Co-Occurrence of NDM-5 and RmtB in a Clinical Isolate of *Escherichia coli* Belonging to CC354 in Latin America

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New Delhi metallo-β-lactamase (NDM)-producing isolates are usually resistant to most β-lactams and other antibiotics as a result of the coexistence of several resistance markers, and they cause a variety of infections associated to high mortality rates. Although NDM-1 is the most prevalent one, other variants are increasing their frequency worldwide. In this study we describe the first clinical isolate of NDM-5- and RmtB-producing *Escherichia coli* in Latin America. *E. coli* (Ec265) was recovered from a urine sample of a female outpatient. Phenotypical and genotypical characterization of resistance markers and conjugation assays were performed. Genetic analysis of Ec265 was achieved by whole genome sequencing. Ec265 belonging to ST9693 (CC354), displayed resistance to most β-lactams (including carbapenems), aminoglycosides (gentamicin and amikacin), and quinolones. Several resistance genes were found, including *blaNDM-5* and *rmtB*, located on a conjugative plasmid. *blaNDM-5* genetic context is similar to others found around the world. Co-transfer of multiple antimicrobial resistance genes represents a particular challenge for treatment in clinical settings, whereas the spread of pathogens resistant to last resort antibiotics should raise an alarm in the healthcare system worldwide.

**Keywords:** metallo-β-lactamase, NDM-5, RmtB, *Escherichia coli*, antibiotic multi-resistance

**INTRODUCTION**

NDM metallo-β-lactamases are carbapenemases capable of hydrolyzing almost all β-lactam antibiotics (except aztreonam), being found in several species of *Enterobacterales*, *Acinetobacter*, and *Pseudomonas*. NDM-producing isolates are usually resistant to most antibiotics due to coexistence of several resistance determinants, and they cause a variety of infections associated with high mortality rates (Mojica et al., 2016). Even though 31 NDM variants have been already
described (https://www.blbd.b, accessed 04.06.21), the NDM-1 variant remains by far the most prevalent worldwide. Substitutions have been observed at 25 of the 270 amino acid positions, of which M154L is present in 11 out of all the distinct variants (Basu, 2020). In contrast to NDM-1 that is widely spread among several Enterobacterales and other Gram-negative bacilli, NDM-5 seems to be more restricted to Escherichia coli isolates (Wu et al., 2019), as it was the first to be reported from a patient in the United Kingdom after previous hospitalization in India (Hornsey et al., 2011). In the Americas, NDM-5 has been only described in clinical isolates in the USA (Mediavilla et al., 2016; Flerlage et al., 2020) and, to the best of our knowledge; it has not been reported in human clinical samples in Latin America, so far. We are aware that the emergence of a novel variant in our geographical region could lead to the replacement or substitution of the prevailing carbapenemases present in the area. On the other hand, RmtB is a 16S ribosomal RNA methyltransferase conferring high-level resistance to aminoglycosides, which has already been disseminated among Enterobacterales worldwide (Doi et al., 2016). This situation has become worrisome due to the loss of one of the most important therapeutic options for treatment of severe infections caused by carbapenem-resistant Enterobacterales (CRE), including those produced by NDM-positive pathogens.

The aim of this study is to report the first clinical isolate of NDM-5- and RmtB-producing E. coli in Latin America and to describe the genetic context of these genes.

MATERIAL AND METHODS

On September 2018, a female outpatient was admitted to Hospital Central de San Isidro “Dr Melchor Ángel Posse” (Buenos Aires Province, Argentina), presenting urinary tract infection symptoms. The patient lived in a retirement nursing home and had not traveled abroad during the prior 12 months. No information about previous antibiotic treatment was recorded at sampling time, nor in the hospital records. A carbapenem-resistant E. coli isolate (Ec265) was recovered from a urine sample. The patient was successfully treated with nitrofurantoin and hydration.

Bacterial identification was carried out by matrix assisted desorption-ionization time of flight mass spectrometry (MALDI-TOF MS) (Bruker Daltonics, Germany).

