 with pHS062105-3 and pECN49-KPC.

Notes: Shown are linear comparison of the three sequenced plasmids pP10159-3, pHS062105-3 and pECN49-KPC (**A**), and that of Tn6296 and its three derivatives from these thee plasmids (**B**). Genes are denoted by arrows. Genes, mobile elements and other features are colored based on function classification. Shading denotes regions of homology (>95% nucleotide identity). Numbers in brackets indicate the nucleotide positions within the corresponding plasmids.

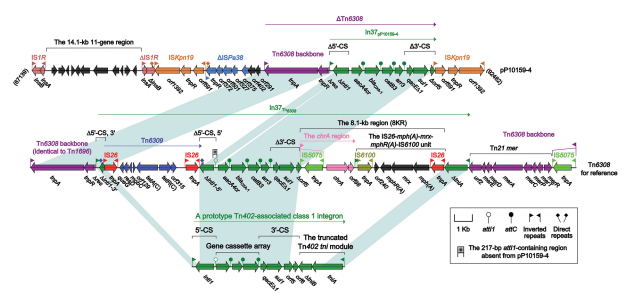


Figure 4 The MDR region from pP10159-4 and comparison with related regions.

Notes: Genes are denoted by arrows. Genes, mobile elements and other features are colored based on function classification. Shading denotes regions of homology (>95% nucleotide identity). Numbers in brackets indicate the nucleotide positions within the corresponding plasmids. Abbreviation: MDR, multi-drug resistant.

jugal transfer genes [found in another type 2 IncC plasmid pSRC119-A/C<sup>27</sup> but not in pR55] in pP10159-5; however, DIER-1 was manifested as the orf546 to sul2 region carrying the floR (florfenicol/chloramphenicol resistance)-ISCR2 region and five putative plasmid maintenance genes in pR55. DIER-2 existed as the orf861 to orf852 region (consisting of 34 putative plasmid maintenance genes) in pR55 and was deleted due to the insertion of ISEc52 at a site between traN and orf147 in pP10159-5; the ISEc52 insertion also resulted in the truncation of traN in pP10159-5. The loss of the orf861 to orf852 region, a part of the plasmid maintenance region as observed in pR55, did not destroy the plasmid maintenance of pP10159-5. A 58.2-kb MDR region (DIER-3) was inserted into rhs2 in pP10159-5 relative to pR55, leading to the truncation of rhs2 as well as the deletion of the downstream orf273-orf243 region. The Tn3-family unit transposon Tn6187 (DIER-4; 32.6 kb in length) was inserted into orf492 in pR55 relative to pP10159-5, splitting orf492 into two separate parts and meanwhile leaving 5-bp direct repeats (DRs; target site duplication signals of transposition) at both ends of Tn6187. The traF to orf186 region (DIER-5; containing not only conjugal transfer but plasmid maintenance genes) was located between uvrD and mobI in pR55, but replaced by a second copy of ISEc52 in pP10159-5; the ISEc52 insertion further led to the truncation of uvrD in pP10159-5. These five DIERs were associated with not only accessory modules but backbone regions, and the insertion and deletion events occurred within the two conjugal transfer regions

*tra1* and *tra2* of pP10159-5 would cause this plasmid to be non-conjugative.

All the resistance genes of pP10159-5 were harbored in the MDR region (Figure 5), which was sequentially organized as IS1R, a 3.6-kb 7-gene region of unknown functions, a 9.6-kb  $bla_{SF0-1}$  ( $\beta$ -lactam resistance) region as observed in pHD0149-2,<sup>28</sup> the *chrA*- and *mph*(A)-carrying 8KR fragment as found in In37 of Tn6308,<sup>22</sup> a 15-kb *aacC2* (aminoglycoside resistance)-*tmrB* (tunicamycin resistance) region that was generated from the insertion of a 7-kb phage-related region at a site between IS*Cfr1* and *bla*<sub>TEM-1</sub> ( $\beta$ -lactamase resistance) of the original *aacC2-tmrB* region as identified in pE11573,<sup>29</sup> a 7.4-kb Tn*21* remnant that was the 3'-terminal IRR to  $\Delta tniA$ region (containing the *mer* locus) of Tn*21*<sup>30</sup> with the disruption of IRR by IS*4321R* and a Tn*1548*-associated region.

