Characterization of an IncFII Plasmid Encoding NDM-1 from Escherichia coli ST131

Rémy A. Bonnin1, Laurent Poirel1, Alessandra Carattoli2, Patrice Nordmann1*

1 Service de Bactériologie-Virologie, INSERM U914 «Emerging Resistance to Antibiotics», Hôpital de Bicêtre, Assistance Publique/Hôpitaux de Paris, Faculté de Médecine et Université Paris-Sud, Paris, France,
2 Department of Infectious, Parasitic and Immuno-Mediated Diseases, Istituto Superiore di Sanità, Rome, Italy

Abstract

Background: The current spread of the gene encoding the metallo-β-lactamase NDM-1 in Enterobacteriaceae is linked to a variety of surrounding genetic structures and plasmid scaffolds.

Methodology: The whole sequence of plasmid PGUE-NDM carrying the blaNDM-1 gene was determined by high-density pyrosequencing and a genomic comparative analysis with other blaNDM-1-negative IncFII was performed.

Principal Findings: Plasmid PGUE-NDM replicating in Escherichia coli confers resistance to many antibiotic molecules including β-lactams, aminoglycosides, trimethoprim, and sulfonamides. It is 87,022 bp in-size and carries the two β-lactamase genes blaNDM-1 and blaOXA-1, together with three aminoglycoside resistance genes aacA4, aadA2, and aacC2. Comparative analysis of the multidrug resistance locus contained a module encompassing the blaNDM-1 gene that is actually conserved among different structures identified in other enterobacterial isolates. This module was constituted by the blaNDM-1 gene, a fragment of insertion sequence ISAb125 and a bleomycin resistance encoding gene.

Significance: This is the first characterized blaNDM-1-carrying IncFII-type plasmid. Such association between the blaNDM-1 gene and an IncFII-type plasmid backbone is extremely worrisome considering that this plasmid type is known to spread efficiently, as exemplified with the worldwide dissemination of blaCTX-M-15-borne IncFII plasmids.

Introduction

Metallo-β-lactamase (MBL) NDM-1 (New Delhi Metallo-β-lactamase) corresponds to one of the latest and most important resistance trait identified in Gram-negative rods [1]. The blaNDM-1 gene has been first identified in the UK, India, and Pakistan, either in various enterobacterial isolates, but also in Acinetobacter sp., Pseudomonas sp., or Vibrio sp. [2–6]. Occurrence of NDM-1 producers in hospitalized patients in the UK was related in many cases with previous hospitalizations in the Indian subcontinent [2]. Additionally, there have been numerous reports of NDM-1-producing Enterobacteriaceae worldwide (infections or colonizations), most often recovered from patients who presented a link with the Indian subcontinent, and in some cases with the Balkans and Middle-East region [6–11].

Five blaNDM-1-bearing plasmids have been fully sequenced, being pHK-NDM (Genbank n’ HQ451074), p271A [JF785549], pNDM-1_Dok01 (AP012208), pNDM10505 [JF503991] and pNDM-KN (JN157804), respectively corresponding to IncI/M-, IncN2-, and IncA/C-type plasmid scaffolds [12–16]. pHK-NDM is an IncI/M 88,803-bp in-size plasmid harboring the blaNDM-1 gene together with the 16S rRNA methylase armA gene, thus conferring high level of resistance to β-lactams and aminoglycosides, and is highly related to plasmids carrying the blaCTX-M-3 gene [12]. Plasmid p271A is 35,947-bp in size and harbors the blaNDM-1 gene as a single resistance gene within an IncN2-type plasmid scaffold on which a new replicase gene was identified [13]. Plasmids pNDM-1_Dok01, pNDM10505 and pNDM-KN belong to the IncA/C broad-host range plasmid family. In addition to the blaNDM-1 gene, those plasmids carry additional resistance genes including a blaCMY-2-like gene, together with an armC or armA 16S RNA methylase gene. Their scaffolds are very similar to those of other IncA/C but blaNDM-1-negative plasmids, known to be responsible for the spread of blaCMY-2-like genes in Enterobacteriaceae in the USA, Canada, and Europe [14,15].

