

1 **A novel oceanic uncultured temperate cyanophage lineage**

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8

9 **Abstract**

10 **Marine cyanobacteria are important contributors to primary production in the**
11 **ocean and their viruses (cyanophages) affect the ocean microbial communities.**
12 **Despite reports of lysogeny in marine cyanobacteria, isolation of lysogenic marine**
13 **cyanophages has not been reported yet. Using assemblies from Red Sea and *Tara***
14 **Oceans metagenomes, we recovered genomes of a novel uncultured marine**
15 **cyanophage lineage, which contain genes coding for common cyanophages proteins,**
16 **as well as the phycobilisome degradation protein NblA, an integrase, and a split DNA**
17 **polymerase. The DNA polymerase forms a monophyletic clade with a DNA**
18 **polymerase from a putative prophage in *Synechococcus* WH8016. The putative**
19 **prophage does not resemble any known cyanophage but shares several genes with**
20 **the newly identified cyanophages. Metagenomic recruitment indicates that the novel**
21 **cyanophages are widespread, albeit at low abundance. This is the first report of a**
22 **putative prophage in a cultured marine *Synechococcus* and the genomic**
23 **characterization of its cyanophage relatives.**

24

25 Introduction

26 Viruses are the most abundant biological entities in the planet (Breitbart, 2012; Paez-
27 Espino et al., 2016). Cyanophages, viruses infecting cyanobacteria, regulate cyanobacterial
28 communities and influence the global nutrient cycles. Viruses also represent major genetic
29 reservoirs and rewire their hosts metabolism (Breitbart, 2012).

30 Viral infections can be lytic or lysogenic. Lytic infections finish with host lysis and
31 virions release, while in lysogenic infections the viral genomes integrate into the host
32 chromosomes as prophages and replicate without virion production (Howard-Varona et al.,
33 2017). Although there are no reports of prophages in cultured cyanobacteria induction of
34 prophages from marine cyanobacterial environmental populations has been studied
35 (McDaniel et al., 2002; Ortmann et al., 2002). Genomic analysis of *Prochlorococcus* phage
36 P-SSP7 suggests that the virus may reproduce in a lysogenic way (Sullivan et al., 2005),
37 additionally a partial P-SSP7-like prophage sequence was identified during a study of
38 environmental *Prochlorococcus* communities from the Pacific Ocean (Malmstrom et al.,
39 2013).

40 We previously screened Bacterial Artificial Chromosome (BAC) libraries for
41 photosystem II (PSII) genes. While some BACs originated from cyanobacteria, other
42 resembled known cyanophage genomes (Zeidner et al., 2005). However, one clone from
43 the Mediterranean Sea (BAC21E04) was notable as it contained numerous ORFs with weak
44 or no homology to anything reported in GenBank, including both non-redundant and
45 environmental-non-redundant entries. BAC21E04 did contain a full length viral-like D1 gene,
46 a partial-length viral-like *talC* transaldolase gene and a putative ribonucleotide reductase
47 (RNR) class II gene (Zeidner et al., 2005). This precluded the possibility to assign affiliation
48 to this BAC at the time.

49 Here we report the identification of a novel lineage of cyanophages related to
50 BAC21E04. These cyanophages possess properties from temperate phages and shares
51 synteny with a putative prophage in *Synechococcus* WH8016.

52

53 **Results and Discussion**

54 To increase our knowledge regarding uncultured cyanophages, we examined
55 metagenomics assemblies from the Red Sea (Philosof et al., 2017) and *Tara* Oceans
56 expedition microbiomes and viromes (Brum et al., 2015; Sunagawa et al., 2015). Four
57 contigs of about 86 kbp representing putative viral genomes related to BAC21E04 were
58 identified (File S1). These contained overlapping terminal regions suggesting that they
59 represent complete, terminally redundant viral Metagenome Assembled Genomes (MAGs)
60 (File S2). The genomic content of the MAGs was compared to 82 genomes from
61 cyanophages infecting *Synechococcus* and *Prochlorococcus* (File S3), identifying 14 564
62 proteins clustered into 2 376 orthologs. Analysis of the shared orthologs between viruses
63 allowed to delineate four clusters of cyanophages corresponding to the viral families
64 *Myoviridae*, *Podoviridae*, *Siphoviridae* and one composed exclusively by the MAGs (Fig.
65 1a), such arrangement remains even when comparing genomic nucleotide distances (Fig.
66 S1). The clustering results suggested that the MAGs presented here belong to a new
67 cyanophage lineage.