Susceptibility testing was performed by disk diffusion test according to the Clinical Laboratory Standards Institute (CLSI) recommendations (CLSI, 2020). Phenotypic screening of metallo-β-lactamases (MBL) was performed by synergy tests using meropenem (10 µg), EDTA (1 µmol), and imipenem (10 µg)-containing disks. Minimal Inhibitory Concentrations (MIC) were determined by Sensititre ARGNF Kit (Thermo Scientific). Results were interpreted according to CLSI guidelines (CLSI, 2020), except colistin and tigecycline for which the EUCAST 2020 breakpoints were considered (https://www.eucast.org, accessed 04.06.21).

Detection and characterization of NDM genes were performed by polymerase chain reaction (PCR), cloning, and sequencing. For blaNDM detection, NDM-F (5´-GGTTTG GCGATCTGTTTTC-3´) and NDM-R (5´-CGGAATGGGCTC ATCAGGATC-3´) primers were used, rendering a 621 bp product. For blaNNDM cloning into a pK19 vector, custom primers including restriction sites were designed (NDM-SalF 5´- TACCGTGCACATGGAAATTGCCCAT-3´ and NDM-EcoR 5´- CGGAATTTCTACGGACGCTGTGTC-3´), and E. coli TOP10 was used as the recipient strain, which was transformed with the recombinant construction. Recombinants were selected on Tryptic Soy Agar containing 30 µg/ml kanamycin and 4 µg/ml meropenem, and checked by disk diffusion tests, followed by blaNNDM amplification using M13_pUC forward (5´-CCCGTACACGAATCAGATGC-3´) and M13_pUC reverse (5´-CAGGAAACAGCTATGAC-3´) primers.

Other β-lactamase genes such as blαCTX-M-type and frequent blαaCM-type were screened by PCR, and detection of the rmtB gene was also performed by PCR (RmtB-F 5´-ACTTTTA CAATCCCTCAATAC-3´ and RmtB-R 5´-AAGTATATAAG GTTCTGTTCG-3´) (Berçot et al., 2011).

Phylogenetic group was carried out by PCR following the Clermont scheme (Clermont et al., 2013).

Plasmid conjugation was performed by a mating-out assay using E. coli J53 (sodium azide resistant) as recipient, and Luria Bertani agar plates supplemented with sodium azide (250 µg/ml) and cefotaxime (2 µg/ml) as selective agents. A REP-ERIC PCR assay was carried out to evaluate the relationship between transconjugant and recipient strains (Versalovic et al., 1991).

Finally, whole genome sequencing (WGS) of the clinical isolate Ec265 was carried out on the Illumina NextSeq platform (San Diego, USA) using a paired-end (PE) library. Briefly, single colonies of the bacteria were grown in 3 ml of lysogeny broth for 18 h at 37°C and the DNA was extracted using a PureLink quick gel extraction kit (Life Technologies, CA, USA). The total genomic DNA was used to library construction with a Nextera DNA Flex kit (Illumina, San Diego, CA, USA). The generated raw reads were initially subjected to quality check using FastQC software (https://www.bioinformatics.babraham.ac.uk/projects/fastqc, accessed 01.17.21), and the paired-reads trimmed to remove adapters and low-quality regions (PHRED quality score below 20) using TrimGalore v0.6.5 (https://github.com/FelixKrueger/TrimGalore, accessed 01.17.21).

Sequence reads were assembled de novo using SPAdes V3.9, and analyzed by using on-line bioinformatic tools (Center for genomic epidemiology, CARD and Pathogen watch). Multilocus sequence typing (MLST) was determined following Achtman and Pasteur schemes (https://pubmlst.org/escherichia/, accessed 01.17.21). Enterobase (https://enterobase.warwick.ac.uk/, accessed April, 2021) was used to determine core genetic relationships among globally disseminated E. coli strains with identical sequence type, as well as to define the clonal complex (CC). To this end, the Ec265 strain was analyzed considering single locus variants (SNV) and double locus variants (DLV). Reads were submitted to EnteroBase under accession number ESC_OA7444AA.
RESULTS
Identification and Susceptibility Profile
Ec265 was identified as Escherichia coli (score value ≥ 2.0, indicating a reliable identification at specie level), displaying a multidrug-resistant profile to penicillins (ampicillin), cephalosporins (cephalotin, cefuroxime, cefoxitin, cefotaxime, ceftazidime, and cefepime), carbapenems (imipenem, meropenem, and ertapenem), aminoglycosides (gentamycin and amikacin), trimethoprim-sulfamethoxazole, tigecycline, and fluoroquinolones (ciprofloxacin and levofloxacin); remaining susceptible to colistin, nitrofurantoin, fosfomycin, tigecycline, and aztreonam. Synergy was observed between EDTA and both meropenem and imipenem disks, suggesting MBL presence. MIC values are shown on Table 1.

