A Multidrug Resistance Plasmid pIMP26, Carrying $\text{bla}_{\text{IMP-26}}$, $\text{fosA5}$, $\text{bla}_{\text{DHA-1}}$, and $\text{qnrB4}$ in Enterobacter cloacae

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IMP-26 was a rare IMP variant with more carbapenem-hydrolyzing activities, which was increasingly reported now in China. This study characterized a transferable multidrug resistance plasmid harboring $\text{bla}_{\text{IMP-26}}$ from one Enterobacter cloacae bloodstream isolate in Shanghai and investigated the genetic environment of resistance genes. The isolate was subjected to antimicrobial susceptibility testing and multilocus sequence typing using broth microdilution method, Etest and PCR. The plasmid was analyzed through conjugation experiments, S1-nuclease pulsed-field gel electrophoresis and hybridization. Whole genome sequencing and sequence analysis was conducted for further investigation of the plasmid. E. cloacae RJ702, belonging to ST528 and carrying $\text{bla}_{\text{IMP-26}}$, $\text{bla}_{\text{DHA-1}}$, $\text{qnrB4}$ and $\text{fosA5}$, was resistant to almost all $\beta$-lactams, but susceptible to quinolones and tigecycline. The transconjugant inherited the multidrug resistance. The resistance genes were located on a 329,420-bp IncHI2 conjugative plasmid pIMP26 (ST1 subtype), which contained $\text{trhK}/\text{trhV}$, $\text{tra}$, $\text{parA}$ and $\text{stbA}$ family operon. The $\text{bla}_{\text{IMP-26}}$ was arranged following $\text{intI1}$. The $\text{bla}_{\text{DHA-1}}$ and $\text{qnrB4}$ cluster was the downstream of $\text{Is}_{\text{CR1}}$, same as that in p505108-MDR. The $\text{fosA5}$ cassette was mediated by $\text{IS}_{4}$. This was the first report on complete nucleotide of a $\text{bla}_{\text{IMP-26}}$-carrying plasmid in E. cloacae in China. Plasmid pIMP26 hosted high phylogenetic mosaicism, transferability and plasticity.

Notoriously, extended and overuse of antibiotics have potentiated globally rapid emergence and spread of carbapenem-resistant Enterobacterales (CRE), posing a serious threat to clinical therapy and infection control. The major driving force for the diversification and dissemination of CRE has been confirmed as the horizontal transfer of plasmid-mediated carbapenem-hydrolyzing enzymes (i.e., carbapenemase) genes, among which the most prevalent and of particular clinical importance were $\text{bla}_{\text{KPC}}$, $\text{bla}_{\text{VIM}}$, $\text{bla}_{\text{IMP}}$, $\text{bla}_{\text{NDM}}$, and $\text{bla}_{\text{OXA-48}}$. $\text{IMP}$, one kind of metallo-$\beta$-lactamases (MBLs), can efficiently inactivate almost $\beta$-lactams except monobactam. $\text{IMP-1}$ was the first transferable MBL detected from Pseudomonas aeruginosa in Japan in 1991. Subsequently, the continuously clinical detection of $\text{bla}_{\text{IMP-1}}$ in different species isolates in Japan, as well as the discovery of $\text{IMP-2}$ in Italy and $\text{IMP-5}$ in Portugal, marked the beginning of the upcoming flourish of IMP MBLs. $\text{IMP-26}$ was first reported as an IMP-4 variant in Singapore in 2010 from a clinical carbapenem-resistant P. aeruginosa isolate by Koh TH et al. However, since then, there have been only sporadic reports on the $\text{IMP-26}$-production in Gram-negative bacilli, especially in Enterobacteriales. Notably, isolates expressing $\text{IMP-26}$ were found significantly more resistant to doripenem and meropenem than that expressing $\text{IMP-1}$.