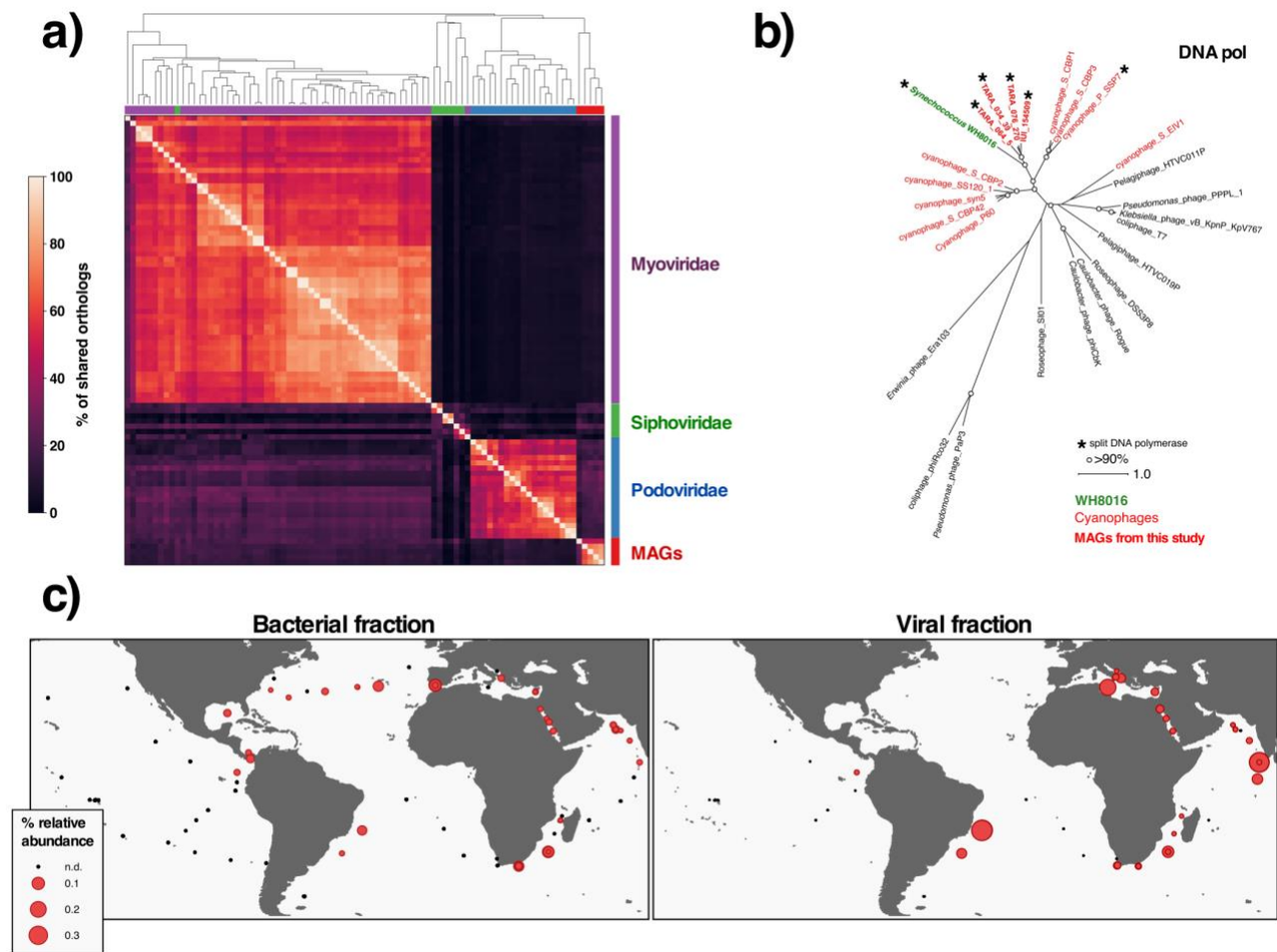
68 Mapping of the *Tara* Oceans metagenomes into the MAGs indicated the presence
69 (>0.01% relative abundance) of these cyanophages in 36 samples from 16 stations with an
70 average of 0.07% (Fig. 1c). While a viral fraction (<0.2 μ m) sample from the South Atlantic
71 (TARA_076) had the highest abundance of the new cyanophages (0.39%), suggesting the
72 presence of cell-free virions from the new family in the environment, two thirds of the
73 samples where the MAGs were detected correspond to the bacterial fraction (0.2–3.0 μ m)
74 probably indicating ongoing infections during sampling (Philosof et al., 2017).

