The Bacteriophage pEp_SNUABM_08 Is a Novel Singleton Siphovirus with High Host Specificity for Erwinia pyrifoliae

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Abstract: Species belonging to the genus Erwinia are predominantly plant pathogens. A number of bacteriophages capable of infecting Erwinia have been used for the control of plant diseases such as fire blight. Public repositories provide the complete genome information for such phages, which includes genomes ranging from 30 kb to 350 kb in size. However, limited information is available regarding bacteriophages belonging to the family Siphoviridae. A novel lytic siphophage, pEp_SNUABM_08, which specifically infects Erwinia pyrifoliae, was isolated from the soil of an affected apple orchard in South Korea. A comprehensive genome analysis was performed using the Erwinia-infecting siphophage. The whole genome of pEp_SNUABM_08 comprised 62,784 bp (GC content, 57.24%) with 79 open reading frames. The genomic characteristics confirmed that pEp_SNUABM_08 is a singleton lytic bacteriophage belonging to the family Siphoviridae, and no closely related phages have been reported thus far. Our study not only characterized a unique phage, but also provides insight into the genetic diversity of Erwinia bacteriophages.

Keywords: singleton; bacteriophage; Siphoviridae; Erwinia pyrifoliae; Erwinia amylovora

1. Introduction

Erwinia pyrifoliae, first isolated in the late 1990s, is regarded as the sole causative agent for blight disease in rosaceous plants in South Korea [1]. It is considered an endemic pathogen in east Asia and has been reported to cause symptoms that are indistinguishable from those observed in the fire blight disease caused by E. amylovora [2]. Owing to the high similarities between these species, comparative studies of the two Erwinia species continue to be published, even several decades after the first identification of E. pyrifoliae [3–6]. South Korea recently witnessed a fire blight outbreak, and the disease control protocol comprised detection, differentiation, and confirmation of pathogens, and eradication of E.-amylovora–infected sites [7,8]. In a few cases, both pathogenic species were detected at one site; however, the molecular techniques used for differentiation between the two pathogens are considered to be time-consuming, which poses challenges in rapid identification [7].

Bacteriophages (phages) infect their host (bacteria) via recognition of specific receptors [9]. The host specificity of phages can be utilized as a biotechnological tool, particularly for therapeutic and diagnostic purposes [10–13]. Several Erwinia phages have exhibited a disease (fire blight) control potency that is comparable to that of antibiotics [14–16]. In addition to therapeutic applications, phages have also been used for diagnosis [17–19].
In a representative trial, a broad-host-range Erwinia phage showed diagnostic potential by exhibiting higher specificity to the pathogen than that demonstrated by a commercial diagnostic strip [20]. A better understanding of phage–host interactions could help to provide valuable insights into disease control and diagnostic strategies.

The complete genomic analysis of phages is a fundamental step for understanding host–phage interactions [21,22]. A number of phages that infect Erwinia have been characterized and sequenced thus far. However, the majority of these phages target E. amylovora, and limited information is available on phages that infect E. pyrifoliae [23–29]. Moreover, the majority of Erwinia phages that have been studied belong to the family Myoviridae [30]. Thus, it is important to focus on the isolation and genomic characterization of Erwinia phages belonging to other families for wider biotechnological applications. In this study, we characterized a highly host-specific Erwinia phage (designated pEp_SNU-ABM_08) isolated from soil samples.

2. Material and Methods

2.1. Phage Isolation

Phage isolation was performed as described previously, with minor modifications [23]. To isolate E. pyrifoliae-infecting phages, 48 samples (31 water samples and 17 soil samples) were obtained from an area located near the region where the blight outbreak was reported in South Korea. The overnight-grown (~18 h) host culture suspension was mixed with the samples for enrichment of phages and cultured for 24 h at 27 °C. A 10-fold serial dilution of enrichment culture was spotted (10 μL) onto a lawn of bacterial culture. Samples positive for lysis zones or plaque(s) were collected to confirm their plaque-forming capacity. The samples were centrifuged, syringe-filtered (0.45 μm), and subjected to a conventional double-layer agar (DLA) assay [23]. The plaques were subcultured five times to obtain a single phage lysate.

2.2. Phage Propagation and Purification

Phage propagation and purification were performed using the DLA assay in accordance with a previously described protocol with minor modifications [23]. The 18 h cultured top agar was collected using SM buffer (100 mM NaCl, 50 mM Tris (pH 7.5), and 10 mM MgSO₄) and centrifuged to isolate phages from the agar matrix, after which it was filtered through a 0.45 μm membrane filter. Precipitation using ammonium sulfate was performed for the purification of phages. The phage lysate was mixed with 2% (v/v) Tween-80, and 35% (w/v) ammonium sulfate, and was incubated for 15 min at 4 °C. The pellicle was resuspended in SM buffer and purified using CsCl gradient ultracentrifugation [23]. The high titer of phage suspension (>10^9 plaque-forming units (PFUs)/mL) dialyzed using 7000 MWCO membrane was stored at 4 °C.

