Sequential Isolation in a Patient of *Raoultella planticola* and *Escherichia coli* Bearing a Novel ISCR1 Element Carrying *bla*<sub>NDM-1</sub>

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## Abstract

**Background:** The gene for New Delhi metallo-β-lactamase 1 (NDM-1) has been reported to be transmitted via plasmids which are easily transferable and capable of wide distribution. We report the isolation of two NDM-1 producing strains and possible in vivo transfer of *bla*<sub>NDM-1</sub> in a patient.

**Methods:** Clinical samples were collected for bacterial culture and antibiotic susceptibility testing from a patient during a 34-day hospitalization. The presence of *bla*<sub>NDM-1</sub> was detected by PCR and sequencing. Plasmids of interest were sequenced. Medical records were reviewed for evidence of association between the administration of antibiotics and the acquisition of the NDM-1 resistance.

**Results:** A NDM-1 positive *Raoultella planticola* was isolated from blood on the ninth day of hospitalization without administration of any carbapenem antibiotics and a NDM-1 positive *Escherichia coli* was isolated from feces on the 29<sup>th</sup> day of hospitalization and eight days after imipenem administration. The *bla*<sub>NDM-1</sub> was carried by a 280 kb plasmid pRpNDM1-1 in *R. planticola* and a 58 kb plasmid pEcNDM1-4 in *E. coli*. The two plasmids shared a 4812 bp NDM-1-IS<sub>CR1</sub> element which was found to be excisable from the plasmid as a free form and transferrable in vitro to a NDM-1 negative plasmid from *E. coli*.

**Conclusion:** *bla*<sub>NDM-1</sub> was embedded in an IS<sub>CR1</sub> complex class 1 integron as a novel 4812 bp NDM-1-IS<sub>CR1</sub> element. The element was found to be able to self excise to become a free form, which may provide a new vehicle for NDM-1 dissemination. This mechanism could greatly accelerate the spread of NDM-1 mediated broad spectrum β-lactam resistance.

## Citation

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## Background

Carbapenems often represent last-resource drugs for Gram negative bacterial infections. The most common mechanism of carbapenem resistance in Gram negative bacteria is the production of carbapenemases, including metallo-β-lactamases (MBLs). The New Delhi metallo-beta-lactamase (NDM-1), a recently identified novel type of MBL can hydrolyze virtually all β-lactams, adding further burden to an already high level of antibiotic resistance. The emergence and global spread of NDM-1 mediated broad spectrum β-lactam resistance has become a major concern worldwide [1,2].

NDM-1 was first identified in a carbapenem-resistant *Klebsiella pneumoniae* strain isolated from urine and an *Escherichia coli* strain isolated from feces of the same patient that was resistant to almost all antibiotics tested except colistin and tigecycline in 2009 in Sweden [3]. NDM-1 positive *K. pneumoniae* and *E. coli* were soon detected in many other countries, including Australia, Canada, Germany, China, France, Japan, the Nordic countries, United States and many others [4–8]. In addition, reported NDM-1 positive bacterial species extended to *Acinetobacter* spp, *Citrobacter freundii*, *Enterobacter cloacae*, *Morganella morgani*, *Providencia* spp, and *Pseudomonas aeruginosa* [9–11]. The world has been alarmed by the wide dissemination of NDM-1 isolates, for which new and effective
antibiotics are currently unavailable. The gene \textit{bla}_{\text{NDM-1}} encoding NDM-1 has been mostly reported to be on plasmids, which could rapidly disseminate and spread between different bacterial species by cell-to-cell transfer of the plasmids. However the plasmids carrying \textit{bla}_{\text{NDM-1}} are different in sizes, ranging from 40 kb to 400 kb \cite{9}, suggesting that direct transfer by plasmid is only one means of NDM-1 dissemination. Multiple mobile genetic elements are also associated with \textit{bla}_{\text{NDM-1}} dissemination \cite{4,11,12}.

The insertion sequence common region 1 (\textit{ISCR1}) is a well recognized system of gene capture. Its role in antibiotic resistance dissemination was first noted by the evidence that a wide variety of antimicrobial determinants are found in the immediate vicinity of \textit{ISCR1} \cite{13,14}. \textit{ISCR1} is capable of mobilizing its upstream sequences by a process of rolling circle transposition \cite{15}. Association of the \textit{ISCR1} element with \textit{bla}_{\text{NDM-1}} has been observed in plasmids pNDM-HK\textit{HV451074} \cite{16}, pMR0211\textit{JN687470} \cite{17} and in \textit{K. pneumoniae} strain 05-506 \cite{3} and \textit{P. aeruginosa} \cite{18}. However the role of the \textit{ISCR1} element in the transfer of the NDM-1 within and between bacterial species is unclear.

