Dissemination of KPC-2-Encoding IncX6 Plasmids Among Multiple Enterobacteriaceae Species in a Single Chinese Hospital

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Forty-five KPC-producing Enterobacteriaceae strains were isolated from multiple departments in a Chinese public hospital from 2014 to 2015. Genome sequencing of four representative strains, namely Proteus mirabilis GN2, Serratia marcescens GN26, Morganella morganii GN28, and Klebsiella aerogenes E20, indicated the presence of bla\textit{KPC}-2-carrying IncX6 plasmids pGN2-KPC, pGN26-KPC, pGN28-KPC, and pE20-KPC in the four strains, respectively. These plasmids were genetically closely related to one another and to the only previously sequenced IncX6 plasmid, pKPC3_SZ. Each of the plasmids carried a single accessory module containing the bla\textit{KPC}-2/3-carrying 1\textit{Tn}_{6296} derivatives. The 1\textit{Tn}_{6292} element from pGN26-KPC also contained \textit{qnrS}, which was absent from all other plasmids. Overall, pKPC3_SZ-like \textit{bla\textit{KPC}}-carrying IncX6 plasmids were detected by PCR in 44.4% of the KPC-producing isolates, which included \textit{K. aerogenes}, \textit{P. mirabilis}, \textit{S. marcescens}, \textit{M. morganii}, \textit{Escherichia coli}, and \textit{Klebsiella pneumoniae}, and were obtained from six different departments of the hospital. Data presented herein provided insights into the genomic diversity and evolution of IncX6 plasmids, as well as the dissemination and epidemiology of \textit{bla\textit{KPC}}-carrying IncX6 plasmids among Enterobacteriaceae in a hospital setting.

Keywords: plasmid, IncX6, genomics, epidemiology, \textit{bla\textit{KPC}}

INTRODUCTION

Klebsiella pneumoniae carbapenamase (KPC), a class A β-lactamase, can hydrolyze almost all β-lactams, including carbapenems (Bush and Fisher, 2011). At least 31 variants (KPC-2 to KPC-32; KPC-1 is essentially identical to KPC-2) of the KPC enzyme have been identified to date. Two Tn3-family unit transposons, Tn4401 and Tn6296, which are genetically divergent from each other, represent the two major prototype genetic platforms carrying \textit{bla\textit{KPC}} genes (Wang et al., 2015). Tn4401 and its derivatives are frequently identified in KPC-encoding plasmids of different incompatibility groups in bacterial isolates from European and American countries, but are rarely

1https://www.ncbi.nlm.nih.gov/pathogens/beta-lactamase-data-resources/
found in isolates from China (Feng et al., 2017). The \( \text{bla}_{\text{KPC}} \) genetic environment in isolates from China is predominantly associated with \( \text{Tn}6296 \) and its derivatives (Wang et al., 2015). Plasmids belonging to incompatibility group X (IncX) are 30–80 kb in size and were initially discovered in the pre-antibiotic era (Datta and Hughes, 1983). IncX plasmids have a narrow host range and are mainly circulated among \textit{Enterobacteriaceae} species (Norman et al., 2008). The backbones of all IncX plasmids have a \( \text{pir}–\text{parA}–\text{hns}–\text{hha}–\text{topB}–\text{pilX} \) (\( \text{rivB} \)–\( \text{actX} \)–\( \text{taxC} \) (\( \text{rlx} \)–\( \text{taxA} \)) (\( \text{dr} \)) organization, but are quite divergent with respect to nucleotide and amino acid sequences similar to \( \text{KPC} \). Comparative genomic analysis has shown that IncX plasmids can be phylogenetically grouped into seven major IncX subgroups, IncX1 to IncX6 (Du et al., 2016), along with another IncX6 subgroup (Bustamante and Iredell, 2017) that is re-designated herein IncX7.

