Draft genome sequence and detailed analysis of *Pantoea eucrina* strain Russ and implication for opportunistic pathogenesis

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**Abstract**

The genus *Pantoea* is a predominant member of host-associated microbiome. We here report on the genomic analysis of *Pantoea eucrina* strain Russ that was isolated from a trashcan at Oklahoma State University, Stillwater, OK. The draft genome of *Pantoea eucrina* strain Russ consists of 3,939,877 bp of DNA with 3,704 protein-coding genes and 134 RNA genes. This is the first report of a genome sequence of a member of *Pantoea eucrina*. Genomic analysis revealed metabolic versatility with genes involved in the metabolism and transport of all amino acids as well as glucose, fructose, mannose, xylose, arabinose, and galactose, suggesting the organism is a versatile heterotroph. The genome also encodes an extensive secretory machinery including types I, II, III, IV, and Vb secretion systems, and several genes for pilus production including the new usher/chaperone system (pfam 05,229). The implications of these systems for opportunistic pathogenesis are discussed.

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**Keywords:** *Pantoea eucrina* Draft genome sequence Detailed annotation Student Initiated Microbial Discovery (SIMD) project Metabolic versatility Secretion systems Pilus

1. Introduction

The strain *Pantoea eucrina* Russ was isolated (by an undergraduate student, BR) from a trashcan surface with frequent human use on Oklahoma State University (OSU) campus in Stillwater, OK. This was part of the Student Initiated Microbial Discovery (SIMD) project at OSU (introduced in [1]). The genus *Pantoea* is a phylogenetically and physiologically diverse genus with members ubiquitously found in host-associated microbiome as plant endophytes, insects symbionts, and members of the human gut microbiomes [2–6]. Endophytic strains range from plant pathogens, plat commensal, to a beneficial strains with growth-promoting effects [7]. *Pantoea* is frequently isolated from the nosocomial environment [8–10] and hence a considerable debate on its role in human infection was recently raised. Genomic analysis of strains belonging to the genus *Pantoea* could potentially contribute majorly to our understanding of opportunistic pathogenesis. Such knowledge can help mitigate the severity of nosocomial infections in immunocompromised patients. Here we report on the first draft genomic sequence, and the detailed annotation and analysis of a *Pantoea eucrina* strain with an emphasis on its pathogenic potential.

2. Genome sequencing information

2.1. Genome project history

The quality draft assembly and annotation were completed in 2015–2016. Table 1 shows the genome project information.

2.2. Growth conditions and genomic DNA preparation

*Pantoea eucrina* Russ was grown overnight at 30 °C on tryptic soy agar plates. Genomic DNA of high sequencing quality was isolated using the MPBio PowerSoil® DNA extraction kit according to manufacturer’s instructions. Negative stain TEM micrographs were obtained using the services of the Oklahoma State University Microscopy Lab. Briefly, the sample was placed on a carbon film TEM grid and allowed to incubate for 2 min, after which the excess liquid was blotted off. Phosphotungstic acid (PTA; 2% w/v) was then added to the grid followed by a 45-sec incubation. Excess PTA was blotted off and the...
grid was allowed to dry before it was visualized using JOEL JEM-2100 transmission electron microscope.

2.3. Genome sequencing and assembly

The genome of Pantoea eucrina Russ was sequenced using the Illumina MiSeq platform at the University of Georgia Genomics Facility using 2X300 paired end chemistry and an average library insert size of 700 bp. The short read de Bruijn graph assembly program Velvet [11] was employed for assembling quality filtered sequence data using the following flags: a kmer value of 101 bp and a minimum contig coverage value of 7 x. The genome project is deposited in GOLD (Genomes On-Line Database) and this Whole Genome Shotgun (WGS) project has been deposited in GenBank under the accession MAYN00000000. The version described in this paper is version MAYN01000000.

2.4. Genome annotation

Using the prokaryotic gene calling software package prodigal [12], a total of 3838 gene models were predicted with average gene size of 931.73 bp. Functional annotation involved a combination of NCBI Blast C+ + homology search, and HHMER 3.0 [13] hmmscan against the PFAM [14] 26.0 database. Additional gene analysis and functional annotation were carried out through the Integrated Microbial Genomes Expert Review (IMG-ER) platform.