Here, we report that we identified two strains of different species (\textit{Raouella planticola} and \textit{E. coli}) carrying NDM-1 isolated 20 days apart from the same patient and showed that \textit{bla}_{\text{NDM-1}} was located on two unrelated plasmids. The \textit{bla}_{\text{NDM-1}} was embedded in an \textit{ISCR1} complex class 1 integron, which is most likely responsible for \textit{bla}_{\text{NDM-1}} transfer between \textit{R. planticola} and \textit{E. coli} in the patient in vivo.

Materials and Methods

Ethics statement

The feces, urine, sputum and blood samples were obtained with the written informed consent from the patient. This study was reviewed and approved by the ethics committee of the National Institute for Communicable Disease Control and Prevention, China CDC, according to the medical research regulations of the Ministry of Health, China. This research was conducted within China.

The Patient

The patient was a 51-year-old farmer from a small village of Gansu Province, who has never visited any cities in China or other countries that have reported NDM-1. On October 1, 2010, the patient was admitted to Lanzhou medical school hospital because of severe multiple left rib fractures as a result of traffic accident. Surgical operation was performed three hours after the accident. A second operation was performed five days later on the 6\textsuperscript{th} of October. The patient was hospitalized for 34 days and discharged on November 6, 2010. The blood, urine, sputum and feces were sampled for isolation of bacteria and antibiotic susceptibility testing as standard patient care except that some samples also were collected for the investigation of NDM-1.

Bacterial isolation, antibiotics resistance testing and molecular typing

Species identification was performed using the Microscan WalkAway 40 S1 identification system (Dade Behring, Deerfield, IL, USA). The 16s rDNA and \textit{tpb\textsubscript{II}} gene sequences were also analyzed to confirm the species identity. Susceptibility of the isolates to antibiotic was determined by the E-test methodology (BioMerieux SA, La Balme-les-Grottes, France). Modified Hodge test was performed to screen for carbapenemases. The results were interpreted according to the 2011 Clinical and Laboratory Standards Institute recommendation \cite{19}. The relationship of NDM-1 positive and negative \textit{E. coli} strains isolated from the same fecal samples was analyzed by PFGE (Pulsed Field Gel Electrophoresis)-\textit{XhoI} digestion method, following the protocol previously described \cite{20}.

PCR primers and detection of \textit{bla}_{\text{NDM-1}}

PCR primers for detecting \textit{bla}_{\text{NDM-1}} and for confirming the genetic environment of the \textit{bla}_{\text{NDM-1}} are listed in Table S1 in File S1. Plasmids were analyzed and sized by the PFGE-S1 nuclease method \cite{21}. Separation of large fragments from restriction enzyme digestion was done with PFGE by using of a CHEF-DR III apparatus (Bio-Rad, Hercules, CA) for 7 h at 6 V/cm and 14°C with initial and final pulse times of 5 s and 20 s, respectively. Southern hybridization was performed with a 560 bp segment of the \textit{bla}_{\text{NDM-1}} gene as a probe using the ECL direct nucleic acid labeling and detection system (GE Healthcare, Buckinghamshire, UK).

Conjugation and transformation of \textit{bla}_{\text{NDM-1}} plasmids

Conjugation experiments were carried out by the solid surface methods with \textit{E. coli} J53 Az as the recipient \cite{4}. Transconjugants were selected on Luria-Bertani agar plates containing sodium azide (100 mg/L) and ampicillin (100 mg/L), confirmed by PCR amplification of the \textit{bla}_{\text{NDM-1}} for NDM-1 positive donors and \textit{bla}_{\text{NDM-1}} gene for NDM-1 negative donors and sequencing of the PCR products. The plasmid DNA of NDM-1 positive strain \textit{E. coli} EcNDM1 was used for transformation and \textit{E. coli} JM109 was used as the recipient.