\( \text{bla}_{\text{KPC}}/2 \)-harboring plasmids have been identified among the IncX3, IncX5, and IncX6 subgroups, including over a dozen plasmids belonging to subgroup IncX3 [e.g., pKpS90 (GenBank accession number \( \text{JX}461340 \)] (Kassis-Chikhani et al., 2013) and pMNCRE44_5 (GenBank accession number \( \text{CP}010881 \)) (Hargreaves et al., 2015), two IncX5 plasmids [pKPC_CAV1492 (GenBank accession number \( \text{CP}011639 \)) and pBR31567 (GenBank accession number \( \text{JX}193302 \))] (Chen et al., 2013), and a single IncX6 plasmid [pKPC3_SZ (GenBank accession number \( \text{KU}032800 \)] (Du et al., 2016)]. Interestingly, \( \text{bla}_{\text{KPC}} \)-harboring plasmids have not been found among the other IncX subgroups.

This study provides evidence for the dissemination of genetically highly similar \( \text{KPC}-2 \)-encoding IncX6 plasmids among at least six \textit{Enterobacteriaceae} species collected in a Chinese public hospital from 2014 to 2015. The complete nucleotide sequences of plasmids pGN2-KPC, pGN26-KPC, pGN28-KPC, and pE20-KPC, extracted from strains belonging to four representative species, were determined to be genetically closely related to the IncX6 reference plasmid pKPC3_SZ. In addition, all five plasmids carried a single accessory region that harbored the \( \text{bla}_{\text{KPC}}/2 \)-gene.

**MATERIALS AND METHODS**

**Bacterial Identification**

Bacterial species identification was performed on the basis of 16S rRNA gene sequencing (Frank et al., 2008). The major plasmid-borne carbapenemase and extended-spectrum \( \beta \)-lactamase genes were screened for by PCR (Chen et al., 2014). All PCR amplicons were sequenced using an ABI 3730 Sequencer (Life Technologies, Carlsbad, CA, United States) using the primers used for PCR.

**Plasmid Transfer**

Plasmid conjugal transfer experiments were carried out using rifampin-resistant \textit{Escherichia coli} strain EC600 (LacZ-, \text{NalR}, \text{RifR}) as the recipient and each of \textit{Proteus mirabilis} GN2, \textit{Serratia marcescens} GN26, \textit{Morganella morganii} GN28, and \textit{Klebsiella aerogenes} E20 as the donor (Feng et al., 2016). Aliquots (3 ml) of overnight cultures of each of the donor and recipient strains were mixed together, harvested, and resuspended in 80 \( \mu \text{l} \) of brain heart infusion broth (BD Biosciences, Franklin Lakes, NJ, United States). The mixture was spotted onto a 1-cm² hydrophilic nylon membrane filter with a 0.45-\( \mu \text{m} \) pore size (Millipore, Billerica, MA, United States) placed onto the surface of a brain heart infusion agar (BD Biosciences, Franklin Lakes, NJ, United States) plate. Plates were incubated for mating at 37°C for 12–18 h. Bacteria were washed from the filter membrane and spotted on Mueller-Hinton agar (BD Biosciences, Franklin Lakes, NJ, United States) plates containing 1 mg/ml rifampin and 2 \( \mu \text{g} \)/ml imipenem to select the transconjugants containing the \( \text{bla}_{\text{KPC}} \) marker.

**Phenotypic Assays**

Activity of Ambler class A/B/D carbapenemases in bacterial cell extracts was determined by a modified CarbaNP test (Feng et al., 2016). Bacterial antimicrobial susceptibility was examined using the broth dilution method, and interpreted as per the Clinical and Laboratory Standards Institute guidelines (CLSI, 2015).

