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To cite this version:
hal-03429615

HAL Id: hal-03429615
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Submitted on 4 Apr 2022

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First Genome Description of Providencia vermicola Isolate Bearing NDM-1 from Blood Culture

David Lupande-Mwenebitu 1,2,*, Mariem Ben Khedher 1,*, Sami Khabthani 1,*, Lalaoui Rym 1, Marie-France Phoba 3,4, Larbi Zakaria Nabti 1,*, Octavie Lunguya-Metila 3,4, Alix Pantel 5,*, Jean-Philippe Lavigne 5,*, Jean-Marc Rolain 1 and Seydina M. Diene 1,2,3,4

1. Introduction

Providencia species are Gram-negative bacteria belonging to the order of Enterobacterales, family of Morganellaceae, and genus Providencia. Their specific power to deaminate...
specific amino acids by oxidation into their corresponding keto and ammonia acids is a particularity that differentiates them from other members of the Enterobacteriaceae family [1,2]. Unlike other bacteria in this family, Providencia species are rarely involved in nosocomial infections [3]. Two species of Providencia, including P. stuartii and P. rettgeri, which are naturally resistant to many antibiotics including colistin and tigecycline, are the most common causes of more than 80% of human clinical infections, mainly urinary tract infections. The other species of this group are P. alcalifaciens, P. burhodogranariea, P. heimbachae, P. rettgeri, P. rustigianii, P. sneebia, P. stuartii, P. thailandensis, P. huaxiensis, and P. vermicola [4–6].

P. vermicola was first isolated from a nematode Steinernema thermophilum collected in soils in India in 2006. Its name means (ver.mi′co.la. L. n. worm; L. suff. -cola of L. n. incola inhabitant; N.L. n. vermicola inhabitant of worms) [7] and is very rarely found as an etiological agent in humans, with only one described case of diarrhea in a 37-year-old patient in India [8]. Providencia species are reported to be found mainly in environments such as water and are mostly involved in infections of birds, fish, and certain insects such as fruit flies [3,9]. Prior to this study, there was no available genome of P. vermicola in the NCBI database. Here, we describe the first complete genome sequence of a clinical multidrug resistant (MDR) P. vermicola isolate from a healthcare facility in Kinshasa, in the Democratic Republic of the Congo, and perform the comparative genomic analysis with the most closely-related species.

2. Materials and Methods

2.1. Bacterial Isolation

Two Gram-negative strains, namely P. vermicola P8538 and Escherichia coli P8540, were isolated in 2017 from blood and urine, respectively, at the KUTH (Kinshasa university teaching hospital) in DR Congo. The KUTH is a government-funded academic tertiary referral hospital in Kinshasa, the capital city of DR Congo. It is the national referral hospital in a country of approximately ninety million people. It has 1000 total beds, of which seven beds are in the intensive care unit (ICU). All isolates were identified first using biochemical tests such as urea, indole, oxidase, citrate, and triple sugar iron (TSI), and were confirmed by Microflex LT MALDI-TOF mass spectrometer (Bruker Daltonics, Bremen, Germany) after being sent to the IHU Méditerranée infection, Marseille.

2.2. Antimicrobial Susceptibility Testing

Susceptibility to 16 antimicrobial agents (i.e., amikacin, amoxycillin, amoxycillin clavulanate, cefepime, ceftixime, cepalothion, colistin sulfate, ciprofloxacin, cotrimoxazole, doxycycline, ertapenem, fosfomycin, gentamicin, imipenem, nitrofurantoin, tazobactam piperacillin; i2a, Montpellier, France) was determined using the disk diffusion method according to the European Committee on Antimicrobial Susceptibility Testing (EUCAST) guidelines (https://www.eucast.org/clinical_breakpoints/, access on 10 February 2021). The minimal inhibition concentration (MIC) of imipenem and colistin were respectively determined by agar dilution in Mueller–Hinton agar (Oxoid, Basingstoke, UK) and by microdilution method using UMIC Colistin kit (Biocentric, Bandol, France) according to CLSI recommendations. This MIC for imipenem was determined after 18–24 h of incubation on Mueller-Hinton plates inoculated with suspensions of isolates at a fixed density (0.5 to 0.6 McFarland standard), using E-test strips (BioMérieux, Marcy l’Etoile, France) according to the manufacturer’s recommendations.