Enterobacter cloacae was one member of the normal intestinal microflora of humans and animals, which has also assumed clinical importance and emerged as a major human pathogen causing hospital-acquired bacteremia, nosocomial pneumonia, urinary tract infections and so on. In the past decade, the emergence of IMP-producing E. cloacae has been extensively reported as a challenge to clinical therapy because of its rapid worldwide transmission. And in China, the most common IMP variants found in E. cloacae were IMP-8 and IMP-4. As for IMP-26-producing E. cloacae, it has been only reported in Chongqing, Shanghai and Beijing worldwide.
Our pilot study firstly reported two IMP-26-producing *E. cloacae* bloodstream isolates in Shanghai\(^1\). Considering the higher carbapenem-hydrolysing activities and emerging reports in China of IMP-26, we subsequently analyzed the transferability and full nucleotide sequence of the corresponding multi-drug-resistance plasmid pIMP26 in this study, which carried several important resistance determinants, such as *blaIMP-26*, *blaIMP-26*, *blaIMP-26*, *aacA4*, *qnrB4* and *fosA5*, conferring resistance to carbapenems, cephalosporins, aminoglycosides and fosfomycin, respectively.

**Methods**

**Isolate and antimicrobial susceptibility testing.** Isolate RJ702 was obtained from the blood of a female patient with uterine malignancy at Ruijin Hospital in April 2013. The carbapenem-resistant isolate was first isolated at day 28 after admission. The previous travel history of the patient was not documented.

The initial species identification of RJ702 was performed using MALDI-TOF MS (bioMérieux, Marcy-l’Étoile, France). The minimum inhibitory concentrations (MICs) of ceftriaxone, ceftazidime, cefotaxime, ceftepime, aztreonam, ciprofloxacin, levofloxacin, amikacin, gentamicin, piperacillin/tazobactam, cepoforazone/sulbactam, trimethoprim/sulfamethoxazole and tigecycline were determined using the broth microdilution method according to guidelines of the Clinical and Laboratory Standards Institute (CLSI M07-A9)\(^2\), while that of meropenem, imipenem and imipenem were determined using the Etest (bioMérieux, Marcy-l’Étoile, France). The susceptibility results were interpreted according to the guidelines of CLSI M07-A9\(^2\), while the breakpoint for tigecycline was according to that of European Committee on Antimicrobial Susceptibility Testing (EUCAST) V6.0\(^3\). *Escherichia coli* ATCC25922 was used as the quality control. PCR was performed to detect the “big five” carbapenemase genes (*blaKPC*, *blaNDM*, *blaIMP*, *blaVIM* and *blaOXA-48*).

**Multilocus typing (MLST).** A MLST scheme was used to assign *E. cloacae* to clonal lineages, including seven housekeeping genes (*dnaA*, *fusA*, *gyrB*, *leuS*, *pyrG*, *rplB*, and *rpoB*) as described by Miyoshi-Akiyama\(^4\). The combination of seven alleles can define the sequence types (STs) on the MLST website (http://pubmlst.org/ecloacae/).

**Plasmid conjugation, S1-nuclease pulsed-field gel electrophoresis (S1-PFGE), and southern hybridization.** The transferability of the resistance genes was assessed in broth culture using *E. coli* J53 Azt (sodiumazide-resistant) as the recipient. The transconjugants were selected on MacConkey agar containing sodiumazide (100 mg/L) and meropenem (2 mg/L) or ceftazidime (1 mg/L). PCR was employed to confirm the existence of *blaIMP-26* DNA plugs of the parental and transconjugant digested with S1-nuclease were prepared and separated by PFGE, and then transferred to positively charged nylon membrane (Roche Applied Science, Germany). The membrane was hybridized with digoxigenin-labeled *blaIMP-26* specific probes.