Whole genome sequencing and sequencing of plasmids

The whole genome of the isolate \textit{R. planticola} was sequenced using a combined strategy of Roche 454 Genome Sequencer FLX System (Roche/454 Life Sciences, Branford, CT, USA) and Illumina sequencing technology (Illumina, San Diego, CA, USA). The Burrows-Wheeler Alignment (BWA) tool was used to map all the Ilumina reads to the scaffolds generated by 454 Newbler. The inter- and intra-scaffold gaps were filled by local assembly of Roche and Illumina reads. The gaps between contigs were closed by targeted PCR amplification and sequencing of the PCR products with BigDye terminator chemistry on an ABI 3730 capillary sequencer.

Plasmid DNA extracted from \textit{E. coli} EcNDM1 was transformed into \textit{E. coli} JM109, and a carbapenem resistant transformant carrying plasmid pEcNDM1-4 named JM109 (pEcNDM1-4) was recovered. Plasmid pEcNDM1-4 extracted from the \textit{E. coli} JM109 (pEcNDM1-4) was used to construct a Tn5 transposon library according to the manufacturer’s recommendations (Epigene, Madison, WI) \cite{22}. Each transposon clone was sequenced using the BigDye terminator chemistry on an ABI 3730 capillary sequencer, first with primer FP-V and primer RP-V to obtain unique sequences flanking the transposon and then by PCR walking sequencing.

The pEcNDM1-4 equivalent plasmid in NDM-1 negative strain \textit{E. coli} EcNDMneg, named pEcNDMneg-4, was sequenced by targeted sequencing. Twenty pairs of overlapping primers were designed according to the sequences of plasmid pEcNDM1-4 (Table S1 in File S1) and used for the amplification of pEcNDMneg-4 as overlapping PCR products which were sequenced.

Nucleotide sequences accession numbers

Sequences of the NDM-1 carrying plasmids were submitted to the GenBank database with accession numbers JX515588,
Kacin, minocycline and linezolid (Figure 1). Nems were used. However the NDM-1 positive sulbactum, ceftizoxime and linezolid were used and no carbapenems were used. Additionally from the same fecal sample a multidrug resistant NDM-1 negative E. coli strain, EcNDMneg, was also isolated. We did not detect any NDM-1 positive bacteria from any other patients in the same ward. Medical records of the patient were reviewed for any evidence of association between administration of antibiotics and dissemination of blaNDM-1. Before the isolation of the NDM-1 positive R. planticola, only mezlocillin/sulbactum, cefizoxime and linezolid were used and no carbapenems were used. However the NDM-1 positive E. coli was isolated after administration of imipenem, piperacillin/tazobactam, amikacin, minocycline and linezolid (Figure 1).

Characterization of strains isolated from the patient
Both RpNDM1 and EcNDM1 were resistant to carbapenem, penicillin and cephalosporin and sensitive to amikacin, colistin sulphate and tigecycline. Additionally, EcNDM1 was also resistant to aztreonam, gentamicin and ciprofloxacin (Table 1). The two E. coli strains EcNDM1 and EcNDMneg had identical resistance patterns with the sole exception that the former was also resistant to carbapenem. PCR sequencing of the blaNDM-1 from both RpNDM1 and EcNDM1 revealed 100% identity with the published blaNDM-1 sequence [3]. PFGE-XbaI analysis of EcNDM1 and EcNDMneg showed identical PFGE patterns (Figure 2b), suggesting that the two strains had the same origin.

The blaNDM-1 location and plasmid analysis
PFGE of S1 digested plasmid DNA and Southern hybridization were performed to examine plasmid profiles and the blaNDM-1 location of E. coli EcNDM1, EcNDMneg and R. planticola RpNDM1. RpNDM1 had two large plasmids, designated as pRpNDM1-1 and pRpNDM1-2 of approximately 280 kb and 240 kb respectively and blaNDM-1 was found to be on the larger plasmid pRpNDM1-1 by Southern hybridization (Figure 2c and 2d). EcNDM1 had five plasmids ranging from 25 kb to 180 kb, which were named as pEcNDM1-1 to pEcNDM1-5 in descending order of their sizes. The blaNDM-1 was found to be on the 60 kb plasmid pEcNDM1-4 by Southern hybridization (Figure 2d). E. coli strain EcNDMneg shared an identical plasmid profile with EcNDM1. The plasmid pEcNDMneg-4 equivalent to plasmid pEcNDM1-4 in EcNDMneg was found to have no hybridization signal with the blaNDM-1 probe (Figure 2c and 2d).

