The Complete Genome Sequence of *Escherichia coli* phage Eco_BIFF

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Abstract

*Escherichia coli* phage Eco_BIFF was isolated from several laboratory stocks of *E. coli* K-12 MG1655 derivatives. The source of the contamination is unknown. Eco_BIFF is a lytic phage that shows effective growth inhibition of *E. coli* K-12. Here, we announce the complete genome sequence of Eco_BIFF, and major findings from its genome annotation.
**Genome Announcement**

*Escherichia coli* K-12 MG1655 is a common laboratory strain used in molecular biology research both as a tool and as a model organism (Blattner et al., 1997). Eco_BIFF was isolated from strain AMD536, a previously described derivative of *Escherichia coli* K-12 strain MG1655 (Cooper et al., 2018). All AMD536 strains infected with Eco_BIFF contained two plasmids: a pBAD30 (Guzman et al., 1995) derivative and a pPRO18 (Lee and Keasling, 2005) derivative. Phage infection was suspected when cells grown in LB at 37°C with aeration lysed in early exponential phase.

*Escherichia* phage Eco_BIFF DNA was isolated from cell lysates of contaminated bacterial cultures using phenol:chloroform:isoamyl alcohol / Sodium dodecyl sulfate (PCI/SDS) DNA extraction (Phagehunting Protocols, [http://phagesdb.org/phagehunters/](http://phagesdb.org/phagehunters/)). A library for DNA sequencing was prepared using the Nextera kit (Illumina). Whole genome sequencing was performed using an Illumina MiSeq Instrument (Wadsworth Center Applied Genomic Technologies Core).

Data generated from the genomic library yielded 262,824 paired-end reads with a read length of 2 x 251 bp. To exclude bacteria sequences, reads were first mapped to the *E. coli* K-12 MG1655 genome reference sequence (U00096.3) using CLC Genomics Workbench. All remaining sequence reads were assembled into a single 49,372-bp contig corresponding to the whole genome of Eco_BIFF. Annotation and comparative analysis were performed using PHASTER (Arndt et al., 2016; Zhou et al., 2011). Rfam was used to search for non-coding RNA genes, structured cis-regulatory elements, and self-splicing RNAs (Kalvari et al., 2018; Nawrocki et al., 2015). The
Eco_BIFF genome has (i) a GC content of 45.26%, (ii) 76 predicted coding regions, and (iii) no predicted non-coding RNA genes, structured cis-regulatory elements or self-splicing RNAs.

The *Escherichia* phage Eco_BIFF genome contains functional genes related to phage architecture and packaging machinery (head protein, portal protein, terminase protein), tail structure for host interaction (tail proteins, tail assembly protein, tail fiber proteins, tail tape measure protein), phage DNA synthesis (ATP-dependent helicase, DNA primase, methylases), and host lysis and degradation (endolysin, holin, exodeoxyribonuclease, polynucleotide kinase/phosphatase, HNH endonucleases and recombination protein). The 23 predicted genes with an assigned function, which are all homologues of genes from related phages, are scattered over the genome and are interspaced with 53 hypothetical proteins, 52 of which are homologous to proteins predicted in related bacteriophage. One hypothetical protein, however, does not share homology to related phages and, therefore, may be an annotation artifact. The most closely related phages to *Escherichia* phage Eco_BIFF include: *Escherichia* phage vB_EcoS_SH2 (KY985004.1), *Escherichia* phage ADB-2 (JX912252.1, Bhensdadia et al., 2013), *Escherichia* phage JMPW2 (KU194205.1), *Shigella* phage SH6 (KX828710.1), and *Enterobacteria* phage T1 (AY216660.1, Roberts et al., 2004). These related phages share >94% nucleotide sequence identity with >86% query coverage, and BLAST E-values of 0 (Search parameters: blastn program, Word Size: 28, expected value: 10, Hitlist size: 100, Match/Mismatch scores: 1, -2, Gapcosts: 0, 2.5, Low Complexity Filter, Filter string: L;m;; Zhang et al., 2000).
Nucleotide Sequence Accession Number. The complete sequence of phage Eco_BIFF can be accessed under the GeneBank accession number MH285980.
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References