**DNA sequencing and genomics analysis.** Genomic DNA of *E. cloacae* RJ702 was isolated using ChargeSwitch® gDNA Mini Bacteria Kit (Life Technologies, Carlsbad, CA, USA) and sequenced by a combination of PacBio RSII (Pacific Biosciences, Menlo Park, CA, USA) and Illumina Hiseq X10 (Illumina, San Diego, CA, USA) sequencing platforms. The assembly was produced firstly using a hybrid *de novo* assembly solution modified by Koren, in which a *de-Brujin* based assembly algorithm and a CLR reads correction algorithm were integrated in “PacBioToCA with Celeras Assembler” pipeline\(^5\). The final assembly generated a circular genome sequence with no gap existed. The precise species identification was established based on average nucleotide identity (ANI) between RJ702 and other type strains of *E. cloacae* subsp. using Orthologous ANI Tool (OAT) recommended by Lee I et al.\(^6\). Annotation of the genomic sequence and alignment with other similar sequences were carried out using the BLAST Ring Image Generator (BRIG)\(^7\) and SnapGene program v4.3.2. Open reading frames (ORFs) were identified using Glimmer version 3.02 (http://cbcb.umd.edu/software/glimmer/). ORFs less than 300-bp were discarded. Insertion elements and resistance genes were identified using ISFinder (https://www.is.biotoul.fr) and ResFinder (https://cge.cbs.dtu.dk/services/ResFinder). PlasmidFinder (https://cge.cbs.dtu.dk/services/PlasmidFinder) and pMLST (https://cge.cbs.dtu.dk/services/pMLST/) were employed to detect and type the plasmids. BLAST (http://blast.ncbi.nlm.nih.gov/Blast.cgi) was used to identify related plasmids carrying *blaIMP-26* to guide PCR-based gap closure and Sanger sequencing to assemble contigs into complete plasmids.

**Nucleotide sequence accession number.** The completely annotated sequence of pIMP26 in *E. cloacae* RJ702 has been deposited in GenBank database (Accession Number: MH399264).

**Ethics approval and informed consent.** The study was approved by Ruijin Hospital Ethics Committee (Shanghai Jiao Tong University School of Medicine), and the Review Board exempted the requirements for informed consent as this study only focused on bacteria.

**Results**

**Precise species identification.** RJ702 was initially identified as *E. cloacae* or *E. asburiae* by MALDI-TOF MS. Additionally, RJ702 showed 95.2874% ANI value with *E. cloacae* ECNIH2 (NZ_CP008823) and only 87.8944% with *E. cloacae* subsp. *E. cloacae* ATCC35953 (NZ_CP011863). The ANI values of RJ702 with *E. cloacae* ATCC13047 (NC_014121) and *E. cloacae* MJ48 (NC_018079) were 89.0906% and 87.9528% respectively. Therefore, RJ702 was precisely identified as *E. cloacae* subsp. *E. cloacae*.

**Antimicrobial resistance and MLST.** *E. cloacae subsp. cloacae* RJ702 belonging to ST528, exhibited resistance to cephalosporins, monobactam, carbapenems, β-lactam/β-lactamase inhibitor combinations (only cepoforazone/sulbactam), aminoglycosides, trimethoprim/sulfamethoxazole, and tigecycline. The transconjugant inherited resistance to these antibiotics (Table 1).
Genome sequencing of RJ702. Whole genome sequencing generated 1,458,457 single reads and 4.70 Gb clean data total bases, which were de novo assembled to 184 contigs (75 > 1,000 bp; N50: 269,528 bp; N90: 45,440 bp). The bases in all contigs of RJ702 was 5.01 Mb with a 54.69% G + C content. The size of chromosome was 4,303,224 bp, and the bases in all contigs of two plasmids in RJ702 were 329,420 bp (pIMP26) and 78,322 bp respectively. PlasmidFinder presented that plasmid pIMP26 hosted two replicons, of which IncHI2 was 327 bp and IncHI2A was 630 bp; while the other plasmid hosted none.