2.3. Transmission Electron Microscopy

The phage suspension (10 μL) was spotted on a copper grid and incubated for 1 min. Excess solution was removed, followed by the conduction of staining with uranyl acetate (2%) for 30 s. The dried grid was examined using the Talos L 120C (FEI, OR, USA) transmission electron microscope (120 kV). Dimensions of the phage virions were analyzed by measuring three independent particles.

2.4. Host Range Analysis

A total of 55 strains including Erwinia spp. (26 E. amylovora strains and 25 E. pyrifoliae strains) isolated from the blight-affected plant tissue were examined for host range analysis using the spot method [23]. The presence of inhibition zone or plaque(s) on the spotted area was considered to denote susceptibility; infectivity was represented as + and −for susceptible and nonsusceptible cases, respectively. The strains that showed an inhibition zone were further assayed using the double-layer method to exclude abortive infection,
using 10-fold serial dilutions of the phage suspension. Genotyping of the Erwinia strains used in the present study was performed using ERIC primers: ERIC_F-3'-ATGTAAA-GCTCCTGGGATTCAC-5', ERIC_R-3'-AAAGTAAGTACTGGGTACCG-5'. PCR reactions were carried out with 500 ng of genomic DNA of the strains and consisted of a denaturation at 95 °C for 3 min, followed by 35 cycles at 94 °C for 1 min, 50 °C for 1 min, and 72 °C for 5 min, and a final extension at 72 °C for 10 min.

2.5. Adsorption and One-Step Growth Curve

The adsorption [31] and one-step growth [32] assays were conducted as described previously. The host strain (exponential phase; ~10⁶ CFU/mL) was mixed with the phage at a multiplicity of infection (MOI) of 0.001 and cultured at 27 °C. For the adsorption assay, samples were collected at 1, 3, 5, 10, 20, 30, 40, 50, and 60 min after incubation. The PFU was measured using the supernatant obtained after centrifugation for the enumeration of unadsorbed phages. For the one-step growth assay, after phage adsorption for 50 min (>95% adsorption rate), unadsorbed phages were discarded after centrifugation. Then, the pellet was resuspended with prewarmed broth and samples were acquired every 20 min for a timeframe of 180 min to analyze the latent period and burst size.

2.6. DNA Extraction and Sequencing

Phage DNA isolation was performed using the phenol–chloroform method [23]. Briefly, 1 mL of the phage suspension (>10¹⁰ PFU/mL) was mixed with 10 U of DNase I and RNase A and incubated for 1 h at 37 °C. After the addition of 0.5 M of ethylenediaminetetraacetic acid for the inhibition of enzymes, the suspension was subjected to treatment with proteinase K for 3 h at 56 °C. Next, DNA was isolated using a phenol:chloroform:isoamylalcohol (25:24:1) solution and precipitated with ethanol. Sequencing and assembly of the isolated DNA were performed using the Illumina HiSeq system (Illumina, San Diego, CA, USA) at Macrogen (Seoul, South Korea).

2.7. Genome Analysis

Open reading frame (ORF) identification was performed using GeneMarkS and the Rapid Annotation using Subsystem Technology (RAST) server [33,34]. The putative function of each ORF was predicted and annotated using protein BLAST. The tRNA, antibiotic resistance-related genes, and virulence-related genes were identified using tRNAscan-SE (v2.0), ResFinder (v3.2), and the VirulenceFinder (v2.0) server, respectively [35–37]. A total of 72 complete genome sequences of Erwinia phage spp. available in the NCBI nucleotide database were included for comparative genome analysis. Comparative genome analysis was performed using Coregene3.5 (blastp threshold: 75), Gepard (word size: 10), and Easyfig (E-value cut off: 0.001) software [38–40]. The phylogenetic tree was generated using Mega X with maximum likelihood method (1000 bootstrap) Virus Classification and Tree Building Online Resource (VICTOR) with the recommended settings for prokaryotic viruses [41,42].

