Abstract: *Vibrio mediterranei* 117-T6 is extensively pathogenic to several *Pyropia* species, leading to the death of conchocelis. In this study, the first *V. mediterranei* phage (named *Vibrio* phage Yong-XC31, abbreviated as Yong-XC31) was isolated. Yong-XC31 is a giant phage containing an icosahedral head about 113 nm in diameter and a contractible tail about 219 nm in length. The latent period of Yong-XC31 is 30 min, and burst size is 64,227. Adsorption rate of Yong-XC31 to *V. mediterranei* can reach 93.8% in 2 min. The phage genome consisted of a linear, double-stranded 290,532 bp DNA molecule with a G + C content of 45.87%. Bioinformatic analyses predicted 318 open reading frames (ORFs), 80 of which had no similarity to protein sequences in current (26 January 2021) public databases. Yong-XC31 shared the highest pair-wise average nucleotide identity (ANI) value of 58.65% (above the ≥95% boundary to define a species) and the highest nucleotide sequence similarity of 11.71% (below the >50% boundary to define a genus) with the closest related phage. In the proteomic tree based on genome-wide sequence similarities, Yong-XC31 and three unclassified giant phages clustered in a monophyletic clade independently between the family Drexlerviridae and Herelleviridae. Results demonstrated Yong-XC31 as a new evolutionary lineage of phage. We propose a new phage family in Caudovirales order. This study provides new insights and fundamental data for the study and application of giant phages.

Keywords: *Vibrio mediterranei*; giant phage; complete genome

1. Introduction

Vibrios are ubiquitous in marine ecosystems living as well-described pathogens of aquatic fauna, for example, *Vibrio anguillarum*, *Vibrio harveyi*, *Vibrio coralliilyticus* and *Vibrio aestuarianus* [1–4]. *Vibrio mediterranei* is a potential emerging pathogen of marine animals such as corals and scallops [5,6]. *V. mediterranei* 117-T6 (CGMCC1.16311) was isolated from the bleached shell-born conchocelis of *Pyropia yezoensis* and was pathogenic to the conchocelis of several *Pyropia* species, including *P. yezoensis* [7,8]. *Pyropia* culture has a long history in China, but there are no effective prevention and control measures for bacterial diseases, which seriously affect the economic income of farmers and the healthy and sustainable development of *Pyropia* industry [8,9]. Global warming has caused an increase in sea surface temperature that has undoubtedly led to the unseasonal outbreaks of Vibrios as well as their increased abundance and virulence in marine environments and aquaculture [10,11]. Therefore, there are a growing need for effective methods for managing bacterial infections.
Viruses, in particular bacteriophages are the most abundant biological entities in the oceans, showing an extremely high, uncharted diversity [12]. The potential of viruses as therapeutic agents to treat infectious diseases has been known for a long time, and they are considered to be important agents and resources as a solution to antibiotic resistance [13]. The phages with double-stranded (ds) DNA genomes larger than 200 kbp are defined as giant or “jumbo” bacteriophages [14]. Giant bacteriophages commonly contain many genes that do not exist in the small genome bacteriophages. For example, some giant phages have several paralogous genes for DNA polymerase and RNA polymerase (RNAP). Importantly, the proteins encoded by these additional genes may replace the function of the host proteins, thereby reducing the dependence of giant phages on their bacterial hosts [15]. In addition, many genes in giant phages are interpreted as coding hypothetical proteins which are not found in small phages, and their biological features are understudied [14]. These genes may be a new resource of proteins for industrial, agricultural, or medical applications in the future [16–19]. However, large phages are not commonly isolated. Limited mobility on semi-solid plates, which prevented formation of visible plaques, may be the possible reason for the rare isolation of giant phage [15,20]. A centrifugal force that is too high, in the centrifugation to remove bacteria and protists, may also be a reason.

In recent years, bacteriophages have been increasingly applied for disease prevention and control in aquaculture [21–23]. To mine potential marine application resources for disease control of Pyropia vibriosis, the first V. mediterranei phage, named Vibrio phage Yong-XC31 (abbreviated as Yong-XC31), was isolated from the coastal water of Meishan island (29°46.989 N 121°57.516 E), Ningbo, China. Characteristics and the complete genome of the phage were analyzed. 

2. Materials and Methods

2.1. Bacterial Strains and Culture Conditions

Vibrio mediterranei strain 117-T6 (China General Microbiological Culture Collection, CGMCC 1.1631) was provided by the Key Laboratory of Marine Biotechnology of Ningbo University [7,8]. V. mediterranei 117-T6 was cultured in NB seawater medium (peptone 10 g, beef extract 3 g, make final volume up to 1 L with filtered seawater, pH 7.2) at 29 °C with shaking at 180 rpm.