2.3. Molecular Mechanisms of Antibiotic Resistance and Whole Genome Sequencing

Real-time PCR and standard PCR were performed to screen for the presence of carbapenem resistance genes including: blaNDM, blaOXA-23, blaVIM, blaOXA-48, and blaKPC. Genomic DNAs (gDNA) of the two carbapenem resistant isolates were extracted using the EZI Advanced XL Biorobot and the tissue DNA kit (Qiagen, Hilden, Germany) with the Bacterial card, according to the manufacturer’s instructions and quantified using NanoDrop 2000 (ThermoFischer, Illkirch, France). Whole genome sequencing was performed using the
MiSeq sequencer (Illumina, San Diego, CA, USA) and the Gridion sequencer (Nanopore, Oxford, UK), according to the Nanopore Template Preparation.

2.4. Bioinformatic Analysis

The genome assemblies were performed using the A5-pipeline on the Illumina reads and using Unicycler for the hybrid assembly, which includes both Illumina and nanopore reads [10,11]. Genome annotation was performed using the prokka (rapid prokaryotic genome annotation) pipeline [12]. Circular genome representations of *P. vermicola* chromosome and plasmid and their comparisons by BlastN with the closest sequences were performed using the CGview Server [13] and locally downloaded EasyFig v2.2 software. Proteome comparison of *P. vermicola* with those *Providencia* species was performed using the “get_homologues.pl” pipeline [14]. Moreover, the average nucleotide identity (ANI) between *P. vermicola* and downloaded genomes was determined using the OrthoANI program (AOT software) [15]. All genes deemed to be candidates of antimicrobial resistance or putative virulence genes were investigated using the ARG-ANNOT database and VFDB with threshold value amino acid alignment ≥70% of the input query sequence to avoid any data extrapolation. PHASTER (PHAge Search Tool Enhanced Release) was used to identify prophage sequences from genomic sequences [16–18].

2.5. Conjugation Experiment

To determine the transferability of carbapenem resistance, a conjugation experiment was performed using *E. coli* J53 (azide-resistant) as the recipient strain, as previously described [19].

3. Results

3.1. Clinical Information and Phenotypic Characterisation of Isolates

The *P. vermicola* P8538 strain was isolated from the blood culture of a 58-year-old patient hospitalized in the ICU for sepsis, whereas the *E. coli* P8540 strain was isolated in the urine of a 26-year-old patient hospitalized in the same ICU, but not at the same time, and who was receiving continuous respiratory assistance. As presented in Table 1, the two isolates were resistant to the majority of the 16 tested antibiotics, with the exception of cefepime, fosfomycin, and cotrimoxazole for the *P. vermicola* P8538 isolate, whereas the *E. coli* P8540 isolate remained susceptible to colistin (Table 1).

3.2. Genome Features

For *P. vermicola* P8538, the genome size was estimated to be 4,432,495-bp and composed of one chromosome 4,280,811-bp in size and 40.80% GC and one plasmid (pPV8538_NDM-1) 151,684-bp in size and 51.93% GC. Genome annotation revealed 4166 predicted genes composed of 3991 CDS, 11 rRNAs, 74 tRNAs, 89 miscRNAs, and 1 tmRNA gene (Table 2).
Table 1. Clinical features, Resistance phenotype, and genotype of *P. vermicola* P8538 and *E. coli* P8540 harboring the *bla*NDM-1 gene.

<table>
<thead>
<tr>
<th>Strain Names</th>
<th>Sex</th>
<th>Age</th>
<th>Sample</th>
<th>Status</th>
<th>Service</th>
<th>Resistance Phenotype</th>
<th>MIC IPM (µg/mL)</th>
<th>Antimicrobial Resistance Genes</th>
</tr>
</thead>
</table>

Table 2. Genome features of *P. vermicola* P8538 and *E. coli* P8540 isolates harboring *bla*NDM-1.

