Complete Genome Sequences of Three *Campylobacter jejuni* Phage-Propagating Strains

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ABSTRACT  Bacteriophage therapy can potentially reduce *Campylobacter jejuni* numbers in livestock, but it requires a detailed understanding of phage-host interactions. *C. jejuni* strains readily infected by certain phages are designated as phage-propagating strains. Here, we report the complete genome sequences of three such strains, NCTC 12660, NCTC 12661, and NCTC 12664.

*Campylobacter jejuni* causes diarrheal disease worldwide, and *C. jejuni* infections arise from consuming and mishandling contaminated poultry (1–3). Phages are being explored as antibiotic alternatives to reduce this burden (4–6). Phages are highly strain specific, so understanding the factors that contribute to this specificity, including capsular polysaccharides (CPSs), flagella (7), and restriction/modification systems (8, 9), can maximize the strain range targeted (10).

*C. jejuni* strains were historically tracked based on phage susceptibility (11, 12). For these typing schemes, each phage was designated a readily infected “phage-propagating” strain. To identify factors governing phage susceptibility in *C. jejuni*, we sequenced the genomes of three *C. jejuni* phage-propagating strains isolated from chickens (12), NCTC 12660, NCTC 12661, and NCTC 12664.

Whole-genome sequencing was performed using the PacBio RS and Illumina MiSeq sequencing platforms. PacBio sequence data were assembled to construct a single closed chromosomal contig for each strain. MiSeq reads were used to validate base calls and to determine the variability at each poly-G tract. Protein-, rRNA-, and tRNA-coding genes were identified as described previously (13). The genome sizes ranged from 1.61 to 1.68 Mb with an average GC content of 30.6%. The three genomes show high similarity to strain NCTC 11168, although NCTC 12660 has at least one small inversion compared to NCTC 11168. These four genomes encode a similar number of genes and pseudogenes, with the genome of NCTC 12660 slightly larger due to the presence of a genomic island. Many of the pseudogenes identified were conserved across all or most of the three strains and NCTC 11168.

We identified several differences in restriction/modification (R/M) and clustered regularly interspaced short palindromic repeats (CRISPR)/CRISPR-associated (Cas) systems between these strains. Relative to the others, NCTC 12661 lacks a type I R/M system, NCTC 12661 and NCTC 12664 lack the type IIG restriction endonuclease (RE) cj1051, NCTC 12661 uniquely encodes a type III R/M system and the type IIG RE (locus tag CJ12661_0039), and the type IV R/M system subunit mcrB is a pseudogene in NCTC 12660. Interestingly, all but NCTC 12664 encode a full type II-C CRISPR/Cas system, with cas9 a pseudogene in NCTC 12664.

CPS variability influences *C. jejuni* phage susceptibility (7, 14), but flagellar glycans play an unknown role (15). Strains NCTC 12661 and NCTC 12664 cluster separately from

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Either scenario could be explained by the many possible recombination, although sequencing or assembly issues could be responsible. The pseD genome variation, within-strain genome variation has been observed (16, 17). We compared our could lead to differences in phage-host interactions. In addition to C. jejuni strains (20). This example highlights the plasticity of C. jejuni genomes.

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REFERENCES