Overall, the blaNDM-1 gene has been more frequently reported onto broad host-range IncA/C-type plasmids, either in clinical or in environmental isolates recovered from the New Delhi area [6]. In addition, some other plasmid scaffolds have been associated with the blaNDM-1 gene including IncF, IncL/M, together with untypeable plasmids [6,8]. In particular, IncF-type plasmids were involved in blaNDM-1 acquisition in isolates recovered in France, India, and Switzerland [8].

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* E-mail: nordmann.patrice@bct.aphp.fr
IncFII-type plasmids are narrow-host range plasmids that are frequently identified among *E. coli* strains [16]. These plasmids are known to be involved in the worldwide dissemination of the *bla*<sub>CTX-M-15</sub> gene in the epidemic ST131 *E. coli* clone [17]. They are characterized by several toxin-antitoxin addiction systems conferring stability during bacterial cell division. The IncFII-type pC15-1a (GenBank n° NC_005327) and pEK516 (EU935738) plasmids carrying the *bla*<sub>CTX-M-15</sub> gene have been fully sequenced [17,19]. In particular, plasmid pEK516 obtained from the UK epidemic CTX-M-15-producing strain D represents the prototypic IncFII-plasmid from the MLST-defined ST131 *E. coli* lineage [18]. Unlike plasmids belonging to other incompatibility groups, the backbones of IncF plasmids exhibit a significant heterogeneity in terms of size and number of replicons [19].

The aim of this study was to characterize in detail an IncFII-type plasmid harboring the *bla*<sub>NDM-1</sub> gene, in order to possibly identify genetic features explaining the successful spread of that gene. That plasmid had been recovered from an *E. coli* isolate belonging to ST131 [20] that corresponds to the main genetic background involved in the worldwide distribution of CTX-M-15 producers [21].

**Results**

**General features**

Our study was initiated by the isolation of a multidrug-resistant *E. coli* strain GUE that had been community-acquired in India [20]. *E. coli* isolate GUE was resistant to most β-lactams (remaining susceptible to aztreonam) and showed reduced susceptibility to carbapenems, minimum inhibitory concentrations (MIC) of imipenem, ertapenem, and meropenem being at 3, 3 and 2 µg/ml, respectively (Table 1). It was also resistant to gentamicin, kanamycin, tobramycin, sulfonamides, tetracycline, and fluoroquinolones, but remained susceptible to amikacin, chloramphenicol, rifampicin, and colistin.

### Table 1. MICs of β-lactams for *E. coli* clinical isolate Gue, *E. coli* J53 pGUE-NDM and *E. coli* J53 reference strains.

| β-lactams         | MIC (µg/ml) | *E. coli* isolate GUE | *E. coli* J53 pGUE-NDM | *E. coli* J53 |
|-------------------|-------------|------------------------|------------------------|--------------|
| Amoxicillin       | >256        | >256                   | 4                      |
| Amoxicillin + CLA*| 256         | >256                   | 4                      |
| Ticarcillin       | >256        | >256                   | 2                      |
| Ticarcillin + CLA*| 256         | 256                    | 2                      |
| Cephalothin       | >256        | >256                   | 4                      |
| Ceftoxatine       | 128         | 128                    | 0.06                   |
| Ceftazidime       | >256        | >256                   | 0.06                   |
| Cefepime          | 32          | 32                     | 0.03                   |
| Aztreonam         | 0.125       | 0.06                   | 0.06                   |
| Meropenem         | 2           | 0.75                   | 0.03                   |
| Ertpenem          | 3           | 0.5                    | 0.03                   |
| Doripenem         | 1.5         | 0.5                    | 0.03                   |
| Imipenem          | 3           | 1.5                    | 0.12                   |

*CLA*: clavulanic acid (4 µg/ml) TZB. doi:10.1371/journal.pone.0034752.t001

Plasmid features

Conjugation assays allowed to transfer the *bla*<sub>NDM-1</sub> gene and identified a ca. 87-kb plasmid named pGUE-NDM. The *E. coli* transconjugant showed resistance to penicillin/inhibitor combinations and broad-spectrum cephalosporins, and reduced susceptibility to carbapenems (Table 1). It was also resistant to kanamycin, gentamicin, tobramycin, trimethoprim, and sulfonamides. Plasmid pGUE-NDM was assigned to the IncFII incompatibility group using the PCR-Based Replicon Typing method. Conjugation frequencies were observed at high rates (2×10<sup>-7</sup> transconjugants per donor cell).

