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Complete genome sequence of a psychrotrophic Pseudarthrobacter sulfonivorans strain Ar51 (CGMCC 4.7316), a novel crude oil and multi benzene compounds degradation strain

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Highlights:

*Pseudarthrobacter sulfonivorans* strain Ar51, a psychotrophic bacterium isolated from the Tibet permafrost of China, can degrade crude oil and multi benzene compounds efficiently in low temperature. Here we report the complete genome sequence of this bacterium. The complete genome sequence of *Pseudarthrobacter sulfonivorans* strain Ar51, consisting of a cycle chromosome with a size of 5.04 Mbp and a cycle plasmid with a size of 12.39kbp. The availability of this genome sequence allows us to investigate the genetic basis of crude oil degradation and adaptation to growth in a nutrient-poor permafrost environment.
Abstract

*Pseudarthrobacter sulfonivorans* strain Ar51, a psychrotrophic bacterium isolated from the Tibet permafrost of China, can degrade crude oil and multi benzene compounds efficiently in low temperature. Here we report the complete genome sequence of this bacterium. The complete genome sequence of *Pseudarthrobacter sulfonivorans* strain Ar51, consisting of a cycle chromosome with a size of 5.04 Mbp and a cycle plasmid with a size of 12.39kbp. The availability of this genome sequence allows us to investigate the genetic basis of crude oil degradation and adaptation to growth in a nutrient-poor permafrost environment.
Arthrobacter are most frequently isolated in soils and play important roles in biogeochemical cycles and decontamination (Unell et al., 2008). A. sulfonivorans, which is a novel methylotrophic strain, has the ability to degrade the dimethyl sulfoxide (DMSO) efficiently (Borodina et al., 2002). Based on phylogenetic groupings, 16S rRNA gene sequence similarities and homogeneity in peptidoglycan types, quinone systems and polar lipid profiles, A. sulfonivorans was reclassified in the genus Pseudarthrobacter gen. nov. Pseudarthrobacter sulfonivorans strain Ar51 was isolated from permafrost of the Tibet in SW China, is a psychrotrophic bacterium that can grow at temperatures between -4°C to 25°C, with an optimum growth temperature of 20°C. Biodegradation experiment has indicated that this strain can utilize the crude oil, alkane and multi benzene compounds efficiently in low temperature. Here we present the complete genome sequence of Pseudarthrobacter sulfonivorans strain Ar51 (previous name is Arthrobacter oxydans Ar51).

The raw sequence was generated using a shotgun approach employing the PacBio® RS II system and whole-genome shotgun technology with the Illumina HiSeq 2000 system yielding ~410× coverage (Roberts et al., 2013). Genomic DNA was sheared to an average fragment size of approximately 20 kb, and subsequently constructed to a SMRTbellTM library with a recovery of 40%. To generate the longest reads possible for this genome, the 20 kb SMRTbell library was size-selected using the BluePippinTM size selection system to remove SMRTbells less than 15 kb. Approximately, 14% of the final SMRTbell library was recovered with an average insert size of approximately 20 kb. The size-selected SMRTbell library was bound with P6 polymerase and sequenced with C4 chemistry and loaded using the one-cell-per-well protocol in the RS II instrument. Three SMRT Cells with 1.45 Gb post-filtering data were used for assembly. De novo assembly was carried out using the Hierarchical Genome Assembly Process (HGAP) version 3 (PacBio DevNet; Pacific Biosciences) workflow, including consensus polishing with Quiver as available in the SMRT® Analysis v 2.3.

The annotation for Pseudarthrobacter sulfonivorans strain Ar51 was conducted with NCBI Prokaryotic Genome Annotation Pipeline (Pruitt et al., 2012). The chromosome of Ar51 consists of 5,043,757 bp, the G+C content is 64.7%, 4390 coding sequence were identified, 12 rRNA genes and 50 tRNA genes. The plasmid of Ar51 consists of 12,394 bp, the G+C content is 56.3%, 3 coding sequence and 5 tRNA genes were identified (Table1). Over 58% (2548) of the predicted open reading frames (ORFs) present in the Ar51 genome could not be assigned a putative function.

Previous studies identified the alkane 1-monoxygenase (alkB) is the key factor to degrade the alkane in the bacteria (Belhaj et al., 2002). Unlike other related alkane biodegrade clusters only contain one copy of alkB, two copies of
alkB genes (AU252_RS14035, AU252_RS11540) were identified in the Ar51 genome. Additionally, the Ar51 also contains multi-aromatic compounds degrade genes, such as biphenyl degradation genes cluster (AU252_RS17305, AU252_RS17310, AU252_RS17315), benzoate degradation genes (benK, AU252_RS22695; benD, AU252_RS22710; AU252_RS22720; AU252_RS22725; AU252_RS22745; benK, AU252_RS00390).

The potential of Pseudarthrobacter sulfonivorans strain Ar51 to produce secondary metabolites was analyzed with the secondary metabolites search tool antiSMASH (Weber et al., 2015), which indicated that 5 putative gene clusters involved in the biosynthesis of different natural products (1 Bacteriocin, 1 typeIII PKS, 1 Siderophore, 2 unspecified cluster) are located in the chromosome and no gene cluster is located in the plasmid. Iron-uptake systems include a unique low-pH-induced iron transporter system and utilization systems. Siderophores are small, high-affinity iron chelating compounds secreted by many microorganisms as a means to acquire sufficient iron for their growth (Neilands, 1995). Ar51 contains a gene cluster to direct synthesis of Desferrioxamine E, a pathway it shares with other Arthrobacter species.

Compared with the psychrotrophic Arthrobacter strain A3, the Ar51 has the stronger ability to synthesis trehalase as the osmotic stress protector (Chen et al., 2011; Sun et al., 2016). In the genome of Ar51, it contains three pathways to synthesis trehalose: the OtsA/B (AU252_RS06465, AU252_RS06470, AU252_RS12615), TreY/TreZ (AU252_RS17210, AU252_RS17210) and TreS (AU252_RS10705, AU252_RS06525) pathways. Glutathione (GSH) is an important antioxidant in plants, animals, fungi, and some bacteria and archaea, preventing damage to important cellular components caused by reactive oxygen species such as free radicals, peroxides, lipid peroxides and heavy metals (Pompella et al., 2003). However, most of Arthrobacter species has no ability to synthesis GSH (Sun et al., 2016). Ar51 contains GSH synthesis pathway, consisting of gamma-glutamyltranspeptidase (AU252_RS22220) and glutathione synthetase (AU252_RS14885). In terms of cold adaption, the genome of this psychrotrophic bacterium has two aquaporin Z genes (AU252_RS16525, AU252_RS18930). Although bacterial aquaporin proteins can theoretically contribute to osmoregulation, studies indicate that they likely function to improve freeze tolerance under rapid-freezing conditions (Tanghe et al., 2006).

The complete genome chromosome and plasmid sequences have been deposited in GenBank database with accession number CP013747.1 and CP013748.1 respectively. This strain has been deposited at the China General Microbiological Culture Collection Center (CGMCC 4.7316).

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Table 1: General genome features of *Pseudarthrobacter sulfonivorans* strain Ar51

<table>
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<th>Feature</th>
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<th>Plasmid</th>
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<tbody>
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<td>Length (bp)</td>
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<tr>
<td>G + C content (%)</td>
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<td>CP013747.1</td>
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