**Abstract**

**Background:** A carbapenem-resistant hypermucoviscous *Klebsiella pneumoniae* isolate was recovered from human sputum.

**Methods:** Whole genome sequencing of this isolate was carried out to reveal its clonal background, antimicrobial resistance determinants and virulence factors. Virulence assays were performed using wax moth larvae. The transfer of *bla*<sub>NDM-5</sub> between bacterial strains was tested using conjugation. 59 genome assemblies of ST29 *K. pneumoniae* and 230 IncX3 plasmids regardless of the carriage of resistance gene were employed for phylogenetic analysis, respectively.

**Results:** The strain carried a virulence plasmid pVir-SCNJ1 bearing the virulence gene *rmpA* and exhibited a high virulence in wax moth. This hypervirulent strain belongs to sequence type 29 and carries *bla*<sub>NDM-5</sub>, which is located on a conjugative plasmid, designated pNDM5-SCNJ1, belonging to type IncX3. pNDM5-SCNJ1 was fully sequenced and shows high similarity with pNDM_MGR194, except some deletion inside the ISaba125 region. Phylogenetic analysis of IncX3 plasmids revealed that although *bla*<sub>NDM-5</sub> can be evolved from *bla*<sub>NDM-1</sub> via point mutations within some IncX3 plasmids, most of *bla*<sub>NDM-5</sub>-carrying IncX3 plasmids probably have acquired *bla*<sub>NDM-5</sub> in multiple events.

**Conclusions:** In this study, we characterized a *bla*<sub>NDM-5</sub>-positive hypervirulent *K. pneumoniae* of sequence type 29 in China. Our results highlight the need for active surveillance on this lineage of carbapenem-resistant *K. pneumoniae*.

**Keywords:** Carbapenem resistance, *bla*<sub>NDM-5</sub>, Hypervirulent, IncX3

**Introduction**

Hypervirulent *Klebsiella pneumoniae* (hvKP) is a worldwide concern due to its capacity to cause life-threatening, community-acquired infections in healthy individuals with high morbidity and mortality [1, 2]. hvKP strains are usually less resistant to most antimicrobials than classic *K. pneumoniae* [3], but the increasing emergence of carbapenemase-producing hypervirulent *K. pneumoniae* (CP-hvKP) compromises options of antimicrobial agents for infection control and drives a global crisis [2, 4]. These CP-hvKP strains are thought to be the result of acquiring plasmid-mediated resistance and virulence markers, either by transferring of resistance plasmids into hvKP strains or virulence plasmids into carbapenem-resistant strains [5]. There have now been several reports of infections caused by carbapenemase-producing hypervirulent *K. pneumoniae* strains [2, 4, 6–9]. These CP-hvKP isolates mainly produce KPC (a
group of serine-lactamases), followed by IMP (a group of metallo-β-lactamases), and they belong to the widely distributed sequence type (ST) 11 [2, 8] and several other STs, e.g., ST25, ST65 [6] and ST36 [4]. Here, we identified an ST29 CR-hvKP clinical strain with K54 serotype carrying blaNDM-5 gene and reported on its characterization.

Materials and methods

Bacterial identification and PCR analysis

The strain SCNJ1 was recovered from the sputum of a patient with an acute bronchiolitis in a hospital of Sichuan Province in November 2018. The initial species identification was performed using the Vitek-2 compact system (bioMérieux, Marcy-l’Étoile, France). A further species confirmation was performed by PCR amplifying of the 16S rRNA gene using the primer pair 27F/1492R [10]. PCR products were purified and then sequenced by Sanger sequencing. The resulting 16S rRNA gene sequences were compared with sequences in GenBank (NCBI) database using BLAST software. The presence of the acquired carbapenemase genes blaKPC, blaNDM, blaGES, blaIMP, blaOXA-48, and blaVIM in this isolate was screened via PCR using primers as previously described [11–14].