**Genomic DNA Sequencing and Sequence Assembly**

Genomic DNA was isolated from \textit{Enterobacteriaceae} isolates GN2, GN26, GN28, and E20 using a Blood and Cell Culture DNA Maxi Kit (Qiagen, Hilden, Germany). Genome sequencing was performed for isolate GN2 using a sheared DNA library with an average size of 15 kb (ranging from 10 to 20 kb) on a PacBio RSII sequencer (Pacific Biosciences, Menlo Park, CA, United States), as well as with a paired-end library with an average insert size of 400 bp (ranging from 150 to 600 kb) on a HiSeq sequencer (Illumina, San Diego, CA, United States). The paired-end short Illumina reads were used to correct the long PacBio reads using proovread (Hackl et al., 2014), then the corrected PacBio reads were assembled de novo using SMARTdenovo².

Genomic DNA from isolates GN26, GN28, and E20 was sequenced from a mate-pair libraries with an average insert size of 5 kb (ranging from 2 to 10 kb) using a MiSeq sequencer (Illumina, San Diego, CA, United States). DNA contigs that were not matched with the reference chromosome sequences of \textit{S. marcescens} (GenBank accession number \( \text{HG}738868 \)), \textit{M. morganii} (GenBank accession number \( \text{CP}023505 \)) or \textit{K. aerogenes} (GenBank accession number \( \text{FO}203355 \)) were assembled based on their contig coverage values using Newbler 2.6 (Nederbragt, 2014). 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 (ORFs) and pseudogenes were predicted using RAST 2.0 (Brettin et al., 2015) combined with BLASTP/BLASTN searches (Boratyn et al., 2013) against the UniProtKB/Swiss-Prot database (Boutet et al., 2016) and the RefSeq database (O’Leary et al., 2016). Resistance genes, mobile elements, and other features were annotated using online

²https://github.com/ruanjue/smartdenovo
databases including CARD (Jia et al., 2017), ResFinder (Zankari et al., 2012), ISfinder (Siguier et al., 2006), and the Tn Number Registry (Roberts et al., 2008). Multiple and pairwise sequence comparisons were performed using MUSCLE 3.8.31 (Edgar, 2004) and BLASTN, respectively. Gene organization diagrams were drawn in Inkscape 0.48.13.

**Nucleotide Sequence Accession Numbers**

The complete sequences of plasmids pE20-KPC, pGN2-KPC, pGN26-KPC, and pGN28-KPC and the draft sequences of the E20, GN2, GN26, and GN28 chromosomes were submitted to GenBank under accession numbers MF156709 to MF156712, CP026722, CP026581, CP026650, and CP026651, respectively.

**RESULTS**

**bla**\textsubscript{KPC}-Carrying Isolates

From 2014 to 2015, a total of 143 carbapenem-resistant *Alcaligenes xylosoxidans* (\(n=6\)), *Acinetobacter baumannii* (\(n=24\)), *Pseudomonas aeruginosa* (\(n=47\)), *Pseudomonas putida* (\(n=5\)), *K. aerogenes* (\(n=6\)), *E. coli* (\(n=2\)), *Enterobacter cloacae* (\(n=2\)), *K. pneumoniae* (\(n=35\)), *P. mirabilis* (\(n=8\)), *S. marcescens* (\(n=5\)), and *M. morganii* (\(n=3\)) isolates were obtained from the 143 different patients with various infections at a Chinese public hospital. Of these carbapenem-resistant isolates, 45 (31.5%) demonstrated class A carbapenemase activity and contained \(bla_{KPC}\) genes, while three isolates (2.1%) had class B carbapenemase activity and carried \(bla_{NDM}\) genes. Carbenapenemase activity and major plasmid-borne carbapenemase genes were not detected in the remaining strains (66.4%). All of the carbapenemase-positive isolates were identified as *Enterobacteriaceae*.

These 45 \(bla_{KPC}\)-carrying isolates consisted of *K. pneumonia* (\(n=31\)), *K. aerogenes* (\(n=6\)), *S. marcescens* (\(n=5\)), and one isolate each of *M. morganii*, *E. coli*, and *P. mirabilis*. In total, 36 of the isolates were recovered from sputum specimens, while the remaining isolates were obtained from urine specimens.