2.2. Antibiotic Susceptibility Test

Antibiotic susceptibility test was performed using antibiological susceptibility discs (Hangzhou Binhe Microorganism Reagent. Co., Ltd., Hangzhou, China), according to the instructions and following the standard of the Clinical & Laboratory Standards Institute (CLSI). 17 types of antibiotic disks were used as follows: penicillin G (6 µg), amoxicillin (10 µg), cephalexin (30 µg), kanamycin (30 µg), gentamicin (10 µg), tobramycin (10 µg), azithromycin (15 µg), aboren (30 µg), chloramphenicol (30 µg), ofloxacin (5 µg), tetracycline (30 µg), doxycycline (30 µg), rifampin (5 µg), trimethoprim/ sulphamethaxazole (1.25/23.75 µg), vancomycin (30 µg), polymyxin (30 µg) and clindamycin (2 µg). Fresh log phase V. mediterranei 117-T6 cultures were spread by sterial swab on Muller-Hinton agar medium, dried at room temperature (about 5 min). Antibiotic discs were then placed on the surface of the medium with triplicates. The plates were incubated at 29 °C for 16 to 18 h. The diameters of the inhibition zones formed were measured.

2.3. Phage Isolation and Morphological Observation

The surface seawater samples used for bacteriophage separation were collected from the seaside of Meishan island (29°46.989 N 121°57.516 E), Ningbo, China on 31 July 2018. The water samples were placed in an ice box and immediately brought back to the laboratory for treatment. After centrifugation at 10,000 × g for 10 min, each 80 mL supernatant was mixed with 40 mL of 3 × NB seawater medium and 2 mL V. mediterranei 117-T6 of log phase (OD600 ≈ 0.6, 1.91 × 10⁹ CFU/mL). The mixtures were cultured at 29 °C with shak-
ing speed of 180 rpm for 3–4 h to enrich the phages, and then centrifuged at 10,000×g for 10 min. The supernatants were filtered through 0.45 μm pore-size filters. Pure phage strain was obtained by three serial single-plaque isolation using the conventional double-layer agar method [24] employing V. mediterranei strain 117-T6 as the host. The bacteriophage was negatively stained with 3% uranium acetate for 20 s and observed under transmission electron microscope (Hitachi H-7650, Tokyo, Japan).

2.4. Thermal and pH Stability

Thermal stability was assessed by exposing aliquots of phage suspension (7.5 × 10^7 PFU/mL, 1 mL, in triplicates) to various temperatures (40, 50, 60 and 70 °C) for 30, 60, 90, 120, 150 and 180 min, respectively. To test the pH stability of Yong-XC31, aliquots of phage suspension (7.5 × 10^7 PFU/mL, 1 mL, in triplicates) were adjusted using NaOH or HCl to various pHs (2–12), withdrawn for 2 h at 29 °C. Titers of the treated and untreated control samples were measured using the double-layer agar plate method. The experiment was repeated three times.

2.5. MOI Selection Experiments and Adsorption Test

In multiplicities of infection (MOIs) selection experiments, 7.5 × 10^5 PFU of Yong-XC31 was mixed with a set of serial dilutions of 117-T6 cell suspensions (7.5 × 10^4 to 7.5 × 10^8 CFU) at MOIs of 0.001, 0.01, 0.1, 1 and 10, respectively, with triplicates. After 10 min of adsorption at 29 °C, the mixtures were centrifuged at 10,000×g for 10 min. The precipitates were suspended in 5 mL NB seawater medium and incubated for 3 h at 29 °C with shaking speed of 180 rpm. Titers in the supernatant of the lysates were measured by using the double-layer agar plate method. The experiment was repeated three times. The MOI with the highest phage production was considered as the optimal one.

To evaluate optimum adsorption time, the phage was mixed with 117-T6 at the optimal MOI of 0.001 (phage-to-bacterium ratio = 7.5 × 10⁵ PFU/7.5 × 10⁸ CFU) with triplicate and incubated at 29 °C with shaking speed of 180 rpm. Samples were taken at 0, 2, 4, 6, 8, 10, 15 and 20 min post-inoculation and centrifuged at 5000×g for 10 min at 4 °C. Phage titers in the supernatant were measured by using the double-layer agar plate method. The experiment was repeated three times.

