Finished Genome Sequence of *Collimonas arenae* Cal35

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We announce the finished genome sequence of soil forest isolate *Collimonas arenae* Cal35, which comprises a 5.6-Mbp chromosome and 41-kb plasmid. The Cal35 genome is the second one published for the bacterial genus *Collimonas* and represents the first opportunity for high-resolution comparison of genome content and synteny among collimonads.

Collimonads are oligotrophic, chitinolytic, rhizosphere-competent soil bacteria with antifungal, mycophagous, and mineral-weathering properties (1). To date, three species of *Collimonas* have been described (2, 3), but the only *Collimonas* genome available is that of the Dutch dune soil isolate *C. fungivorans* Ter331 (CFTer331, GenBank accession no. CP002745 for the chromosome and EU315244 for plasmid pTer331). The CFTer331 genome has been instrumental in describing the *Collimonas* chitinolytic system (4), the genes underlying the production of antifungal polynylene-like compounds called collimycins (5), the molecular interactions with model fungus *Aspergillus niger* (6), and the phenotypic variation among other collimonads isolated from the same dune soil as Ter331 (7). However, with only one *Collimonas* genome sequence available to date, intrageneric comparison at single-nucleotide resolution has not yet been feasible. Here, we report the finished genome sequence of *Collimonas arenae* Cal35 (CCal35), which was recovered from a forest soil in California (8).

Genomic DNA of CCal35 was extracted from an overnight culture using a DNeasy Blood and Tissue kit (QIAGEN, Valencia, CA). A 10-kb PacBio RS II-compatible library was constructed by the UC Davis Genome Center and sequenced on the 3 single-molecule real-time (SMRT) cells using P4-C2 chemistry. De novo assembly was performed with the help of SMRT Analysis software v2.2.0 (Pacific Biosciences) featuring HGAP 2 (9), and subsequent correction with quiver in addition to Gepard v1.30 (10) to reveal two circular replicons: a 5,603,532-bp chromosome (*G*+*C* content 56.15%; 85.54× coverage) and a 41,440-bp plasmid (*G*+*C* content 50.54%; 73.21× coverage) which we designated pCal35. Gene prediction by RAST (11) revealed 5,019 coding sequences, 54 tRNA genes, and 3 rRNA operons (5S, 16S, 23S) on the chromosome, in addition to 57 coding sequences on pCal35.

The chromosome of CCal35 appears mostly syntenic with that of CFTer331, with the exception of one 2.75-Mbp inversion centered on the origin of replication. CCal35 shares 69% of its genes with Ter331, of which 80% are >80% identical at the predicted amino acid level. The chitinolytic system of CCal35 is very similar to that of CFTer331, but 35 possesses at least two additional chitinase genes. Clearly missing from the CCal35 genome is gene cluster K which is responsible for the production of collimycins (5–7). Interestingly, in CFTer331, this gene cluster is located near one of the chromosome inversion points. Unrelated to pTer331 (12), plasmid pCal35 carries several genes with high coding similarity to plasmids from plant-pathogenic species of *Xanthomonas* (for example, pXAC64, pXCV38, pXcB) and *Ralstonia* (pRSC35). In addition, it harbors a stretch of DNA with high similarity to plasmid 2 from *Polaromonas* JS666 and coding for tRNA 2-selenouridine synthase, alkane 1-monooxygenase, and a LuxR-type regulator.

### Nucleotide sequence accession numbers

The Cal35 chromosome and plasmid sequences are available under GenBank accession numbers CP009962 and CP009963, respectively.

### ACKNOWLEDGMENTS

This work was funded by the University of California at Davis, the California Tomato Research Institute, and the Ministry of Science and Technology (NSC-103-2911-I-005-301, NSC-102-2911-I-005-301) and the Ministry of Education (the ATU plan) of Taiwan, Republic of China.

Assembly and annotation of the CCal35 genome sequence were performed by V.C.L.d.J. and J.J.W. (in the lab of J.H.J.L., visiting from the UC Davis Genome Center). Interpretation of the annotation data was done by J.H.J.L., who also drafted the manuscript with inputs, edits, and final approval by all other authors. This is NIOO-KNAW publication number 5741.

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