Antimicrobial susceptibility tests

In vitro susceptibility tests of cefepime, piperacillin-tazobactam, ampicillin, ampicillin/sulbactam, cefotetan, ceftriaxone, cefazolin, nitrofurantoin, ceftazidime, tobramycin, ciprofloxacin, trimethoprim/sulfamethoxazole, aztreonam, amikacin, gentamicin and levofloxacin were performed using Vitek-2 system. The minimum inhibitory concentrations (MICs) of imipenem, meropenem and colistin against the isolate were determined using the microdilution broth method following recommendations of the Clinical Laboratory Standards Institute (CLSI) [15]. Breakpoints of colistin were defined by the European Committee on Antimicrobial Susceptibility Testing (EUCAST) (http://www.eucast.org/), otherwise, we applied those defined by the CLSI.

String test

String test was performed by stretching a mucoviscous string from the colony using a standard bacteriologic loop as described previously [16]. Strains that formed viscous strings > 5 mm in length were defined as hypermucoviscous.

Conjugation

Conjugation experiments were carried out using broth-based methods with the azide-resistant Escherichia coli strain J53 as the recipient and transconjugants were selected using 2 μg/ml meropenem plus 150 μg/ml sodium azide. The presence of blaNDM-5 in transconjugants was confirmed by PCR and sequencing.

Virulence assay

The virulence potential of the SCNJ1 strain was assessed using wax moth (Galleria mellonella) larvae weighing 250 to 350 mg (Tianjin Huiyude Biotech Company, Tianjin, China) with method described previously [17]. Overnight cultures of K. pneumoniae strains were adjusted with phosphate-buffered saline (PBS) to concentrations of 1 × 10⁴ CFU/ml, 1 × 10⁵ CFU/ml, 1 × 10⁶ CFU/ml, 1 × 10⁷ CFU/ml after being washed with PBS. 10 μl of inoculum was injected into the hemocoel of sixteen larvae using a 25-μl Hamilton syringe via the last left proleg. The larvae were then incubated at 37°C and the number of live larvae was counted at 12 h intervals for 3 days. Two blaKPC-2-carrying carbapenem-resistant K. pneumoniae clinical isolates of ST11:K47, KPNJ2 and KPLZ1050, without rmpA and rmpA2, were used as the control.

Genome sequencing and analysis

The strain was subjected to whole genomic sequencing using an Illumina HiSeq 2000 system with the 150-bp paired-end approach and 150× coverage. Reads were trimmed using Trimmomatic [18]. Draft genome was then assembled using the SPAdes program [19]. Annotation was carried out using Prokka [20]. Sequence type and capsular type of this strain were determined using the assembled contigs to query the Multi-Locus Sequence Typing (MLST) v 2.0 (https://cge.cbs.dtu.dk/services/MLST/) and wzc genotyping system as previously described [21], respectively. In addition, the wzi genotyping system [22] and KLeborate (https://github.com/katholt/Kleborate/) were employed to confirm the sequence type and capsular type. Clonal complexes (CCs) were determined by using eBURST v3 based on K. pneumoniae MLST data (https://eburst.mlst.net ). Virulence genes were identified using the Virulence Factors Database (VFDB) available at http://www.mgc.ac.cn/VFs/main.htm. Antimicrobial resistance genes were identified using the ResFinder v3.1 software of the Center for Genomic Epidemiology (CGE, http://genomicepidemiology.org/).

Plasmids pVir-SCNJ1 and pNDM5-SCNJ1 were completely circularized with gaps between the contigs closed by PCR and respective ampiclons sequenced using Sanger sequencing, respectively. Identification of plasmid incompatibility types were performed on complete sequences of plasmids via the online service PlasmidFinder v2.0 at CGE (https://cge.cbs.dtu.dk/services/PlasmidFinder/). The annotations of the plasmid sequences were conducted using the RAST tools and
edited manually [23]. Sequence alignment of \( \text{bla}_{\text{NDM-5}} \)-carrying plasmids was performed using BLAST and visualized with EasyFig v 2.2.3 [24]. Alignments with highly homologous complete plasmid sequences of \( \text{pVir-SCNJ1} \) available in NCBI were performed by using the BRIG tool [25]. The circular map of \( \text{pNDM5-SCNJ1} \) was also generated using BRIG [25].