Resistance Markers and Mobilization
The E. coli strain Ec265, belonging to phylogenetic group F, was positive for bla\textsubscript{NDM} gene detection. PCR for bla\textsubscript{CTX-M} and bla\textsubscript{OXA} genes rendered negative results. The bla\textsubscript{NDM} gene was successfully transferred to E. coli J53 showing that this resistance marker was located on a conjugative plasmid. Resistance to gentamicin and amikacin was co-transferred in the conjugation assay (Table 1). Presence of the rmtB gene was detected in both Ec265 and the transconjugant (Tc265). REP-ERIC PCR discarded any clonal relationship between them. Recombinant pK19 plasmid harboring the bla\textsubscript{NDM} gene was sequenced and the bla\textsubscript{NDM-5} gene was confirmed. Recombinant strain also displayed a resistant phenotype against meropenem (8 µg/ml) and imipenem (>16 µg/ml).

Because of the unusual variant of bla\textsubscript{NDM} for our region, and the resulting resistance in the clinical isolate, a deep genomic analysis of Ec265 was performed by WGS.

Whole Genome Sequencing
WGS analysis revealed a 5 171 045-bp genome size, with 50.5% GC content, 166 contigs (>200 bp), and N50 value of 162 218 bp (GenBank accession no. JACXX1000000000). Resistance analysis of Ec265 predicted several acquired antimicrobial resistance genes, such as rmtB, aac(3)-IId, adaA2, aph(3’’)-Ib, aph(6)-Id (aminoglycoside resistance), dfrA12 and dfrA17 (trimethoprim resistance), bla\textsubscript{NDM-5} and bla\textsubscript{TEM-1B} (β-lactam resistance), sul1 and sul2 (sulfonamide resistance), ermA(B) and mhp(A) (macrolide resistance), tet(B) (tetracycline resistance) and chromosomal point mutations in parE (I355T), parC (S80I, D84G), and gyrA (S83L, D87N), involved in fluoroquinolones resistance (Table 2).

Ec265 belongs to Fim type H58 CH type 88-58 (fimH58 and fumC88), and the serotype predicted was O153:H34. Some acquired virulence factors were identified, such as air (enteroaggregative immunoglobulin repeat protein), eilA (Salmonella HiiA homolog), gad (glutamate decarboxylase), lpf\textsubscript{A} (long polar fimbriae), and mcmA (micronic M part of colicin H) (Table 2).

According to PlasmidFinder (https://cge.cbs.dtu.dk/services/PlasmidFinder/, accessed 01.17.21), Ec265 harbors different replicon types: IncFIA, IncFIB, IncFII, and IncQ1; however, when Pathogenwatch software (https://pathogen.watch/, accessed 01.17.21) was used, Col and Col156 replicon types were also detected (Table 2). There are several replicon types reported to be associated with bla\textsubscript{NDM-5} carrying plasmids in Enterobacteriales worldwide, being IncX3 the most common, followed by IncFIB, IncFII, and IncFIA (Wu et al., 2019). Two of these frequent replicons were detected in Ec265 (IncFII and IncFIB). The potential location of bla\textsubscript{NDM-5} in an IncFII-

| Antibiotic | MIC (µg/ml) |
|------------|-------------|
| Ampicillin | >16/4/16    |
| Ampicillin/sulbactam | >16/8/8/16/8 |
| Amoxicillin/clavulanic acid | >16/8/16/8 |
| Cephalothin | >16/16    |
| Cefotaxime | >16/16    |
| Ceftazidime | >32/32 |
| Cefepime | >16/16    |
| Amoxicillin | >8/8     |
| Nitrofurantoin | >32/32 |
| Ciprofloxacin | >16/16 |
| Levofloxacin | >4/0.5 |
| Gentamicin | >8       |
| Amikacin | >32/32    |
| Polymyxin | >32/32    |
| Chloramphenicol | 16/16 |
| Trimethoprim/sulfamethoxazole | >2/38/2/38 |
| Colistin | ≤1       |

