



2 Title

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- 3 Draft genome sequence of Phascolarctobacterium faecium DSM 14760 isolated
- 4 from koala feces in Australia

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- 16 evelynvaldovinos@icloud.com.
- 17 Abstract
- 18 Phascolarctobacterium faecium was originally isolated from koala feces in 1992. Here, we sequence and
- analyze the type strain, with a length of 2,317,131 bp, 27 contigs, and average G+C content of 43.73%,
- and three virulence factors that are potential health hazards to humans..
- 21 Announcement
- 22 Introduction
- 23 Phascolarctobacterium faecium is a spiral-shaped, gram-negative, anaerobic bacterium (1) that produce
- short-chain fatty acids acetate and propionate (2) and are associated with hosts' metabolic state and
- 25 mood (3), The bacteria can be used as a biomarker for colorectal cancers due to its high prevalence in
- patients (4). The 16S rRNA sequence can be found in NCBI under the accession number X72865. P.
- 27 faecium is found in the gut microbiome of koalas (2) and humans (3). The type of strain was isolated
- feces in Australia in 1992 (5). Feces were emulsified in 0.25-strength Ringer solution serially diluted 10-

- 29 fold, plated onto Wilkins-Chalgren Anaerobe agar (Oxoid) with 5% defibrinated horse blood, incubated
- for 5 days at 37°C in 10% CO2, 10% H2, and 80% N2 by Bio-Bag (6). P. faecium was sequenced as part of
- 31 the Genomic Encyclopedia of Type Strains, Phase IV, with the goal of sequencing genomes valuable for
- metagenomic binning, taxonomic classification, and comparative biology (7).

33 Methods and related outcomes

- 34 Details on organism growth and DNA isolation to be provided by DSMZ. Draft genome of
- 35 Phascolarctobacterium faecium DSM 14760 was sequenced at the DOE Joint Genome Institute (JGI)
- using the Illumina HiSeq-2000 1TB platform, generated 8,845,394 raw reads totaling for 1,335,654,494
- 37 bp. Raw data was filtered using BBTools per SOP 1061, which removed about 200 bp from each contig.
- 38 The final drafts had a total of 27 contigs in 21 scaffolds, totaling 2,317,131 bp in size. The sequencing
- 39 FASTQ file, using the BZZZN library, showed 8,787,470 reads. Pair-end sequence processed 151 bp reads.
- 40 The genome assembly was 100% complete with 0.23% contamination using the software, SPAdes,
- 41 v3.10.1, with parameters —phred-offset 33 —cov-cutoff auto –t 16 –m 115 —careful –k 25,55,95 —12
- 42 (8). The Standard JGI Microbial Genome Annotation Pipeline was used for annotation (9).

43 Results

- 44 The genomic features of Phascolarctobacterium faecium are listed in Table 1. The highest
- 45 similarity detected was draft type strain P. faecium JCM 14760 genome (Assembly accession
- 46 number GCF 009719105.1; 23 contigs). Next closest relatives were Acidaminococcus
- 47 fermentans DSM 20731 and Acidaminococcus massiliensis Marseille-P2828 T, with respective
- 48 (dDDH-d4) similarities of 39.5% and 31.5%. These relationships are depicted in the phylogenetic
- 49 tree in Figure 1.
- 50 The Comprehensive Antibiotic Resistance Database (CARD) website version 4.0.0 compares our bacterial
- 51 genome sequences to determine their antibiotic resistance through their AMR genes (10). There is
- 52 resistance to the disinfecting agents fluoroquinolones, tetracycline, and vancomycin, which are the van
- 53 genes. The CRISPRCasFinder website version 1.12 identified two CRISPR localization regions in the
- 54 genome with evidence (11). AntiSMASH 7.0 identified two secondary metabolite regions in the genome:
- NZ_QLTS01000002.1 (RiPP-like) and NZ_QLTS01000006.1 (ranthipeptide) (12). Using the Virulence
- 56 Factor Database (VFDB) software version 2.0, there are three main virulence factors that can be a
- 57 possible health hazard to humans (13). Factors are based on stress response, biofilm formation, and
- 58 virulence gene activation.

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Table 1: Genomic Features of *Phascolarctobacterium faecium* DSM 14760.

Feature:	Finding:
Length (bp)	2,317,131
No. of Scaffolds	21
No. of Genes	2248

No. of Protein coding genes	2176
No. of rRNA genes	8
No. of tRNA genes	54
GC content %	43.73
Scaffold N50 (bp)	195,470
Average fold coverage	576.43

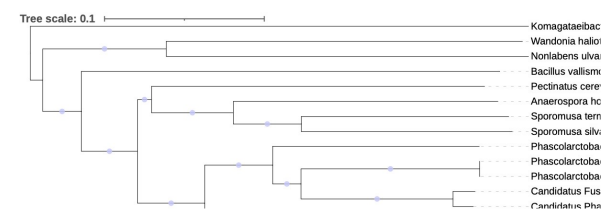


Figure 1. The tree above was inferred with FastME 2.1.6.1 [7] from whole-proteome-based GBDP distances. The branch lengths are scaled via GBDP distance formula *d5*. Branch values are GBDP pseudo-bootstrap support values > 60 % from 100 replications, with an average branch support of 100.0 %. The tree was midpoint-rooted [8].

Data availability statement

- The raw reads have been deposited in the NCBI SRA under the accession number
- 68 X72865.

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- 73 05CH11231Links to an external site.

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