The influence of temperature and pH on the adsorption of the phage was monitored. The phage was mixed with 117-T6 at the optimal MOI of 0.001 as above at 29 °C at the pH of 4, 5, 6, 7, 8 and 9, respectively, with triplicates, and then centrifuged at 5000×g for 10 min at 4 °C. Phage titers in the supernatant were measured by using the double-layer agar plate method. The experiment was repeated three times.

2.6. One-Step Growth Experiment

Phage Yong-XC31 was mixed with 117-T6 (1.36 × 10⁸ CFU/mL) at a MOI of 0.001 with triplicate and allowed to adsorb for 2 min at 29 °C. Then, centrifuged at 6000×g for 10 min. Pellets containing the infected cells were washed twice and re-suspended in 40 mL of fresh NB seawater medium, incubated at 29 °C with shaking at 220 rpm. Samples were taken at 0, 10, 20, 30, 40, 50, 60, 90, 120, 150, 180 and 210 min, respectively, centrifuged at 5000×g for 10 min at 4 °C, and then phage titers in the supernatant were immediately determined by the double-layer agar method. Titer measurements were conducted in triplicate.

2.7. Host Range Determination

The host range of Yong-XC31 was tested against 34 bacterial strains including the indicator host (Table 1) using the spot test according to the references [25,26]. Cultures of each bacterial strain (10^⁸ CFU/mL) were mixed with melted 0.7% agar (43 °C) NB seawater medium, respectively, with triplicates and poured on a 1.5% solid agar to make double layer agar plates. After solidification, 5 μL of phage suspensions (10⁶ PFU/mL) were
spotted on each plate. The plates were cultured at 29 °C for 8 h. It was considered as positive for clear lysis zones or plaques on the bacteria lawn.

Table 1. Bacterial strains used in the host range assay and infection results.

| No. Bacteria Strains | Lytic Ability | No. Bacteria Strains | Lytic Ability |
|----------------------|---------------|----------------------|---------------|
| Vibrio mediterranei 117-T6 | + | Escherichia coli DH5α | - |
| Vibrio hispanicus XXY | - | Citrobacter freundii | - |
| Vibrio fluvialis XXY1 | - | Shewanella putrefaciens | - |
| Vibrio unifascis XXY | - | Streptococcus iniae | - |
| Vibrio fluvialis XXY2 | - | Alternate pseudomonas XY | - |
| Vibrio parahaemolyticus 1A08161 | - | Shigella dysenteriae | - |
| Vibrio parahaemolyticus 1A11655 | - | Proteus vulgaris | - |
| Vibrio alginolyticus LDF | - | Proteus mirabilis | - |
| Vibrio alginolyticus XY | - | Aeromonas hydrophila | - |
| Vibrio alginolyticus WY | - | Shigella sonnei | - |
| Vibrio alginolyticus SZT | - | Pseudomonas aeruginosa LDF | - |
| Vibrio harveyi 1-5 | - | Pseudomonas aeruginosa SZT | - |
| Vibrio harveyi LDF | - | Aeromonas sobria ATCC4979 | - |
| Vibrio pacinii XY | - | Aeromonas sobria ATCC4979 | - |
| Vibrio anguillarum | - | Pseudoalteromonas isaakenskiana | - |
| Aeromonas hiraeolus XXY | - | Salmonella paratyphi B | - |
| Edwardsiella tarda SZT | - | Enterobacter cloacae | - |
| Edwardsiella tarda LW | - | Marinomonas sp.XY | - |
| Enterobacter sakazakii | - | Pseudomonas sp. XY | - |

(+) representative infection, (-) representative non-infection.

2.8. Genome Extraction and Sequencing

Genomic DNA of phage Yong-XC31 was extracted utilizing the modified method of standard phenol-chloroform extraction [27]. DNase I and RNase A (TransGen Biotech, Beijing, China) to a final concentration of 1 ug/mL were added to the purified phage Yong-XC31 stock solution. The mixture was incubated at 37 °C for 2 h to remove contaminating bacterial DNA and RNA. DNase I was deactivated by incubating the solution for 15 min at 80 °C. After adding EDTA to a final concentration of 20 mM, proteinase K at 50 ug/mL, and sodium dodecyl sulfate at 0.5% (w/v), the mixture was incubated for 1 h at 56 °C. An equal volume of phenol was added to extract the viral DNA, followed by centrifugation at 10,000 × g for 10 min. The aqueous layer was moved to a fresh tube, to which an equal volume of phenol–chloroform-isoamyl alcohol (25:24:1) was added and mixed, and then centrifuged at 10,000 × g for 10 min. The aqueous layer was mixed with isovolumic chloroform and centrifuged 10,000 × g for 5 min. The aqueous phase was added with an equal volume of isopropanol, stored at −20 °C for 3 h, and centrifuged at 4 °C at 13,000 × g for 20 min. The precipitated DNA was washed with 75% ethanol. The obtained Yong-XC31 DNA was resuspended in deionized water and stored at −20 °C for further experiments.

