Genetic Acquisition of NDM Gene Offers Sustainability among Clinical Isolates of *Pseudomonas aeruginosa* in Clinical Settings

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Abstract

New Delhi metallo β-lactamases are one of the most significant emerging resistance determinants towards carbapenem drugs. Their persistence and adaptability often depends on their genetic environment and linkage. This study reports a unique and novel arrangement of *bla*NDM-1 gene within clinical isolates of *Pseudomonas aeruginosa* from a tertiary referral hospital in north India. Three NDM positive clonally unrelated clinical isolates of *P. aeruginosa* were recovered from hospital patients. Association of integron with *bla*NDM-1 and presence of gene cassettes were assessed by PCR. Genetic linkage of NDM gene with IS*Ab*125 was determined and in negative cases linkage in upstream region was mapped by inverse PCR. In which only one isolate’s NDM gene was linked with *bla*NDM-1 and presence of gene cassettes were assessed by PCR. Genetic linkage of NDM gene with IS*Ab*125 was determined and in negative cases linkage in upstream region was mapped by inverse PCR followed by sequencing analysis. In continuation significance of this novel linkage was further analyzed wherein promoter site detected by Softberry BPROM software and activity were assessed by cloning succeeding semi-quantitative RT-PCR indicating the higher expression level of NDM gene. This study concluded out that the unique genetic makeup of NDM gene with DNA-dependent-RNA-polymerase favours adaptability to the host in hospital environment against huge antibiotic pressure.

Introduction

The New Delhi Metallo-β-lactamase (NDM) has emerged as a major carbapenemase with rapid dissemination worldwide [1]. Clinical isolates harbouring NDM gene are often referred as superbugs. The NDM gene has been identified usually in enterobacteriaceae and recently in *Pseudomonas* spp. [2] which expresses resistance towards carbapenems and represents a significant threat for clinicians. This situation has been further complicated by the association...
of NDM gene with other resistant determinants [3]. Most importantly, however they represent an important stage in the evolution of antibiotic era.

Transfer of the \textit{bla}\textsubscript{NDM} among promiscuous plasmids is the major dissemination route including clonal outbreaks. Noticeably, \textit{bla}\textsubscript{NDM} has been frequently identified from unrelated gram negative bacilli, harboured by different plasmid types [4]. However, the mechanism of multiresistance trait of NDM positive isolates remained wanted and speculated that it could have been captured from original chromosomal location by mobile genetic elements. It has been reported that the \textit{bla}\textsubscript{NDM} can be carried by different plasmid types (IncA/C, IncF, IncL/M, or untypable) and occasionally found to be chromosomally integrated [5]. The sequence of few plasmids carrying \textit{bla}\textsubscript{NDM} are now available and reveals its inimitable genetic features such as association with insertion sequences and transposons for high mobility, acquisition of other multiresistance regions (\textit{aadB}, \textit{dfrA12}, \textit{bla}\textsubscript{OXA-30}, \textit{aacA4}) including additional important enzymes which make it more versatile [6].

In this study we have presented a unique genetic makeup of \textit{bla}\textsubscript{NDM} gene among clinical isolates along with other resistance factors. However, this novel finding suggests the insertion of \textit{bla}\textsubscript{NDM} within DNA directed RNA polymerase gene, thus favouring its survivability within hospital.

**Materials and Methods**

**Bacterial isolates & Carbapenem susceptibility**

A total of 105 consecutive, non-duplicates, carbapenem non-susceptible isolates of \textit{P.aeruginosa} were collected from indoor patients of Sir SunderLal Hospital, Banaras Hindu University, Varanasi, India, during March 2011 to September 2011. Identification of organisms was done by the conventional methods [7].

The metallo β-lactamase status of the strains was established by the Imipenem—Ethylene diamine tetra-acetic acid (EDTA) disc potentiation method [8]. A previously confirmed and sequenced clinical isolate of \textit{E. coli} harbouring \textit{bla}\textsubscript{NDM} was taken as positive control [9] and \textit{E. coli} ATCC 25922 was used as negative control.