2.5. Comparative genomics

We compared the genome of Pantoea eucrina strain Russ to 22 closely related genomes (IMG Ids: 648276708 (Pantoea sp. ab), 649633081 (Pantoea sp. At-9b), 2511231025 (Pantoea sp. YR343), 2511231035 ((Pantoea sp. GM01), 2519899784 (Pantoea sp. SC1), 2545824509 (Pantoea sp. GL120224-02), 2551306469 (Pantoea sp. A4), 2551306543 (Pantoea sp. B40), 2582581300 (Pantoea sp. 9140), 2602041550 (Pantoea sp. AS-PWVM4), 2602042078 (Pantoea sp. 9133), 2609460089 (Pantoea sp. IMH), 2616644925 (Pantoea sp. 3.5.1), 2617271108 (Pantoea sp. FFS), 2619619082 (Pantoea sp. SL1_5M), 2627853687 (Pantoea sp. MBL3J), 2627853912 (Pantoea sp. SM3), 2630968876 (Pantoea sp. PSH11), 2630968889 (Pantoea sp. PSH12), 263641558 (Pantoea sp. BL1), 2643221431 (Pantoea sp. isolate 98), 2651869657 (Pantoea sp. RIT-P-B)) using the “Genome clustering” function on the IMG-ER analysis platform based on the KEGG profile. We also used principal component analysis to compare the genomes based on several genomic features including the genome size, the number of genes, the number of transporters identified, the GC content, the number of non-coding bases, the number of genes belonging to COG categories, as well as the number of genes belonging to each COG category. The PCA analysis was conducted using the “princomp” function in the labdsv library of R [15]. The results were visualized using a biplot, where genomes were represented by stars and genomic features or COG categories used for comparison were represented by arrows.

Table 1

| MIGS ID | Property | Term       |
|---------|----------|------------|
| MIGS 31 | Finishing quality | Draft      |
| MIGS 28 | Libraries used     | Illumina X200 paired end chemistry |
| MIGS 29 | Sequencing platforms | Illumina Miseq |
| MIGS 31.2 | Fold coverage | 300x |
| MIGS 30 | Assemblers       | Velvet 2.0 |
| MIGS 32 | Gene calling method | Prodigal, IMG-ER |
| Genbank ID | MAYN00000000 |
| GenBank date of release | July 2016 |
| GOLD ID | Gp0126378 |
| BIOPROJECT | PRJNA327384 |
| MIGS 13 | Project relevance | Environmental |

3. Results and discussion

3.1. Classification and features

Cells of P. eucrina strain Russ are Gram-negative, motile rods that were arranged in singles (Fig. 1). Colonies on TSA agar were yellow.

Within the genus Pantoea, 24 species are described with widely published names P. agglomerans type strain ATCC 27155, P. allii type strain BD390, P. ananatis type strain ATCC 33244, P. anthophila type strain BD871, P. breneri type strain BD873, P. calida type strain 1400/07, P. citreus type strain BD875, P. coffeiphila type strain DSM 28482, P. conspicua type strain BD805, P. cyripedii type strain ATCC 29267, P. deleyi type strain BD767, P. dispersa type strain ATCC 14589, P. eucalypti type strain BD769, P. eucrina type strain BD872, P. gaviniae type strain A18/07, P. intestinalis type strain DSM 28113, P. punctata type strain BD876, P. rodasii type strain BD943, P. rwandensis type strain BD944, P. septica type strain BD874, P. stewartii type strain ATCC 8199, P. terrea type strain BD877, P. theicola type strain DSM 29122, P. vagans type strain BD765, and P. wallisia type strain BD946. Strain Russ shares 96.6% with P. agglomerans, 96.3% with P. allii, 97% with P. ananatis, 97% with P. anthophila, 96.6% with P. breneri, 96.4% with P. calida, 94.9% with P. citreus, 97.9% with P. coffeiphila, 95.7% with P. cyripedii, 96.8% with P. dispersa, 98.5% with P. eucalipti, 100% with P. eucrina, 96.9% with P. gaviniae, 95.5% with P. intestinalis, 96.2% with P. punctata, 97.5% with P. rodasii, 97.5% with P. rwandensis, 97.8% with P. septica, 97.8% with P. stewartii, 96.6% with P. terrea, 96.3% with P. theicola, 96.4% with P. vagans, and 98.5% with P. wallisia in the Pantoea genus. Phylogenetic analysis based on the 16s rRNA gene placed Pantoea eucrina strain Russ in the same node with the Pantoea eucrina strains IHB B 10086, C7, and CT194 (Table 2, and Fig. 2).