The NEBNext Ultra II DNA Library Prep Kit for Illumina (#E7645) was used for constructing genomic library. Genome sequencing of the phage was performed using the Illumina MiSeq (SanDiego, CA, USA) sequencing platform to obtain 2 × 300 bp paired-end reads. Low-quality (Q-value < 20) reads and adapters were filtered out using fastp. SPAdes 3.13.0 software (http://cab.spbu.ru/software/spades/) was utilized to assemble the trimmed reads. Phage genome termini were analyzed using our proposed method [28].

2.9. Genome Annotation and Taxonomic Analyses

ORFs prediction was initially conducted with RAST (http://www.rast.nmpdr.org), and then identified with HMMER and HHpred web server [29,30]. All predicted ORFs were manually verified using the BLAST tool of NCBI (https://blast.ncbi.nlm.nih.gov/). tRNAscan-SE was used to search for tRNA genes (http://lowelab.ucsc.edu/tRNAscan-SE/; [31]), and the antibiotic resistance genes and virulence factors were searched in CARD database (http://arpcard.mcmaster.ca) and VFDB database (http://www.mgc.ac.cn/VFs/main.htm), respectively.
Nucleotide sequence comparisons were firstly conducted using NCBI BLASTn [32]. The pair-wise average nucleotide identity (ANI) values between the giant phage Yong-XC31 and the phages with the 10 top highest homology (Table 2) in BLASTn comparison (e-value < 10\(^{-5}\)) were confirmed using OrthoANI [33]. The in silico DNA–DNA hybridization (isDDH) values between Yong-XC31 and these closely related giant phages were performed using GGDC (formula 2) [34]. The percentage of conserved proteins (POCP) values between Yong-XC31 and these related giant phages were calculated as described previously [35]. Genome comparison of Yong-XC31 and the closest related phage BONAISHI was done by a genome comparison visualizer, Easyfig [36]. To estimate the nucleotide sequence similarity between Yong-XC31 and other phages in current (26 January 2021) public databases, the Pairwise Sequence Comparison (PASC) classification tool [37] (https://www.ncbi.nlm.nih.gov/sutils/pasc/viridty.cgi) was used. ViPTree online [38] (available at https://www.genome.jp/viptree/) was used to generate a proteomic tree based on genome wide sequence similarities, computed by tBLASTx, gathering 56 classified phages of the nine families of Caudovirales, Yong-XC31 and 10 giant phages in Table 2.

Table 2. Average nucleotide identity (ANI), in silico DNA–DNA hybridization (isDDH) and percentage of conserved proteins (POCP) values between *Vibrio* phage Yong-XC31 and the giant phages with the 10 top highest homology in BLASTn comparison (e-value < 10\(^{-5}\)).

| Strain                | Accession no. | ANI, % | isDDH, % | POCP, % |
|-----------------------|---------------|--------|----------|---------|
| *Vibrio* phage BONAISHI | MH595538      | 58.65  | 0        | 6.774   |
| *Aeromonas* phage phiA55  | HM452126      | 0      | 0        | 0.606   |
| *Aeromonas* phage CC2    | JX123262      | 0      | 0        | 0.268   |
| *Aeromonas* phage AS-yj   | MF498774      | 0      | 0        | 0.278   |
| *Aeromonas* phage AS-szw | MF498773      | 0      | 0        | 0.273   |
| *Aeromonas* phage AS-zj   | MF448540      | 0      | 0        | 0.272   |
| *Aeromonas* phage AS-sw   | MF498775      | 0      | 0        | 0.271   |
| *Pseudomonas* phage Phabio | MF042360     | 0      | 0        | 0.26    |
| *Vibrio* phage 2.275.O._10N.286.54.E11 | MG592671 | 0      | 0        | 0.245   |
| *Vibrio* phage 2 TSL-2019     | MF063068      | 0      | 0        | 0       |

2.10. Genome Sequence Accession Number

The complete genome sequence of *Vibrio* phage Yong-XC31 is available from GenBank under nucleotide accession number MK308674. The phage has been deposited in China General Microbiological Culture Collection Center under number CGMCC No. 17098.

