



Complete Genome Sequences of Six BI Cluster *Streptomyces* Bacteriophages, HotFries, Moozy, Rainydai, RavenPuff, Scap1, and SenditCS

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ABSTRACT Six double-stranded DNA *Streptomyces* bacteriophages, HotFries, Moozy, RavenPuff, Scap1, Rainydai, and SenditCS, were isolated using the phytopathogen *Streptomyces scabiei* as a host. These phages have been identified as *Siphoviridae* and members of cluster BI by genomic analysis.

B acteriophages HotFries, Moozy, RavenPuff, Rainydai, SenditCS, and Scap1 were isolated by direct plating of processed soil samples collected around Baltimore, Maryland, and Kottayam, India (see Table 1), onto lawns of *Streptomyces scabiei* RL-34 (ATCC 49173), a phytopathogen responsible for common scab disease (1). The samples were collected and isolated by undergraduate researchers as part of the SEA-PHAGES program at the University of Maryland Baltimore County (2).

The phages were plaque purified on lawns of *S. scabiei* grown on supplemented nutrient agar (3) with Trypticase soy soft agar overlays incubated for 24 to 48 h at 30°C, and a high-titer crude lysate was harvested as described (4). Examination by transmission electron microscopy revealed the phages to be *Siphoviridae*, with icosahedral capsids with an average width of 54 nm (standard deviation [SD], \pm 5 nm) and flexible tails with an average length of 254 nm (SD, \pm 15 nm). The host range of the phages was tested by spotting diluted crude lysate on lawns of *Streptomyces* spp. (5). All six phages infected *Streptomyces mirabilis* NRRL B-2400 and *Streptomyces neyagawaensis* ISP 5588 at similar efficiencies of plating (EOP). Moozy, Rainydai, and RavenPuff also infected *Streptomyces* SC 2364 but at a reduced EOP. SenditCS, Scap1, and HotFries lysed *S. azureus* SC 2364 but did not produce infectious particles. RavenPuff demonstrated the broadest host range, infecting *Streptomyces bobili* IMRU 3310, *S. bottropensis* ISP-5262, *S. diastatochromogenes* IFO 3337, and *S. griseus* subsp. *griseus* (ATCC 10137).

Phage DNA was isolated using the Wizard DNA clean-up system (Promega). Sequencing libraries were prepared by the Pittsburgh Bacteriophage Institute from genomic DNA using an NEB Ultra II FS kit with dual-indexed barcoding. Forty-eight libraries were then pooled and run on an Illumina MiSeq instrument, yielding at least 80,000 single-end 150-base reads for each genome. These reads were then assembled using Newbler version 2.9 with default settings, and in each case, the assembled reads yielded a single phage contig which was checked for completeness, accuracy, and phage genomic termini by using Consed version 29 (6). Phage genomes averaged $2,549 \times$ coverage and were identified as linear with 9-bp 3' sticky overhangs. Genome annotation was completed using DNA Master (7).

All six bacteriophages were classified as members of cluster BI based on nucleotide conservation, genomic synteny, and phylogenetic analysis (8). HotFries, Moozy, Raven-Puff, and Scap1 are in subcluster BI2 and have an average genome length of 43,503 bp (SD, \pm 305 bp) and a GC content of 61.0% (SD, \pm 0.2%). Rainydai and SenditCS are in subcluster BI4 and have an average genome length of 56,878 bp (SD, \pm 1,153 bp) and

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	Genome	9-bp 3' overhang	% GC	No. of			GenBank
Phage name	size (bp)	sequence	content	ORFs ^a	Origin	Cluster	accession no.
HotFries	43,699	5'-CGCCGCCCT	60.9	57	Owings Mills, MD	BI2	MH155869
Moozy	43,545	5'-CGCCGCCCT	61.2	58	Olney, MD	BI2	MH155872
RavenPuff	43,709	5'-CGCCGCCCT	60.9	59	Catonsville, MD	BI2	MH155878
Scap1	43,060	5'-CGCCGCCCT	60.9	55	Kottayam, India	BI2	MF975637
Rainydai	57,623	5'-CGCCCGCCT	58.1	91	Rosedale, MD	BI4	MH155877
SenditCS	55,993	5'-CGCCCGCCT	58.2	86	Ellicott City, MD	BI4	MH155880

TABLE 1 Properties of six cluster BI Streptomyces phages

^aORFs, open reading frames.

a GC content of 58.2% (SD, \pm 0.1%). Between 55 and 59 protein-coding genes were identified in the Bl2 phages, while 86 and 91 protein-coding genes were identified in the Bl4 phages (see Table 1). No tRNA genes were identified. The BI phages displayed low average nucleotide identity (ANI) between each other (0.76 [SD, \pm 0.12]), although the subcluster ANIs ranged from 0.88 to 0.98 (9). For additional information, see the Actinobacteriophages Database (10).

Data availability. The GenBank accession numbers for the genome sequences reported here are provided in Table 1. The raw sequencing reads are available in the SRA under the accession number SRP159070.

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Members of the 2017–2018 UMBC Phage Hunters class are listed at http://phages .umbc.edu/home/class-lists/2017-18/.

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