| TABLE 1 | MIC values for Ec265, transconjugant, and recipient strains. |
|----------|---------------------------------------------------------------|
| Antibiotic | MIC (µg/ml) |
| Ec265 | Tc265 | E. coli J53 |
|-----------------|-----------------|-----------------|
| Ampicillin | >16 | >16 | ≤8 |
| Ampicillin/sulbactam | >16/8 | >16/8 | ≤8/4 |
| Amoxicillin/clavulanic acid | >16/8 | >16/8 | ≤8/4 |
| Cephalothin | >32 | >32 | 16 |
| Cefotaxime | >16 | >16 | ≤4 |
| Ceftazidime | >32 | >32 | ≤1 |
| Cefepime | >16 | >16 | ≤2 |
| Aztreonam | <8 | <8 | ≤8 |
| Imipenem | 8 | 8 | ≤1 |
| Meropenem | >16 | >16 | ≤1 |
| Ertapenem | >2 | >2 | ≤1 |
| Gentamicin | >8 | >8 | ≤4 |
| Amikacin | >32 | >32 | ≤8 |
| Nalidixic acid | >16 | ≤16 | ≤16 |
| Ciprofloxacin | >2 | 0.12 | 0.12 |
| Levofloxacin | >4 | 0.5 | 0.5 |
| Nitrofurantoin | ≤32 | ≤32 | ≤32 |
| Fosfomycin | ≤32 | ≤32 | ≤32 |
| Chloramphenicol | 16 | 16 | ≤8 |
| Trimethoprim/sulfamethoxazole | >2/38 | >2/38 | ≤2/38 |
| Colistin | ≤1 | ≤1 | ≤1 |

| TABLE 2 | Genomic characteristics of Ec265 isolate. |
|----------|----------------------------------------|
| Characteristics | Escherichia coli Ec265 |
| Genome data | 5 171 045 |
| % GC content | 50.5 |
| N50 (bp) | 162 218 |
| Resistome Antibiotics | bla\textsubscript{NDM-5}, bla\textsubscript{TEM-1B} |
| Aminoglycosides | rrtB, aac(3)-IId, adaA2, aph(3’’)-Ib, aph (6)-Id |
| Trimethoprim | dfrA12, dfrA17 |
| Sulfonamides | sul1, sul2 |
| Macrolides | ermA(B), mhp(A) |
| Tetracycline | tet(B) |
| Chromosomal point mutations | |
| Fluoroquinolones | parE (I355T), parC (S80I, D84G), gyrA (S83L, D87N) |
| Acquired virulence factors | air, eIA, gad, lpf\textsubscript{A}, mcmA |
| Plasmids | IncFIA, IncFIB, IncFII, IncQ1, Col, and Col156 |
| GenBank accession number | JACXX1000000000 |
type plasmid is of concern considering its known capacity of efficient spread among bacteria (Bonnin et al., 2012).

The contig in which blaNDM-5 was found according to the assembly (contig 54, 10 926 bp) was annotated with RAST software in order to determine its genetic context. A truncated insertion sequence ISAba125 was found upstream of blaNDM-5, whereas bleomycin resistance gene (bleMBL), that encodes a bleomycin resistance protein (BRP), was observed downstream (Figure 1B). These two features are shared with other common genetic contexts of blaNDM (Wu et al., 2019). Some of the elements of NDM-5 genetic context (ISAba125 truncated sequence, IS91-family transposase, and Tn21 Urf2) code for transposase products, that may have had a role in the mobilization of this gene from other genetic platforms. Genes encoding BRP and NDM are co-expressed from the same promoter (Dortet et al., 2017), and could be co-selected either by bleomycin-like molecules (cancer treatment drugs) or carbapenems. Further downstream of bleMBL, there are a set of several genes, including trpF (encoding a phosphoribosyl anthranilate isomerase), dsbD (encoding a protein disulfide reductase), and a typical class 1 Integron containing sul1, qacE, aadA2, and dfrA12 resistance genes. Finally, the genetic region containing bleNDM-5 and class 1 Integron is flanked by two copies of IS26 transposase that delimit a small resistance island.

