Complete Genome Sequences of *Paenibacillus larvae* Phages Halcyone, Heath, Scottie, and Unity from Las Vegas, Nevada

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ABSTRACT  We present the complete genome sequences of four phages that infect *Paenibacillus larvae*, the causative agent of American foulbrood disease in honeybees. The phages were isolated from beehives and beeswax products from Las Vegas, Nevada. The genomes are 50 to 55 kbp long and use the “direct terminal repeats” DNA-packaging strategy.

American foulbrood, currently the most destructive bacterial disease affecting the honeybee, *Apis mellifera*, is caused by the Gram-positive bacterium *Paenibacillus larvae* (1). There is growing interest in phages that infect and lyse *P. larvae*, as antibiotic resistance is now widespread (2). There are currently 26 complete *P. larvae* phage genome sequences in the literature and more in the process of being published (3–8). Here, we present the complete genome sequences of phages Halcyone and Heath, isolated from soil underneath healthy hives in Gilcrease Orchards in North Las Vegas, Nevada; phage Unity, isolated from material inside a beehive at the University of Nevada Las Vegas (UNLV); and phage Scottie, isolated from commercial hand cream (Burt’s Bees) purchased in the Las Vegas area.

The phages were amplified using *P. larvae* NRRL 2605, an ERIC I genotype strain, and plated on modified brain heart infusion agar with soft agar. Phage DNA was purified with phenol-chloroform extraction at the University of Nevada Las Vegas (UNLV) and sequenced at Brigham Young University with Illumina HiSeq 2500 sequencing with 250-bp paired-end reads. The genome sequences were assembled with Geneious v. 10.2.2 (Biomatters, Auckland, New Zealand) with medium-low sensitivity/fast and manually annotated using DNA Master (9) by students in the course BIOL 209X Phage Discovery at UNLV.

Each phage’s GenBank accession number, isolation source, and assembly results are shown in Table 1. All 4 phages are in the family *Siphoviridae* with linear double-stranded DNA (dsDNA) genomes and use the “direct terminal repeats” (DTR) DNA-packaging strategy (10, 11). The DTR sequence of each phage was identified using Pile-up Analysis Using Starts & Ends (PAUSE) (https://cpt.tamu.edu/computer-resources/pause/) and Geneious, looking for a sharply delimited region with double coverage depth (11). Halcyone, Heath, and Scottie have identical DTR sequences 377 bp long, whereas Unity has a different DTR sequence 378 bp long. The genomes were oriented by setting the first base of the DTR sequence to be the first base of the genome.

The genomes for Halcyone, Heath, and Scottie are 55 kbp long, which is at the maximum of the range of the *P. larvae* phage genome length (3–8), and the genome of Unity is 50 kbp long. A multiple alignment of genome sequences with ClustalW shows that Halcyone and Heath are very closely related to each other, while Scottie is
TABLE 1 GenBank accession numbers and genome assembly results for *Paenibacillus larvae* phages

<table>
<thead>
<tr>
<th>Phage name</th>
<th>GenBank accession no.</th>
<th>Isolation source</th>
<th>Genome assembly results</th>
<th>Genome length (bp)</th>
<th>Avg coverage (X)</th>
<th>GC content (%)</th>
<th>DTR sequence length (bp)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Halcyone</td>
<td>MH460827</td>
<td>Soil</td>
<td></td>
<td>55,560</td>
<td>83</td>
<td>48.6</td>
<td>377</td>
</tr>
<tr>
<td>Heath</td>
<td>MH460826</td>
<td>Soil</td>
<td></td>
<td>55,560</td>
<td>222</td>
<td>48.6</td>
<td>377</td>
</tr>
<tr>
<td>Scottie</td>
<td>MH460825</td>
<td>Hand cream</td>
<td></td>
<td>55,990</td>
<td>173</td>
<td>48.6</td>
<td>377</td>
</tr>
<tr>
<td>Unity</td>
<td>MH460824</td>
<td>Beehive</td>
<td></td>
<td>50,316</td>
<td>320</td>
<td>49.1</td>
<td>378</td>
</tr>
</tbody>
</table>

a little more distant and Unity is more distant still; the difference is largely due to a 5-kbp region missing in Unity but present in the other three phages. All four phages encode a large terminase, a portal protein, a major tail protein, two tail assembly proteins, a tail tape measure protein, and an N-acetylmuramoyl-l-alanine amidase. The tail assembly proteins appear to have a predicted translational frameshift similar to those of the G and G-T genes in phage Lambda (12, 13), located in the 3’ region of the upstream tail assembly protein (gp14). We tentatively identify the heptanucleotide slippery sequence as “TAAAAAA.” Current work is ongoing to identify more *P. larvae* phage protein functions and provide a comparative genomic analysis of *P. larvae* phages.

Data availability. The GenBank accession numbers for the four complete *Paenibacillus larvae* phage genome sequences are listed in Table 1.

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REFERENCES