All MBL positive isolates were suspended in 1mL of buffered peptone water supplemented with 30% glycerol (peptone glycerol) and were kept at −80°C. The isolates were also stored in minimal media (10% Peptone, 5% Sodium Chloride and 0.8% Agar) as stab culture. ERIC-PCR was performed for genotyping by using their respective primers to determine clonal relatedness of the isolates [10].

**Ethics Statement**

This work and obtained samples specifically for this study has been ethically approved by the chairperson of the ethical committee of Institute of Medical Sciences, Banaras Hindu University, Varanasi; Ref. No.-Dean/2012-13/114.

**Genotypic detection of \textit{bla}\textsubscript{NDM} and their association with mobile genetic elements**

Genomic DNA for PCR was extracted by using QIAamp DNA mini kit (Qiagen, Germany). Genotypic detection of \textit{bla}\textsubscript{NDM} was performed by PCR assay in all the MBL positive isolates. Reaction conditions and primers used for PCR amplification were as described earlier [11]. Consecutively, the presence of integrons were examined by PCR, using primers to amplify a 160 bp fragment for class 1 \textit{integrase}, and 288 bp fragment for class 2 \textit{integrase}. The primers, PCR conditions and reaction mixtures used were as described previously [12]. To find the
Genetic association of \( \text{bla}_{\text{NDM}} \) gene with integrons, PCR was performed using forward primer of the conserved region (5’CS) of integron gene and reverse primer of the characterized \( \text{bla}_{\text{NDM}} \) gene [12]. Integrons contain multiresistant regions known as gene cassettes which were determined by 59 base elements PCR (Table 1)[13].

Insertion sequences (IS), one of the major genetic elements in transposition of antibiotic resistance genes. In order to assess the linkage of NDM gene with the insertion sequence in our study, PCR analysis was performed by using forward primer of IS\(_{\text{Aba125}}\) (Acinetobacter specific), and reverse primer of NDM gene. The reaction mixture and running conditions were as described earlier [14].

Detection of Novel linkage by inverse PCR

In order to determine the genetic environment surrounding the \( \text{bla}_{\text{NDM-1}} \) gene (apart from their linkage with IS\(_{\text{Aba125}}\) and integrons) inverse PCR was performed. DNA was extracted from the three strains and digested with Sau3AI (Biolabs, US). DNA fragments obtained were then autoligated at 16°C with T4 DNA ligase in higher dilution to allow self circularization. The fragment of DNA containing the \( \text{bla}_{\text{NDM-1}} \) gene was used as a template for an inverse PCR with primers designed from the \( \text{bla}_{\text{NDM-1}} \) gene sequence. Inv1F as forward and Inv1R as reverse primer [Table 1] were used for amplification. Reaction mixture was approximately 5 ng of template DNA, 10 pmol of each primer, 200 \( \mu \text{M} \) deoxynucleoside triphosphate (dNTP) mix, 2 mM MgCl2 and of Taq DNA polymerase in the reaction buffer supplied with the enzyme. Reaction conditions were: initial denaturation at 94°C for 3 min followed by 32 cycles at 94°C for 30 s, 58°C for 45 s, 72°C for 1 min 25 s, and final extension at 72°C for 8 min.

Once the sequence of amplified product of inverse PCR was obtained, the linkage was confirmed by using POL’F 5’ACGTCAGAGCGATGAGACG-3’ as forward primer and Inv1R 5’ATTCGTCGGCGGATGAGG-3’ as reverse primer to detect 5’ region of target site, followed by second PCR reaction with NDM 3PF 5’GATCGTGATGAGGCCATTCGC-3’ as forward primer and POL’R 5’GACCTGGAACTGACCGTACG-3’ as a reverse primer to confirm 3’ target region. To further establish this linkage, the PCR reaction was performed using POL’F as forward and POL’R as reverse primer among NDM positive isolates.