<table>
<thead>
<tr>
<th>Features</th>
<th><em>P. vermicola</em> P8538</th>
<th><em>E. coli</em> P8540</th>
</tr>
</thead>
<tbody>
<tr>
<td>Genome size</td>
<td>4,432,495-bp</td>
<td>4,809,673-bp</td>
</tr>
<tr>
<td>% GC content</td>
<td>41.1%</td>
<td>50.9%</td>
</tr>
<tr>
<td>No. of contigs</td>
<td>2</td>
<td>210</td>
</tr>
<tr>
<td>N50</td>
<td>184,648-bp</td>
<td>80,966-bp</td>
</tr>
<tr>
<td>No. of predicted genes</td>
<td>4166</td>
<td>4951</td>
</tr>
<tr>
<td>No. of CDS</td>
<td>3991</td>
<td>4553</td>
</tr>
<tr>
<td>No. of predicted tRNAs</td>
<td>74</td>
<td>83</td>
</tr>
<tr>
<td>No. of predicted rRNA</td>
<td>11</td>
<td>13</td>
</tr>
<tr>
<td>No. of predicted tmRNA</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>No. of predicted miscRNAs</td>
<td>89</td>
<td>303</td>
</tr>
<tr>
<td>No. of phage sequences</td>
<td>5</td>
<td>2</td>
</tr>
<tr>
<td>Sequence Type (ST)</td>
<td>-</td>
<td>1421</td>
</tr>
</tbody>
</table>

As shown in Figure 1, genome comparison with 10 published *Providencia* genomes revealed multiple mobile genetic elements (MGEs) in the *P. vermicola* P8538 chromosome. Indeed, three complete prophage regions were identified from the PHASTER database, including Phage_Entero_mEp460 “54.42-Kb, 39.96% GC”, Phage_Salmo_Fels_2 “59.9-Kb, 41.21%GC”, and Phage_Escher_HK639 “41.7-Kb, 42.31%GC”, respectively. Additionally, a complete siderophore biosynthesis NRPS (39.62-Kb, 33.92%GC) composed of 19 genes can be identified (Figure 1). As presented in Supplementary Figure S1, this siderophore NRPS was only identified from the NCBI database in three genomes including *P. stuartii* PRV00010, *Morganella morganii* VGH116, and *Salmonella enterica* 2014AM-3158 with average % homology of 96%, 44.37%, and 33.89%, respectively (Supplementary Table S1). The identified Type-I-F CRISPR-cas3 system (10.9-Kb, 43.63% GC) appeared to be specific to the *P. vermicola* P8538 isolate since it was not identified in any of the 10 compared genomes (Figure 1). Interestingly, a complete Type VI secretion system (T6SS) encoding gene was harbored in the *P. vermicola* P8538 chromosome and was partially detected in the compared genomes (Figure 1).

However, in addition to the MGEs identified from the *P. vermicola* P8538 chromosome, a complete and recombinant plasmid harboring the New Delhi metallo-β-lactamase-1 (*bla*NDM-1) gene has been identified from the genome sequences. As presented in Figure 2, this plasmid was mainly characterized by the presence of multiple plasmid conjugative transfer genes (14 *tra* genes), a toxin/antitoxin *higAB* system, a transposon-containing-NDM-1, a glutathione detoxification system, and four other antibiotic resistance genes.

### 3.3. Genome Comparison with Closely Related Species

As shown in Figure 3, the whole-proteome-based phylogenetic tree and pairwise comparison of *P. vermicola* P8538 with 10 other *Providencia* species revealed that our *P. vermicola* was more closely related to *P. stuartii* genomes than those of the two recently published *P. vermicola* genomes. It appeared clear that the published *P. vermicola* G1 was wrongly identified and should be reidentified as *P. rettgeri* (Figure 3). Indeed, *P. vermicola* P8538 shared between 90.79% and 97.61% of proteome homology with *P. stuartii* and only 88.6% homology with *P. vermicola* LLDR26 (Figure 3). Interestingly, *P. vermicola* P8538 exhibited only 81% of ANI with published *P. vermicola* LLDR26 and 77.24% of ANI with *P. vermicola* G1 (Figure 3).

This result was also confirmed based on the RpoB % aa identity which was 99.11% with that of *P. stuartii*, 98.96% *P. vermicola* LLDR26, and only 98.29% with *P. vermicola* G1 (Figure 3).

In addition to this evidence, as shown on Figure 4, the OrthoANI analysis revealed 82.95% and 82.89% average nucleotide identity of P8538 genome with *P. stuartii* AR_0026 and *P. stuartii* MRSN2154 genomes, respectively. Interestingly, *P. vermicola* P8538 exhibited only 81% of ANI with published *P. vermicola* LLDR26 and 77.24% of ANI with *P. vermicola* G1. These results clearly suggested a wrong identification of the latter. Indeed, *P. vermicola* G1 showed 99.24% of ANI with *P. rettgeri* 151.
Figure 1. Circular map of the *P. vermicola* P8538 chromosome and its genomic comparison with the nine closest *Providencia* genomes.