Backbones of pIMP26. Plasmid pIMP26 was a 329,420-bp closed circular DNA sequence with an average G + C content of 48.24% (Fig. 2). It hosted two repB replicons, and both belonged to IncHI2 prototype (ST1 subtype). BLAST searches indicated the backbone regions of pIMP26 highly similar to a 324,503-bp IncHI2 plasmid pIMP26. It shared high similarity with p505108-MDR from a blaTEM-1B producing E. cloacae type). BLAST searches indicated the backbone regions of pIMP26 highly similar to a 324,503-bp IncHI2 plasmid pIMP26 (Fig. 2). Annotation of the finished sequence data revealed that pIMP26 contained 381 ORFs, including repB, traB, trhK/trhV (for stabilizing mating pairs during plasmid conjugation), four transfer gene clusters (locus 1-4), and two IS elements (IS6100, IS6100-cr1), transposons (Tn3 family, etc.) and integrons (intI1, intI1), and contained multiple resistance genes (blaIMP-26, fosA5, qnrB4, aac(6’)-Ib3, aac(6’)-IIc, aacA4, aph(6)-IId, strA, mhp(A), ere(A), catA2, tet(D), dfrA18, blaSHV-12), two copies of blaTEM-1BP, and five copies of sul1). According to BLAST searches (Fig. 3), the blaIMP-26 region was sequentially arranged as intI1, blaIMP-26, ltrA, qacEΔ1 and sul1 (position: 2567–7765), same as the blaIMP-26 cluster in pIMP-4 from an IMP-4-producing E. cloacae isolate in Taiwan (87% query coverage and 99% nucleotide identity) (Fig. 2). Annotation of the finished sequence data revealed that pIMP26 contained 381 ORFs, including repB (position: 32558–33613 and 47056–47931, for plasmid replication initiation), trhK/trhV (for stabilizing mating pairs during plasmid conjugation), four transfer gene clusters (locus traB, traN, traA and traM, encoding the conjugative apparatus), the parA family operon (for the replication/partitioning system with repB), and the sbcA family operon (for plasmid stability).

Resistances regions in pIMP26. Plasmid pIMP26 was rich in mobile genetic elements, including IS elements (IS4, IS6, IS26, ISCR1, etc.), transposons (Tn3 family, etc.) and integrons (intI1, intI1), and contained multiple resistance genes (blaIMP-26, fosA5, qnrB4, aac(6’)-Ib3, aac(6’)-IIc, aacA4, aph(6)-IId, strA, mhp(A), ere(A), catA2, tet(D), dfrA18, blaSHV-12), two copies of blaTEM-1BP and five copies of sul1). According to BLAST searches (Fig. 3), the blaIMP-26 region was sequentially arranged as intI1, blaIMP-26, ltrA, qacEΔ1 and sul1 (position: 2567–7765), same as the blaIMP-26 cluster in pIMP-4 from an IMP-4-producing E. cloacae isolate in Taiwan (87% query coverage and 99% nucleotide identity) (Fig. 2). Annotation of the finished sequence data revealed that pIMP26 contained 381 ORFs, including repB (position: 32558–33613 and 47056–47931, for plasmid replication initiation), trhK/trhV (for stabilizing mating pairs during plasmid conjugation), four transfer gene clusters (locus traB, traN, traA and traM, encoding the conjugative apparatus), the parA family operon (for the replication/partitioning system with repB), and the sbcA family operon (for plasmid stability).

Conjugation, S1-PFGE, and southern hybridization. The transconjugant RJ702-1 was obtained by plasmid conjugation experiments. S1-PFGE revealed that RJ702 harbored two plasmids (~320-kb and ~70-kb), and RJ702-1 inherited both. Southern hybridization analysis revealed blaIMP-26 located on the ~320-kb plasmid (pIMP26) (Fig. 1).
Discussion

The undesirable antibiotic resistance (especially carbapenem-resistance) has appeared and disseminated rapidly in Gram-negative bacilli, which was attributed largely to the acquisition of multiple resistance genes by horizontal plasmid-mediated genes transfer. Our study was to map the genetic environment of a novel multi-drug-resistance plasmid pIMP26, in order to provide a new insight for the potential spread of blaIMP-26 and fosA5 or correlations between genetic diagnosis and clinical treatment.

