Complete Genome Sequence of *Mycobacterium ulcerans* subsp. *shinshuense*

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*Mycobacterium ulcerans* subsp. *shinshuense* produces mycolactone and causes Buruli ulcer. Here, we report the complete sequence of its genome, which comprises a 5.9-Mb chromosome and a 166-kb plasmid (pShT-P). The sequence will represent the essential data for future phylogenetic and comparative genome studies of mycolactone-producing mycobacteria.

Received 4 August 2016 Accepted 8 August 2016 Published 29 September 2016

Citation Yoshida M, Nakanaga K, Ogura Y, Toyoda A, Ooka T, Kazumi Y, Mitarai S, Ishii N, Hayashi Y, Hoshino Y. 2016. Complete genome sequence of *Mycobacterium ulcerans* subsp. *shinshuense*. Genome Announc 4(5):e01050-16. doi:10.1128/genomeA.01050-16.

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*Mycobacterium ulcerans* is a causative agent of Buruli ulcer (1). Buruli ulcer is characterized as a chronic, indolent, necrotizing disease of skin and soft tissue, which produces ulcers followed by scar formation (2). Its first reported case in Japan occurred in 1980 (3, 4). The causative agent was isolated and identified as *M. ulcerans* subsp. *shinshuense* because it was closely related to *M. ulcerans*, although certain distinct differences were observed (3–5). The first clinical isolate (ATCC 33728T) is a reference strain of *M. ulcerans* subsp. *shinshuense* (6). Here, we report the complete genome sequence of the chromosome and a large plasmid of ATCC 33728T.

The strain was grown on Middlebrook 7H10 agar medium. DNA was purified with a high pure PCR template preparation kit (Roche Diagnostics). The genome sequence was determined using the 454 GS FLX Titanium system (Roche Diagnostics). A total of 889,811 single-end and 115,770 paired-end (8-kb insert) reads were assembled with the GS Assembler software version 2.6 into three scaffolds containing 259 gaps. All gaps, except for one large gap in the plasmid, were closed by sequencing of gap-spanning PCR products using an ABI3130xl DNA sequencer (Applied Biosystems). Illumina 150 × 2 paired-end reads (1,249,780 reads) obtained by a MiSeq sequencer (Illumina) and PacBio reads (47,671 subreads over 15.0 kb) obtained by an RS II system were used for sequence- and assemble-error correction and closing of a large gap in the plasmid. Automated annotation was carried out with the Microbial Genome Annotation Pipeline (MiGAP) (http://www.migap.org). Manual curation of the automatic annotation was performed using the in silico molecular cloning (In Silico Biology).

The chromosome of ATCC 33728T is 5,899,681 bp in length with a 65.64% G+C content, and the plasmid pShT-P is 166,617 bp in length with a 62.76% G+C content. The average nucleotide identities were 98.36% to *M. ulcerans* (strain Ag99) and 99.10% to *M. liflandii* (strain 128FXT), indicating their close genetic relatedness and supporting its taxonomic position as a subspecies of *M. ulcerans* (7). The chromosome contains 5,015 predicted protein coding sequences (CDSs), which is much more than those of the *M. ulcerans* strain Ag99 chromosome (4,160). Plasmid pShT-P contains 72 CDSs, whereas 81 CDSs were present in the large plasmid of *M. ulcerans* (8). The total number of insertion sequences is 233 (12 on the plasmid), which is less than that of *M. ulcerans* (324), but more than that of *M. marinum* (50 in strain M) (9). Notably, the number of IS2606 is significantly less than that of *M. ulcerans* (2 versus 91). The draft genome sequence of ATCC 33728 was previously determined (10), but it contains no plasmid sequence. This is probably due to the deletion of plasmid pShT-P, which frequently occurs during *in vitro* cultivation (10, 11). The genome sequence of *Mycobacterium ulcerans* subsp. *shinshuense* ATCC 33728T will represent the essential data for future phylogenetic and comparative genome studies and will be useful for better understanding the evolution of mycolactone-producing mycobacteria.

**Accession number(s).** This complete genome sequence has been deposited at DDBJ/ENA/GenBank under the accession no. AP017624 to AP017625.

**FUNDING INFORMATION**

This work, including the efforts of Kazue Nakanaga, Yoshitoshi Ogura, Atsushi Toyoda, Tadasuke Ooka, Tetsuya Hayashi, and Yoshihiko Hoshino, was funded by Ministry of Education, Culture, Sports, Science, and Technology (MEXT) (221S0002). This work, including the efforts of Yoshihiko Hoshino and Kazue Nakanaga, was funded by Japan Society for the Promotion of Science (JSPS). This work, including the efforts of Mitsunori Yoshida, Norihisa Ishii, and Yoshihiko Hoshino, was funded by Japan Agency for Medical Research and Development (AMED).

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