3. Results

3.1. Antibiotic Susceptibility of *V. mediterranei* 117-T6

*V. mediterranei* 117-T6 was resistant to 4 of 17 tested antibiotics, which were penicillin G, aboren, polymyxin and clindamycin, respectively (Figure 1). It was intermediate to azithromycin, vancomycin and gentamicin. Though, causing inhibition zones on *V. mediterranei* 117-T6 lawn, amoxicillin, rifampicin, tobramycin, tetracycline, doxycycline, chloramphenicol, ofloxacin and selectrin were not effective inhibitors to *V. mediterranei* 117-T6 as bacterial colonies existing in the inhibition zone of these antibiotics. Bacterium resistant to three or more antimicrobials was defined as multidrug resistant (MDR) [39,40]. *V. mediterranei* 117-T6 is a typical multidrug resistant bacterium.

3.2. Phage Morphology

*Vibrio* phage Yong-XC31 produced transparent circular plaques with clear and regular edges on *V. mediterranei* lawns (Figure 2a). Yong-XC31 is large in size having a head with icosahedral approximately spherical structure, about 113 nm in diameter, and a contractible tail, about 219 nm in length (Figure 2b).
Aeromonas phage AS-yj MF498774 0 0 0.278 
Aeromonas phage AS-szw MF498773 0 0 0.272 
V. mediterranei 117 T6 0.245 
V. mediterranei 117 T6 0.271 

Figure 1. Results of antibiotic susceptibility test. The size of the inhibition zone indicates the susceptibility of bacteria to antibiotics following the standard of the Clinical & Laboratory Standards Institute (CLSI).

Figure 2. Morphology of the plaque and *Vibrio* phage Yong-XC31. (a) plaques produced by Yong-XC31 on *V. mediterranei* lawn; (b) electron micrograph of Yong-XC31. Bar represents 100 nm.

3.3. Thermal and pH Stability

Temperature and pH stability provide more latent capacity with regard to phage storage, transport and potential applications. In the thermal and pH stability assays, Yong-XC31 was very stable at 40 °C maintaining constant titer for over 3 h, relatively stable at 50 °C and 60 °C (Figure 3a). The phage was stable at pH 5 to 8. Although the titer declined dramatically at pH 2, 10 and 11 (Figure 3b), Yong-XC31 can tolerate a pH ranging from 2–4 and 9–10 for at least two hours without complete loss of infectivity.

3.4. Optimal MOI and Factors Influencing Adsorption

Among all the tested MOIs, the optimal MOI is 0.001, when mixing $7.5 \times 10^5$ PFU of Yong-XC31 with $7.5 \times 10^8$ CFU 117-T6 host cells (Figure 4a).
Figure 3. Thermostability (a) and pH stability (b) of *Vibrio* phage Yong-XC31. All the experiments were conducted in triplicates.

Figure 4. Multiplicities of infection (MOIs) curve (a), temporary adsorption kinetics of *Vibrio* phage Yong-XC31 (b), influence of temperature on adsorption kinetics (c), and influence of pH on adsorption kinetics (d). All the experiments were conducted with triplicates.
Adsorption is a key stage in virus recognition of a sensitive host cell. Adsorption of Yong-XC31 to *V. mediterranei* 117-T6 is very efficient. The adsorption rate can reach 93.8% in 2 min at 29 °C (Figure 4b). External conditions such as temperature and pH are influential to phage adsorption, in turn, affect potential applications. From 0 to 50 °C, the adsorption rate of Yong-XC31 to *V. mediterranei* 117-T6 increases with the increase of temperature (Figure 4c). From pH 4–8, the adsorption rate increases with the increase of pH, while decreased at pH 9 (Figure 4d).

### 3.5. One-Step Growth Curve

The one-step growth curve of Yong-XC31 showed a latent period of about 30 min. There are two generally accepted methods for calculating phage burst size. The average burst size of Yong-XC31 was 64,227 calculated as the ratio of mean yield of phage particles liberated to the mean phage particles that infected the bacteria in the latent period (Figure 5) referring to reference [41], and 2 PFU/cell calculated as the ration of the final count of liberated phage particles to the initial count of infected bacterial cells at the beginning of the latent period (Figure 5) referring to reference [24].