Firstly, the backbone of pIMP26 was blasted with different plasmids in BLAST. The origins of functional modules in pIMP26, such as multiple antibiotic resistance determinants, stably conjugal transfer (tra and trh family), mobile elements and plasmid maintenance (stb family) (Fig. 2), represented a strong transferability, stability and plasticity of this plasmid. IncHI2 was one of the most prevalent broad-host-range plasmid families carrying different resistance determinants simultaneously in Enterobacterales. As previously reported on E. cloacae, most β-lactamase-encoding genes (blaSHV-12, blaCTX-M-15, blaNDM-1, blaIMP-4, etc.) were also located on IncHI2 plasmids (subtype ST1) of 290~340-kb in size, and our study also fit it. It should be noted that the similar backbone shared by pIMP26 and other plasmids (Fig. 2) in clinical isolates of E. cloacae, K. pneumoniae and S. enterica from different areas strongly suggested that inter-species genetic exchange also occurred, thus broadening the host range and dissemination of combined cargo genes. Besides, pIMP26 contained a wide variety of transposable elements carrying known antibiotic resistance genes. Tn3 family transposon was the medium of TEM genes and fosA5 was also located in Tn3 in pIMP26 (Figs 2 and 3). The archetype of Tn3 was known as some of the earliest unit transposons identified in Gram-negative bacilli. Tn3 family members demonstrated transposition immunity, but homologous and/or res-mediated recombination between related elements can occur, creating hybrid elements. And this would explain multiple Tn3-mediated resistance elements in pIMP26 in this study. However, further study is definitely needed to characterize the mechanisms behind the transfer or recombination of Tn3.

IMP-26, firstly found in P. aeruginosa in Singapore, was differed from IMP-4 at position 145 (G to T change); the translated amino acid sequence differed from IMP-4 at residue 49 (phenylalanine for valine). Blast searches indicated that the genetic structure surrounding blaIMP-26 has only revealed in a study from Vietnam up to now, containing intI1-blaIMP-26-qacG-aac(6′)-Ib-orf3-orf4 (Fig. 2), and our study was the first time focusing on the complete nucleotide of the plasmid carrying blaIMP-26. Interesting was the blaIMP-26 region in pIMP26 different from that found in Vietnam (though both located on intI1); but same as the blaIMP-4 cluster of pIMP-4 in

Figure 1. (A) The S1-PFGE profile of E. cloacae RJ702 and its transconjugant RJ702-1. M, Salmonella enterica serotype Braenderup H9812 was digested with XbaI as a molecular size marker. (B) The Southern blotting profile of E. cloacae RJ702 and its transconjugant RJ702-1 with blaIMP-26 specific probes.
Shanghai (Genbank ID: FJ384365) (Fig. 3). It prompted that the \textit{bla}_{IMP-26} detected in our study maybe originated from \textit{bla}_{IMP-4} or the genetic mutation may occur during transfer of \textit{bla}_{IMP-26} cassette.

The prevalence and dissemination of \textit{fosA5} have probably been underestimated\textsuperscript{36}. Previous study once found IS10 playing an important role in the mobilization of \textit{fosA5}\textsuperscript{27}. However, the upper half consistent with pHKU1 in

\begin{figure}
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\includegraphics[width=\textwidth]{figure2}
\caption{Circular map of plasmid pIMP26. The two inner circles represented the G+C content plotted against the average G+C content of 48.24\% (black circle) and GC skew information (green and purple circles). Circles in different colors represented different plasmids (details in the legend), and the Genbank numbers were as follows: pIMP26 (MH399264), p505108-MDR (KY978628), pCNR48 (LT994835), pEC-IMPQ (EU855788), pIMP4-SEM1 (FJ384365), and pGMI14-002 (CP028197). The location of discussed resistance genes and \textit{intI} were also demonstrated on the outer cyan-blue circle. The annotation of the genetic components were added manually using the Microsoft PowerPoint 2016 program.}
\end{figure}