Phylogenetic analysis

All assembled \( \text{K. pneumoniae} \) genomes (n = 6823; accessed by March 1, 2019) were retrieved from GenBank. MLST typing was performed using the script (https://github.com/tseemann/mlst). A total of 59 assemblies of ST29 \( \text{K. pneumoniae} \) were included and aligned with that of strain SCNJ1 using CSI Phylogeny 1.4 (https://cge.cbs.dtu.dk/services/CSIPhylogeny/) (Additional file 1: Table S2). Gubbins (version 2.3.4) was used to remove single nucleotide polymorphisms (SNPs) on recombination sites [26]. The filtered SNPs were then used as input for inferring a phylogenetic tree using RAxML with the GTRGAMMA model and 1000 bootstraps [27]. ABRicate (https://github.com/tseemann/abricate) was used to identify antimicrobial resistance genes in these genomes and the capsular typing of \( \text{K. pneumoniae} \) was performed with the \text{wzc} genotyping system.

The sequence of all available IncX3 plasmids regardless of the carriage of resistance gene (n = 230; accessed by March 24, 2019) were retrieved from the GenBank (Additional file 1: Table S4). Orthogroups were identified using OrthoFinder [28] and used for multiple sequence alignments (MSA) with MAFFT [29]. The species tree was inferred from the concatenated MSA using FastTree [30]. The STRIDE algorithm (Specie Tree Root Inference from Duplication Events) was used to root the species tree in OrthoFinder.

Results and discussion

Antimicrobial susceptibility test showed that the \( \text{K. pneumoniae} \) SCNJ1 strain was resistant to imipenem (MIC, \( \geq 256 \mu g/ml \)) and meropenem (MIC, \( \geq 256 \mu g/ml \)) cefepime (MIC, 16 \( \mu g/ml \)), piperacillin-tazobactam (MIC, \( \geq 128 \mu g/ml \)), ampicillin (MIC, \( \geq 32 \mu g/ml \)), ampicillin/sulbactam (MIC, \( \geq 32 \mu g/ml \)), amoxicillin (MIC, \( \geq 64 \mu g/ml \)), cefotaxime (MIC, \( \geq 64 \mu g/ml \)), cefazolin (MIC, \( \geq 64 \mu g/ml \)), nitrofurantoin (MIC, 128 \( \mu g/ml \)) and ceftazidine (MIC, \( \geq 64 \mu g/ml \)), but was susceptible to colistin (MIC, 2 \( \mu g/ml \)), tobramycin (MIC, \( \leq 1 \mu g/ml \)), ciprofloxacin (MIC, \( \leq 0.25 \mu g/ml \)), trimethoprim/sulfamethoxazole (MIC, \( \leq 20 \mu g/ml \)), aztreonam (MIC, \( \leq 1 \mu g/ml \)), amikacin (MIC, \( \leq 2 \mu g/ml \)), gentamicin (MIC, \( \leq 1 \mu g/ml \)) and levofloxacin (MIC, \( \leq 0.25 \mu g/ml \)). Strain SCNJ1 showed hypermucoviscosity phenotype as evidenced by forming a viscous string about 35 mm, which is beyond the > 5 mm to define hypermucoviscous.

PCR and sequencing showed that \( \text{bla}_{\text{NDM-5}} \) was the only carbapenemase-encoding gene carried by the strain SCNJ1. NDM-5, a variant of NDM (New Delhi Metallo-\( \beta \)-lactamase), was first identified in an \( \text{E. coli} \) ST648 isolate (EC045) in the UK in 2011 from a patient with a recent hospitalization history in India [31]. Although \( \text{bla}_{\text{NDM-5}} \) has been widely found in \( \text{K. pneumoniae} \) strains since its first discovery (Additional file 1: Table S1), \( \text{bla}_{\text{NDM-5}} \)-carrying hypermucoviscous \( \text{K. pneumoniae} \) remains uncommon. We found one publication that described a \( \text{bla}_{\text{NDM-5}} \)-positive \( \text{K. pneumoniae} \) isolate (K2/ST14) in China in 2015 [32], which was speculated to be hypermucoviscous on the basis of genome analysis. However, no experimental data was included.