Compared to other Pantoea species with sequenced genomes, strain Russ shares 93% 16s rRNA gene similarity with representatives of Pantoea dispersa, 97% similarity with representatives of Pantoea stewartii, and 96% similarity with representatives of Pantoea ananatis.

3.2. Genome properties

The genome assembly process produced a contig N50 of 2633,372 bp with a total genome size of 3,939,877 bp. The GC content was 55.88%. One hundred and thirty four RNA genes were identified in the genome including 11 ribosomal RNA and 75 tRNA genes. The ribosomal RNA operon showed a typical bacterial organization with genes for 5S, 16S, and 23S rRNA and tRNAs tRNA^ile and tRNA^A^l^s. Of the...
Table 2
Classification and general features of Pantoea eucrina Russ [25].

| MIGS ID | Property | Term | Evidence code |
|---------|----------|------|---------------|
| Classification | Domain Bacteria | TAS [26] |
| | Phylum Proteobacteria | TAS [26] |
| | Class Gammaproteobacteria | TAS [26] |
| | Order Enterobacteriales | TAS [26] |
| | Family Enterobacteriaceae | TAS [26] |
| | Genus Pantoea | TAS [26] |
| | Species eucrina | TAS [26] |
| | Strain: Russ | TAS [26] |
| Gram stain | Negative | TAS [26] |
| Cell shape | Rod | TAS [26] |
| Motility | Motile | TAS [26] |
| Sporulation | Non-sporing | TAS [26] |
| Temperature range | Mesophile | TAS [26] |
| Optimum temperature | 28 °C | TAS [26] |
| pH range; optimum Carbon source | d-glucose, d-fructose, D-galactose, trehalose, D-mannose, cellulose, L-arabinose, glycerol, inositol, Dsaccharate, cis-aconitate, D-glucuronate, D-galacturonate, N-acetylglucosamine, D-glucose, DL-lactate, L-histidine, L-arabitol, L-glutamate, L-alanine and l-serine succrose, maltotriose, maltose, D-arabitol, L-arabitol, xylitol, D-mannitol, adonitol and citrate. | TAS [26] |

3838 detected genes, 3704 genes (96.51%) were protein-coding, of which 80.22% had a function prediction, 74.65% represented a COG functional category, and 8.1% were predicted to have a signal peptide. Using PSORT [16], we classified proteins as 41% cytoplasmic, 0.72% extracellular, and 30.3% associated with the membrane. Based on the presence of 139 single copy genes [17], the genome is predicted to be 81.3% complete. Genome statistics are shown in Table 3. The distribution of genes into COG functional categories is shown in Table 4.

