## Complete Genome of *Lactococcus lactis* subsp. *cremoris* UC509.9, Host for a Model Lactococcal P335 Bacteriophage

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Here, we report the complete genome of *Lactococcus lactis* subsp. *cremoris* UC509.9, an Irish dairy starter. The circular chromosome of *L. lactis* UC509.9 represents the smallest among those of the sequenced lactococcal strains, while its large complement of eight plasmids appears to be a reflection of its adaptation to the dairy environment.

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Lactococcus lactis strains are used extensively worldwide for the production of fermented dairy products. Bacteriophage (phage) attack during this fermentation process can lead to slow or failed fermentations and is therefore of major economic concern (1). *L. lactis* subsp. *cremoris* UC509 is an Irish cheddar starter strain and is the lysogenic host of the model P335-type phage Tuc2009 (2–6). *L. lactis* UC509.9, whose genome sequence is presented here, is a prophage-cured Tuc2009-sensitive derivative of UC509 (7).

While lactococcal phages are subject to intensive scientific scrutiny, the specific interactions with their hosts are poorly understood. To further our understanding regarding the molecular interplay between Tuc2009 and its host, we sequenced the genome of L. lactis UC509.9. Sequencing was performed by Agencourt Bioscience (Beverly, MA) and Macrogen (Seoul, Republic of Korea) using a combination of 454 sequencing of a 3-kb fragment library using Roche standard procedures and of Sanger sequencing of a 36-kb insert library followed by homopolymer tract correction using Illumina sequencing. Initial sequence assembly was performed using GSassembler (Roche). Gap closure and quality improvements were performed by Sanger sequencing of gapclosing PCR products as suggested by Projector 2 (8) with the Staden package (9). Homopolymer tract single nucleotide polymorphisms (SNPs) were detected and corrected using Robust Variant detection (ROVAR) (V. de Jager, B. Renckens, R. J. Siezen, and S. A. F. T. van Hijum, unpublished data [https://trac.nbic.nl /rovar/]) applied to Illumina sequencing data as described previously (10), resulting in a >200-fold coverage of the genome. Putative protein-encoding genes were identified using Prodigal version 2.0 (11). The results were inspected using Artemis (12), with manual checking and editing using BLASTP, Pfam (13), Kyoto Encyclopedia of Genes and Genomes (KEGG) (14), and Clusters of Orthologous Groups (COG) databases (15).

The complete genome of *L. lactis* UC509.9 consists of a single circular chromosome of 2,250,427 bp (35.88% G+C content)

plus eight plasmids: pCIS1 (4,263 bp), pCIS2 (5,961 bp), pCIS3 (6,159 bp), pCIS4 (7,045 bp), pCIS5 (11,676 bp), pCIS6 (40,285 bp), pCIS7 (53,051 bp), and pCIS8 (80,592 bp). The L. lactis UC509.9 genome is predicted to contain 2,066 proteinencoding genes, of which 168 are pseudogenes. Forty-three of these 168 pseudogenes are identical to those found in L. lactis subsp. cremoris SK11 (GenBank accession no. CP000425.1). The genome of L. lactis UC509.9 contains 104 transposase-encoding genes involving a total of 106,746 bp, including 42 copies of IS182 and 29 copies of IS981. The combination of the smallest lactococcal chromosome identified so far and the high number of transposons and pseudogenes suggests that the genome has undergone significant genome decay while adapting to the nutrient-rich dairy environment. A region of approximately 11 kb in size not present in other L. lactis genomes appears to be an integrated plasmid that includes the restriction-modification system ScrFII (16). The L. lactis UC509.9 plasmid complement encodes various traits for adaptation to the dairy environment, such as lactose and casein metabolism.

Nucleotide sequence accession numbers. The complete chromosome and plasmid complement of *L. lactis* subsp. *cremoris* UC509.9 were deposited in GenBank under accession no. CP003157 (chromosome), CP003165 (pCIS1), CP003164 (pCIS2), CP003163 (pCIS3), CP003162 (pCIS4), CP003161 (pCIS5), CP003160 (pCIS6), CP003159 (pCIS7), and CP003158 (pCIS8).

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