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Figure 2. Comparison of the *P. vermicola* pPV8538_NMD-1 plasmid with *E. coli* pEC8540 plasmid and the three closest plasmids retrieved from the NCBI database.

Figure 3. Proteome Pairwise comparison and the *RpoB* phylogenetic tree of *Providencia* genomes.
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Figure 4. Average nucleotide identity comparison of *P. vermicola* P8538 with *Providencia* species using the OrthoANI software.

3.4. Resistome

Regarding the antibiotic susceptibility phenotype of the *P. vermicola* P8538 isolate (Table 1), the resistome analysis confirmed the observed β-lactam resistance phenotype by the presence of the *bla*CMY-6, and *bla*NDM-1 genes from the plasmid pPV8538_NDM-1. Moreover, other antibiotic resistance genes were found, including *rmtC* and *aacA*4 conferring resistance to aminoglycosides, *catA*1 for phenicol resistance, *sul*1 and *aac6'-Ib-cr* for resistance to sulfonamides and quinolones, respectively (Table 1). Interestingly, the Tn3-NDM-1 transposon (21,774-bp, 59.40% GC) on the plasmid was the vehicle of four resistance genes, namely *bla*NDM-1, *rmtC*, *Sul*1, and *aacA*4 (Figure 2). A total of 20 virulence associated genes from the *P. vermicola* P8538 were identified and are presented in Supplementary Table S2.
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3.5. Genomic Analysis of the *E. coli* P8540 Isolate

Interestingly, during this study, our investigation of the potential spread of *bla*NDM-1 in this hospital revealed a positive *E. coli* P8540 strain isolated from the urine of a 26-year-old hospitalized patient. This strain was subjected to whole genome sequencing and resulted in a genome assembled into 210 contigs with a size of 4,809,673-bp and 50.9% GC content. The details of the genome features are presented in Table 2. MLTS analysis reveals a type ST1412 clone and plasmid finder analysis detected five plasmid replicons which were classified as col (MG828), IncA/A2, IncFIA/H11, IncI, and IncR. Interestingly, the same pPV8538_NDM-1 plasmid identified in the *P. vermicola* P8538 isolate was detected and identified from this *E. coli* P8540 isolate (pEC8540), suggesting a conjugation transfer event of this plasmid between these two Enterobacteriaceae species (Figure 2). Unfortunately, our in vitro experiment to transfer by conjugation the pPV8538_NDM-1 plasmid into the *E. coli* J53 (azide-resistant) strain was unsuccessful after three repeated assays.

Resistome analysis of *E. coli* P8540 confirmed the presence of the *bla*NDM-1, *bla*TEM-1B, *bla*SHV-12, and *bla*CTX-M-88 genes, conferring the observed β-lactam resistance of this isolate. Moreover, the presence of multiple genes conferring resistance to aminoglycosides (n = 12), phenicols (n = 4), sulfamide/trimethoprim (n = 2), macrolides (n = 1), quinolones (n = 18), and tetracyclines (n = 8) were identified (Table 2).

4. Discussion

In this paper we report the first complete genome sequences of the *P. vermicola* species. The genome analysis and comparative genomics of this clinical MDR *P. vermicola* P8538 isolate revealed significant genomic variations compared to other *Providencia* species. This may indicate the ability of this bacterium to colonize several hostile environments, given the presence of several MGEs in the genome, which is well documented in the literature [3,20,21]. We identified a conjugative and recombinant plasmid harboring antimicrobial resistance genes which was identified in two different pathogenic Enterobacterial species (*E. coli* and *P. vermicola*) from the same hospital, suggestive of a spread of the MDR plasmid within this healthcare setting. Thus, the existence of a conjugative plasmid harboring the NDM-1 enzyme in this hospital appeared to be a serious concern for infection and prevention control measures.