3.3. Insights from the genome sequence

Genome analysis of Pantoea eucrina Russ identified a microorganism with a typical Gram-negative cell wall structure. The genome also suggests that the cell envelope contains the polar lipids phosphatidylglycerol, phosphatidylcholine, phosphatidylethanolamine, phosphatidylserine, and cardiolipin since genes for their biosynthesis were identified in the genome. Strain Russ genome also encodes for a complete flagellar assembly, in agreement with the isolate’s electron micrograph, as well as a type I pilus system belonging to the newly identified usher/chaperone system [18]. This system was first characterized in Acinetobacter baumannii and is linked to the early steps in biofilm formation [18]. The presence of genes encoding this system in Pantoea eucrina combined with the observation that it also possesses flagella imply a possible role of these genes to its virulence especially in nosocomial settings by allowing it to establish biofilms. When compared against the virulence factor database [19], the genome of Pantoea eucrina Russ showed 1077 virulence factor hits (29% of the protein-coding genes). These included Type I, Type II, Type III, Type IV, and Type Vb secretion systems. Most of these secretion systems have been linked to virulence in Gram-negative organisms [20], and could potentially contribute to opportunistic pathogenesis. Aside from their potential to be opportunistic pathogens, Pantoea endophytic strains range in their relationship with plant hosts from pathogenic to beneficial growth-promoters or bioprotectors [7]. Previous research suggested a relationship between the presence of virulence-associated genes on mobile elements and the pathogenicity of the strain towards plants [7]. Even though strain Russ was not isolated from a plant host, we sought to examine the possibility of its potential pathogenicity, or lack thereof, towards plants. Among 11 possible transposases identified in the genome, one putative transposase (IMG gene ID: 2650202385) is present in a cluster (with one other transposase and two insertion elements proteins) downstream from a sucrose utilization cluster and a tellurite resistance cluster. Since sucrose is a predominant disaccharide in higher plants tissues [21], the capability to degrade sucrose would be highly beneficial for plant endophytes. Some plants are also known to accumulate tellurium [22], which would warrant a mechanism of tellurium resistance in the endophytic bacteria affiliated with such plant hosts. The presence of sucrose utilization gene cluster, as well as a tellurium resistance gene cluster upstream from transposases and transposable elements in the genome of strain Russ might suggest its potential for plant pathogenesis. However, this claim requires further investigation.

Further analysis of KEGG pathways identified almost compete to complete catabolic pathways for utilization of glucose, fructose, mannose, xylose, arabinose and galactose, and all amino acids as carbon and energy sources. The genome also suggests the capability of xanthine degradation to glycine as well as uracil degradation to 3-hydroxypropionate, both of which indicate the capability to utilize purines and pyrimidines as energy sources. The genome encodes a complete TCA cycle and electron transport chain with F-type ATPase subunits confirming the aerobic nature of the microorganism. Facultative anaerobiosis is also suggested by the genome based on the presence of genes encoding for enzymes involved in lactate, acetate and formate fermentation were identified. Genomic analysis suggested auxotrophy for VitB12 and Riboflavin. However comparison of the protein-coding genes against the transporter database [23] identified several ABC and secondary transporters that could potentially be used for the import of such molecules.

3.4. Insights from comparative genomics

When the genome of Pantoea eucrina Russ was compared to 22 closely related genomes based on their KEGG profiles, the genome clustered with Pantoea sp. PSNH1 previously isolated from patients in a hospital setting and shown to carry several plasmids with antibiotic resistance cassettes [24] (Fig. 3A). This genomic similarity was confirmed when several genomic features (including the genome size, the number of genes, the number of transporters identified, the GC content, the number of non-coding bases, the number of genes belonging to COG categories, as well as the number of genes belonging to each COG category) were used to compare Pantoea eucrina Russ genome to the 22 other closely related genomes. The Russ genome was shown to cluster with the genomes of strains PSNH1, PSNH2, IMH, and B40 based on the lower number of genes belonging to the COG categories E, K, G, R, and P in these genomes (Fig. 3B).
virulence factor database identified 1077 genes in the genome with possible associated function including type I, II, III, IV, and Vb secretion systems, most of which could potentially contribute to opportunistic pathogenesis that was previously reported for members of the genus Pantoea. Comparative genomics using general genomic features as well as the KEGG function profile clustered the Russ genome with Pantoea strains previously isolated from hospital settings and shown to harbor antibiotic resistance-encoding plasmids.

4. Conclusions

This study presents the first draft genome sequence and functional annotation of a member of the genus Pantoea. The genome of Pantoea eucrina strain Russ revealed extensive sugar and amino acid degradation machinery, as well as the potential to use purines and pyrimidines as carbon and energy sources. Type 1 pili belonging to the new usher/chaperone system (pfam 05,229), and possession of flagella could contribute to the capability to form biofilms. Comparison to the virulence factor database identified 1077 genes in the genome with potential virulence-associated function including type I, II, III, IV, and Vb secretion systems, most of which could potentially contribute to opportunistic pathogenesis that was previously reported for members of the genus Pantoea. Comparative genomics using general genomic features as well as the KEGG function profile clustered the Russ genome with Pantoea strains previously isolated from hospital settings and shown to harbor antibiotic resistance-encoding plasmids.