**Sequence analysis of plasmid pGUE-NDM**

Whole plasmid sequencing identified plasmid pGUE-NDM to be 87,022 bp in-size with a GC content of 53% and 47 open reading frames (ORF) (Figure 1 and Table S1). BLAST analysis of the complete nucleotide sequence was performed in comparison with the reference IncFII plasmid pC15-1a (Genbank n°AY458016) and with plasmid pEK516 UK (Genbank n° EU935738). Comparative DNA sequence analysis confirmed that pGUE-NDM possessed an IncFII-type backbone and exhibited a significant synteny within the two scaffolds, with the exception of regions containing accessory genes (Table S1).

The backbone of pGUE-NDM included a complete array of genes involved in replication, conjugation and partition. The replication operon was composed by four ORFs exhibiting high identity with IncFII-specific replicase genes. The toxin/antitoxin addiction systems system *pemJ/pemK* and *hok/mok* were identified as previously described for other FI-plasmids (Figure 1), together with the *parB*, *parC* genes encoding proteins involved in plasmid stability (Figure 1 and Table S1). A complete transfer operon (locus *tra-ribo*) was identified being involved in the plasmid dissemination.

The *bla*<sub>NDM-1</sub> gene was localized in a multidrug resistance (MDR) region of 20,181 bp. This region was bracketed by two copies of insertion sequence IS26 in opposite orientations creating an IS26-made compound transposon that was not bracketed by a target site duplication. Immediately upstream of the *bla*<sub>NDM-1</sub> gene, a remnant of IS<sub>Aba125</sub> insertion sequence was identified (Figure 2). This truncated IS<sub>Aba125</sub> contained the ~35 promoter sequence leading to the *bla*<sub>NDM-1</sub> gene expression [13]. Upstream of the IS<sub>Aba125</sub>, two IS26 were identified bracketing the *aacC2* aminoglycoside resistance gene. Then, three gene cassettes being part of a remnant of a class 1 integron, namely *aacC4*, *bla*<sub>NDM-1</sub>, and a truncated *catB4* genes were identified, that latter gene being bracketed by another IS26 element (Figure 2). Downstream of the *bla*<sub>NDM-1</sub> gene, the *bla*<sub>MBL</sub> gene encoding resistance to bleomycin was identified followed by a truncated phosphoribosylanthranilate isomerase gene (*AisO*), and then by a truncated twin-arginine translocation pathway signal protein gene (*AisFt*), as previously found on other plasmid scaffolds (Figure 3). Then, the IS<sub>CR1</sub> insertion sequence was identified [22], followed by a class 1 integron structure containing three gene cassettes, namely *aadA2* encoding resistance to streptomycin and spectinomycin, *orfE* of unknown function, and *dfrA12* encoding resistance to trimethoprim, and then the class 1 integrase gene. Finally, an additional copy of IS26 truncating the *Tn1721* transposase gene was identified (Figure 2). Analysis of surrounding sequences of each IS26 elements revealed that none of them was bracketed by direct repeat (DR) sequences. Similarly, no DR was identified that would suggest that some of these elements may form an IS26-made transposons. This result reinforced the hypothesis of successive homologous recombination events at the origin of the global structure of this MDR region.
Evolution of the blaNDM-1 genetic context

In order to get further insights into the blaNDM-1 gene acquisition, a comparison of the genetic structures previously identified with that identified on plasmid pGUE-NDM was performed (Figure 3). Interestingly, a common module that we termed NDM module was systematically identified. This module was composed of the ISAba125 fragment containing the 235 promoter region, the blaNDM-1 gene, the bleomycin resistance gene bleMBL, and a truncated phosphoribosylanthranilate isomerase. Recently, composite transposon Tn125 has been identified in several A. baumannii isolates [23,24]. This 10,099-bp in-size transposon of was made of two copies of ISba125 bracketing a 7,925-bp fragment and was inserted into the chromosone of different strains (Figure 3). Transposon Tn125 also contained the NDM module, together with the groES and groEL genes and the ISCR21 insertion sequence (Figure 3). It seems that the NDM module has integrated the pGUE-NDM backbone by a recombination event mediated by IS26 elements. Overall, a series of IS26-mediated recombination events has likely been at the origin of the formation of the large resistance gene array identified on plasmid pGUE-NDM.