Tn1548 was a IS26-flanked composite transposon lacking DRs at its ends, and had an IS26-In27-ISCR1- $\Delta$ ISEc28-armA (aminoglycoside resistance)-ISEc29-msr(E)-mph(E) (macrolide resistance)-orf543-repAciN-IS26 structure. There were a number of Tn1548-associated elements with the replacement of In27 by different class 1 integrons,<sup>31</sup> eg, In0 carrying no gene cassette array in pP10159-5 (Figure 5).

Remarkably, pP10159-5/pR55 belonged to the same incompatibility group and were genetically very closed related with respect to their plasmid backbones, but they carried totally different profiles of accessory modules that were inserted at different sites of the plasmid backbones; a similar observation was also found for pP10159-4/pNDM-CIT.

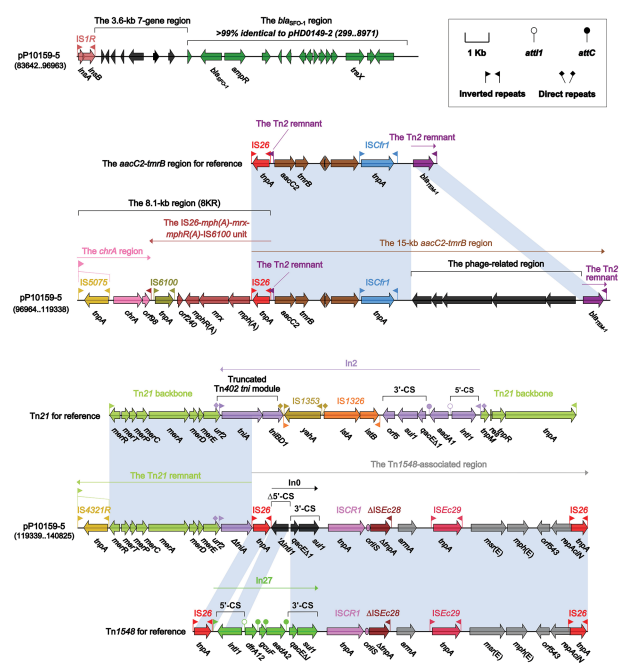


Figure 5 The MDR region from pP10159-5 and comparison with related regions.

**Notes:** Genes are denoted by arrows. Genes, mobile elements and other features are colored based on function classification. Shading denotes regions of homology (>95% nucleotide identity). Numbers in brackets indicate the nucleotide positions within the corresponding plasmids. **Abbreviation:** MDR, multi-drug resistant.

## Discussion

*C. freundii* isolates are resistant to cephalosporins due to inducible expression of chromosomally encoded AmpC  $\beta$ -lactamase,<sup>32</sup> but in general still remain susceptible to carbapenems. *C. freundii* can persist in the hosts for long periods, which likely facilitates the acquisition and accumulation of various resistance determinants under

high selective pressure within the hospital environments. Carbapenem-resistant *C. freundii* isolates expressing plasmid-encoding carbapenemases, such as KPC,<sup>33</sup> NDM,<sup>34</sup> and VIM,<sup>35</sup> have emerged during the past decade and, moreover, the coproduction of two or more carbapenemases, eg, KPC-2 + NDM-1<sup>36</sup> and NDM-1 + VIM-4 + OXA-181,<sup>37</sup> have also been identified in *C. freundii*, making the corresponding isolates highly resistant to  $\beta$ -lactams including carbapenems.

Coexistence of two to four different resistance plasmids has been observed in *C. freundii*.<sup>36,38,39</sup> We recently determined the complete nucleotide sequences of four plasmids p112298-KPC (belonging to an unknown incompatibility group; accession number KP987215),<sup>36</sup> p112298-NDM (IncX3 type; KP987216),<sup>36</sup> p112298-catA (IncHI2; KY270851) and p112298-tetA (type 1 IncC)<sup>40</sup> coexisting in a clinical *C. freundii* isolate 112,298. In this follow-up study, high-throughput genomic sequencing disclosed the co-occurrence of five resistance plasmids pP10159-1, pP10159-2, pP10159-3, pP10159-4 and pP10159-5 in a clinical *C. freundii* isolate P10159. This is the first report of co-occurrence of five resistance plasmids of different incomparability groups, with determination of their complete sequences, coexisting in a bacterial isolate.