Draft genome sequences of SCNJ1 was assembled into 29 contigs (28 were \( > 1000 \) bp in length), which comprises 5,474,953 bp, with a 57.29% GC content. SCNJ1 was assigned to capsular type \( \text{wzi}115-\text{K54} \) and sequence type ST29 (\( \text{gapA-infB-mdh-phiE-rpoB-tonB} \) allele number 2–3–2-3-6-4-4). K54 is a hypervirulent member of \( \text{K. pneumoniae} \) [16] and has been described in several previous publications as linked to ST29 [33–37]. To date, \( \text{K. pneumoniae} \) strains with the ST29 group has a worldwide distribution and has been found carrying a variety of carbapenem genes, including \( \text{bla}_{\text{NDM-1}} \) [38–40], \( \text{bla}_{\text{KPC}} \) [41], \( \text{bla}_{\text{OXA-48}} \) [42] and \( \text{bla}_{\text{OXA-181}} \) [38], as well as several extended-spectrum \( \beta \)-lactamases (ESBLs) genes [43, 44] in various countries. However, the currently available evidence is insufficient to demonstrate whether ST29, a member of CC29, is an epidemic clone mediating the spread of specific and clinically relevant antibiotic resistance genes.

The gene \( \text{bla}_{\text{NDM-5}} \) is described for the first time in a strain of \( \text{K. pneumoniae} \) ST29 in our work, as demonstrated by the phylogenetic tree based on filtered SNPs of all available ST29 \( \text{K. pneumoniae} \) strains (Fig. 1). The phylogenetic analysis also showed that strain SCNJ1 was clustered with four isolates recovered in China and was closest to strain SCLZ15–011 (GCA_001630805, carrying no carbapenemase gene, recovered in 2016 in China) with 198 SNPs difference (Fig. 1). It should be noted that sporadic cases due to ST29 \( \text{K. pneumoniae} \) were frequently detected, mainly from liver abscess patients [37, 43, 45, 46], and multidrug-resistant ST29 hvKP strains have been reported in different locations of China [5, 33, 47]. This highlights the need to monitor the epidemiology of the ST29 clones of \( \text{K. pneumoniae} \) isolates in China.

We found that strain SCNJ1 harbored genes encoding regulators of the mucoid phenotype (\( \text{rmpA} \)), aerobactin
(iucABCD and iutA), ent siderophore (entABCDEFS, fepABCDG), salmochelin (iroBCDEN), yersiniabactin (fyuA, irp1, irp2 and ybtAEPQSTUX) and type 3 fimbriae (mrkABCDFHIJ) etc. These genes are frequently associated with hypervirulence phenotype of K. pneumoniae [16, 48]. The rmpA, iutA, iucABCD and iroBCDN genes were carried by a 211,807-bp plasmid, designated pVir-SCNJ1. The rmpA2 (another regulator of mucoid phenotype) gene on the pVir-SCNJ1 plasmid was truncated, due to a frameshift mutation introducing an internal stop codon. pVir-SCNJ1 was an IncHI1/IncFIB-type plasmid and was 99.71% identical to the known virulence plasmid pLVPK (219,385 bp, GenBank accession no. NC_005249) at 93% coverage [49] (Fig. 2). It was notable that pVir-SCNJ1 was highly similar (99% coverage and 99.99% identity, Fig. 2) to the recently-identified plasmid pL22–1 (212, 635 bp, GenBank accession no. NZ_CP031258) that recovered from a Klebsiella quasipneumoniae strain L22, which suggests that the pLVPK-like plasmid has the potential to mediate inter- and intra-species transfer of virulence genes [50]. Our virulence assays showed that survival of G. mellonella was 0% with strains SCNJ1, while survival was 56.2 and 50.0% with the control strains KPLZ1050 and KPNJ2, at an inoculum of 1 × 10^5 cfu/mL at 72 h after infection, respectively (Fig. 3, Additional file 1 :Table S3). These findings suggest that strain SCNJ1 was hypervirulent.

