Complete Nucleotide Sequence of KPC-3-Encoding Plasmid pKpQIL in the Epidemic Klebsiella pneumoniae Sequence Type 258

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We have determined the entire DNA sequence of plasmid pKpQIL, the blaKPC-3-carrying plasmid harbored by the carbapenem-resistant Klebsiella pneumoniae clone sequence type 258 (ST 258) in Israel. pKpQIL is a 113,637-bp, self-transmissible plasmid that belongs to the incompatibility group IncFII. It consists of a large backbone of a pKPN4-like plasmid and carries the blaKPC-3-containing Tn4401a transposon of a pNYC-like plasmid.

Bacterial plasmids are extrachromosomal genetic elements that play a crucial role in the acquisition and dissemination of antibiotic resistance determinants through inter- and intraspecies horizontal gene transfer. Resistance to carbapenems in Enterobacteriaceae does not occur naturally but rather arises by the acquisition of various β-lactamases encoded by transferrable plasmids (15).

During the last decade, carbapenem-resistant KPC-producing Enterobacteriaceae strains, particularly Klebsiella pneumoniae, have emerged and spread globally (14). These KPC-producing strains harbor plasmids encoding KPC-type carbapenemases. Several KPC-encoding plasmids from K. pneumoniae were partially or fully sequenced, including blaKPC-2 and blaKPC-3-carrying plasmids from the United States (6) and blaKPC-2-carrying plasmids from Greece, Colombia (12), and China (19). These plasmids differed in size and harbored the transposon Tn4401, proven to be involved with the dissemination of blaKPC (12, 14). This element has been found entirely or in part in all of the blaKPC-encoding plasmid sequences to date and has been found in one of two isoforms, a or b, that differ by a 100-bp deletion upstream of the blaKPC gene (6). A variant of this transposon, harboring ISKpn8, was reported from China (19).

In 2006, an extremely drug-resistant KPC-3-producing K. pneumoniae strain emerged in Israel (8), causing a nationwide clonal outbreak. This strain is genetically related to K. pneumoniae outbreak isolates from various regions in the United States (13), identified as sequence type 258 (ST 258) (7). Soon thereafter, this sequence type was found to have spread further geographically to China and several countries within the Middle East, Europe, and South America (1, 4, 11, 20). The K. pneumoniae ST 258 strain isolated in Israel is susceptible only to gentamicin and colistin (8), thereby limiting therapeutic options and thus posing a clinical and public health threat (18).

Clinical isolates of K. pneumoniae ST 258 studied during the last 4 years in Israel have been found to harbor an identical blaKPC-3-encoding plasmid of about 105 kb (13), designated pKpQIL (9). This plasmid was shown to be self-conjugative and exclusively responsible for carbapenem resistance in these isolates. pKpQIL contains the Tn4401 element and TEM-1 gene (9). In this paper, we report the complete sequence and analysis of pKpQIL.

Plasmid pKpQIL was purified from a representative clinical isolate of carbapenem-resistant KPC-3-producing K. pneumoniae ST 258, isolate 557. This isolate harbored two plasmids: pKpQIL and an additional 240-kb plasmid (9). Plasmid DNA was purified using a NucleoBond PC 100 plasmid midi kit (Macherey-Nagel GmbH, Düren, Germany) and transformed into Escherichia coli GeneHogs (Invitrogen Corp., Dorset, United Kingdom). Transformants harboring the blaKPC gene were identified by PCR and selected as described previously (8). pKpQIL was isolated and subjected to a complete DNA sequence analysis using “454 massively parallel DNA sequencing” (Dyn G. S. Ltd., Caesarea, Israel). The resulting sequences were assembled to eight contigs using the 454 Newbler assembler software (10). Sequence gaps on the plasmid were closed by PCR and sequencing. GeneMark (http://exon.biology.gatech.edu/gmhm2_prok.cgi) was used to predict the putative open reading frames (ORFs). Overlapping and closely clustered ORFs were manually inspected. The G+C content of the plasmid was identified using the GC calculator (http://www.siencebuddies.org/science-fair-projects/project_ideas/Genom _GC_Calculator.shtml). The nucleotide acid and deduced protein sequences were analyzed at the NCBI website (http://www.ncbi.nlm.nih.gov/).

The entire DNA sequence of pKpQIL is composed of 113,637 bp forming a circular plasmid (Fig. 1) with a G+C content of 53.9%. One hundred thirty-six open reading frames were identified. The products of 67 ORFs showed a high similarity to proteins of known functions. Sequence analysis of pKpQIL showed a high similarity with two K. pneumoniae plasmids. Seventy-five percent of the sequence was highly similar to that of plasmid pKPN4 (GenBank accession no. CP000649), and 12% of pKpQIL was highly similar to the partially sequenced plasmid pNYC (GenBank accession no. EU176011).
FIG. 1. Genetic map of the full-length, 113-kb pKpQIL plasmid. The arrows indicate gene location and orientation. Predicted ORFs with known functions are indicated by black arrows. ORFs of hypothetical proteins are indicated by white arrows. Boundaries of Tn4401 are marked with a bracket. The plasmid map was drawn using Redasoft visual cloning software. The black arc inside the circle shows the region in common with plasmid pKPN4 of *K. pneumoniae* MGH 78578, and the gray arc indicates the region in common with plasmid pNYC of *K. pneumoniae* strain YC.