There is the increasing prevalence of MDR C. freundii strains, which resulted from co-selection of genes encoding resistance to multiple antimicrobial classes, thereby leaving few or no options of antimicrobial treatment.<sup>36,38,39</sup> In this study, the five plasmids from C. freundii P10159 contain a total of 24 different genes or gene loci involved in the resistance to  $\beta$ -lactams including carbapenems, aminoglycosides, quinolones, macrolides, phenicols, rifampicin, sulfonamides, tunicamycin, bleomycin, quaternary ammonium compounds, chromate, mercury and tellurium, which are associated with several mobile elements including insertion sequences, integrons and transposons. Notably, the existence of redundant resistance genes made the P10159 strain extremely highly resistant to the corresponding classes of antibiotics, including carbapenems (*bla*<sub>NDM-1</sub>, *bla*<sub>IMP-4</sub> and *bla*<sub>KPC-2</sub>), aminoglycosides (armA and aacC2), quinolones (qnrS1 and aacA4cr), macrolides [mph(A) and mph(E)] and sulfonamides (two copies of sul1).

The co-selection and maintenance of multiple plasmids and an extremely large number of resistance genes in a single bacterial isolate reflect active and complex horizontal genetic transfer events that have taken place under selective pressures associated with many kinds of antibiotic molecules or toxic compounds.

## Acknowledgment

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## Disclosure

The authors report no conflicts of interest in this work.

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# Supplementary material

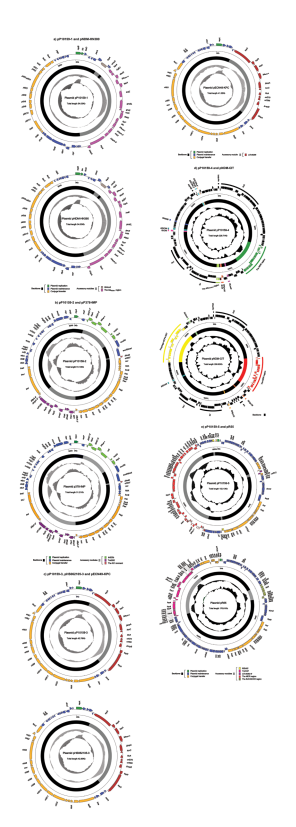


Figure SI Plasmid schematic maps.

Notes: The nine plasmids pP10159-1 and pNDM-HN380 (**A**), pP10159-2 and pP378-IMP (**B**), pP10159-3, pHS062105-3 and pECN49-KPC (**C**), pP10159-4 and pNDM-CIT (**D**), and pP10159-5 and pR55 (**E**) are included in the comparative analysis. Genes are denoted by arrows, and the backbone and accessory module regions are highlighted in black and color, respectively. The innermost circle presents GC-skew [(G-C)/(G + C)], with a window size of 500 bp and a step size of 20 bp. The next-to-innermost circle presents GC content.

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Figure S2 Linear comparison of pP10159-4 and pNDM-CIT sequences.

Notes: Genes are denoted by arrows. Genes, mobile elements and other features are colored based on function classification. Shading denotes regions of homology (>95% nucleotide identity).

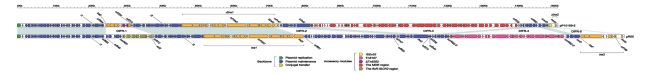


Figure S3 Linear comparison of pP10159-5 and pR55 sequences.

Notes: Genes are denoted by arrows. Genes, mobile elements and other features are colored based on function classification. Shading denotes regions of homology (>95% nucleotide identity).

Table SI Results of modified CARBA-NP test

Bacterial strain	Substrat	te	Detecting carbapenemase activity			
	I	II	111	IV	v	(Ambler class)
P10159	Red	Yellow	Yellow	Yellow	Red	A + B
NDM-EC600	Red	Yellow	Yellow	Red	Red	В
NDM-TOPI0	Red	Yellow	Yellow	Red	Red	В
IMP-EC600	Red	Yellow	Yellow	Red	Red	В
IMP-TOP10	Red	Yellow	Yellow	Red	Red	В
KPC-EC600	Red	Yellow	Red	Yellow	Red	Α
KPC-TOP10	Red	Yellow	Red	Yellow	Red	Α
TOPIO	Red	Red	Red	Red	Red	-
EC600	Red	Red	Red	Red	Red	_

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