\begin{figure}
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\includegraphics[width=\textwidth]{figure3}
\caption{Plasmid accessory resistance regions. The comparison of linear DNA against the corresponding regions in different plasmids. The resistance genes were indicated by orange arrows and the insertion sequences are indicated by purple arrows. Shading regions denoted regions of homologous (>95\% nucleotide identity).}
\end{figure}
pIMP26 indicated that IS4 might also related to its mobilization\(^\text{36}\) (Fig. 3). Plasmid carrying bla\(_{OHEA}\) was usually reported also carrying qnrB4, bla\(_{DSTV-1}\)\(^\text{38}\). This suggested that the cassette in common of qnrB4 and bla\(_{OHEA}\)\(^\text{1}\) (Fig. 3) (including that in pIMP26) was derived from the same immediate ancestor. The qnrB4-bla\(_{OHEA}\)-containing region of pIMP26 was located after the 3’ conserved sequence (3’-CS) of intI1 (Fig. 3), containing aac(6’)-Ic, qacE\(_\Delta I\) and sulI. Besides, an insertion sequence common region 1 (ISCR1) was identified downstream of sulI. ISCR1 could mobilize the nearby sequence and a truncated 3’-CS from one integron to the 3’-CS of another integron through rolling-circle transposition, and provide a promoter for the expression of nearby genes\(^\text{39}\); this may lead to the co-carriage of multiple resistant genes in one plasmid and the multi-drug resistance of clinical isolates.

Interestingly, our study showed the qnrB4- and aac(6’)-Ib3-harboring RJ702 susceptible to quinolones (MIC = 0.5 or 0.25). We speculated that it was due to the absence of other mechanisms of chromosomal resistance (e.g., alterations in type II topoisomerases) in RJ702 other than plasmid-mediated quinolone resistant (PMQR) genes. Researchers found that PMQR mechanism caused only low-level quinolone-resistance on its own, which may not exceed the clinical breakpoints of susceptibility for quinolones but facilitated selections of higher-level resistance and posed threats to the treatment of infections by microorganisms hosting PMQR genes\(^\text{40}\), which could validate our speculation and underline the necessity of monitoring on PMQR genes.

This is the first report on the entire structure of bla\(_{IMP-26}\)-carring plasmid. To some extent, our study evidenced the increasing clinical significance of IncHI2 replicons as resistance genes’ reservoirs and provided insights on the possibilities of further spread in China and highlighted the needs for intensive surveillance and precautions.

Conclusions

We firstly reported here the complete nucleotide sequence of a plasmid carrying bla\(_{IMP-26}\), which was an IncHI2 replicon simultaneously encoding multidrug resistance determinants, including β-lactam (bla\(_{IMP-26}\) blad\(_{HHA}\)-Ib3, blad\(_{DSTV-1}\), aminoglycoside (aac(6’)-Ic, aacA4, aph(6)-IIc, etc.), fluoroquinolone (qnrB4, aac(6’)-Ib3) and fosfomycin (fosA5) resistance genes. New genetic context of fosA5 was also characterized. The novel plasmid with multi-insertion of different resistant components and stable inheritance emphasized controlled use of clinical antibiotics to prevent selective pressure aggravating the emergence and dissemination of multi-drug resistance.

Data Availability

The datasets used and/or analyzed during the current study are available from the corresponding author on reasonable request.

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Conception and design of study: L.H. and J.S., S.W. Acquisition of data (laboratory or clinical): S.W., K.Z., S.X., F.G. and X.L. Analysis and interpretation of data: S.W., K.Z. and L.X. Contribution of reagents/materials/analysis tools: L.H. and Y.N. Drafting of article and/or critical revision: S.W., K.Z., L.H. and J.S. Final approval of the submitted manuscript: All.

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