![One-step growth curve](image)

**Figure 5.** One-step growth curve. All the experiments were conducted at MOI = 0.001 with triplicates.

### 3.6. Host Range Determination

To test Yong-XC31 host range, 34 different bacterial strains (Table 1) were used employing spot test method. Among these bacterial strains, only *V. mediterranei* 117-T6 was found to be susceptible to phage Yong-XC31. Yong-XC31 showed strict host specificity, which may be favorable for the application of the phage as it is difficult to change the normal flora.
3.7. Genome Analysis of Yong-XC31

Yong-XC31 is a typical giant phage at genome level. Complete genome sequencing was conducted using next-generation sequencing (NGS) with an average read length of 284.6 bases. 81.66% of the sequencing reads were matched to the complete genome (133,281 out of 163,212 reads) with an average sequencing depth of 130.6-fold. The complete genome of Yong-XC31 is 290,532 bp long with a G + C content of 45.87%. Yong-XC31 genome is unique sharing the highest BLASTn homology with the most related phage BONAISHI (sequence ID: MH595538, query cover as low as 1%, identity 70.62%). 25.2% predicted Yong-XC31 ORFs might encode novel proteins as they have no similarity to protein sequences in current public databases (26 January 2021).

No tRNA gene was found in the genome. Among the 318 ORFs of Yong-XC31, 17.3% were predicted functions, 82.7% predicted as hypothetical proteins. No ORFs associated with virulence factors, toxins or antibiotic resistance genes were found among the annotated ones within Yong-XC31 genome. The predicted ORFs could be classified into five functional categories, including DNA replication/regulation, bacteriophage packaging, bacteriophage structure, lysis and hypothetical protein (Figure 6, Supplement Table S1).

6 ORFs were annotated as structural and assembly genes (Supplement Table S1). 15 ORFs were annotated to be involved in DNA replication, recombination, and repair. 2 ORFs encode homing endonucleases of HNH family (ORF68 and ORF141), one of which is an intron-encoded homing endonuclease (ORF141) located between genes encoding subunits of RNAP.

As mentioned in the introduction, giant bacteriophages commonly contain many genes that do not exist in the small genome bacteriophages. Notably, Yong-XC31 harbours multiple enzyme genes related to bacteriolyis. At least 5 ORFs predicted to play a role in host cell lysis, including a peptidase (family M48, ORF29), a permuted papain-like amidase

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**Figure 6.** Genome map of *Vibrio* phage Yong-XC31. The arrow indicates the coding protein sequence, the protein function is distinguished by color, red stands for lysis, yellow for packaging, blue for DNA replication/regulation and green for structure, gray indicates hypothetical protein.
enzyme (ORF175), a glycosyl hydrolases (family 18, ORF295), a peptidoglycan hydrolase LytM (ORF143) and a lysin (ORF302). Remarkably, the lysin has a modular structure comprising an N-terminal glycoside hydrolase 19 domain, and a C-terminal peptidoglycan binding domain (PGBD).

As introduction described, some giant phages contain additional genes which may replace the function of the host proteins, thereby reducing the dependence of giant phages on their bacterial host. Noticeably, Yong-XC31 harbours eight paralogous genes (ORF34, ORF46, ORF47, ORF69, ORF136, ORF140, ORF142, ORF153) for RNAP. In particular, Yong-XC31 harbours genes (ORF265 and ORF271) encoding thymidylate synthase (dTMP synthase) and thymidylate kinase (TdT kinase) responsible for the de novo biosynthesis of thymidylate (dTMP) and as the salvage enzyme which leads to the production of dTMP even in presence of dTMP synthase inhibition [42]. Yong-XC31 also harbours a gene (ORF278) encoding tRNA nucleotidyltransferase/poly(A) polymerase participating translation, ribosomal structure and biogenesis [43].

3.8. Taxonomy

Yong-XC31 is a novel phage. As described above, Yong-XC31 shared the highest BLASTn identity (70.62%) with the closest related phage BONAISHI, with the query cover as low as 1%. Only 8 genes were found to be conserved in Yong-XC31 and Vibrio phage BONAISHI by the comparison of Yong-XC31 and Vibrio phage BONAISHI (Figure 7).

Figure 7. Genome comparison of Vibrio phage Yong-XC31 and the most closely related phage BONAISHI. Strain-specific protein coding genes are shown in light blue, protein coding genes conserved in Yong-XC31 and Vibrio phage BONAISHI are shown in yellow.

Further, the genome of Yong-XC31 shared only 0–58.65% ANI values with the giant phages with the 10 top highest homology in BLASTn comparison (e-value < 10^{-5}) (Table 2), which are substantially below the ≥95% ANI boundaries to define a species [44,45]. The isDDH values between Yong-XC31 and these giant phages all were 0, lower than the 70% cut off to define a species [34]. The POCP values between Yong-XC31 and giant phage species were 0.26–6.77%, much lower than the genus boundary cut-off of 50%.