Conjugation was performed with EcNDM1, EcNDMneg and RpNDM1 respectively as donor and E. coli J53 Az2 as recipient. Both pEcNDM1-4 and pEcNDMneg-4 were found to be a conjugal plasmid with a relatively high transfer frequency at approximately 1 x 10^{-2} per donor cell. However pRpNDM1-1 failed to transfer into E. coli J53 Az2 by conjugation.

Sequence analysis of plasmids pRpNDM1-1, pEcNDM1-4 and pEcNDMneg-4
Plasmid pRpNDM1-1 was sequenced by whole genome sequencing of RpNDM1 using a combination of Roche 454 sequencing with 283 867 paired-end reads generated for a 18.48-fold coverage and Illumina sequencing with 15 713 532 75-bases paired end reads for an approximately 205.7-fold coverage. Most of the Roche and Illumina reads were assembled into 252 contigs and 23 scaffolds. Gaps were closed and genome structure confirmed by targeted PCR and Sanger sequencing. The chromosome and the two plasmids of RpNDM1 were completely assembled. The blaNDM-1 carrying plasmid pRpNDM1-1 was 277 682 bp in size and belongs to the IncH group by sequence homology analysis [23, 24]. The plasmid encodes 261 coding sequences (CDSs), of which 136 encode proteins with homology to proteins of known functions and the remaining 125 (47.9%) encode hypothetical proteins. The blaNDM-1 was found within a 39 394 bp antibiotic resistance region, starting from a Trn3-like 38 bp invert repeat (IR) upstream of merR (encoding putative transcriptional regulator) at bp 176 794 to the end of the interrupted 38 bp IR of Trn5036 at bp 216 377 and was flanked by a 5 bp direct repeat, characteristic of Trn3-family transposons. Besides blaNDM-1, six antibiotic resistance genes, aacA4 (resistance to aminoglycoside), catB8 (resistance to chloramphenicol), sul1 (resistance to sulfonamides), aadA2 (resistance to aminoglycoside), blacTX-M-9 (resistance to lactam antibiotics), arr3 (resistance to rifampin) and a truncated dfrA27 (resistance to trimethoprim) with a 113 bp deletion in the middle of the gene were also found in this region (Figure 3a).

pRpNDM1-1 was found to share 86% of its sequence by length with Klebsiella oxytoca strain E718 plasmid pKOX-R1 (GenBank accession number CP003684), showing three segments of colinearity, although genetic arrangement of the segments was slightly different (Figure S2 in File S1). In comparison to pKOX-R1, pRpNDM1-1 carries an additional 32 395 bp antibiotic resistance region, starting from downstream of IS26 at position bp 177 651 to upstream of class 1 integron 5'CS at position bp 21 0027, carrying 7 antibiotic resistance genes including blaNDM-1. Interestingly, Klebsiella oxytoca strain E718, a NDM-1 carrying strain, isolated from Taiwan harbored another plasmid, pKOX_NDM1 in addition to pKOX-R1. The blaNDM-1 was found to be located on pKOX_NDM1, which had little similarity with pRpNDM1-1 except for a 100% identical segment of 1294 bp carrying 3'-IS900-1-blacTX-M-1-blaNDM-1-3', which indicates that the NDM-1 acquisition by plasmid pRpNDM1-1 has a complex history (Figure S2 in File S1).