The 45 isolates came from 10 different hospital departments: 25 from the Intensive Care Unit, 6 from the Department of Gerontology, 5 from the Department of Respiratory Medicine, 2 from the Department of Neurology, 2 from the Department of Urology, and 1 each from the Department of General Surgery, the Department of Neurosurgery, the Emergency Department, the Department of Endocrinology, and the Department of Traditional Chinese Medicine (Supplementary Table S1). Thirty-six (80.0%) of the 45 isolates carried one or more \(\beta\)-lactamase genes \([bla_{TEM}, bla_{SHV}, bla_{CTX-M-1}\text{G} \text{ (Group)}, bla_{CTX-M-9G}, bla_{OXA-1} \text{ and } bla_{OXA-2}\text{]}\) in addition to \(bla_{KPC}\).

**pKPC3\textsubscript{SZ}-Like IncX6 Plasmids From \(bla_{KPC-2}\)-Carrying Isolates**

Four \(bla_{KPC}\)-positive isolates, *P. mirabilis* GN2, *S. marcescens* GN26, *M. morganii* GN28, and *K. aerogenes* E20, were arbitrarily selected for genome sequencing. GN2 was isolated from the urine specimens of an elderly female with urinary tract infection in 2014, while GN26 and GN28 (in 2015) and E20 (in 2014) were isolated from sputum specimens from three different elderly males suffering from pulmonary infections. These four patients were admitted to the hospital because of primary diseases consisting of myocardial infarction, cerebral infarction sequelae, cerebral contusion and pneumonia, respectively, and developed the above hospital-acquired infections during hospitalization.

GN2, GN26, GN28, and E20 each contained an IncX6 plasmid, designated pGN2-KPC, pGN26-KPC, pGN28-KPC, and pE20-KPC, respectively. These plasmids were 45.6–46.3 kb in length, with 62–65 predicted ORFs (Table 1). The modular structure of each plasmid was divided into the backbone regions along with a single accessory module, which was defined as an acquired DNA region associated with mobile elements, and was inserted into the backbone (Figure 1 and Supplementary Figure S1). A total of three resistance genes were identified: \(bla_{KPC-2}\) was located in all four plasmids, while \(\Delta{bla_{TEM-1}}\) was identified in pGN2-KPC, pGN26-KPC, and pGN28-KPC, and \(qnrS1\) was found in pGN26-KPC. All these resistance genes were located in the accessory modules.

All four \(bla_{KPC-2}\)-carrying plasmids could be transferred to *E. coli* EC600 via conjugation, generating the corresponding

| Category                      | pGN2-KPC | pGN26-KPC | pGN28-KPC | pE20-KPC | pKPC3\textsubscript{SZ} |
|-------------------------------|----------|-----------|-----------|----------|------------------------|
| Total length (bp)             | 46,320   | 46,292    | 46,123    | 45,579   | 43,333                 |
| Total number of ORFs          | 65       | 65        | 63        | 62       | 61                     |
| Mean G+C content, %           | 47.9     | 47.7      | 47.9      | 47.9     | 47.7                   |
| Length of the backbone (bp)   | 32,849   | 32,720    | 32,652    | 32,722   | 32,721                 |
| Accessory module(s)           | \(\Delta{bla_{TEM-1}}\) | \(\Delta{bla_{TEM-1}}\) | \(\Delta{bla_{TEM-1}}\) | \(\Delta{bla_{TEM-1}}\) | \(\Delta{bla_{TEM-1}}\) |
| [resistance gene(s) harbored] | \(bla_{KPC-2}\) region \((bla_{KPC-2} \text{ and } \Delta{bla_{TEM-1}})\) | \(bla_{KPC-2}\) region \((bla_{KPC-2} \text{ and } \Delta{bla_{TEM-1}})\) | \(bla_{KPC-2}\) region \((bla_{KPC-2} \text{ and } \Delta{bla_{TEM-1}})\) | \(bla_{KPC-2}\) region \((bla_{KPC-2} \text{ and } \Delta{bla_{TEM-1}})\) | \(bla_{KPC-3}\) region \((bla_{KPC-3} \text{ and } \Delta{bla_{TEM-1}})\) |

