

Salmonella enterica subsp. enterica serovar Enteritidis strain SE74 chromosome, complete genome

GenBank: CP050723.1

[FASTA Graphics](#)

LOCUS CP050723 438 bp DNA linear BCT 06-APR-2020

DEFINITION Salmonella enterica subsp. enterica serovar Enteritidis strain SE74

chromosome, complete genome.

ACCESSION [CP050723](#) REGION: 995854..996291

VERSION CP050723.1

DBLINK BioProject: [PRJNA615288](#)
BioSample: [SAMN14450169](#)

KEYWORDS .

SOURCE Salmonella enterica subsp. enterica serovar Enteritidis
(Salmonella
enteritidis)

ORGANISM [Salmonella enterica subsp. enterica serovar Enteritidis](#)
Bacteria; Proteobacteria; Gammaproteobacteria; Enterobacterales;
Enterobacteriaceae; Salmonella.

REFERENCE 1 (bases 1 to 438)

AUTHORS Chen,Z.

TITLE Genomic analyses of multidrug-resistant Salmonella Indiana,
Typhimurium, and Enteritidis isolates using MinION and MiSeq
sequencing technologies

JOURNAL Unpublished

REFERENCE 2 (bases 1 to 438)

AUTHORS Chen,Z.

TITLE Direct Submission

JOURNAL Submitted (26-MAR-2020) Joint Institute for Food Safety and
Applied
Nutrition, University of Maryland, 5145 Campus Drive, Patapsco
Building, Suite 2134, College Park, MD 20742, USA

COMMENT The annotation was added by the NCBI Prokaryotic Genome
Annotation

Pipeline (PGAP). Information about PGAP can be found here:
https://www.ncbi.nlm.nih.gov/genome/annotation_prok/

##Genome-Assembly-Data-START##

Assembly Method :: Unicycler v. 0.4.8

Assembly Name :: SE74

Genome Representation :: Full

Expected Final Version :: Yes

Genome Coverage :: 97x

Sequencing Technology :: Illumina MiSeq; Oxford Nanopore MiniION


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Control and Prevention"
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KTI"

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ORIGIN

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