75 All four MAGs contained a cyanophage-like ribonucleotide reductase (RNR) class II
76 gene (Fig. S2), a *talC* transaldolase gene (Fig. S2), and an exonuclease (*exo*). Three
77 contained tRNA genes, an integrase gene, and a D1 gene with viral signatures (Sharon et
78 al., 2007). Also, for the first time in marine cyanophages genes coding for the phycobilisome

79 degradation protein NblA were detected in each of the newly identified cyanophages, with
80 three of the MAGs carrying an additional copy of *nblA*. Functional *nblA* genes can be found
81 in freshwater cyanophages (Gao et al., 2012), however their presence in marine viruses is
82 intriguing as no sequenced marine cyanobacteria carries the gene (Supp Info). Besides of
83 the terminase large subunit (TerL), portal, major capsid protein (MCP), and putative tail
84 proteins identified based on structural predictions, no other known phage structural proteins
85 could be classified supporting the classification of these cyanophages into a new family (File
86 S4).

87 The new cyanophages also encode a split DNA polymerase, which formed a new
88 cluster within the T7-like cyanophage DNA polymerase family (Fig. 1b). Unexpectedly, a
89 split DNA polymerase from *Synechococcus* WH8016 also clustered with the new phage
90 family. Examination of the vicinity of the DNA polymerase genes in the genome of WH8016
91 revealed the existence of a genomic island bearing a possible prophage of approximately
92 100 kbp (Fig. 2) with distinct nucleotide usage. The putative prophage does not resemble
93 any known cultured marine cyanophage genome at the nucleotide level, but do shares
94 several genes with BAC21E04 and other cyanophages including the newly identified viral
95 family (split DNA polymerase genes, integrase, phage/plasmid related protein; Fig. 2). The
96 putative host site-specific attachment site (*attB*) for WH8016 could be located between a
97 tRNA-Pro located upstream the integrase in the borders of the prophage-like area, however
98 the region does not contain detectable known viral structural protein coding genes.

99 Genomic evidence for a partial prophage in *Prochlorococcus* cells was recently
100 observed using single cell genomics (an ~8 kbp fragment resembling part of the podophage
101 P-SSP7 genome (Malmstrom et al., 2013)). Based on our observations we suggest that the
102 newly identified marine phages form a new cyanophage family related to the identified
103 putative prophage in *Synechococcus* WH8016.