2.8. Proteome Analysis

Mass spectrometric analysis of the structural proteins of phage pEp_SNUABM_08 was performed as described [23]. The phage proteins prepared by implementing five rounds of freeze–thaw cycles were separated using 12% SDS-PAGE. Subsequently, the proteins spanning the entire lane were in-gel digested with trypsin (100 ng/µL). The resultant proteins derived from pEp_SNUABM_08 were identified by using a nano-high-resolution LC-MS/MS spectrometer Q Exactive Hybrid Quadrupole-Orbitrap (Thermo Scientific, MA, USA) equipped with the Dionex U 3000 RSLC nano HPLC system.
3. Results

3.1. Biological Analysis of pEp_SNUABM_08

The *Erwinia* phage pEp_SNUABM_08 was isolated from the soil of a blight-affected apple orchard (located on Jecheon, Chungcheongbuk province) using *E. pyrifoliae* (KACC13945) as an indicator strain. Morphological analysis revealed that the phage belonged to the Siphovirus family. The phage pEp_SNUABM_08 possessed an icosahedral head (62 ± 4 nm in diameter) and a long noncontractile tail (190 ± 12 nm in length) (Figure 1).

![Transmission electron microscopy result showing the structure of *Erwinia pyrifoliae* phage pEp_SNUABM_08. Scale bar: 100 nm.](image)

The host range assay was performed against 51 *Erwinia* strains, including *E. amylovora* and *E. pyrifoliae*, isolated from blight-affected plant tissues around South Korea in 1999, 2019, and 2020, and four other strains belonging to three bacterial species (Table 1). The phage pEp_SNUABM_08 showed highly specific infectivity. Of 26 *E. amylovora* strains, phage pEp_SNUABM_08 could infect only two strains, YKB 14748 and RA0030, isolated from Chungcheongbuk and Gyeonggi provinces. Of 25 *E. pyrifoliae* strains, the phage pEp_SNUABM_08 mainly infected strains isolated from Gangwon province (11/15 strains) having different clonal lineages, and 1 strain isolated from Gyeongsangbuk province, which showed a geographical spread (Figure S1). Lysis from without was not observed and plaques were obtained for all susceptible strains in case of low-concentration phage suspension(s).

**Table 1.** Host range of *Erwinia* phage pEp_SNUABM_08.

| Species              | Strain  | Year | Province          | Infectivity |
|----------------------|---------|------|-------------------|-------------|
| *Erwinia amylovora*  | YKB 14715 | 2019 | Chungcheongbuk    | −           |
|                      | YKB 14740 | 2019 | Chungcheongbuk    | −           |
|                      | YKB 14742 | 2019 | Chungcheongbuk    | −           |
|                      | YKB 14748 | 2019 | Chungcheongbuk    | +           |
|                      | YKB 14750 | 2019 | Chungcheongbuk    | −           |
|                      | RA0030    | 2020 | Gyeonggi          | +           |
|                      | RA0031    | 2020 | Gyeonggi          | −           |
|                      | RA0032    | 2020 | Gyeonggi          | −           |
|                      | RA0033    | 2020 | Gyeonggi          | −           |
The phage pEp_SNUABM_08 showed a low adsorption rate, which reached 70%, 90%, and 95% at 10, 30, and 40 min, respectively (Figure 2A). The first burst of phages commenced at 40 min (latent period) after inoculation with the burst size of 20 PFU/cell and the second burst was observed at 100 min after inoculation (Figure 2B).
3.2. Protein Analysis of pEp_SNUABM_08

Upon analyzing the profiles of structural proteins of pEp_SNUABM_08, we identified 12 structural proteins including 10 generally identified structural proteins including head-to-tail joining protein (gp12), portal protein (gp13), major capsid protein (gp16), tape measure protein (gp24), and two distinct proteins (Ig-like domain-containing protein; gp21, and head decoration protein; gp15), which were also predicted in the genome analysis (Table 2).

Table 2. Profiles of LC-MS/MS-detected structural proteins ofErwinia phage pEp_SNUABM_08.

| Protein | Putative Function                        | Molecular Weight (kDa) | Number of Identified Peptides | Coverage (%) |
|---------|------------------------------------------|------------------------|-----------------------------|--------------|
| gp12    | Head-to-tail joining protein             | 9.6                    | 11                          | 28           |
| gp13    | Portal protein                           | 61.2                   | 100                         | 25           |
| gp15    | Head decoration protein                  | 14.2                   | 2                           | 19           |
| gp16    | Major capsid protein                     | 39.9                   | 90                          | 36           |
| gp21    | Ig-like domain-containing protein        | 41.9                   | 58                          | 11           |
| gp22    | Putative tail assembly chaperone         | 17.3                   | 127                         | 40           |
| gp24    | Tape measure protein                     | 148.4                  | 528                         | 48           |
| gp25    | Distal tail protein                      | 64.7                   | 258                         | 21           |
| gp29    | Putative tail protein                    | 139.1                  | 218                         | 25           |
| gp30    | Putative tail fiber protein              | 23.7                   | 5                           | 10           |
| gp38    | Virion protein                           | 25.2                   | 106                         | 38           |
| gp51    | Tail fiber protein                       | 9.2                    | 10                          | 47           |

