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Genome Sequences of Three *Lactobacillus* Species Strains of the Stomach of the White-Footed Deermouse (*Peromyscus leucopus*)

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ABSTRACT Three colony types of *Lactobacillus* were isolated from the stomach of LL colony stock *Peromyscus leucopus* deermice, a reservoir for several human zoonoses. Genome sequences revealed two isolates to be new strains of *Lactobacillus animalis* and *Lactobacillus reuteri*. The third was distinct from known species and was provisionally designated *Lactobacillus* sp. strain LL6.

Peromyscus leucopus, a widely distributed, abundant mammal in North America, is an important reservoir for agents of tick-borne zoonoses, including Lyme disease, babesiosis, anaplasmosis, and a viral encephalitis (1). The genus *Peromyscus* clusters with hamsters and voles rather than with murids such as *Mus musculus* (2). *P. leucopus* is the target for field-delivered oral vaccines (3), but little is known about the gastro-intestinal (GI) microbiota of this or other *Peromyscus* species.

During study of the GI tract of animals of *P. leucopus* LL stock (Peromyscus Genetic Stock Center, University of South Carolina), we identified three types of *Lactobacillus* in the stomachs, ceca, and feces of both females and males. The organisms were isolated from stomach contents on Rogosa SL agar (Sigma-Aldrich) plates in a candle jar at 37°C and were distinguished by colony morphologies. Single colonies of each were cultivated in Difco MRS broth (Becton, Dickinson). High-molecular-weight genomic DNA was extracted from harvested cells with a Qiagen Genomic-tip 100/G kit.

Their genomes were sequenced by two methods, (i) for long reads on an Oxford Nanopore Technology (ONT) MinION Mk1B instrument with a ligation sequencing kit, R9.4.1 flow cell, MinKNOW v. 19.6.8 for primary data acquisition, and Guppy v. 3.2.4 for base calling with default settings and (ii) for short reads on an Illumina MiSeq instrument with paired-end v2 microchemistry and 150 cycles. The library was constructed using the NEXTflex rapid DNA-seq kit (Bioo Scientific), the quality of sequencing reads was analyzed using FastQC (Babraham Bioinformatics), and reads were trimmed of Phred scores of <15 and corrected for poor-quality bases using Trimmomatic (4).

A hybrid assembly of each genome was carried out with Unicycler v. 0.4.7 (5) with default settings and 16 threads on the high-performance computing cluster of the University of California Irvine. Complete, circularly permuted chromosomes were obtained for two of the isolates (LL1 and LL7), and two contigs of 1.75 Mb and 0.32 Mb, respectively, were obtained for the third (LL6) (Table 1). Annotation was provided by the NCBI Prokaryotic Genome Annotation Pipeline (6). By the criterion of 16S ribosomal sequence identity of \geq 99%, isolates LL1 and LL7 were identified as *Lactobacillus animalis* and *Lactobacillus reuteri*, respectively. Other sequences of these organisms confirmed those classifications.

L. reuteri LL7 had another replicon (Table 1) that had coding sequences for a Rep_3 (pfam01051) plasmid replication initiation protein and multiple transposases. This is consistent with a type of megaplasmid observed in other lactobacilli (7).

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Received 8 August 2019 Accepted 10 September 2019 Published 3 October 2019 TABLE 1 Resources from this study, sequencing statistics, and characteristics of chromosomes and plasmids of three Lactobacillus spp.

		BioProiect BioSample		SRA accession no. ONT reads	No. of ONT reads	Mean covera	Mean coverage (×) Size (bp)	Size (bp)		G+C content (%)	(%	GenBank accession no.	
Species	Strain	accession no.	Strain accession no. accession no. (Illumina, ONT) (×106)/N ₅₀ (kb) ONT Illumina Chromosome Plasmid Chromosome Plasmid Chromosome(s)	(Illumina, ONT)	$(\times 10^6)/N_{50}$ (kł	DNT (c	Illumina	Chromosome	Plasmid	Chromosome	Plasmid	Chromosome(s)	Plasmid
Lactobacillus animalis LL1 PRJNA533837 SAMN11468794 SRX5831611, 2.31/6.11 SRX5831612	LL1	PRJNA533837	SAMN11468794	SRX5831611, SRX5831612	2.31/6.11	2,098	2,098 182	2,280,577 4,016 41.0	4,016	41.0	37.1	37.1 CP039849	MK858222
Lactobacillus reuteri LL7 PRJNA554696 SAMN12280509 SRX6494060, SRX6494061	LL7	PRJNA554696	SAMN12280509	SRX6494060, SRX6494061	1.52/0.87	423	218	2,205,740	178,977 38.9		38.1	CP041676	CP041677
Lactobacillus sp.	9TT	PRJNA554698	LL6 PRJNA554698 SAMN12280527 SRX6492358, SRX6492357	SRX6492358, SRX6492357	2.73/6.21	2,601 278		2,067,236	6,969	33.5	36.4	VLLR01000001 (contig 1), VLLR01000003 VLLR01000002 (contig 1)	VLLR01000003

Isolate LL6 could be not be identified with a known *Lactobacillus* species in the genome database. Over its full-length 16S rRNA gene sequence, the closest organism was *Lactobacillus amylolyticus* at 95.6% nucleotide identity. The identities of LL6's coding sequences for the core genome genes *recA*, *rpoB*, *gyrA*, and *rplB* were no more than 83%, 87%, 81%, and 94%, respectively, with any other *Lactobacillus* sp. These distances suggest that this represents a previously unknown taxon, provisionally designated *Lactobacillus* sp. strain LL6.

Data availability. The strains described here are available from the corresponding author. Table 1 provides the publicly available resources from this study with hyperlinks to repositories.

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