The plasmid pEcNDM1-4 was sequenced using the EZ-Tn5 transposition system. The genetic structure and the sequence of the plasmid were further confirmed by PCR and sequencing of the PCR products using 19 pairs of overlapping primers. The plasmid was 58 228 bp and was found to share a backbone of approximately 51 kb with plasmid pKp96 [25] at 99% DNA sequence identity including the complete array of genes for replication, plasmid transfer, partition and stabilization (Figure S1 in File S1). The backbone of pEcNDM1-4 was interspersed by four insertions. Insertion-1 (bp 4384 to bp 7924) was comprised of blaCTX-M-9 (resistance to lactam antibiotics), catB3 (resistance to chloropenicillin), a truncated arr3 and a truncated dfrA27. The truncated dfrA27 is unusual as it contained a 113 bp deletion in the middle of the gene and is identical to that found in plasmid pRpNDM1-1. Insertion-2 (bp 11 394 to bp 16 257) was comprised of a truncated blacTX-M-1 (resistance to bleomycin), blacTX-M-1, a truncated IS900-1-sul1 and an intact ISCR1. Insertion-3 (bp 17 091 to bp 17 909) was comprised of a mobile element IS26. Insertion-4 (bp 35 885 to bp 36 170) was comprised of a partial mobile element IS26. Insertion-4 (bp 35 885 to bp 36 170) was comprised of a partial mobile element IS26. Insertion-4 (bp 35 885 to bp 36 170) was comprised of a partial mobile element IS26.
for 2 pairs of primers, 17 pairs of the primers successfully amplified products and the sequences obtained from sequencing the PCR products were identical to that of pEcNDM1-4. We found that pEcNDMneg-4 was completely co-linear and identical to pEcNDM1-4 except for the region where the 2 pairs of primers failed. The gap was closed using new primers and the plasmid pEcNDMneg-4 was 53,416 bp and was 100% identical to plasmid pEcNDM1-4 except for a 4,812 bp deletion which contained a truncated \textit{ble} \textit{MBL}, \textit{bla} \textit{NDM-1}, a truncated \textit{IS} \textit{Aba125}, the class 1 \textit{integron} \textit{3}\textit{CS (qacE1/sul1)} and an \textit{IS} \textit{CR1} element. This 4,812 bp \textit{NDM-1-IS} \textit{CR1} element is discussed in detail below.

### Comparison of the two \textit{bla}\textit{NDM-1} carrying plasmids, pRpNDM1-1 and pEcNDM1-4

The two \textit{bla}\textit{NDM-1} carrying plasmids, pRpNDM1-1 and pEcNDM1-4, were unrelated in plasmid backbone and genetic arrangement, except for a 9,332 bp-segment between bp 6,874 and bp 16,205 in pEcNDM1-4 and between bp 210,030 and bp 200,699 in pRpNDM1-1 which was 100% identical (Figure 3b). The 9,332 bp-segment consisted of a truncated \textit{arr3}, an unusually truncated \textit{dfrA27,3} CS, \textit{IS} \textit{CR1}, a truncated \textit{amikacin}, \textit{bla} \textit{NDM-1}, a truncated \textit{IS} \textit{Aba125} and a second copy of \textit{3}\textit{CS} and an \textit{IS} \textit{CR1} with a truncated \textit{qacE1}. This segment was located in a complex class 1 integron in both pRpNDM1-1 and pEcNDM1-4. In comparison to the complex class 1 integron in pRpNDM1-1, the complex class 1 integron in pEcNDM1-4 contained a 218 bp deletion in the 5' \textit{CS} and immediately downstream of the 5' \textit{CS} an additional gene cassette array \textit{aacA4-bla} \textit{OXA-30-bla} \textit{catB3} which is prevalent in \textit{Enterobacteriaceae} plasmids, such as plasmid pCF1-1 (Figure 3a).

The discovery and characterization of the 4,812 bp \textit{NDM-1-ISR1} element

When the plasmid DNA extracted from JM109 (pEcNDM1-4) was probed with the \textit{bla} \textit{NDM-1} probe, a 5 kb band (Figure 2f,
Table 1. Antimicrobial susceptibility patterns of R. planticola KpNDM1, E. coli EcNDMneg, E. coli EcNDM1-4 and their transformants.

| Antibiotics | Minimum inhibitory concentration (mg/L) |
|-------------|----------------------------------------|
|             | R. planticola | E. coli | E. coli | E. coli | E. coli |
|             | RpNDM1 | EcNDM1 | EcNDMneg | EcNDMneg (4812) | JM109 (pEcNDM1-4) | JM109 |
| TZP         | >256  | >256 | 16 | 32 | >256 | 0.75 |
| CTX         | >256  | >256 | >256 | >256 | >256 | 0.25 |
| ATM         | 0.094 | 32   | 32   | 32   | 0.064 | 0.064 |
| GEN         | 4     | 16   | 16   | 16   | 0.125 | 0.094 |
| AMK         | 3     | 6    | 6    | 6    | 1.5   | 0.5   |
| CIP         | 0.75  | >32  | >32  | >32  | 0.5   | 0.032 |
| IMP         | >32   | >32  | 0.19 | 16   | 16    | 0.19  |
| MEM         | >32   | >32  | 0.23 | 16   | 16    | 0.016 |
| CST         | 0.5   | 0.75 | 0.5  | 0.5  | 0.125 | 0.125 |
| TGC         | 0.75  | 0.38 | 0.38 | 0.38 | 0.38  | 0.25  |
| MHT*        | positive | positive | negative | positive | positive | negative |

TZP: piperacillin/tazobactam; CTX: cefotaxime; ATM: aztreonam; GEN: gentamicin; AMK: amikacin; CIP: ciprofloxacin; IPM: imipenem; MEM: meropenem; CST: colistin; TGC: tigecycline.