Southern Hybridization

Simultaneously, to validate our study Southern blotting was performed on agarose gel by in-gel hybridization [15] with the \( \text{bla}_{\text{NDM}} \) probe labelled with DIG HIGH PRIME LABELING MIX.

### Table 1. List of primers used in the study.

| Primers | Target | Sequence (5’–3’) | Reference |
|---------|--------|------------------|-----------|
| HS287 F HS286 R | 59be association | GGGATCCGCGCTGCTANCTCVRRCGTAGASC GGGATCCGCGCTGCTANCTCVRRCGTAGASC | 12 |
| Inv 1F Inv 1R | To amplify flanking region | ATGGAAACTGGCGACCAACGG AATCGTGCGCGGCGATGACCACCA | Present study |
| POL’F POL’R | To amplify flanking region | ACGTCAGAGCGATGAGACG GACCTGGAACTGACGTACG | Present study |
| NDM 3P | 3’ end of linkage with DNA directed RNA Polymerase reverse | GATCGTGATGACCGCATTCGCC | Present study |
| Native Pm Mutated Pm NDMc | To amplify RNA polymerase promoter along with integrated NDM gene | AACCTGATTGTCGAGCCTCTACTC CTGGTCTACTCCAGTAAAACA* CATCGAAATCGCGGATGGAG | Present study |

*underlined regions are mutated bases

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(ROCHE, Germany) detection Kit. The digoxigenin-labeled \textit{bla\textsubscript{NDM}} specific probe was prepared using primers (Forward NDM 5’GGGCAGTCGCTTCCAACGGT 3’ and Reverse 5’CGACCGGCAGGTTGATCTCC 3’) that amplify a 130 bp region of the \textit{bla\textsubscript{NDM}} gene. Total genomic DNA was digested by HindIII for fragmentation of DNA which is followed by the transfer to nylon membrane (Hybond N, Amersham, UK) and then hybridised with prepared \textit{bla\textsubscript{NDM}} specific probe. Detection was performed by using an NBT color detection kit (ROCHE, Germany). Promoter activity was assessed by designing two sets of primers; in first set forward primer (RNAPROM) was designed from upstream region of the RNA polymerase promoter whereas in the second set the primer with desired mutation was generated (MUTPROM) in the promoter region. For both the sets the reverse primer used was NDMc [Table 1]. The PCR amplicons were sequenced to confirm and cloned using p\textsuperscript{GEM-\textsubscript{T}} vector (Promega, Germany) in E. coli JM107. Further, RNA was isolated from both the constructs using RNeasy\textsuperscript{R} Mini Kit (Qiagen Hilden, Germany) and cDNA was prepared using QuantiTect Reverse Transcription kit (Qiagen, Hilden, Germany). Thereafter, Semi quantitative reverse transcriptase PCR was performed to determine the expression of \textit{bla\textsubscript{NDM}} in both the constructs.

**Transferability & plasmid profiling**

Horizontal transferability of \textit{bla\textsubscript{NDM}} was investigated by transformation assay. Conjugation experiment was carried out between clinical isolates as donors and streptomycin resistance E. coli recipient strain B (Genei, Bangalore, India). Overnight culture of bacteria were diluted in Luria Bertani broth (Hi-Media, Mumbai, India) and was grown at 37°C till the O.D. of the recipient and donor culture reached 0.8–0.9 at A\textsubscript{600}. Donor and recipient cells were mixed at 1:5 donor-to-recipient ratios and transconjugants were selected on imipenem (0.25mg/L) + streptomycin (1000mg/L) agar plates. Transformation was carried out using E. coli JM107 as recipient. Transformants were further selected on imipenem (0.25mg/L) containing LB agar plates.

For the detection of incompatibility group type of plasmid in transformants with \textit{bla\textsubscript{NDM}} as well as in \textit{bla\textsubscript{NDM}} harbouring donor strains, PCR based replicon typing was carried out targeting 18 different replicon types [16]. Plasmid stability and fitness was also assessed by serial passage of \textit{bla\textsubscript{NDM}} positive isolates on LB broth without any antibiotic pressure. After each passage the isolates were tested for the presence of \textit{bla\textsubscript{NDM}} PCR assay.

