Draft genome sequence of Ochrobactrum pseudogrignonense strain CDB2, a highly efficient arsenate-resistant soil bacterium from arsenic-contaminated cattle dip sites

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Publication Details
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Abstract
We report the 4.97-Mb draft genome sequence of a highly efficient arsenate-resistant bacterium, Ochrobactrum sp. strain CDB2. It contains a novel arsenic resistance (ars) operon (arsR-arsC1-ACR3-arsC2-arsH-mfs) and two non-operon-associated ars genes, arsC3 and arsB. The genome information will aid in the understanding of the arsenic resistance mechanism of this and other bacterial species.

Keywords
ochrobactrum, sequence, sites, genome, dip, draft, cattle, contaminated, arsenic, bacterium, soil, resistant, arsenate, efficient, highly, cdb2, strain, pseudogrignonense

Disciplines
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Draft Genome Sequence of *Ochrobactrum pseudogrignonense* Strain CDB2, a Highly Efficient Arsenate-Resistant Soil Bacterium from Arsenic-Contaminated Cattle Dip Sites

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We report the 4.97-Mb draft genome sequence of a highly efficient arsenate-resistant bacterium, *Ochrobactrum* sp. strain CDB2. It contains a novel arsenic resistance (ars) operon (arsR-arsC1-ACR3-arsC2-arsH- mfs) and two non-operon-associated *ars* genes, *arsC3* and *arsB*. The genome information will aid in the understanding of the arsenic resistance mechanism of this and other bacterial species.

Analysis revealed a novel *ars* operon (arsR-arsC1-ACR3-arsC2-arsH-mfs). Interestingly, two arsenate reductase genes coexist in it: the deduced protein ArsC1 is highly homologous to the ArsC of *Saccharomyces cerevisiae* (4), while ArsC2 more closely matched the *Escherichia coli* reductase (5). The last gene (mfs) is predicted to specify a protein belonging to the superfamily of major facilitator transporters while sharing only limited homology to known ArsB and ACR3 proteins. The function of this transmembrane transporter warrants investigation. We noticed that such an operon also exists in the *Ochrobactrum anthropi* strain ATCC 49188 genome (accession no. NC_009668.1, locus 521,109 to 525,495). A similar operon (lacking *mfs* has been identified in *Ochrobactrum tritici* strain SCI24 (6).

Two other putative *ars* genes, *arsC3* (which, like *arsC2*, also codes for a glutaredoxin-dependent arsenate reductase) and *arsB*, were also found in the genome. However, rather than being located in *ars* operons, each of them presents independently of other correlated *ars* genes. The existence of genes for three arsenate reductases and three transporters in the same genome may explain the remarkably high resistance of this bacterium to arsenate (1).

**Nucleotide sequence accession numbers.** The draft annotated genome sequence has been deposited at DDBJ/EMBL/GenBank under the accession number AKVI00000000. The version described in this paper is the first version, AKVI01000000. The draft genome sequence has been deposited at DDBJ/EMBL/GenBank under the accession number AKVI00000000.

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**REFERENCES**


A *Ochrobactrum* strain (CDB2) isolated from arsenic-contaminated cattle tick dip sites in northeastern New South Wales, Australia, exhibited high resistance to arsenical compounds, especially arsenate (1). Obtaining the full genome sequence of this organism would help us to understand the mechanism of arsenic resistance of this bacterium and related species.

A pair-end library was constructed and pyrosequencing was performed using a Roche GS FLX+ sequencer at the John Curtin School of Medical Research, the Australian National University, Canberra. After assembling the raw data by using the GS de novo Assembler (Roche, version 2.6.3), we used the RAST (Rapid Annotations using Subsystems Technology) server (2) for draft genome annotation, subsystem classification, and G+C content calculation. The putative *ars* genes were further analyzed with BLASTp and HHpred (http://toolkit.tuebingen.mpg.de/hhpred).

The NCBI Prokaryotic Genomes Automatic Annotation Pipeline (PGAAP) (http://www.ncbi.nlm.nih.gov/genomes/static/Pipeline.html) was used to produce GenBank format data.

The draft genome of *Ochrobactrum* sp. strain CDB2 comprises 4,971,228 bases assembled from a 20-fold coverage of raw data and has a G+C content of 53.6%. It contained 147 contigs, with the longest consisting of 350,235 bases while the median length of all contigs was 25,233 bases. We found 4,868 coding sequences (CDSs) and 54 tRNA tRNA genes in the constructed genome.

Our previous identification at the genus level was based on an analysis of a partial 16S rRNA gene sequence (1,341 bases) and a fatty acid composition profile (1). With the full 16S sequence obtained, a new search showed a 100% match with a 1,387-bp 16S sequence from *Ochrobactrum pseudogrignonense* strain CCUG 30717T (accession no. AM422371) (3). The nucleotide sequences of three other genes, *groEL*, *gyrB*, and *recA* (accession no. FM863822, FM863816, and AM422877, respectively), also matched perfectly, except for one base in *groEL*, between CDB2 and *Ochrobactrum pseudogrignonense* CCUG 30717T. These identified CDB2 as a strain of *Ochrobactrum pseudogrignonense*.