MHT*: Modified hodge test was performed and interpreted according to 2011 CLSI recommendation.

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labelled b) hybridized with the blonDM-1 probe at low intensity although this band cannot be seen on the agarose gel (Figure 2e), in addition to the heavy smear band that corresponds to the blonDM-1 carrying 58 kb plasmid. The 5 kb band was unexpected, suggesting that a free form of a 5 kb DNA molecule carrying blonDM-1 exists in the cell.

Using primer pair C4-F and C5-R designed to amplify the sequence between arr3 and qnrA3 (Figure 2a), two fragments of approximately 5 kb and 10 kb were amplified from pEcNDM1-4 (Figure 2g, Lane 1) and one fragment of approximately 5 kb was amplified from pEcNDMneg-4 (Figure 2g, Lane 2). The latter was much more intense, although similar amounts of templates were used for the 2 PCR amplifications. Cloning and sequencing of the 5 kb and 10 kb agarose gel purified fragments from the pEcNDM1-4 PCR amplification revealed that the 10 kb fragment corresponds to the region between arr3 to qnrA3 (see Figure 2a) and the 5 kb fragment corresponds to the same region but is devoid of the 4812 bp NDM-1-IS element. We also sequenced the 5 kb PCR product from pEcNDMneg-4. The sequences of the two 5 kb fragments were 100% identical. These results suggest that the 4812 bp NDM-1-IS element can be excised from the plasmid.

The 5 kb band observed in the Southern hybridization of undigested plasmid DNA extract described above must be the 4812 bp NDM-1-IS element, providing further evidence of the existence of a free form of the 4812 bp NDM-1-IS element.

Four pairs of primers, L1F/R to L4F/R, facing diverging directions were designed to confirm the structure of the 4812 bp NDM-1-IS element. An approximately 5 kb fragment was amplified from EcNDM1 using each of the four pairs of primers (Figure 2g). Sequencing of these PCR products showed that they all corresponded to the 4812 bp NDM-1-IS element, albeit the sequences started from different positions as the primers anneal at different sites. These PCR experiments confirmed the presence of the 4812 bp NDM-1-IS element. However the PCR results cannot determine whether the NDM-1-IS element is in a circular form or in tandem multiple copies since the diverging primer pairs can amplify a product in both scenarios.

We then used restriction mapping to determine whether there is more than one copy of the NDM-1-IS element on the plasmid. There is a Nod site at bp 12 238 located in the 4812 bp NDM-1-IS element and a Nod site upstream at bp 22 118 (Figure 2a). When the plasmid pEcNDM1-4 was digested by Nod and probed with the blonDM-1 probe, a 10 kb hybridization band was observed as expected (Figure 2e and 2f). A 5 kb hybridization band was also observed and was approximately double of the intensity of the 10 kb hybridization band, suggesting that there were double copies of the 5 kb band relative to the 10 kb band. We then used 2 more restriction enzymes, ClaI and BsaXI, to further determine whether there is a single copy or multiple copies of the NDM-1-IS element. The ClaI (restriction sites flanking the blonDM-1 gene at bp 4619 and bp 18 518) and BsaXI (restriction sites flanking the blonDM-1 gene at bp 6738 and bp 16 939) digestions were expected to generate 13 kb and 10 kb fragments respectively. However when plasmid pEcNDM1-4 was digested by ClaI and BsaXI and probed with the blonDM-1 probe, a 23 kb band and a 20 kb band were hybridized with the blonDM-1 probe respectively, which is 10 kb larger than expected (Figure 2f). The extra 10 kb indicates that there are 2 more copies of the NDM-1-IS element present in tandem. However this result is inconsistent with the C4-F/C5-R PCR results which only detected a 5 kb and a 10 kb band corresponding to 0 or 1 copy of the NDM-1-IS element. The discrepancy can be explained by that the C4-F/C5-R PCR failed to amplify the 20 kb amplicon that contains 3 copies of the NDM-1-IS element because of the length limit of the PCR amplification. This also explains the large difference in band intensity observed between pEcNDM1-4 and pEcNDMneg-4 in the C4-F/C5-R PCR described above. There must be a very small proportion of the pEcNDM1-4 carrying 0 or 1 copy of the NDM-1-IS element with the majority carrying 3 copies. Therefore the pEcNDM1-4 size is expected to vary from 33 416 kb (0 copy of the NDM-1-IS element) to 67 852 kb (3 copies of the NDM-1-IS element),...
which may also partly explain the smear observed in Figure 2e and Figure 2f of the total plasmid extract. Note that the assembled pEcNDM1-4 sequence has only one copy of the NDM-1-ISCR1 element with a size of 58,228 bp, due to the limitation of sequence assembly that cannot resolve the repeats.
Considering all of the results above, the 4812 bp NDM-1-IS\textsubscript{CR1} element exists in two forms in EcNDM1 as illustrated in Figure 2a. One was located on the plasmid predominantly in three tandem copies, while the other was an extra-plasmid and extra-chromosomal free form. The latter must have been released from the plasmid in small quantities during replication, which may have resulted in some plasmids containing aberrant copies of the 4812 bp NDM-1-IS\textsubscript{CR1} element.