Conjugation assays showed that strain SCNJ1 transferred a plasmid carrying bla<sub>NDM-5</sub> to E. coli J53 at a frequency of 10<sup>−6</sup> (transconjugant/recipient) by mating, suggesting that bla<sub>NDM-5</sub> was carried on a self-transmissible plasmid, which was assigned pNDM5-SCNJ1. In addition to the bla<sub>NDM-5</sub>, strain SCNJ1 had a few chromosomal resistance genes, including the ESBL gene bla<sub>SHV-187</sub>, fluoroquinolone-resistance genes oqxA and oqxB, and fosfomycin-resistance gene fosA.

pNDM5-SCNJ1 was a 45,255-bp IncX3 plasmid, with an average GC content of 46.83% and had no other known antimicrobial resistance genes except bla<sub>NDM-5</sub>. pNDM5-SCNJ1 consists of a 30-kb backbone comprising several sets of genes (pir and bis encoding replication initiation protein, parA for plasmid partitioning, hns and topB for maintenance and a gene cluster responsible for conjugation) and a genetic load region with high GC content between the
resolvase and the hns gene, which are typical of IncX3 plasmids (Fig. 4a). BLASTn revealed that the sequence of pNDM5-SCNJ1 was highly similar (100% coverage and 99.99% identity) to the plasmid pNDM_MGR194 (GenBank accession no. KF220657) recovered from a K. pneumoniae isolate in India, as well as a number of previously described IncX3 plasmids carrying blaNDM-5 in China.

In the genetic load region of pNDM5-SCNJ1 (Fig. 4b), the umuD gene was split into two fragments at the
nucleotide position 336 bp by the blaNDM-5-containing structure (IS26-ctuA1-tat-trpF-blaMBL-blaNDM-5-IS5Aba125-IS5−ISAbAba125-IS3000−ATn2), resulting in a pair of 3-bp direct repeats (TGT). In such a genetic context, an IncX3 plasmid pEC14_35 (GenBank accession no. JN935899) without any antibiotic-resistance gene, which was isolated from a patient in the USA in 1989, was likely to be the ancestral vector. It is also likely that pNDM5-SCNJ1 has diverged recently from blaNDM-1-positive plasmids pBJ01 (GenBank accession no. JX296013) by sequential mutations (Fig. 4b). In the subsequent genetic variant, the ISAba125 was truncated by the insertion of IS5 element (at 166 bp upstream blaNDM-5 start codon) and a 4-bp flanking direct repeats (CTAA). CTAA was identified. Comparisons of the genetic contexts of blaNDM-5 in pNDM_MGR194, pP744T, pTK1044 and pNDM-SCNJ1 showed that the remnant of ISAba125 was conserved, but the length of the remnant of ISAba125 between IS3000 and IS5 differed (pNDM_MGR19: 1002 bp, pP744T: 404 bp, pNDM-SCNJ1: 112 bp, pTK1044: 0 bp), suggesting that IS5 has inserted into ISAba125 at the same position in these plasmids and that gene deletions caused by homologous recombination plays a possible role in the formation of diversified ΔIS-Aba125 region.

IncX3 plasmids are narrow-host-range vectors of the Enterobacteriaceae [51, 52]. Searches on IncX3 plasmids in NCBI showed that they were recovered from various species of Enterobacteriaceae (Additional file 1: Table S4), including *E. coli*, *K. pneumoniae*, *Citrobacter freundii*, *Enterobacter cloacae*, *Klebsiella oxytoca*, *Enterobacter hormaechei*, *Salmonella enterica*, *Kluyvera intermedia*, *Morganella morganii*, *Citrobacter freundii*, *Enterobacter cloacae*, *Klebsiella oxytoca*, *Enterobacter hormaechei*, *Salmonella enterica*, *Kluyvera intermedia*, *Morganella morganii*, *Raoultella planticola* and *Raoultella ornithinolytica* from different countries, suggesting a wide distribution of IncX3 plasmids. Three kinds of carbapenemase genes were found to be carried by IncX3 plasmids, including *blaNDM*, *blaOXA-181* and *blaKPC*. The carriage rate of *blaNDM* was significantly higher (*n* = 150, 64.94%) than those of *blaKPC* (*n* = 18, 7.79%) and *blaOXA-181* (*n* = 17, 7.35%) (Additional file 1: Table S4). Of note, IncX3 plasmids were found to carry many different *blaNDM* alleles, including NDM-1, 4, 5, 6, 7, 13, 19, 20, 21, which were mainly recovered from China, confirming that IncX3 plasmids function as a common vehicle in facilitating the rapid dissemination of NDM-type MBLs among Enterobacteriaceae in China.