\(pGN2\text{-KPC}, pGN26\text{-KPC}, pGN28\text{-KPC}, \text{ and } pE20\text{-KPC}\) were fully sequenced in this work, while \(pKPC3\text{\textsubscript{SZ}}\) was derived from GenBank. Genetic comparison of these five sequenced plasmids was interpreted in the main text.
**bla**<sub>KPC</sub>-positive *E. coli* transconjugants GN2-KPC-EC600, GN26-KPC-EC600, GN28-KPC-EC600, and E20-KPC-EC600. Class A carbapenemase activity was detected for all transconjugants, and resulted from the production of the KPC-2 enzyme. Both the wild-type and transconjugant strains were resistant to ampicillin, cefepime, meropenem, and aztreonam. Moreover, GN26 was resistant to ciprofloxacin, but its transconjugant was intermediately resistant to this drug due to the presence of *qnrS1* known to mediate the low-level resistance to fluoroquinolones (Table 2). In conclusion, each of GN2, GN26, GN28, and E20 harbored a conjugative **bla**<sub>KPC</sub>-carrying IncX6 plasmid, which accounted for the carbapenem resistance phenotype.

Based on the complete sequences of the five IncX6 plasmids (pGN2-KPC, pGN26-KPC, pGN28-KPC, pE20-KPC, and pKPC3_SZ), a total of nine genes were arbitrarily selected to screen for the prevalence of pKPC3_SZ-like IncX6 plasmids among the 45 **bla**<sub>KPC</sub>-positive isolates. Of these nine selected genes, eight [replication initiation: repA (replication initiation protein); maintenance: parA (partitioning ATPase), topB (type III topoisomerase), dnaJ (molecular chaperone), and ftsH (cell division protein); conjugal transfer: tivB3-4 (P-type type IV secretion, inner-membrane component of translocation channel and ATPase), tivB6 (P-type type IV secretion, inner-membrane component of translocation channel), and tivB10 (P-type type IV secretion, outer-membrane component of translocation channel)] were from backbone regions, while the remaining one was quinolone-resistance gene *qnrS1*. PCR analysis and amplicon sequencing showed that all eight backbone genes were present in 24 isolates, including 11 *K. pneumoniae* isolates, all 6 *K. aerogenes* isolates, 4 *S. marcescens* isolates, and 1 isolate

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**TABLE 2 | Antimicrobial drug susceptibility profiles.**