In addition, the Bacterial and Archaeal Viruses Subcommittee (BAVS) within the International Committee on the Taxonomy of Viruses (ICTV), which holds the responsibility of classifying new prokaryotic viruses, recently redefined a genus as a cohesive group of viruses sharing a >50% of high degree of nucleotide sequence similarity [45,46]. Yong-XC31 shared the highest nucleotide sequence similarity, as low as 11.71%, with the closest related phage in PASC [37] search. It’s much below the >50% boundaries to define a genus. Summarily, these results indicate the status of Yong-XC31 as a taxonomically unique phage presenting a novel monophyletic genus.

Most jumbo phages were attributed to the Myoviridae family. Yet, in the proteomic tree constructed based on genome wide sequence similarities (Figure 8), Yong-XC31 and three unclassified giant phages clustered in a monophyletic clade independently between the family Drexlerviridae and Herelleviridae. These four giant phages harbor genomes of 242,446–309,157 bp. A new phage family is proposed in Caudovirales order.
Figure 8. The proteomic tree is generated using ViPTree online based on genome-wide similarities determined by tBLASTx. 56 type species of nine families belonging to the order Caudovirales, *Vibrio* phage Yong-XC31, and 10 giant phages in Table 2 were included in the analysis. Bacteriophage family assignments according to the official ICTV classification are provided with different color bars. No color bar is marked to unclassified giant phages. Red star refers to Yong-XC31. The sizes or size ranges of the giant phage genomes are shown in red font on the right.

4. Discussion

*Pyropia* is an important cultivated seaweed in Northeast Asia [7, 47]. In recent years, increase in the prevalence of diseases and pests has caused a subsequent reduction in their quantity and commercial value [47, 48]. Among these diseases, yellow spot disease (YSD) is a destructive disease of the conchocelis sporeling culture of *Pyropia* [7, 47, 48]. *V. mediterranei* 117-T6 (CGMCC1.16311) was pathogenic to the YSD of conchocelis of *Pyropia* [7, 47, 48].
**Pyropia** species [8]. Traditional antibiotic therapy has no stable effect on YSD [49]. In this study, antibiotic susceptibility test demonstrated that *V. mediterranei* 117-T6 was multidrug resistant, which accord with the ineffective antibiotic treatment. Therefore, the development of new tools and strategies to control pathogens and treat diseased *Pyropia* is becoming a most important issue.

Phages are currently emerging as potential treatments for multidrug resistant bacterial infections. Yet, literature search yielded no information about *V. mediterranei* phage. In this study, the first *V. mediterranei* phage was isolated and identified as a novel giant phage representing a novel phage genus. Proteomic tree indicates that Yong-XC31 together with giant *Vibrio* phage BONAISHI, *Vibrio* phage 2 TSL-2019 and *Pseudomonas* phage Phabio represent a new evolutionary lineage independently between the family Drexleviridae and Herelleviridae. The four giant phages all infect Gram-negative bacteria, contain genome of 242,446–309,157 bp with only an average 1/4 of the predicted genes being assigned functions. A new family is proposed here in Caudovirales. In the novel proposed family, Yong-XC31 presents a monophyletic genus in consideration of the pair-wise ANI, isDDH, POCP and PASC values.

The therapeutic value of a candidate bacteriophage relies on the characterization of viral properties such as stability, growth kinetics, host range and viral yield [50]. Results of this study indicated that the temperature and pH stability of Yong-XC31 was good, which is conducive to its application to control *V. mediterranei* infections in complex environment. Results also demonstrated other good properties of Yong-XC31 including a very low optimal MOI (0.001), efficient adsorption to host (reach 93.8% adsorption rate in only 2 min), a short latent period (30 min), a high burst size (64,227) and strict host specificity. Additionally, our small-scale laboratory experiments proved that Yong-XC31 could protect both the free-living conchocelis (FLC) and the shell-borne conchocelis (SBC) of *Pyropia* from harm caused by *V. mediterranei* [49]. These results suggest significant application potential of Yong-XC31 in *Pyropia* production.

Thus, far, only fourteen jumbo phages infecting *Vibrio* bacteria have been characterized. Among them, Yong-XC31 is most related to the unclassified phage BONAISHI, which is strictly lytic for several strains of *V. coralliilyticus* pathogenic to coral [50]. Yong-XC31 and BONAISHI share several special characteristics. Their genomes are both about of 290 kb, contain more than one paralogous gene encoding RNAPs and do not harbour identified tRNA gene.

