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The coexistence of two <i>bla</i> _{NDM-5} genes on an IncF plasmid
as revealed by nanopore sequencing
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19 Abstract

In a carbapenem-resistant *Escherichia coli* clinical isolate of sequence type 167, two copies of bla_{NDM-5} were found on a 144,225-bp IncF self-transmissible plasmid of the F36:A4:B- type. <u>Both</u> bla_{NDM-5} genes were located in 11,065-bp regions flanked by two copies of IS26. The two regions were identical in sequence but were present at different locations on the plasmid, suggesting a duplication of the same region. This study highlights the complex genetic contexts of bla_{NDM-5} . 26 New Delhi metallo-β-lactamase (NDM) is a type of carbapenem-hydrolysing enzymes 27 (carbapenemases) with the ability to hydrolyze all β -lactams except monobactams (1), 28 representing a serious challenge for treatment of bacterial infections, infection control 29 and public health. Up to now To date, there are 21 variants of NDM, among which with NDM-5 is one of the most common variants encountered in the Enterobacteriaceae 30 (2-5). The NDM-5-encoding gene, *bla*_{NDM-5}, usually exists in a single copy on 31 32 plasmids. However, we have found the peculiar presence of two copies of *bla*_{NDM-5} on 33 a single plasmid within an *Escherichia coli* clinical isolate, which is reported here.

34

35 E. coli strain SCEC020007 was recovered from urine of a female outpatient with 36 urinary tract infection in October 2016 in China. The strain was resistant to amikacin 37 (MIC, >512 µg/ml), ceftazidime (>512 µg/ml), ceftazidime-avibactam (>512/4 µg/ml), ciprofloxacin (256 µg/ml), imipenem (64 µg/ml), meropenem (256 µg/ml), 38 39 piperacillin-tazobactam (>512/4 µg/ml) trimethoprim-sulfamethoxazole and 40 (128/2,432 µg/ml), but was susceptible to aztreonam (8 µg/ml), colistin (2 µg/ml) and tigecycline (0.25 µg/ml) as determined using the broth dilution method of the Clinical 41 Laboratory Standards Institute (6). As there are no breakpoints of colistin and 42 43 tigecycline from CLSI, those defined by EUCAST (http://www.eucast.org/) were 44 applied.

45

A draft genome sequence of the strain was generated on the Illumina HiSeq X10 platform, which generated 5,557,833 clean reads and 1.67 Gb clean bases. A total of 113 contigs (102 >1,000 bp; *N50* 126,680 bp) with a 50.76% GC content were *de novo* assembled using SPAdes (7). Strain SCEC020007 belonged to phylogenetic group A as determined using PCR as described previously (8) and sequence type 51 167 (ST167) as determined using the genomic sequence to query the E. coli multi-locus 52 sequence typing database 53 (http://enterobase.warwick.ac.uk/species/index/ecoli). Antimicrobial resistance genes 54 were identified from genome sequences using the ABRicate program (https://github.com/tseemann/abricate) 55 the ResFinder database to query 56 (http://genomicepidemiology.org/). Strain SCEC020007 had 9 antimicrobial resistance genes mediating resistance to aminoglycosides (aadA2, aadA5, rmtB), 57 58 β -lactams (*bla*_{NDM-5} and *bla*_{TEM-1}), tetracycline (*tet(A*)), sulphonamides (*sul1*) and 59 trimethoprim (*dfrA12* and *dfrA17*). Plasmid replicon types within strain SCEC020007 were determined using by the PlasmidFinder tool at http://genomicepidemiology.org/. 60 61 Surprisingly, strain SCEC020007 had an IncFIA, an IncFII and an IncB/O/K/Z replicon 62 but no IncX3, which is the common replicon type of plasmids associated with *bla*_{NDM-5}. 63

To untangle the genetic context of *bla*_{NDM-5}, strain SCEC020007 was subjected to 64 65 sequencing using the long-read real-time MinION Sequencer (Nanopore, Oxford, UK). The A de novo hybrid assembly of both short Illumina reads and long MinION reads 66 was performed constructed using Unicycler v0.4.3 (9) under conservative mode for 67 an increased accuracy. The C complete circular contigs generated were then 68 69 corrected using Plion v1.22 (10) with Illumina reads for several rounds until no 70 change was detected. The hybrid assembly of Illumina and MinION reads revealed 71 that strain SCEC020007 had a 4.8-Mb circular chromosome, a 144,225-bp plasmid containing an IncFIA and a FII replicons (designated pNDM5_020007) and an 72 84,952-bp plasmid with an IncB/O/K/Z replicon (designated pBOKZ_020007). 73 74 Surprisingly, there were two copies of *bla*_{NDM-5} in strain SCEC020007, both of which were present on pNDM5_020007. Both *bla*_{NDM-5} genes were located in 11,065-bp 75