| Antibiotics          | GN2       | GN2-KPC-EC600 | GN26      | GN26-KPC-EC600 | GN28      | GN28-KPC-EC600 | E20       | E20-KPC-EC600 | EC600     |
|----------------------|-----------|---------------|-----------|---------------|-----------|---------------|-----------|---------------|-----------|
| Ampicillin           | > 1024/R  | > 1024/R      | > 1024/R  | > 1024/R      | > 1024/R  | > 1024/R      | > 1024/R  | > 1024/R      | < 4/S     |
| Cefepime             | > 256/R   | 64/R          | > 256/R   | 16/R          | 256/R     | 128/R         | > 256/R   | 32/R          | < 2/S     |
| Meropenem            | 16/R      | 4/R           | 32/R      | 8/R           | 16/R      | 4/R           | 32/R      | 8/R           | < 1/S     |
| Aztreonam            | > 512/R   | 512/R         | > 512/R   | > 512/R       | > 512/R   | > 512/R       | > 512/R   | > 512/R       | < 4/S     |
| Amikacin             | < 8/S     | < 8/S         | < 8/S     | < 8/S         | < 8/S     | < 8/S         | < 8/S     | > 1024/R      | > 8/S     |
| Tetracycline         | 64/R      | < 1/S         | 64/R      | < 1/S         | 64/R      | < 1/S         | 64/R      | < 1/S         | < 1/S     |
| Ciprofloxacin        | 16/R      | < 1/S         | 4/R       | 2/R           | 64/R      | < 1/S         | 128/R     | < 1/S         | < 1/S     |
| Nitrofurantoin       | 128/R     | 16/S          | > 128/R   | 16/S          | 64/S      | 16/S          | 128/R     | 16/S          | 8/S       |
| Trimethoprim         | > 32/R    | < 0.25/S      | < 0.25/S  | < 0.25/S      | < 0.25/S  | < 0.25/S      | > 32/R    | < 0.25/S      | < 0.25/S  |
| Sulfamethoxazole     | > 608/R   | < 4.75/S      | < 4.75/S  | < 4.75/S      | < 4.75/S  | < 4.75/S      | > 608/R   | < 4.75/S      | < 4.75/S  |
| Tigecycline          | < 1/S     | < 1/S         | < 1/S     | < 1/S         | < 1/S     | < 1/S         | < 1/S     | < 1/S         | < 1/S     |
| Colistin             | 128/R     | < 1/S         | > 128/R   | < 1/S         | < 1/S     | < 1/S         | < 1/S     | < 1/S         | < 1/S     |

S, sensitive; R, resistant; I, intermediately resistant.

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**FIGURE 1 | Linear comparison of IncX6 plasmids.** 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).
each of *M. morgani*, *E. coli*, and *P. mirabilis*, indicating that these isolates harbored IncX6 plasmids. All five replication and maintenance genes, but none of the three conjugal transfer genes, were detected in another *S. marcescens* isolate, probably indicating that this isolate contained an IncX6 plasmid that had lost the conjugal transfer genes. None of the eight selected genes were detected in the remaining 20 isolates, signifying that these isolates did not carry IncX6 plasmids. *qnrS1* was detected in 26 *K. pneumoniae* isolates, 6 *K. aerogenes* isolates, and 4 *S. marcescens* isolates, denoting coexistence of *bla*KPC and *qnrS1* in these isolates.

**Genomic Comparison of IncX6 Plasmids**

pGN2-KPC, pGN26-KPC, pGN28-KPC, and pE20-KPC showed the highest sequence identity to the IncX6 reference plasmid pKPC3_SZ ([Du et al., 2016]), with >92% query coverage and >99% nucleotide identity. The major backbone genes or gene loci included *repA* and its iterons (replication initiation), *parA* and *topB–hha–hns* (maintenance), and *rlx, dtr, tivB, cpl*, and *eex* (conjugal transfer). *repA* coded for the IncX6-specific replication initiation protein and was not identified in any other available sequences. A 253-bp region containing seven imperfect GGGTTTATATCCCGATA direct repeats was located 73-bp upstream of *repA*, and may function as iterons that bind the RepA protein. *ParA* was the partitioning ATPase responsible for plasmid segregation and stability ([Schumacher, 2008]), however, centromere-binding protein ParB and its binding sites *parC* could not be located. The gene expression modulation (gmn) region, composed of *topB, hha* (transcriptional regulator), and *hns* (histone-like DNA-binding protein), was involved in plasmid maintenance ([Norman et al., 2008]). The conjugal transfer region was composed of a complete set of P-type conjugative DNA transfer genes, including *rlx* and *dtr* (DNA transfer; encoding relaxase Rlx and an auxiliary protein, Dtr), *tivB1–tivB11* (encoding P-type IV secretion system elaborating the pilus for mating pair formation), *cpl* (encoding a coupling protein that links DNA transfer and mating pair formation), and *eex* (entry exclusion preventing nucleoprotein transport between donors) ([De La Cruz et al., 2010; Chen et al., 2013; Thomas et al., 2017]).