Some specific MGEs have been identified in this particular *P. vermicola* P8538 isolate, including a glutathione detoxification system from the plasmid pPV8538_NDM-1 which was detected by BlastN from the NCBI database in very few bacterial plasmids from *Salmonella enterica*, *Proteus mirabilis*, *Serratia marcescens*, *Enterobacter hormaechei*, *Klebsiella quasipneumoniae*, and *michiganensis*. As reported in the literature, this system is involved in the glutathione-dependent process of formaldehyde detoxification [22,23]. Moreover, we identified from the P8538 chromosome a NRPS cluster for siderophore biosynthesis which has only been identified in three other enterobacterial genomes, namely *P. stuartii* PRV00010, *M. morganii* VGH116, and *S. enterica* 2014AM-3158, and was absent in the two genomes of *P. vermicola* recently deposited in the NCBI database. As reported, siderophore systems are low molecular weight molecules which are widespread in the
bacterial and fungal world, with more than 200 biosynthetic and diverse types. They play the role of capturing, solubilizing and delivering essential Fe(III) ions in the cytoplasm [24] and are involved in the growth and development of microorganisms but also in bacterial virulence, as described in E. coli in urinary tract infections [25]. They are also involved in bacterial dissemination by induction of inflammation in the lungs [26,27]. The T6SS acts as a virulence factor in the majority of proteobacteria with the ability to attack eukaryotic and prokaryotic target cells through a complex process, secreting toxic effectors through a contact mechanism into neighboring bacteria or eukaryotic cells, causing cell lysis or growth arrest [28,29]. This complex process involves the transport of proteins through a contractile bacteriophage-like tail structure [30,31]. Six secretion systems have thus far been identified and are referred to as Type I to Type VI (T1SS to T6SS). T6SS was first discovered in Vibrio cholerae and Pseudomonas aeruginosa in 2006, and several studies have subsequently demonstrated its presence in many Gram-negative bacteria, including many human and animal pathogenic strains [32]. Indeed, studies have shown that certain T6SS subunits have structural homologies with other subunits of the bacteriophage T4, including the main tail protein and its injection syringe. It has therefore been established that T6SS is phylogenetically and structurally very close to the bacteriophage T4 [32]. Thus, we believe that the integration of several exogenous sequences, including bacteriophages and T6SS, may play a role in the adaptation and survival of P. vermicola which evolves in endosymbiosis in Steinernema thermophilum, a nematode in which P. vermicola develops [33].

5. Conclusions

This study highlights the emerging threat of blaNDM-1 dissemination in Kinshasa. To the best of our knowledge, this study describes, for the first time in the Democratic Republic of the Congo, the blaNDM-1 gene, in a bacterial genus of Enterobacteriaceae and in a rare species (P. vermicola), about which very little is known in Africa in general and nothing is known on the genomic level for this species. The fact that P. vermicola has been widely described as a nematode endosymbiont, nematodes which infect and kill fish and insects, some of which are used as food in certain environments such as Kinshasa, gives rise to speculation about the role that the habit of humans eating insects might play in the transmission of this bacterium. In addition, S. thermophilum, a nematode in which P. vermicola develops, is not recognized as a human pathogen. It cannot be excluded that this microorganism could be found in other parasites, in this case causing bacteria–parasite co-infections in humans. The identification of these NDM-1-producing isolates, which are also resistant to several other antibiotics and shared through the same plasmid with another isolate (P8540) in the same healthcare facility, confirms the existence of mobile genetic element exchanges among the circulating isolates within the University Hospital of Kinshasa. Therefore, it is urgent to improve surveillance and clinical practices to reduce or prevent the spread of resistance.

Supplementary Materials: The following are available online at https://www.mdpi.com/article/10.3390/microorganisms9081751/s1, Figure S1: BlastP comparison of the P. vermicola P8538 Siderophore biosynthesis NRPS cluster with that from P. stuartii, M. morganii, and Salmonella enterica. Protein annotation and the BlastP results are presented in Supplementary Table S1: BlastP comparison of the Siderophore biosynthesis NRPS cluster of P. vermicola P8538 with that of P. stuartii, M. morganii, and Salmonella enterica. Supplementary Table S2: Virulence factor genes identified in P. vermicola P8538 isolate. Whole genome sequences accession: The whole genome sequence of the clinical P. vermicola P8538 isolate was deposited at GenBank under the numbers CP048796 and CP048797.

Funding: The French government supported this work under the “Investissements d’avenir” programme (reference: Mediterranee Infection 10-IAHU-03).

Institutional Review Board Statement: This study was conducted in accordance with existing ethical guidelines and was approved by the Institutional Review Board of the Catholic University of Bukavu (UCB/CIES/NC/08/2019). The study was conducted in accordance with the principles of the Declaration of Helsinki. Anonymous and codified clinical patient data were used with strains collected and stored; informed consent was no longer required.

Informed Consent Statement: Not applicable.

Data Availability Statement: Not applicable.

Acknowledgments: We are very grateful to the IHU Mediterranée-Infection, Marseille, France for financial support, and TradOnline for English correction of the manuscript.

Conflicts of Interest: None of the authors report any conflict of interest or financial disclosures.

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