3.3. Genomic Analysis of pEp_SNUABM_08

pEp_SNUABM_08 harbors a 62,715 bp dsDNA genome (GC content, 57.24%). Seventy-nine ORFs were detected in the genome; however, tRNA genes were not detected. Among the predicted ORFs, 48 were located on the positive strand and 31 were located on the negative strand (Figure 3). The predicted ORFs were categorized into the following four groups based on their function: nucleotide regulation (e.g., Cas4-like protein, putative exonuclease), structure and packaging (e.g., putative tail fiber protein, terminase small/large subunit), lysis (e.g., lysis protein A like protein, Rz like protein), and hypothetical proteins. The majority of the ORFs were found to be for hypothetical proteins (57 ORFs; 72.15%); lysogeny-, antibiotic-resistance-, and bacterial-virulence-related genes were not detected (Table S1).
Figure 3. Complete genome map of pEp_SNUABM_08. The ORFs were categorized and color-coded based on their function; structure and packaging, blue; nucleotide regulation, yellow; lysis, red; hypothetical protein, gray. Scale = base pair.

The predicted genes showed best matches with those of phages classified under the genus Chivirus with a relatively low identity (23–76%). The genome organization of pEp_SNUABM_08 was slightly similar to that of the well-known lytic bacteriophage Chi; however, a strong homology could not be observed for the last section of the genome (~37 kb; Figure S2). As expected, the complete nucleotide sequence of pEp_SNUABM_08 did not form a cluster with phages of the family Siphoviridae, which infect bacterial strains of class Gammaproteobacteria (Burkholderia, Erwinia, Escherichia, Klebsiella, Pantoea, Pseudomonas, and Salmonella), as shown in the phylogenetic tree (Figure 4), and did not exhibit the generation of a strong line in the dot plot analysis (Figure 5).
Figure 4. Whole-genome phylogenetic analysis of Siphoviridae phages infecting members of the class Gammaproteobacteria. The pEP_SNUABM_08 phage isolated in the present study is highlighted using a black arrow (▶). The genera (Chivirus, magenta; unclassified, mosaic; Casadabanvirus, cyan; Abidjanvirus, light green; Nickievirus, brown; Serutavirus, dark green; Vidquintavirus, orange; Sashavirus, turquoise; Kilunavirus, crimson; Septimavirus, sky blue; Kagunavirus, bisque; Jershevirus, violet; Dhillonvirus, kakhi; Murrayvirus, petal; Stanboltvirus, gray; Detrovirus, white; Hendrixvirus, chartreuse; Lambdavirus, black; Feofaniavirus, chrome yellow) and host (Burkholderia, red; Erwinia, orange; Escherichia, yellow; Klebsiella, green; Pantoea, blue; Pseudomonas, dark blue; Salmonella, purple) are indicated using respective colors.
A core gene analysis was performed to conduct a more detailed comparation. The replication strategy of Erwinia phage pEp_SNUABM_08 was similar to that of Chi, as the major component phage proteins (41/79; 51.9%) encoding structural, packaging, nucleotide regulation, or lysis proteins were conserved. In contrast, the hypothetical proteins encoded by the last section of their genomes were unique to each phage. The proteins are considered to endow pEp_SNUABM_08 with the singleton status. Moreover, phylogenetic analysis among Erwinia phages showed the existence of a sole cluster of pEp_SNUABM_08 (Figure 6). Only one core protein was shared among Siphoviridae phages infecting
Erwinia spp. (pEp_SNUABM_08, 49, 59, Midgardsorm38), namely the tape measure protein.