The backbones of these five plasmids displayed only two major modular differences (Figure 1): (i) a 128-bp duplication in *cpl* of pGN2-KPC resulted in frameshift mutation, turning *cpl* into a pseudogene but retaining the conjugal transfer ability of pGN2-KPC, and (ii) a 71-bp deletion within *orf393* (coding for an XRE-family Helix-turn-helix protein) was identified in pGN28-KPC, again causing the hypothetical gene *orf393* to become a pseudogene.

The accessory modules of pGN2-KPC, pGN26-KPC, pGN28-KPC, pE20-KPC, and pKPC3_SZ were named the *bla*KPC regions (Figure 1), and were highly similar to one another (Figure 2). The *bla*KPC regions from pGN2-KPC, pGN28-KPC, and pE20-KPC comprised a ΔTn6296 derivative and an ISKpn19 element, while that from pGN26-KPC consisted of a ΔTn6296 derivative and an ISKpn19-containing ΔTn6292 derivative. The *bla*KPC region from pKPC3_SZ contained only a ΔTn6296 derivative (Figure 2).

Tn6296 was originally identified in plasmid pKP048 from *K. pneumoniae* ([Jiang et al., 2010]). It was generated by the insertion of the core *bla*KPC-2 genetic platform (Tn6376- *bla*KPC-2–ΔISKpn6–korC–orf6–klcA–ΔrepB) into the *mcp* (methyl-accepting chemotaxis protein) gene of the cryptic transposon Tn1722, truncating *mcp* and splitting Tn1722 into ΔTn1722-5’ [IRL (inverted repeat left)–tnpAR–res] and ΔTn1722-3’ [Δmcpc–IRR (inverted repeat right)]. The Tn3-family unit transposon Tn6292, as observed in pIMP-HZ1 from *K. pneumoniae*, carried a core quinolone resistance genetic platform, *qnrS1–ΔISecl2*, and contained an ISKpn19 insertion within *tnpA* ([Feng et al., 2016; Liang et al., 2017]).

The ΔTn6296 derivatives from these five plasmids were slightly different from each other, with deletions and insertions relative to the prototype Tn6296 (Figure 2 and Supplementary Table S2). First, ΔTn1722-5’ was lost from all five ΔTn6296 elements. Second, a 70-bp deletion within *tnpA* of Tn6376 and a 70-bp deletion within Δ*tnpA* of ΔISKpn6 were found in pE20-KPC and pGN26-KPC, respectively, leading to frameshift mutations of these two coding regions. Third, the insertion of a 624-bp Δ*blaTEM-1*-containing region between ISKpn27 and *bla*KPC-2/3 was identified in pGN26-KPC, pGN22-KPC, pGN28-KPC, and pKPC3_SZ. Two promoters, consisting of the intrinsic P1 promoter and an upstream Tn6376-provided P2 promoter, were found to govern the *bla*KPC-2 expression of Tn6296 ([Wang et al., 2015]). The insertion of the Δ*blaTEM-1*-containing region resulted in the loss of IRL-Tn6376 and P1 (Supplementary Figure S2), leaving P2 as the only promoter for *bla*KPC expression. Finally, the 3’-terminal regions of these five ΔTn6296 derivatives were truncated in different formats: (i) ΔTn1722-3’ was absent in each of pE20-KPC, pGN22-KPC, and pGN28-KPC due to connection of ISKpn19; (ii) in pGN26-KPC, the *qnrS1*-containing ΔTn6292 element (6.4 kb in length) was connected with a 1.6-kb ΔTn6376 remnant with deletion of IRL–*tnpA*–res and truncation of *tnpR*, and the introduction of ΔTn6292 into pGN26-KPC probably resulted from homologous recombination between Tn6292 and a pre-existing ISKpn19 element (as observed in pE20-KPC, pGN22-KPC, and pGN28-KPC), with ISKpn19 acting as the common region necessary for recombination; and (iii) ΔTn1722-3’ was also lost in pKPC3_SZ, although none of the ISKpn19-related elements or any other regions were found to be adjacent to the 3’-end of ΔTn6296.