Ec265 blaNDM-5 context is the same as the one observed in a clinical strain of E. coli from France containing blaNDM-1 (Figures 1A, B). Furthermore, considering the blaNDM-5 contexts of greater identity corresponding to clinical strains from other countries around the world (Canada, Myanmar, and Italy), they did not show the presence of Urf2 and aldA elements (Figures 1B, C). All these strains mentioned above contain IncFII-type plasmids harboring blaNDM-1 or blaNDM-5.

Two genes codifying a proton antiporter (cdu2) and the chaperonin GroEL were located downstream of rmtB (contig 87). This 1.68 kb-arrangement was 100% identical to other rmtB-containing Enterobacteriales (GenBank accession nos. CP050367, MN061455, MN007141) recovered from clinical samples.

According to Achtman scheme, Ec265 belongs to ST9693 (allelic profile: adk-85, fumC-88, gyrB-78, icsA-29, mdh-59, purA-5, recA-62) and according to Pasteur database, it belongs to ST39 (allelic profile: dinB-13, icsA-39, pabB-11, polB-16, putP-12, trpA-25, trpB-8, uidA-19). Ec265 reported in this study do not belong to STs commonly associated to E. coli NDM-producing isolates elsewhere (such as ST101, ST167, ST131, ST405, ST410, and ST648) (Dadashi et al., 2019). EnteroBase software, which works with Achtman scheme (7 MLST genes), showed that ST9693 belongs to CC354 and to date, there was only one more E. coli ST9693 isolate record from Norway (April 2021). The analysis of closely related STs, such as SLVs, displayed a total of 477 E. coli isolates, of which 457 (96%) belonged to the international ST354. The inclusion of DLVs rendered 31 more isolates (a total of 508), grouped in 29 different STs belonging to CC354.

ST354 has been previously reported in carbapenem-resistant E. coli clinical isolates harboring NDM-5 (Zhang et al., 2016; Aung et al., 2018) and KPC (Zouh et al., 2018) in Asian countries meanwhile only in KPC-producing E. coli ST2287, a SLV of ST354 complex, in USA (Chavda et al., 2016). Although ST354 has been associated with E. coli isolates resistant to several antibiotics, recovered from humans (Dadashi et al., 2019) and animals (Guo et al., 2015; Zhuge et al., 2020), it is not considered a high-risk clone.

**DISCUSSION**

To our best knowledge, even if NDM-5-producing clinical isolates have been reported in other countries around the
world, so far in Latin America, essentially $bla_{NDM-1}$ has been reported (Wu et al., 2019), while $bla_{NDM-5}$ was only described in Brazil in an Enterobacter bugandensis isolate from an environmental sample (Matteoli et al., 2020). This work would be the first report of a clinical isolate of carbapenem-resistant E. coli carrying NDM-5 in our country and region, that also displays resistance to amikacin and fluoroquinolones. It is important to highlight that previously, the replacement of ESBLs in Argentina (CTX-M-2 by CTX-M-15 hegemony shift) happened in an unnoticed way until detected by searching for specific resistance markers instead of general resistance mechanisms (Sennati et al., 2012) showing the significance of reporting novel variants to understand the changing epidemiology of countries and regions. Besides, we demonstrated $bla_{NDM-5}$ localization in a conjugative plasmid, raising an alert about the potential dissemination of this resistance marker to a high-risk clone, in addition to the fact that this NDM variant display higher levels of resistance when compared to NDM-1 (Hornsey et al., 2011).

The co-transfer of multiple antimicrobial resistance genes represents a particular challenge for clinical treatment in health care settings, and the spread of isolates resistant to last resort antibiotics, such as carbapenems and amikacin, should be a global warning in public health that deserves close monitoring.

**DATA AVAILABILITY STATEMENT**

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found below: https://www.ncbi.nlm.nih.gov/genbank/, JACXX1000000000.

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GG and JDC conceived and designed the experiments. AC, BG, RF-E, GG, and JDC wrote the manuscript. AC, RF-E, and JDC analyzed the data. FG supplied the clinical isolate and performed the phenotype characterization. AC performed the experiments. NL and BF performed WGS. All authors contributed to the article and approved the submitted version.

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**Conflict of Interest:** 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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