76 regions flanked by two copies of IS26 and the two regions were identical in sequence but were present at different locations on pNDM5 020007 (Figure 1), suggesting that 77 the 11,065-bp region is duplicated. The presence of the two *bla*_{NDM-5} genes and their 78 locations on pNDM5 020007 were confirmed by PCR. The 11,065-bp region 79 contained a complex class 1 integron with a *dfrA17-aadA5* cassette array and ISCR1 80 81 (insertion sequence common region 1), which is truncated by IS26 at its 5' conserved segment, a 69-bp remnant of ctuA1 (encoding an ion tolerant protein), dsbC 82 83 (encoding an oxidoreductase), *trpF* (encoding a phosphoribosylanthranilate 84 isomerase), *ble* (mediating bleomycin resistance), *bla*_{NDM-5}, a truncated ISAba125 85 and a truncated ISEcp1/ISEc9 element (Figure 1). The co-existence of two bla_{NDM-5} 86 genes has been reported before but the co-existence of two bla_{NDM-1} genes has 87 been described previously (11, 12). Two tandem copies of *bla*_{NDM-1} genes have been found in the chromosome of an ST167 E. coli in China (11) and a Pseudomonas 88 aeruginosa strain in Serbia (12). In both cases, the tandem copies of bla_{NDM-1} are 89 90 associated with ISCR1 but not IS26. It is known that ISCR1 uses the rolling circle 91 mechanism for transposition and may generate tandem duplication of its mobilized sequence via homologous recombination (13). However, the duplication of the 92 93 11,065-bp region carrying *bla*_{NDM-5} on pNDM5_020007 is not tandem, suggesting that the duplication might not result from the action of ISCR1 but could be mediated by 94 95 IS26. The exact mechanism for the duplication of such a large region warrants further 96 studies.

97

Assembly based on Illumina reads alone generated only a single contig containing *bla*_{NDM-5} and was unable to reveal that there <u>were</u> actually <u>were</u>-two identical copies of
the same contig. This imposes difficulties for completing the *bla*_{NDM-5}-carrying plasmid

sequence by convention<u>al</u> methods including PCR and Sanger sequencing to close
gaps between contigs. By contrast, MinION sequencing <u>is_was_able</u> to resolve the
copy numbers of genes and contigs<u>and their exact position on the plasmid relative to</u>
<u>each other</u>.

105

106 Plasmid multi-locus sequence typing (pMLST) was performed using the pMLST tool 107 (https://cge.cbs.dtu.dk/services/pMLST/). pNDM5_020007 belongs to the F36:A4:B-108 type. pNDM5 020007 was has closest similarity (97% coverage and 99% identity) to a 149.5-kb unnamed plasmid (GenBank accession no. CP023871) from E. coli strain 109 110 FDAARGOS 434, which was recovered from a human rectal swab in British 111 Colombia, Canada, in 2014. This unnamed plasmid also carries *bla*_{NDM-5} (a single 112 copy) and belongs to the F36:A4:B- type. Backbones of pNDM5_020007 and the 113 unnamed plasmid of strain FDAARGOS_434 are almost identical, suggesting that 114 they might have originated from a common plasmid. Conjugation experiments were 115 carried out in broth and on filters with the azide-resistant E. coli strain J53 as the 116 recipient. pNDM5_020007 was able to be transferred by conjugation, suggesting that 117 it is self-transmissible.

118

In conclusion, we identified the presence of two *bla*_{NDM-5} genes on an F36:A4:Bself-transmissible plasmid. The co-existence of two *bla*_{NDM-5} genes was due to the duplication of an IS26-bracketed region containing IS*CR1*.

122

Nucleotide sequence accession numbers. The complete sequence of
 pBOKZ_020007, pNDM5_020007 and the chromosome of strain SCEC020007 has

125 been deposited into GenBank under the accession no. CP025625, CP025626 and

126 CP025627, respectively.

127

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- 133

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177

178 Figure legend

Figure 1. pNDM5_020007 and the genetic context of bla_{NDM-5} . The two 11,065-bp bla_{NDM-5} -containing regions bracketed by IS26 are indicated by orange circles in the map of pNDM5_020007 and are shown in detail at the bottom. Δ represents truncated genes or mobile genetic elements.