In summary, complex transposition and homologous recombination events, particularly those involving the three prototype mobile elements Tn6296, Tn6292, and ISKpn19, occurred to promote the assembly and mobilization of the *bla*KPC regions in these plasmids.

**DISCUSSION**

IncX6 plasmid backbones have very limited sequence identity (<18% BLAST coverage and <84% nucleotide sequence identity) to those of other subgroups. Indeed, dramatic genetic diversities are presented among different IncX subgroups. Nevertheless, IncX6 plasmids contain the core IncX backbone makers responsible for
FIGURE 2 | The bla_{KPC} regions from IncX6 plasmids, and comparison with related regions. 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).
Previously sequenced IncX plasmids mostly belong to the IncX1–IncX4 subgroups, with very few representatives of IncX5–IncX7 plasmids. Currently, only five IncX6 plasmids have been fully sequenced, including pKPC3_SZ (Du et al., 2016) and the pGN2-KPC, pGN26-KPC, pGN28-KPC, and pE20-KPC plasmids sequenced in the current study. The five plasmids all originate from clinical isolates belonging to different Enterobacteriaceae species, namely E. cloacae, P. mirabilis, S. marcescens, M. morgani, and K. aerogenes, respectively, all of which come from China. Each of these five IncX6 plasmids contains a single accessory module containing two or three resistance genes, with all five carrying plasmids appear to be an important vehicle for dissemination of KPC-2-Encoding IncX6 Plasmids in a Hospital.

BlaoKPC, pKPC3_SZ, pGN2-KPC, and pGN28-KPC harboring of these IncX6 plasmids encode multi-drug resistance. IncX6 plasmids to various geographic areas to understand the contribution of IncX6 plasmids to blaKPC epidemiology among Enterobacteriaceae isolates.

**AUTHOR CONTRIBUTIONS**

DZ and BoL: conception and design of the study. BiL, JF, DZ, BoL, ZZ, ZY, QJ, PW, XC, BG, JH, PM, WW, WC, YT and JW: acquisition of data. BiL, JF, DZ, and BoL: analysis and interpretation of data. BiL, JF, DZ, and BoL: drafting the article. BiL, JF, DZ, BoL, ZZ, ZY, QJ, PW, XC, BG, JH, PM, WW, WC, YT, and JW: critical revision. All authors read and approved the final manuscript.

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**SUPPLEMENTARY MATERIAL**

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmicb.2018.00478/full#supplementary-material

**FIGURE S1** | Schematic maps of IncX6 plasmids. Genes are denoted by arrows, and the backbone and accessory module regions are highlighted in black and gray, respectively. The innermost circle presents GC skew ([G-C]/[G+C]), with a window size of 600 bp and a step size of 20 bp. The next-to-innermost circle presents GC content.

**FIGURE S2** | Alignment of promoter-proximal regions of blaKPC-2. The 898–354 bp upstream sequences together with the start codon of the blaKPC-2 genes from IncX6 plasmids and Tn6296 are aligned by MUSCLE. Shown are core promoter regions, −35 and −10 elements, transcription starts, Shine-Dalgarno (SD) sequences for ribosome recognition and translation starts.

**TABLE S1** | KPC-producing Enterobacteriaceae strains.

**TABLE S2** | Major features of ΔTn6296 derivatives compared to Tn6296.

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plasmid replication initiation (*repA* and *bis*), maintenance (*parA*, *hns–hha–topB*, *releB*, and *dnaJ*), and conjugal transfer (*rlx*, *dtf*, *tivB*, *cpl*, *eex*, and *actX*).
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**Conflict of Interest Statement:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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