



## Complete Genome Sequence of *Staphylococcus aureus* FCFHV36, a Methicillin-Resistant Strain Heterogeneously Resistant to Vancomycin

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We report here the sequence of the entire chromosome of *Staphylococcus aureus* strain FCFHV36, a methicillin-resistant strain heterogeneously intermediate to vancomycin, bearing a type II staphylococcal chromosome cassette *mec* element (SCC*mec*), belonging to multilocus sequence type (MLST) 105, and isolated from a vertebra of a patient with osteomyelitis.

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We present here the sequence of the entire chromosome of *Staphylococcus aureus* strain FCFHV36, recovered from a vertebral biopsy sample from a patient diagnosed with community-acquired osteomyelitis and under medical care in a hospital in the state of Santa Catarina, Brazil. FCFHV36 presents an MIC of vancomycin of 2  $\mu$ g/ml, which classifies it as being susceptible to this antibiotic. The patient, however, did not respond to therapy with vancomycin. A heterogeneous resistance (heterogeneous vancomycin-intermediate *S. aureus* [hVISA]) profile was detected by the population analysis profile-area under the curve (PAP-AUC) technique (1), meaning that cell subpopulations of this strain present higher MICs than those of the overall cell population. The PAP-AUC ratio of FCFHV36 to hVISA type strain Mu3 was 1.02.

The total genomic DNA of FCFHV36 was used to construct a paired-end (PE) library and a mate-paired (MP) library, which were separately sequenced using the MiSeq platform (Illumina, Inc.). A total of 4,981,028 75-bp-long reads were generated for the paired-end library, and a total of 1,181,570 250-bp-long reads were generated for the mate-paired library, which had a mean insert distance of 2,700 bp. Both libraries were simultaneously used as input for de novo assembly using the A5 pipeline (2), generating 46 contigs (sum, 2.84 Mbp; N<sub>50</sub>, 147.9 kbp; max length, 385.9 kbp). In order to build a scaffold, the contigs were ordered by synteny against a reference chromosome using Gepard (3). The reference genome chosen was the publicly available genome presenting the most similar k-mer spectrum to the contigs, as determined using KmerFinder (4), which was S. aureus JH1 (NCBI GenBank accession no. NC\_009632). Contigs pertaining to plasmids were separated. The contig order was verified by aligning mate-paired reads against the scaffold and verifying the existence of mate pairs straddling the gap close to the mean insert distance using Geneious 7 (5). The correct position of contigs without synteny to the JH1 reference genome was also determined by matepaired distance information. Gaps were then filled with GapFiller

using reads from the paired-end library. After manual curation of gaps, the final circularized chromosome was annotated with Prokka (6), and features were manually curated by blasting against the Gen-Bank nr database. Insertion sequences were found using the ISfinder database (7) and annotated manually using Artemis (8).

The chromosome of FCFHV36 carries 2,619 protein-coding sequences, 7 pseudogenes, 58 tRNA genes, and 16 rRNA genes. The *mecA* gene, which confers resistance to most  $\beta$ -lactams, is carried by a type II staphylococcal chromosome cassette *mec* element (SCC*mec*). *In silico* multilocus sequence typing (MLST) was able to attribute sequence type 105 (ST105) to the strain. Comparative genomics between this sequence and vancomycin-susceptible, VISA, and other hVISA stains will help determine the polymorphisms that correlate with decreased vancomycin susceptiblity in *S. aureus*, which is hard to detect in the clinical laboratory.

**Nucleotide sequence accession number.** The complete genome sequence of *S. aureus* strain FCFHV36 has been deposited in DDBJ/EMBL/GenBank under the accession no. CP011147.

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We declare no conflicts of interest.

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