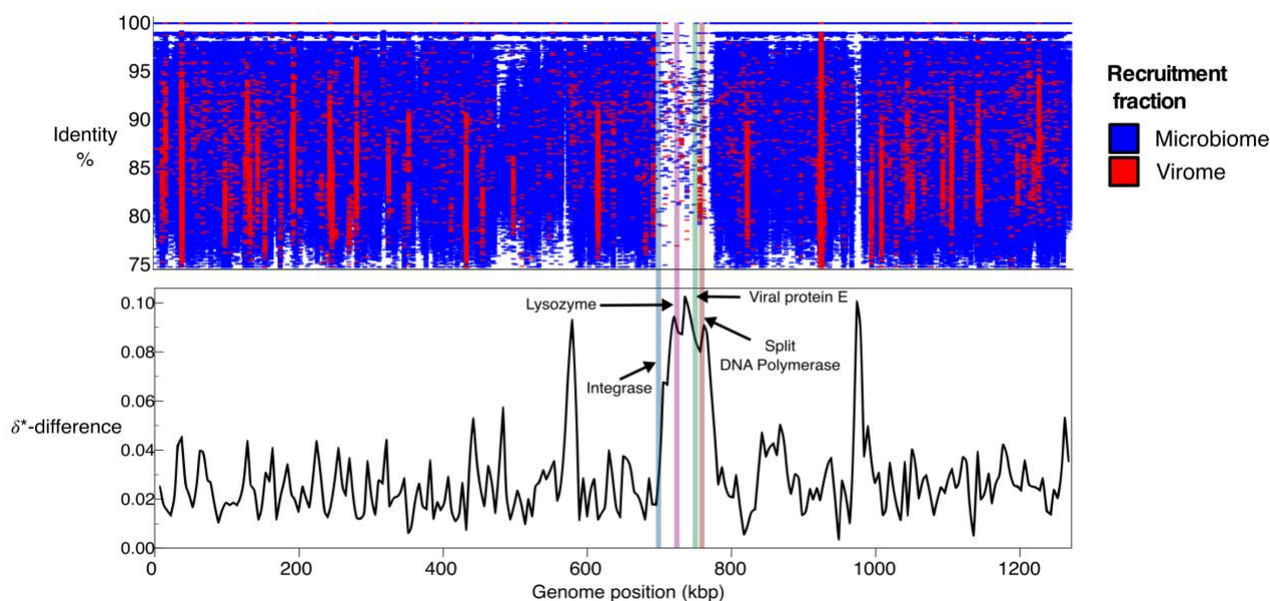


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105 **Figure 1. a)** Comparison of Cyanophages genomes clustering using percentage of shared
 106 orthologs. Heat map representation of protein orthologs shared by cyanophages, scaled
 107 between 0 (darker) and 100% (lighter). Dendrogram tips are colored according to the
 108 taxonomic affiliation of the cyanophages using the following key: *Myoviridae* (purple),
 109 *Podoviridae* (blue), *Siphoviridae* (green), and the MAGs identified in this study (red).
 110 Clustering identifies four large groups corresponding to the taxonomic families from the
 111 cyanophages (Full details in the Supplemental information). **b)** Phylogenetic protein tree of
 112 DNA Polymerase. The newly identified MAGs and *Synechococcus* WH 8016 are shown in
 113 bold red. Cyanophage and cyanobacteria sequences are shown in red and green,
 114 respectively. Split DNA polymerases are marked with an asterisk. The MAGs form a
 115 monophyletic clade with a split DNA polymerase from a putative prophage in
 116 *Synechococcus* WH8016. **c)** Distribution and relative abundance of the newly identified
 117 cyanophages in Tara Oceans metagenomes. Each red circle is scaled to represent the

118 relative abundance of the MAGs in each station. Black points indicate stations where the
119 MAGs could not be detected (<0.01%).

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Figure 2. A putative prophage in the genome of *Synechococcus* WH8016. Top: Recruitment
123 of *Tara* Oceans reads from 331 metagenomes onto *Synechococcus* WH8016 genome.
124 Each recruited read is drawn according to the position of the genome where it aligned and
125 the nucleotide identity to the recruitment area. Reads are colored according to the sample
126 fraction of origin, red for virome (<0.2 μm) and blue for microbiome (0.2–3.0 μm). Bottom:
127 Karlin signature (δ^*) difference, the difference of the relative abundance of dinucleotides
128 between a sliding window and the whole sequence (Karlin et al., 1998), was calculated using
129 a window size of 10 000 bp and a step size of 5 000 bp. Colored bars spanning both panels
130 indicate position of some of the shared proteins in synteny with the newly identified
131 cyanophages.

132

133 **Materials and methods**

134 A detailed methodology can be found in the Supplemental Information.

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136 **Data and code availability**

137 The sequence of the MAGs can be found in File S1. The rest of the sequences are deposited
138 at GenBank as follows: BAC21E04 [AY713439-AY713444], *Synechococcus* WH8016
139 [AGIK01000001]. File S3 contains the accessions for the collection of cyanophages.
140 Supporting data can be retrieved from <https://osf.io/q3khc/> while the Jupyter Notebooks and
141 scripts used are publicly available at https://github.com/BejaLab/TM_Cyanophages.

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143 **Author Contributions**

144 J.F.-U. and O.B conceived the project. J.F.-U., A.P., I.S., and O.B. performed bioinformatic
145 analyses. J.F.-U. and O.B. wrote the manuscript with contributions from all authors to data
146 analysis, figure generation, and the final manuscript.

147

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154 **Conflict of interest**

155 The authors declare no conflict of interest.

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