**Detection of other resistant determinants**

To investigate the presence of other resistant determinants along with NDM, isolates were further tested for the co-existence of other MBL genes such as \textit{bla\textsubscript{IMP}} and \textit{bla\textsubscript{VIM}}, as well as ESBLs, AmpC and class D carbapenemase (OXA-48, OXA-58, OXA-23 and OXA-198) genes by multiplex PCR [17–23]. In addition, the study of efflux pump activity of the strains was phenotypically detected by double disc synergy test using meropenem and CCCP (carbonyl cyanide m-chlorophenylhydrazone) [24]. Consequently the expression of the Mex-efflux system was determined by quantitative real time PCR assay. PA01 strain was used as control [25].

**Antibiotic susceptibility testing**

Antimicrobial sensitivity testing was performed on Mueller-Hinton agar (Hi-Media, Mumbai, India) plates by Kirby Bauer disc diffusion method and interpreted as per CLSI recommendations [26]. The antibiotic tested were amikacin (30μg), gentamicin (10μg), netilmicin (30μg), tobramycin (10μg), ceftazidime (30μg), ciprofloxacin (5μg), imipenem (10μg), meropenem (10μg), piperacillin/tazobactum (100/10μg) and polymyxin B (300μg) (Hi-Media, Mumbai, India). MICs of all the isolates were determined by the agar dilution method against
cefotaxime, ceftazidime, ceftriaxone (Hi-Media, Mumbai, India), cefepime (Alembic Ltd., Vadodara, India), aztreonam (Aristo Pharmaceuticals Ltd., Mumbai, India), imipenem (United Biotech, Solan, India), meropenem (AstraZeneca Pharmaceuticals Ltd., Bangalore, India), tigecycline (Taj Pharmaceuticals Ltd., Mumbai, India), polymyxin (Celon laboratories Ltd., Andhra Pradesh, India).

*Escherichia coli* ATCC 25922 was used as a control.

### Sequencing analysis

All the PCR amplicons were purified using the QIAquick Gel Extraction kit (QIAGEN Inc., Valencia, CA) and were subjected to DNA sequencing (Merck, Bangalore, India). Sequences were analyzed using the BLAST suite of programs. ([http://www.ncbi.nlm.nih.gov/BLAST/](http://www.ncbi.nlm.nih.gov/BLAST/)). Further, promoter sites were also determined by using Soft Berry BPROM software ([http://linux1.softberry.com/berry.phtml?topic=bprom&group=programs&subgroup=gfindb](http://linux1.softberry.com/berry.phtml?topic=bprom&group=programs&subgroup=gfindb)).

### Results

#### Genetic context of *bla*<sub>NDM</sub>

Among the 105 isolates, 38 strains exhibited MBL activity by Imipenem EDTA disc potentiation method, of which three nonclonal isolates (PA6, PA38 and PA47) were found to harbour *bla*<sub>NDM-1</sub> ([Table 2](#tab2)) and were carrying class 1 integron too. While out of the 35 MBL positive but *bla*<sub>NDM</sub> negative isolates, 7 and 3 isolates were harbouring VIM and IMP gene respectively. Remaining phenotypically MBL positive isolates did not showed any amplification with the targeted MBL gene primers (IMP, VIM, NDM). Further, class 1 integron was detected among the 99/105 (94.28%) isolates. None of the study isolates were found carrying class 2 integrase.

On performing the ERIC PCR for all the 105 isolates a total of 46 different clonal types were observed. However two non MBL strains were found to be clonal with PA47.

