







## Complete Genome Sequences of Two Escherichia Phages Isolated from Wastewater in Finland

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ABSTRACT Escherichia phages vB\_EcoM-fFiEco06 and vB\_EcoM-fHoEco02 were found to have 167,076-bp and 167,064-bp genomes, respectively. They are members of genus T4virus, and they are 99.96% identical to each other. The host ranges of the phages are different, probably due to a few differences in their tail protein amino acid sequences.

hage vB\_EcoM-fFiEco06 (fFiEco06) was isolated from a mixture of municipal wastewater samples collected from several towns in Finland using clinical Escherichia coli strain 123738 (obtained from HUSLAB, Helsinki, Finland) as a host. Phage vB\_EcoMfHoEco02 (fHoEco02) was isolated from a hospital wastewater sample collected in Helsinki, Finland, using the same host. Both phages form clear ~2-mm-diameter plaques and show the morphology of a typical member of the Myoviridae when examined with transmission electron microscopy. The phages had rather narrow host ranges. Out of 200 clinical E. coli strains tested, fFiEco06 infected 9 strains and fHoEco02 infected 17 strains, indicating that even though the two phages were isolated using the same host, their host specificities are slightly different.

Phage DNA was isolated with the Invisorb Spin virus DNA minikit (Stratec Biomedical). Next-generation sequencing was performed at the Institute for Molecular Medicine Finland (FIMM) Technology Centre Sequencing Unit. DNA libraries were constructed with the Nextera DNA sample preparation kit, and paired-end sequencing was done using a MiSeq PE300 sequencer (Illumina, San Diego, CA, USA) with a 300-nucleotide (nt) read length. Reads were assembled with the A5-miseq integrated pipeline for de novo assembly of microbial genomes (1). The pipeline yielded 167,076-bp and 167,064-bp contigs for fFiEco06 and fHoEco02, respectively, with 271and 145-fold median coverage depths. The G+C content of both genomes was 35.3%.

Phage genomes were aligned pairwise to each other and to the enterobacterial phage T4 sequence (GenBank accession number NC\_000866), the type virus of genus Tavirus in viral family Myoviridae and subfamily Tevenvirinae, with EMBOSS Stretcher (2). fFiEco06 and fHoEco02 were 99.96% identical to each other, and they showed 82.17% and 82.32% identity to T4, respectively. The two phages can thus be considered representatives of the same species and as members of T4virus. The genomes were annotated using Rapid Annotations using Subsystems Technology (RAST) (3-5), BLASTP (6), and tRNAscan-SE version 2.0 (7). fFiEco06 and fHoEco02 had 274 and 272 proteincoding genes, respectively, and both phages had 10 tRNA genes. The difference in the number of protein-coding sequences (CDSs) between the two phages results from small indels in the fHoEco02 genome that cause premature stop of CDSs corresponding to fFiEco06 FE6\_028 and FE6\_212, coding for small outer capsid (Soc) protein and a hypothetical protein, respectively. In addition to these differences, there is a 9-bp

Received 3 April 2018 Accepted 29 April 2018 **Published** 31 May 2018

Citation Kiljunen S, Wicklund A, Skurnik M. 2018. Complete genome seguences of two Escherichia phages isolated from wastewater in Finland. Genome Announc 6:e00401-18. https://doi.org/10.1128/genomeA.00401-18.

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deletion in the fHoEco02 genome just before the start codon of HE2\_107, coding for the rl.1 conserved hypothetical protein, which may influence its translation.

Even though the sequences of fFiEco06 and fHoEco02 were almost identical, the host specificities of the two phages were different. Therefore, we made pairwise comparisons of the amino acid sequences of their tail proteins with EMBOSS Stretcher. There were 1- to 2-amino-acid differences between CDSs coding for tail sheath monomers (FE6\_175 and HE2\_174), long tail fiber proximal subunits (FE6\_253 and HE2\_251), long tail fiber distal subunits (FE6\_256 and HE2\_254), and tail fiber adhesins (FE6\_257 and HE2\_255), which probably explain the different host ranges of the phages.

Accession number(s). The genomic sequences of vB\_EcoM-fFiEco06 and vB\_EcoMfHoEco02 have been deposited in GenBank under the accession numbers MG781190 and MG781191, respectively.

## **ACKNOWLEDGMENTS**

Helsinki University Hospital special state subsidy for health science research grants and the Jane & Aatos Erkko Foundation are acknowledged for funding (to M.S.).

The funders had no role in the study design, data collection and interpretation, or the decision to submit the work for publication.

Pentti Kuusela and Anu Kantele are acknowledged for the clinical E. coli strains and Sheetal Patpatia for laboratory assistance.

S.K., A.W., and M.S. designed the work, A.W. performed the experiments, S.K. analyzed the data, and S.K. and M.S. wrote the paper.

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