The 4812 bp NDM-1-IS\textsubscript{CR1} element was transferable by transformation to \textit{E. coli} EcNDMneg.

The transferability of the 4812 bp NDM1-IS\textsubscript{CR1} element was examined by using \textit{E. coli} strain EcNDMneg as a host which is sensitive to imipenem. The 4812 bp NDM-1-IS\textsubscript{CR1} element was amplified by PCR using primer pair L5-F/R (Table S1 in File S1) and the PCR product was ligated by T4 DNA ligase to become circular. The PCR product was introduced into \textit{E. coli} EcNDMneg by electroporation, and a carbapenem-resistant transformant named EcNDMneg (4812) was found. The susceptibility profile of EcNDMneg (4812) was identical to that of strain EcNDM1 (Table 1). Targeted PCR sequencing found that the 4812 bp NDM-1-IS\textsubscript{CR1} element precisely inserted downstream of the existing IS\textsubscript{CR1} in pEcNDM1-4 to recreate an identical gene arrangement to pEcNDM1-4.

### Discussion

This case of isolating two NDM-1 positive strains of different species, \textit{R. planticola} and \textit{E. coli} isolated from the same patient provided an interesting scenario of emergence of NDM-1 within the same patient \textit{in vivo}. The NDM-1 positive \textit{R. planticola} strain was isolated when only mezlocillin/sulbactum, ceftriaxime and linezolid were used, suggesting that the emerging of NDM-1 was not due to administration of carbapenem antibiotics in this case. However, the NDM-1 positive \textit{E. coli} was isolated after administration of imipenem which may have allowed the selection of an \textit{E. coli} strain that possessed NDM-1. The 9332 bp fragment carrying an unusually truncated \textit{dfrA27} gene was shared by the two plasmids, suggesting an additional link between the two \textit{bla}_{NDM-1} carrying plasmids pRpNDM-1 and pEcNDM1-4. The latter further harboured an additional gene cassette array, \textit{aacA4-cr-bla\textsubscript{OXA-30/catB3}}, which is highly prevalent in \textit{Enterobacteriaceae}, with a 218 bp deletion in the 5' CS. This deletion would most likely have been generated when the gene cassette array was inserted in pEcNDM1-4. Therefore the likely process is that \textit{E. coli} acquired \textit{bla}_{NDM-1} from \textit{R. planticola} and additionally acquired the gene cassette array \textit{aacA4-bla\textsubscript{OXA-30/catB3}} from another \textit{Enterobacteriaceae} plasmid, such as pCF1-1. The alternative is acquisition of \textit{bla}_{NDM-1} by \textit{R. planticola} from \textit{E. coli}, which is also possible but less likely since acquisition of a 218 bp region to undo the deletion in

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Figure 3. Schematic representation of the DNA sequences surrounding the \textit{bla}_{NDM-1} in plasmid pEcNDM1-4, pRpNDM1-1 and homologous sequences in plasmid pCF1-1 to illustrate the relationship of the three plasmids. (a): Illustration of the adjacent genes of NDM-1-IS\textsubscript{CR1} of pRpNDM1-1, pEcNDM1-4 and homologous sequences in plasmid pCF1-1. Arrows represent ORFs and their direction of transcription; Filled triangles show the same 218 bp insertion in pCF1-1 and in pRpNDM1-1, compared with pEcNDM1-4; Open rectangle represent hypothetical proteins between \textit{bla\textsubscript{CTX-M-9}} and IS\textsubscript{26} in plasmid pRpNDM1-1; Pink shades across the plasmids indicates more than 99% identity between sequences. (b): MUMer-based genomic display between pEcNDM1-4 and pRpNDM1-1. The corresponding homologous sequences represented in (a) were highlighted in red lines.