However, among the three NDM harbouring isolates 59b<sup>e</sup> PCR amplification and their sequencing revealed presence of dihydrofolate reductase (*dhfr*) and aminoglycoside acetyl transferase (*aac(6)’*) within the gene cassette. While analysing the upstream linkage of NDM gene with ISA<sub>ba125</sub>, isolate PA6 formed a band of 850 bp thus showing the association between them which was confirmed by sequencing. However, in case of PA38 and PA47 inverse PCR result gave a novel finding that *bla*<sub>NDM</sub> gene got integrated within DNA dependent RNA polymerase of host genome. This was further confirmed by designing primers which could amplify both 5’ and 3’ region of the insert ([Fig. 1](#fig1)). This insertion was further verified and supported by the fact that the amplified product of higher basepair with Pol’F and Pol’R could be observed in PA38 and PA47 while expected size of amplification was evident with PA6 ([Figs. 2 and 3](#fig2 and #fig3)).

### Table 2. Clinical details of the isolates harbouring NDM gene.

| ID no. | Isolate          | Patient age/Sex | Clinical specimen | Ward/OPD/ICU | Current diagnosis | Carbapenem Susceptibility | Other susceptible antibiotics tested *in vitro* | Coproduction of Class C | Coproduction of Class C-NDM or other MBL genes |
|--------|------------------|-----------------|-------------------|--------------|------------------|---------------------------|-----------------------------------------------|-------------------------|---------------------------------------------|
| PA6    | *P. aeruginosa*   | 24year/female   | Tissue            | Burn         | sepsis           | R                         | Polymyxin B                                  | CMY-2                   | NDM                                        |
| PA38   | *P. aeruginosa*   | 25 year/female  | Central tip       | FICU         | VAP              | R                         | Polymyxin B                                  | ACT, CMY-2              | NDM                                        |
| PA47   | *P. aeruginosa*   | 29 year/female  | Pus               | Ortho        | sepsis           | R                         | Polymyxin B                                  | DHA, ACT                | NDM, IMP, VIM                              |

OPD, outpatient department; ICU, Intensive care unit; VAP, ventilator-associated pneumonia; FICU, Female intensive care unit R, resistant; Class C-Plasmid mediated AmpC β-lactamases.

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was also observed that the recombinant \( \text{bla}_{\text{NDM}} \) used promoter region of DNA dependent RNA polymerase for its expression (Fig. 4). Further, Semi quantitative reverse transcriptase PCR also establishes DNA dependent RNA polymerase promoter activity for higher expression of NDM gene (Fig. 5).

Results of Southern hybridization were evident to suggest genetic background of the \( \text{bla}_{\text{NDM}} \). Plasmids of all three isolates were transferred to the nylon membrane while only PA6 hybridized with NDM-1 specific probe. Whereas when the same experiment was done with total DNA of all the test isolates, it was observed that all of them hybridized with the probe showing identical banding pattern with PA38 and PA47. Transformation and conjugation was successful only with PA6, where transconjugant showed to carry \( \text{bla}_{\text{NDM}} \) within the 30–50kb plasmid. However, all the three isolates were carrying the plasmid of IncT type. The plasmid was highly unstable and was lost after six consecutive passages without any antibiotic in LB broth.

Coproduction of other resistant determinants

All the three isolates were negative for ESBLs but coproducing AmpC \( \beta \)-lactamase, while PA47 was carrying \( \text{bla}_{\text{IMP}} \) and \( \text{bla}_{\text{VIM}} \) along with NDM gene. All of them were harbouring class D carbapenemase as well (Table 2). On performing the quantitative real time PCR to assess the level of transcription in MexAB-OprM efflux pump system, it was observed that the transcription level was 16-fold higher while under inducing condition when compared with PA01.