The region in common between pKpQIL and pKPN4 consists of 86,610 bp (positions 26360 to 112970 on pKpQIL). This region shows 99% identity with plasmid pKPN4 at both the sequence and the gene organization level. This plasmid is one of five plasmids harbored by multidrug-resistant *K. pneumoniae* MGH 78578 (ATCC 700721) that was isolated from the sputum of a 66-year-old male pneumonia patient hospitalized in Massachusetts General Hospital in 1994 (http://www.expasy.ch/sprot/hmap/KLEP7.html). The common coding region of these two plasmids consists of genes responsible for plasmid maintenance, transmission, and antibiotic resistance (Fig. 1).

The genes identified on pKpQIL that are responsible for plasmid maintenance and stability, as in other enteric plasmids (2), include plasmid segregation genes (*parA* and *parB*), type I restriction modification systems, and mediators of plasmid stability (*stbA* and *stbB*). The *tra* gene region, involved in plasmid transfer via conjugation, makes up approximately 35 kb (30% of the plasmid length) of both plasmids, supporting our previous findings that pKpQIL is self-transmissible (9). An additional common region between pKpQIL and pKPN4 carries antibiotic resistance genes, including the *aadA* gene, *bla*OXA-9, and *bla*TEM-1 and is similarly positioned in the transposon Tn331, as reported previously (17). Sequence analysis of this region showed two main modifications. The first is the truncation of *aadA* being a result of the IS26 insertion at position 682 of the *aadA* gene. A second is a nonsense mutation (A→G) at position 350 of the *bla*OXA-9 gene, leading to inactivation of this gene. The presence of *bla*OXA-9 and *bla*TEM-1 together with
bla_{KPC-3} was previously reported in other K. pneumoniae isolates (5, 16, 21).

From positions 4828 to 18898 (14,070 bp), pKpQIL shows 99% identity with the plasmid pNYC of the K. pneumoniae strain YC (12). This region contains the 10-kb bla_{KPC-3}-carrying Tn4401 transposon responsible for carbapenem resistance that has been previously characterized (6, 12). The Tn4401 transposon of pKpQIL has a 100-bp deletion and is therefore isofrom a, as in agreement with previous findings (7).

The remaining part of pKpQIL (overall, 12,954 bp) includes 4,828 bp upstream of Tn4401 and 8,126 bp downstream of Tn4401. The 4,828 bp show a 94% homology to the plasmid pKp91 of K. pneumoniae 342 (GenBank accession number NC_011281). This region includes four hypothetical proteins, an endonuclease, and the origin of replication of pKpQIL was found to belong to the IncFII-like incompatibility group, based on the RepA sequence (3). RepA showed 89% amino acid identity with the IncFII RepA from Salmonella enterica plasmid pSLT (GenBank accession number AE006471). Previously reported IncFII plasmids were also identified to carry antibiotic resistance genes, including CTX-M- and SHV-type extended-spectrum β-lactamase (ESBL)-encoding genes, plasmid-mediated quinolone resistance genes, or ampC genes (2). pKpQIL is the first bla_{KPC-3}-carrying plasmid found to belong to the IncFII group. PCR-based replicon typing (PBRT) applied by Carattoli et al. to select bla_{KPC-3}-carrying plasmids from different outbreak strains resulted in negative results for all the replicons (2). Other bla_{KPC-3}-carrying plasmids analyzed in the same study belonged to other replicon types, such as plasmid 9 of K. pneumoniae from the United States (GenBank accession number FJ223607), found to belong to the IncN-like type (2), and plasmid pBC633 of K. pneumoniae strain KN633 from Colombia (GenBank accession number EU176012), which had a CoE-like replication origin (2).

pKpQIL was found to carry two replicons. In addition to the IncFII-like replicon mentioned previously, another origin of replication (RepA, position 53179 to 54075) exists on the common part of pKpQIL and pKP24 and shows 100% homology to RepA of pKP24. These findings correlate to other known IncF plasmids that often carry more than one replicon and are known as the low-copy-number plasmids described by Carattoli et al. (2). The Inc-F-like plasmids are known to be limited in their host range to the genera of Enterobacteriaceae (2). The KPC-encoding plasmids from K. pneumoniae that have been fully sequenced are plasmid 9 (GenBank accession number FJ223607), plasmid 12 (GenBank accession number FJ223605), and plasmid 15S (GenBank accession number FJ223606). Other than the common Tn4401 region, necessary for the spread of the bla_{KPC-3} gene (14), these KPC-encoding plasmids are different in their gene organization and in their antibiotic resistance features. The majority of these plasmids (pKP048 from China [19], pNGR from Greece, and pNYC from the United States [12]) were demonstrated to be self-conjugative, as was pKPQIL (9), while others, such as pBC633 (GenBank accession number EU176012), were not (12).

Previous findings indicate that all bla_{KPC-3}-carrying plasmids investigated harbor the Tn4401 structure that is located on different plasmid backbones. The complete sequencing of pKpQIL presented in the current study suggests that it was formed as a result of the plasmid rearrangement of a pKPN4-like and pNYC-like plasmid. Additional studies on KPC-encoding plasmids harbored by K. pneumoniae ST 258 from other geographical areas and plasmid comparison studies are warranted in order to reveal the mechanisms driving the success of certain KPC-carrying plasmids compared to others.

Nucleotide sequence accession number. The complete sequence of pKpQIL is available from the GenBank database under accession number GU595196.

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REFERENCES