Genomic Sequencing of *Orientia tsutsugamushi* Strain Karp, an Assembly Comparable to the Genome Size of the Strain Ikeda

Hsiao-Mei Liao, Chien-Chung Chao, Haiyan Lei, Bingjie Li, Shien Tsai, Guo-Chuian Hung, Wei-Mei Ching, Shyh-Ching Lo

Division of Cellular and Gene Therapies, Office of Cellular, Tissue and Gene Therapies, Tissue Microbiology Laboratory, Center for Biologics Evaluation and Research, U.S. Food and Drug Administration, Silver Spring, Maryland, USA;
Naval Medical Research Center, Silver Spring, Maryland, USA;
Uniformed Services University of the Health Sciences, Bethesda, Maryland, USA

* Present address: Haiyan Lei, Pediatric Oncology Branch, National Cancer Institute, Bethesda, Maryland, USA.
H.-ML. and C.-C.C. contributed equally to this work.

*Orientia tsutsugamushi*, an intracellular bacterium, belongs to the family *Rickettsiaceae*. This study presents the draft genome sequence of strain Karp, with 2.0 Mb as the size of the completed genome. This nearly finished draft genome sequence was annotated with the RAST server and the contents compared to those of the other strains.

**Results and Discussion**

The first version of draft genome sequence of strain Karp was deposited in GenBank under accession no. LYMA00000000. O. tsutsugamushi is an obligate intracellular organism causing scrub typhus. The disease in endemic triangle influences more than one billion people in the world (1). The genome size among the strains is varied; only two completed genomes, those of strain Boryong and Ikeda, have been published (2, 3). The genome of *O. tsutsugamushi* contains near 40% identical repeats, which troubles the sequence assembly. In this study, we combined a cloned-based method and high-throughput sequencing to assemble the draft genome of strain Karp, which was originally isolated from Papua New Guinea (4, 5).

Clone-based whole-genome shotgun sequencing generated the reads by sequencing bacterial artificial chromosome (BAC) clones with the Sanger method. The assembled reads formed 116 contigs before high-throughput whole-genome sequencing (HTGS) was available; the contig length ranged from 1,153 to 1,617 bp. The raw reads of HTGS were used to join the contigs from the Sanger read assembly, and the original 116 contigs were reduced to 99. HTGS was conducted using the Illumina MiSeq platform with the 2 × 250 bp paired-end mode. The trimmed raw reads were de novo assembled into an original 1,011 contigs using CLC Genomics Workbench 9.0, with an average coverage of 73 ×. The contigs from HTGS were first mapped to three references: (i) the contig set from the Sanger assembly, (ii) strain Boryong (accession no. NC_009488.1), and (iii) strain Ikeda (accession no. NC_010793.1), and subjected to a BLAST search against the nt/nr database to eliminate contigs from background DNA contaminations. The raw reads retrieved from filtered contigs were de novo assembled and mapped again to the three references, plus a published Karp draft genome (GenBank accession number NZ_LANM00000000.1) for the secondary background cleaning. All retained HTGS contigs of strain Karp can be mapped to at least one of the references, and five contigs from HTGS (combined length, 11,304 bp) were added to the original 99 Sanger contigs (2,011,605 bp) to form the draft genome of 2,022,909 bp (30.41% G+C content). The 104 (99 + 5) contigs of Karp were aligned to the Ikeda complete genome using the CONTIGuator software (6).

The Karp genome was annotated using RAST server version 2.0 (7), with 2,089 coding sequences (CDSs), 1,090 transcribed from the positive strand, and 999 transcribed from the negative strand; 2,052 of these were categorized into 185 subsystems, and 37 were RNAs. A SEED Viewer sequence comparison (8) showed that 1,604 genes of strain Boryong and 1,951 genes of strain Ikeda could be found in the Karp draft genome. The HTGS was also conducted to sequence the other two *O. tsutsugamushi* strains, AFSC4 and AFSC7, without the information about the repetitive sequences assembled from cloned-based sequencing. The SEED Viewer functional comparison (8) revealed that two genes presenting in both AFSC4 and AFSC7 were absent in Karp, even though it possesses more CDSs in its genome (data not shown). Previous studies showed that strain Karp was sensitive to antibiotics, whereas AFSC4 was insensitive (9), and AFSC7 had similar internal observations. A completed draft genome of AFSC4 and AFSC7 for delicate comparison with the Karp genome may provide the possible targets for investigation of microbial drug resistance mechanisms.

**Accession number(s).** The first version of draft genome sequences of *Orientia tsutsugamushi* strain Karp was deposited in GenBank under accession no. LYMA00000000.

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REFERENCES