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pRpNDM1-1 is less probable. There is also a possibility that blaNDM-1 was gained from another source by one or both species independently.

**ISCR1** element was first observed in class 1 integrons In6 and In7 in 1995 [26]. Since then, **ISCR1** elements were shown to have a close association with trimethoprim, quinolone, aminoglycoside resistance genes and several β-lactamase genes. Two copies of **ISCR1** with a 3’CS upstream flanking different subtypes of qnr genes were also reported in China [27], France [17] and other countries [28]. **blaNDM-1** has always been found to be carried on a structure consisting of a partial ISAb125 and a **bla** gene with **blaNDM-1** sandwiched between them [3]. **ISCR1** elements adjacent to the conserved ISAb125-blaNDM1-bla*QAC* element have been reported [3,16,18]. However, this is the first time to report that **blaNDM-1** was flanked by an **ISCR1** element on both sides. The conserved 1596 bp ISAb125-blaNDM1-bla*QAC* region was flanked by a copy of the integron 3’CS qacE1/sul and **ISCR1** on both sides (Figure 3a) in the two NDM-1 carrying plasmids of different species *R. planticola* and *E. coli*.

**ISCR1** is a known means of mobilizing its adjacent sequence via rolling circle transposition. Partridge and Hall [14] and Toleman et al. [15] demonstrated that rolling circle replication of the **ISCR1** element can produce circular modules that include the **ISCR1** and its upstream adjacent antibiotic resistance genes. These circular modules can then be rescued by recombination between homologous fragments. The 4812 bp NDM-1-**ISCR1** element may be mobilized by rolling circle transposition. Our isolation of two nearly identical *E. coli* strains from the same fecal sample with only one being NDM-1 positive presented an interesting scenario for the acquisition of **blaNDM-1** and provided a vehicle for experimental transfer of NDM-1 antibiotic resistance. Our *in vitro* transformation experiment using a synthesized circular 4812 bp NDM-1-**ISCR1** element showed that the element successfully transformed EcNDMneg containing plasmid pEcNDMneg-4 to a NDM-1 positive strain with the precise insertion of the element into the **ISCR1** downstream in pEcNDMneg-4 to create an identical gene arrangement to pEcNDM1-4, demonstrating its mobility via **ISCR1**. However, *in vitro* transformation is artificial. There are likely other mechanisms of transfer.

The advantage for the **blaNDM-1** gene embedding in a class 1 integron **ISCR1** complex is that one of the most widely used mechanisms for the spread of antibiotics resistance across species is harnessed for NDM-1 dissemination. Therefore in addition to clonal expansion of a NDM-1 positive strain, NDM-1 can spread by multiple means including inter-strain spread within and between species through plasmids, transposons and the NDM-1-**ISCR1** element. Given that strains carrying class 1 integrons or **ISCR1** complex class 1 integrons are common among bacteria [13,15,29], this is potentially a highly effective means for the spread of **blaNDM-1**.

In conclusion, **blaNDM-1** was found to be embedded in an **ISCR1** complex class 1 integron, which is most likely responsible for the **blaNDM-1** transfer between *R. planticola* and *E. coli* in a patient *in vivo*. The novel 4812 bp NDM-1-**ISCR1** element which can self excise to become a free form could provide a new vehicle for NDM-1 dissemination.

**Supporting Information**

**Figure S1**  Figure S1, Figure S2 and Table S1. (DOC)

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**Author Contributions**

Conceived and designed the experiments: JL RJ JX. Performed the experiments: JL YX MY XC XB Yan Wang QS HZ QM QC AZ CY XL. Analyzed the data: JL RJ BL WL Yiting Wang. Contributed reagents/materials/analysis tools: JL JX BL. Wrote the paper: JL RJ JX.

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