Letter to the editor

Genomic features of a clinical ESBL-producing and colistin-resistant hypermucoviscous K. quasipneumoniae subsp. similipneumoniae from Brazil

Dear Editor:

Klebsiella pneumoniae is widely recognized as a significant threat to human health, probably associated to the emergence of hypervirulent and pan-resistant strains associated with hospital infections. Klebsiella quasipneumoniae is a novel, recently described, bacterial species which can be as virulent as other K. pneumoniae strains, causing invasive infections and resulting in high mortality rates. In this report, we report the first draft genome of a clinical ESBL-producing, colistin-resistant, highly virulent and hypermucoviscous K. quasipneumoniae subsp. similipneumoniae isolate in Brazil.

The KP121 strain was isolated from the rectal mucosa of a patient in a teaching hospital in Southeastern Brazil, on July 28, 2015. It was formerly characterized as K. pneumoniae at the hospital by the Vitek-2® system (bioMérieux, Brazil). Antimicrobial susceptibility tests were performed by Microflex TM matrix-assisted laser desorption/ionization time-of-flight mass spectrometry (MALDI-TOF MS; Bruker Daltonics, Bremen, Germany), remaining sensitive to ciprofloxacin and tigecycline. Colistin susceptibility was confirmed by the broth microdilution method according to EUCAST guidelines, and >2 μg/mL was considered resistant to colistin. It is important to note that, 13 days prior to sample isolation, the patient was colonized at the same site by a KPC-producing K. pneumoniae (positive blaKPC gene), both from different clones assessed by PFGE. At the time of KP121 strain isolation, the patient was on ceftriaxone, cefepime, and vancomycin due to other hospital infections, and was discharged 14 days after.

Total genomic DNA of the strain was subjected to whole genome sequencing (WGS) using Illumina NextSeq 500 sequencer (Illumina, San Diego, CA). Sequence reads were de novo assembled using the Velvett pipeline (version 5.0.1) and Geneious (version R9). This assembly was submitted for annotation using the NCBI Prokaryotic Genome Annotation Pipeline, and pairwise alignment was performed by BLASTn homology searches (http://blast.ncbi.nlm.nih.gov). Multilocus sequence typing (MLST) (https://cge.cbs.dtu.dk/services/MLST/), Resfinder (https://cge.cbs.dtu.dk/services/MLST/) and PlasmidFinder (https://cge.cbs.dtu.dk/services/PlasmidFinder/) were used to determine the sequence type, antibiotic resistance genes, and plasmid types present in the isolate. The BLASTn tool and the pairwise alignment of Geneious also allowed comparisons with the genomic DNA of the NTUH-K2044 strain. Were examined chromosomal mutations in genes encoding PhoP/Q, PmrA/B and CrrA/B two-component systems and mutations or insertions in mgrB.

Analyses revealed that the strain belonged to the sequencing type (ST) 1484. KP 121 was identified as K. quasipneumoniae subsp. similipneumoniae based on two criteria: (i) a high pairwise average nucleotide identity (ANI) of 98.98% with the reference strain KP142 (CP023478.1); and (ii) presence of the blaKPC-B-2 gene (Table 1). The strain was placed in the KPIib phylgroup which further supported subspecies classification. This strain presented the wzi sequence (wzi allele: 251) assigned to the (K-loci)-type KL16 (http://bigd.db.web.pasteur.fr/klebsiella).

Although K. pneumoniae is the most clinically relevant species within the genus, the emergence and description of a closely related highly virulent subspecies, hypermucoviscous and with extensive drug-resistant characteristics is significant, as it may be another important resistance and virulence gene reservoir. In epidemiological terms, plasmids from the incompatibility group IncH12 were found which, according to the literature, are associated with multidrug-resistant (MDR) isolates.

Although it belongs to the capsular type 16, the hyper-mucoviscosity characteristic of this strain was confirmed by the phenotypic test according to Catalán-Nájera et al.
which evidenced ≤5 mm strings. In addition, the strain presented virulence genes encoding the aerobactin (iutA), type 1 fimbriae (fimABCDEFGHJK), type 3 fimbriae (mrk-ABCDEFHJ), Escherichia coli common pilus (ecpA), outer core lipopolysaccharide (wabG), enterobactin (entE), urease synthesis associated genes (ureADE), allABCDRS, KfuABC, ugaE and traT. In addition, this strain also produces a moderate level of biofilm.³

Previously published studies have assessed healthcare-associated infections (HAI) epidemiology caused by MDR microorganisms, indicating that these infections are highly prevalent and difficult to control.³ The description of a non-common, highly resistant strain makes this issue even more troubling. The strain described here is colistin-resistant, but none of the classical mutations responsible for this phenomenon was detected. However, other mutations not yet described in the literature were identified (S363I in pmrB, L424F in phoQ, Q99R; Q239H; Q287; L295K in crvB). Although we did not do experiments to confirm the relationship of these mutations with resistance to colistin, this may be related to this phenotype. Another strain of *K. quasipneumoniae* subsp. *similipneumoniae* was recently described in Brazil, which although susceptible to colistin presented the KPC gene, reinforcing the idea that strains of this lineage are an important reservoir of resistance genes.⁶ Unfortunately, the increased use of polymyxins has led to reports of colistin-resistant isolates worldwide, particularly in Brazil.³ Although this strain does not have any carbapenemase gene, it is resistant to carbapenems, as well as to aminoglycosides (mediated by aph(3)’-Ia, ant(2)’-Ia and adaA2), fluoroquinolones (mediated by qnrA1, qnrA2 and qnrB) and fosfomycin (mediated by fosA). In addition and more seriously, this strain presents the *blaCTXM-9* gene, conferring an ESBL phenotype.

To the best of our knowledge, this is the first report of a highly virulent and hypermucoviscous colistin-resistant *K. quasipneumoniae*, which may represent a new challenge for clinical management and public health concern in Brazil. The description of this strain sets up a dark storm in hospital settings, due to the real possibility of a new resistance and virulence genes reservoir, which represents a challenge for the development of more effective surveillance and prevention programs.

### Nucleotide sequence accession number

The sequence of the KP121 strain was deposited in GenBank with accession number QXXP000000000.

### Ethical approval

Not required.

### Funding

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### Conflicts of interest

The authors declare no conflicts of interest.

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### REFERENCES


### Table 1 - Average nucleotide identity of KP121 genome against *Klebsiella* genomes.

<table>
<thead>
<tr>
<th>Query <em>Klebsiella</em> genome</th>
<th>Size (nucleotides)</th>
<th>DNA (G+C) content (%)</th>
<th>ANI of KP121 against query genome</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>K. quasipneumoniae</em> subsp. <em>quasipneumoniae</em> UCICRE (K535451)</td>
<td>5,263,297</td>
<td>58.16</td>
<td>96.71%</td>
</tr>
<tr>
<td><em>K. quasipneumoniae</em> subsp. <em>similipneumoniae</em> CP023478</td>
<td>5,217,300</td>
<td>57.85</td>
<td>98.98%</td>
</tr>
<tr>
<td><em>K. pneumoniae</em> subsp. <em>pneumoniae</em> KPC42 (NZ_QOD00000000)</td>
<td>5,315,950</td>
<td>57.37</td>
<td>93.90%</td>
</tr>
<tr>
<td><em>K. variicola</em> at-22 (NC_013850)</td>
<td>5,458,505</td>
<td>57.58</td>
<td>93.60%</td>
</tr>
</tbody>
</table>

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