Discussion

The blaNDM-1 gene is now identified worldwide and it has been speculated that particular genetic features could be at the origin of that wide diffusion. After the characterization of IncA/C, IncN2, and IncL/M plasmids bearing the blaNDM-1 gene, our study characterized an IncFII plasmid which backbone is known to be a major vehicle for dissemination of the blaCTX-M-15 gene [17]. It might be hypothesized that the endemicity of the blaCTX-M-15-positive and IncFII plasmid in the Indian subcontinent has created a favorable context for hosting and therefore disseminating the blaNDM-1 gene. That finding is particularly threatening when considering the explosive diffusion of the blaCTX-M-15 gene that has been witnessed worldwide during the last decade.

Analysis of the genetic features of plasmid pGUE-NDM did not identify any particular element that would positively or negatively interfere into its spreading potency such as addiction systems, partitioning systems or virulence factors in comparison to others IncFII-type plasmids. Indeed, the backbone part of the plasmid (being replication system, maintenance systems and transfer systems) corresponded to one previously identified. However, it appeared that acquisition of the blaNDM-1-containing module resulted from a series of IS26-related recombination events. Our study further underlines that the current dissemination of the blaNDM-1 gene is associated to a variety of genetic backgrounds.

Materials and Methods

Antimicrobial agents and MIC determinations

Susceptibility testing was performed by disk diffusion assay (Sanofi-Diagnostic Pasteur, Marnes-la-Coquette, France) and interpreted according to CLSI [25]. The MICs for carbapenems were determined by Etest (AB Biodisk, Solna, Sweden) on Mueller-Hinton agar plates at 37°C. The production of MBLs was evaluated using Etest, combining imipenem and EDTA as recommended by the manufacturer (AB biolMérieux).
Plasmid identification and preparation

Plasmid pGUE-NDM was assigned to the IncFII incompatibility group using the PCR-Based Replicon Typing (PBRT) method [26]. Conjugation experiments were performed as previously described using azide-resistant *E. coli* strain J53 [27]. Transconjugants were selected on Trypticase-Soy media containing 100 µg/ml of ticarcillin and 100 µg/ml of sodium azide.

High-density pyrosequencing and sequence assembly

Plasmid DNA was isolated from the *E. coli* transconjugant using Qiagen Maxiprep kit (Qiagen, Courtaboeuf, France). The complete sequencing work flow of the Illumina Genome Analyzer IIX system (Illumina Inc., San Diego, CA) was performed by the DNAVision company (Gosselles, Belgium) and is described at www.dnavision.com.

Genome assembly and annotation

Reads from each sample were trimmed to remove poor quality sequence using the following procedure: keeping only reads for which the quality of all the fifty first bases is greater or equal to ten in both reads; for each read trimming the tail of the sequence from the first position for which the quality is smaller than ten; trimming poly-A sequence artifacts; and trimming adapter sequences. Then, the assemblies were carried out using Velvet2 assembler [28] in order to produce contigs from Illumina GAIIX reads. A total of ca. 12,000 high-quality reads were derived from the pGUE-NDM library, of which 482 contig were obtained using Velvet2 assembler including 14 contigs *de novo*. Contigs were considered as DNA contamination if showing at least 90% similarity with *E. coli* K12 substrain MG1655 genomic DNA sequences (NCBI accession number NC_000913.2) using the BLASTn algorithm. Among the 482 contigs, 430 were considered as DNA contaminations. A single contiguous sequence was obtained from the 52 contigs obtained using PCR-based gap closure et PCR combinations to close gaps and verify the position of each contig.

Genome comparison

The BLASTp algorithm was used to search for protein similarities by using as a reference the *E. coli* K12 chromosome sequence. The criterion used to evaluate the deduced amino acid sequence homology was >50% similarity at the amino acid level and >50% coverage of protein length.

Nucleotide sequence accession numbers

The pGUE-NDM plasmid sequence was submitted to the GenBank database and can be found under accession number JQ364967.

Transparency declaration

None to declare.

Supporting Information

Table S1 Orfs identified in pGUE-NDM. (DOC)

Author Contributions

Conceived and designed the experiments: RAB LP PN. Performed the experiments: RAB. Analyzed the data: RAB LP AC. Contributed reagents/materials/analysis tools: LP PN. Wrote the paper: RAB LP PN.
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