Antimicrobial susceptibility

On observing the antimicrobial susceptibility of these NDM producers, the isolates showed resistance towards all the tested antibiotics namely; cephalosporins, aztreonam, carbapenems, aminoglycoside and fluoroquinolones but were susceptible against polymyxin B. MIC of carbapenem was much higher for PA38 and PA47 compare to PA6 (Table 3).
Discussion

Among the carbapenemases, the NDM gene has gained particular attention due to its global dissemination and multidrug resistance phenotype [27]. In the study, we found that all the three blaNDM harbouring strains were highly resistant to the antibiotics tested except polymyxin. Although the MBL enzymes donot affect monobactam, the co-existence of different AmpC genes such as blaEBC, blaDHA and blaCIT in these isolates were found to confer resistance to monobactam [14, 28]. Besides harbouring blaNDM the isolates were also carrying class D carbapenemases along with intrinsic mechanism. Similar genotype was earlier reported from India [29]. However, it leaves scope for future studies to address the exact role of six different carbapenem resistant determinants in a single isolate (PA47) when exposed to carbapenems. The study could demonstrate that blaNDM-1 in PA6 is plasmid mediated which was transferable by conjugation. However, same incompatibility type plasmid (lacking blaNDM-1) was present in other two isolates (PA38 and PA47). So, this plasmid could play the role of carrier for their acquisition of resistant determinant which later got integrated within the host genome.

![Figure 3. DNA directed RNA polymerase gene amplification in three NDM harbouring isolates. Lane 1-PA6 showing expected amplification of the intact gene; Lane 2 & 3-PA38 & 47 showing amplification of the gene along with the NDM insert (amplified product was comparatively of higher basepair than PA6).](image1)

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![Figure 4. Promoter sequence for NDM gene as determined by Softberry BPROM software. (A) Promoter sequence of blaNDM within RNA polymerase gene in P. aeruginosa strain (AF047025) and (B) NDM promoter in A. baumannii associated with some insertion sequence (HQ857107). The −35 and −10 motifs of promoter are in boldface type whereas ORF sequence in red color font.](image2)

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In the previous study it was reported that the NDM gene has originated from *Acinetobacter baumannii* and is linked with IS_Aba125 in the upstream region. However, during horizontal transfer the gene got excised from the *Acinetobacter* DNA along with whole or truncated portion of IS_Aba125 [30]. Therefore *Acinetobacter* specific primer was used for the study and it was observed that only PA6 showed the presence of this insertion sequence in the upstream area. But acquisition of *bla*NDM within the DNA dependent RNA polymerase of host chromosome is quite unique and not reported previously.

The new genetic makeup of *bla*NDM under the control of RNA polymerase promoter might be responsible for higher level of expression, which could probably be the explanation for high MIC of PA38 and PA47.

The clinical challenge posed by *bla*NDM is currently higher worldwide due to its potential transferability. As our findings also suggest that the unique genetic makeup of *bla*NDM is for make an ease to endure the isolate within hospital environment to counter increasing antibiotic pressure. This is an important consideration that *Pseudomonas* with recombinant *bla*NDM is crucial for the hospital infection management and therapeutic options, otherwise this new genetic adaptability will lead in to more serious public health implications. From the study it cannot be truly predicted whether this recombination was a by chance event or a true event since

**Table 3. MIC (mg/L) of *P. aeruginosa* against cephalosporins and carbapenems.**

| Isolate ID | CTX | CAZ | CEP | CFT | ERTA | IMP | MER |
|------------|-----|-----|-----|-----|------|-----|-----|
| PA6        | ≥512| ≥512| ≥512| ≥512| 256  | 128 | 64  |
| PA38       | ≥512| ≥512| ≥512| ≥512| 512  | 512 | 512 |
| PA47       | ≥512| ≥512| ≥512| ≥512| 512  | 512 | 512 |

MIC, Minimum Inhibitory Concentration; CTX, Cefotaxime; CAZ, Ceftazidime; CEP, Cefepime; CFT Ceftriaxone; ERTA, Ertapenem; IMP, Imipenem; MER, Meropenem.

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the sample size was too small. Therefore, further investigation is advocated to establish this fact.

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Author Contributions
Conceived and designed the experiments: SM SU AB. Performed the experiments: SM SU AB APM DC. Analyzed the data: SM AB. Contributed reagents/materials/analysis tools: SM SU AB APM DC MRS. Wrote the paper: SM AB.

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