Phylogenetic analysis based on concatenated MSA of IncX3 plasmids revealed that pNDM5-SCNJ1 was closely related (100% coverage, 99.99% identity) to plasmid pQDE2-NDM (GenBank accession no. MH917280), which also carried *blaNDM-5* that was recovered from a *K. pneumoniae* isolate in Shandong, China, in 2015 (Fig. 5). The phylogenetic tree also showed that most of *blaNDM-5*-carrying IncX3 plasmids are tightly clustered with each other and formed a relatively distinct clade, with only sporadic ones clustered into clades with *blaNDM-1*-carrying plasmids, suggesting that although *blaNDM-5* is likely to evolve from *blaNDM-1* via point mutations on some IncX3 plasmids, most of IncX3 plasmids probably have acquired *blaNDM-5* in multiple events.
Conclusion
In conclusion, our work identified an ST29 CP-hvKP carrying the carbapenemase gene \( \text{bla}_{\text{NDM-5}} \) and provided additional evidence of the rapid dissemination of \( \text{bla}_{\text{NDM-5}} \) by pNDM-MGR194-like plasmid among Enterobacteriaceae in China. The association of the epidemic IncX3 plasmid carrying \( \text{bla}_{\text{NDM-5}} \) with a hypervirulent \( K. \) pneumoniae lineage, ST29/K54 in this case, is quite worrisome and may pose a great threat to humans. More extensive surveillance and effective action to control its further dissemination are urgently required.

Additional file

Additional file 1: Table S1. Background information on the \( \text{bla}_{\text{NDM-5}} \)-positive \( K. \) pneumoniae isolates. Table S2. ST29 \( K. \) pneumoniae strains with genome sequences available in the GenBank. Table S3. Survival (number of larvae) of \( G. \) mellonello after infection by \( K. \) pneumoniae strain SCNJ1. Table S4. The names, host species, accession numbers, carbapenemase genes and locations of IncX3 plasmids. (DOCX 109 kb)

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Authors’ contributions
LZ designed the experiments. GW, CL, FZ and WC performed the experiments. SN, YM and JZ analyzed the data. YL analyzed the data and wrote the manuscript. YY wrote the manuscript. YC edited the original draft. All authors read and approved the final manuscript.

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Availability of data and materials
Draft whole-genome sequences of the SCNJ1 strain has been deposited into GenBank under accession no. MK715436 and MK715437, respectively.

Ethics approval and consent to participate
The current study was approved by the Ethics Committee of Southwest Medical University (No.201903–194) and were carried out in accordance with the approved guidelines. Written informed consent was exempted, since this retrospective study mainly focused on bacteria and patient intervention was not required.

Consent for publication
Not applicable.

Competing interests
The authors declare that they have no competing interests.

Author details
1Department of Laboratory Medicine, The First People’s Hospital of Neijiang, Neijiang, Sichuan, China. 2Department of Immunology, School of Basic Medical Sciences, Southwest Medical University, Luzhou, Sichuan, China. 3Department of Pathogenic Biology, School of Basic Medical Sciences, Southwest Medical University, No.1 Section 1, Xiang Lin Road, Longmatan District, Luzhou 646000, Sichuan, China. 4Department of Population Medicine and Diagnostic Sciences, College of Veterinary Medicine, Cornell University, Ithaca, NY, USA.

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