Authors’ contributions

FM, BR, RR, MBC, and NY contributed to the analysis. FM, WDH, DPF, and NY wrote the manuscript. BR, CB, and RAH performed the lab experiments.

Table 3

| Attribute              | Value  | % of total |
|------------------------|--------|------------|
| Genome size (bp)       | 3,939,877 | 100%       |
| DNA coding (bp)        | 3,459,667 | 87.81%     |
| DNA G+C (bp)           | 2,205,503 | 55.98%     |
| DNA scaffolds          | 8      | 100%       |
| Total genes            | 3838   | 100%       |
| Protein coding genes   | 3704   | 96.51%     |
| RNA genes              | 134    | 3.49%      |
| Pseudo genes           | 0      |            |
| Genes in internal clusters | 829    | 21.60%     |
| Genes with function prediction | 3079   | 80.22%     |
| Genes assigned to COGs | 2865   | 74.65%     |
| Genes with Pfam domains | 3267   | 85.12%     |
| Genes with signal peptides | 312    | 8.10%      |
| Genes with transmembrane helices | 853    | 22.23%     |
| CRISPR repeats         | 0      |            |

The total is based on the total number of protein coding genes in the genome.

Table 4

| Code | Value | % of total | Description                                                      |
|------|-------|------------|------------------------------------------------------------------|
| J    | 242   | 7.5%       | Translation, ribosomal structure and biogenesis                  |
| A    | 1     | 0.03%      | RNA processing and modification                                  |
| K    | 258   | 8%         | Transcription                                                    |
| L    | 124   | 3.84%      | Replication, recombination and repair                            |
| B    | 0     | 0%         | Chromatin structure and dynamics                                  |
| D    | 44    | 1.36%      | Cell cycle control, cell division, chromosome partitioning       |
| V    | 67    | 2.08%      | Defense mechanisms                                               |
| T    | 186   | 5.77%      | Signal transduction mechanisms                                   |
| M    | 233   | 7.19%      | Cell wall/membrane biogenesis                                    |
| N    | 77    | 2.30%      | Cell motility                                                    |
| U    | 39    | 1.21%      | Intracellular trafficking and secretion                          |
| O    | 120   | 3.72%      | Posttranslational modification, protein turnover, chaperones     |
| C    | 176   | 5.46%      | Energy production and conversion                                 |
| G    | 312   | 9.07%      | Carbohydrate transport and metabolism                            |
| E    | 315   | 9.76%      | Amino acid transport and metabolism                               |
| F    | 95    | 2.94%      | Nucleotide transport and metabolism                               |
| H    | 177   | 5.49%      | Coenzyme transport and metabolism                                 |
| I    | 97    | 3.01%      | Lipid transport and metabolism                                   |
| P    | 198   | 6.14%      | Inorganic ion transport and metabolism                            |
| Q    | 43    | 1.33%      | Secondary metabolites biosynthesis, transport and catabolism      |
| R    | 232   | 7.19%      | General function prediction only                                  |
| S    | 165   | 5.11%      | Function unknown                                                 |
| -    | 1640  | -          | Not in COGs                                                      |

The total is based on the total number of protein coding genes in the genome.

Competing interests

All authors declare no competing interests.

Transparency document

The Transparency document associated with this article can be found, in online version.

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Pantoea eucrina strain Russ was isolated and selected for sequencing as part of a Howard Hughes Medical Institute funded project at Oklahoma State University. The project aims at improving undergraduate student persistence through authentic laboratory research. During a
Fig. 3. Comparative genomics of Pantoea eucrina strain Russ and 21 closely related genomes. (A) KEGG profile clustering of the genomes compared in this study. (B) PCA biplot of the genomic features and COG category distribution in the genomes compared. Genomes are represented by stars, where the strain name is depicted. Arrows represent genomic features or COG categories used for comparison. The arrow directions follow the maximal abundance, and their lengths are proportional to the maximal rate of change between genomes. The first two components explained 75% of variation.
two-semester long effort, undergraduate students isolate an environmental strain, and extract its genomic DNA. The genome is then sequenced and analyzed by undergraduate students as part of an upper division microbial genomics class. The current genome was analyzed by a team of undergraduate (BR, RR), and graduate (FM) students.

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