### Family
- **M**. **Myoviridae**
- **P**. **Podoviridae**
- **A**. **A cidoviridae**
- **C**. **C asiciviridae**
- **A**. **Autographiviridae**

### Genus
- **M**. **Machanivirinae**
- **E**. **Erissovirinae**
- **W**. **Wellingtovirinae**
- **D**. **Derbiocovirinae**
- **A**. **Agencovirinae**
- **J**. **Jepetovirinae**
- **R**. **Rasinsivirinae**
- **J**. **Johncovirinae**
- **K**. **Kvinsivirinae**
- **F**. **Fussetovirinae**
- **C**. **Cohnicovirinae**
- **E**. **Erocentovirinae**
- **R**. **Rovinovirinae**
- **S**. **Sassovirinae**
- **Y**. **Yokoshimavirinae**
- **E**. **Elnavirinae**
- **D**. **Deriavirinae**
- **K**. **Kvingsavirinae**
- **E**. **Erocentovirinae**
- **U**. **Unclassified**

### Host
- **E.** **sauvagae**
- **E.** **tasseolitella**
- **E.** **tamarisci**
- **E.** **persicina**
- **E.** **horticola**
- **E.** **pyrifolia**

**Figure 6.** Phylogenetic analysis of Erwinia phages conducted using the VICTOR software. The pEp_SNUABM_08 phage isolated in the present study is highlighted using a black arrow (▶). The numbers above the branches represent Genome-BLAST Distance Phylogeny (GBDP) pseudo-bootstrap support values based on the conduction of 100 replications.
The structure of the tail tip protein (gp29) of pEp_SNUABM_08 comprised an N-terminal baseplate binding domain and a C-terminal oligosaccharide binding domain (Figure 7A). The N-terminal domain showed homology with phage-originated domains such as T5 baseplate hub (Q6QGE9), Lambda Tip attachment protein J (P03749), and phage-tail 3 (PF13550.7). However, the C-terminal of gp29 matched with the binding domain of the gene transfer agent of Rhodobacter capsulatus (6TEH). Additionally, gp29 of pEp_SNUABM_08 did not show clustering with other closely related phages, as shown in Figure 7B.

**Figure 7.** Schematic representation of the tail tip protein (gp29) of pEp_SNUABM_08 (A). The N-terminal (baseplate binding domain), and C-terminal (oligosaccharide binding domain) are colored in blue and red, respectively. Phylogeny analysis of the tail tip protein of pEp_SNUABM_08 (▶) and related phages (B).

### 4. Discussion

The first step in this study was the isolation of phages with different host ranges and analysis of their genome sequences. In this study, we isolated the *Erwinia* phage pEp_SNUABM_08. Although most reported *Erwinia* phages exhibit morphological characteristics similar to those of myoviruses or podoviruses, pEp_SNUABM_08 is a distinct siphovirus-shaped virion. It exhibited high host specificity against the *E. pyrifoliae* strains (Table 1) compared to the myophages or podophages that were isolated from South Korea. Nevertheless, pEp_SNUABM_08 may possess biocontrol and diagnostic potential for black shoot disease, which has been previously reported in Gangwon province, South Korea [23,43,44].

The host specificity of pEp_SNUABM_08 may be associated with its atypical tail protein, which is responsible for host recognition [45,46]. In particular, it has been elucidated that the host receptor-binding protein (RBP) is located at the tip of the tail fiber protein, which is located in the C-terminal end of the protein [47,48]. The arrangement of tail fiber genes in the lytic phage pEp_SNUABM_08 is similar to that observed in phage lambda, as they harbor the lambda-like tail fiber domains, namely tail completion protein Z (gp19), tail tube terminator protein U (gp20), tail tip protein M (gp25), tail tip protein L (gp26), tail tip assembly protein I (gp27), and tail tip attachment protein J (gp29) [49]. The tail tip attachment protein J is considered to play a major role in host recognition, and the tail tip
protein of pEp_SNUABM_08 (gp29) also comprises the N-terminal baseplate binding domain, C-terminal RBP, and oligosaccharide binding domain (Figure 7A). However, the C-terminal domain of pEp_SNUABM_08 did not exhibit strong homology with any of the phage-originated RBPs, and gp29 was solely clustered among closely related phage tail proteins (Figure 7B). This single clustering characteristic may explain the host specificity of pEp_SNUABM_08.

Many years have passed after its initial isolation, and phages that are closely related to phage pEp_SNUABM_08 have not been reported. In addition, the majority (72%) of its genome encoded hypothetical proteins; many proteins did not exhibit homology with other proteins available in the database. The distinct biological and genomic characteristics of *Erwinia* pEp_SNUABM_08 are considered to originate from its singleton status. Further studies focusing on structural and functional characterization of this virus may provide insights into novel phage–host interaction mechanisms, especially for *Erwinia* phages.

**Supplementary Materials:** The following are available online at www.mdpi.com/article/10.3390/v13071231/s1, Table S1: The profile of ORFs in *Erwinia* phage pEp_SNUABM_08; Figure S1: ERIC-PCR genotyping of *Erwinia* spp. used in this study for host range analysis; Figure S2: Comparative whole genome analysis of *Erwinia* phage pEp_SNUABM_08 and *Salmonella* phage Chi.

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