Most phages rely on RNAPs of a bacterial host to transcribe their genes [51]. A strategy used by some phages is to depend on their own single-subunit RNAPs to transcribe a subset of viral genes [52]. An even more radical strategy is used by some giant phage, not need to rely on the host RNAP, yet only relies on self-encoded phage multi-subunit RNAPs: virion-associated RNAPs and early expressed RNAPs [14]. These RNAPs can perform two functions. The virion-associated RNAP is injected into the host cell together with phage DNA and transcribes early phage genes [53]. The early expressed RNAP formed by putative RNAP subunits which would transcribe viral genes expressed in the middle and late stages of infection [54]. This study finds that Yong-XC31 contains five genes encoding virion-associated RNAPs (ORF34, ORF136, ORF140, ORF142, ORF153) and three early expressed RNAPs (ORF46, ORF47, ORF69). This may be beneficial to its efficient infection. In addition, the absence of detectable tRNA in phage BONAISHI genome suggests that it is well adapted to the translation machinery of its hosts, which is a critical process for efficient phage propagation [50]. Similarly, we did not find any tRNA in the genome of Yong-XC31.

As mentioned above, alternative therapies must be developed to mitigate the sharp increase in antibiotic resistance. Besides phage itself, novel antimicrobial strategies include enzyme-based antibiotics (“enzybiotics”) such as phage lytic enzymes. Yong-XC31 harbors multiple (at least eight) enzyme genes related to bacteriolysis. These enzyme genes may be potential beneficial resources.
Very little is known about the processes of host–phage interaction in marine environments. Bailey et al., found that *Synechococcus* phage S-PM2 contain genes encode homologs of the key photosystem II reaction center core polypeptides (D1 and D2) and proposed that this might play an active role in protecting their hosts against photoinhibition, thereby ensuring an energy supply for replication by preventing the deleterious effects on host cell integrity seen during acute photoinhibition [55]. Phages infecting marine picocyanobacteria often carry a *psbA* gene, which encodes a homolog to the photosynthetic reaction center protein, D1. Bragg et al., proposed that phage encoded D1 may help to maintain photosynthesis during the lytic cycle, which in turn could bolster the production of deoxynucleoside triphosphates (dNTPs) for phage genome replication [56]. They examined the contribution of phage *psbA* expression to phage genome replication under constant low irradiance and predicted that phage *psbA* expression could lead to an increase in the number of phage genomes produced during a lytic cycle of between 2.5 and 4.5% [56]. In this study, Yong-XC31 possesses unusual genes rarely present in other phages. In particular, Yong-XC31 contains an ORF encoding Methyl-accepting chemotaxis protein (MCP) and a gene encoding EIIB belonging phosphotransferase system. As the predominant chemoreceptor and signal transducer in bacteria and archaea, MCPs also termed transducer-like proteins (Tlps), serve as sensors in bacterial chemotactic signaling [57]. MCPs sense intracellular and environmental signals, and relay them to the downstream signaling pathways in the cytoplasm. Then, bacteria utilize the well-known chemotactic responses to move towards factors that favor survival [58]. Further studies are needed to find whether the expression of the viral methyl-accepting chemotaxis protein gene may enhance the host survival and persistence in the complex environment, thus profiting the phage itself. The phosphoenol phosphotransferase system (PTS) is a multi-component signal transduction cascade that regulates diverse aspects of bacterial cellular physiology in response to the availability of high-energy sugars in the environment [59]. In bacteria, there are often many different EII's. EII's are responsible for the phosphorylation of the carbohydrate as well as its transportation across the bacterial membrane [60]. The significance of gene encoding EIIB in Yong-XC31 remains to be studied.

5. Conclusions

In conclusion, *Vibrio* phage Yong-XC31 was isolated and characterized as a new potential biocontrol strategy to control *Vibrio mediterranei*. A total of 80 orphan ORFs in Yong-XC31 may be a new resource of applications. Giant phage Yong-XC31 has very unique genome sequence. Blast search results, ANI values, isDDH values, POCP values, nucleotide sequence similarity estimated via PASC and proteomic tree demonstrated Yong-XC31 as a singleton phage distantly related to previously sequenced bacteriophages, representing a new evolutionary lineage of phage. We propose a new phage family with Yong-XC31 as the representative specie of a genus. This study expands the diversity of phages.

6. Patents

Lihua Xu; Dengfeng Li; Jing Fang et al., A virulent phage vB_VmeM-Yong XC31 of *Vibrio mediterranei* and its application (ZL201910792955.8).

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