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Provirophages in the *Bigelowiella* genome bear testimony to past encounters with giant viruses

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Virophages are recently discovered double-stranded DNA virus satellites that prey on giant viruses (nucleocytoplasmic large DNA viruses; NCLDVs), which are themselves parasites of unicellular eukaryotes. This coupled parasitism can result in the indirect control of eukaryotic cell mortality by virophages. However, the details of such tripartite relationships remain largely unexplored. We have discovered ~300 predicted genes of putative virophage origin in the nuclear genome of the unicellular alga *Bigelowiella natans*. Physical clustering of these genes indicates that virophage genomes are integrated into the *B. natans* genome. Virophage inserts show high levels of similarity and synteny between each other, indicating that they are closely related. Virophage genes are transcribed not only in the sequenced *B. natans* strain but also in other *Bigelowiella* isolates, suggesting that transcriptionally active virophage inserts are widespread in *Bigelowiella* populations. Evidence that *B. natans* is also a host to NCLDV members is provided by the identification of NCLDV inserts in its genome. These putative large DNA viruses may be infected by *B. natans* virophages. We also identify four repeated elements sharing structural and genetic similarities with transposons—a class of mobile elements first discovered in giant viruses—that were probably independently inserted in the *B. natans* genome. We argue that endogenized provirophages may be beneficial to both the virophage and *B. natans* by (i) increasing the chances for the virophage to coinfect the host cell with an NCLDV prey and (ii) defending the host cell against fatal NCLDV infections.

virophage | nucleocytoplasmic large DNA virus | microbial community | endogenous virus | Maverick/polinton

Sputnik was first described in 2008 as a new class of small icosahedral viruses with an ~20-kb circular double-stranded DNA genome (1). Sputnik is a satellite virus, because its replication depends upon proteins produced by the nucleocytoplasmic large DNA virus [NCLDV; also giant virus or proposed order *Megavirales* (2)] *Acanthamoeba polyphaga* Mimivirus (APMV; *Mimiviridae*) and replicates in APMV viral factories. Sputnik was shown to inhibit replication of its helper virus and thus acted as a parasite of that virus. In analogy to the term bacteriophage it was called a virophage, but this designation has been challenged (3). Three additional virophages infecting members of the *Mimiviridae*, e.g., Sputnik 2, Rio Negro, and Zamilon, were subsequently reported (4–6). Virophages that prey on giant viruses that infect heterotrophic nanoflagellates and microalgae have also been discovered, including Organic Lake virophage 1 [OLV1 (7)], Mavirus (8), and a virophage of the *Phaeocystis globosa* virus (PgVV) (9), yet the classification of the latter as a virophage sensu stricto is uncertain. In addition, complete or near-complete virophage genomes have been assembled from environmental DNA: Yellowstone Lake virophages 1–7 (YSLV1–7) and Ace Lake Mavirus (ALM) (10, 11).

Overall, virophage genomes have similar sizes (~18–28 kb) and low G+C content (~27–39%) and are related to Sputnik by genetic and structural homologies (12). Among the 20–34 protein-coding sequences predicted in virophage genomes, the putative core gene set comprises six genes encoding the FtsK-HerA family DNA-packaging ATPase (ATPase), primase-superfamily 3 (S3) helicase,

cysteine protease (PRO), and zinc-ribbon domain (ZnR) as well as major and minor capsid proteins (MCPs and mCPs, respectively) (12). In addition, genes encoding two different families of integrases have been identified in several virophages: A putative rve integrase was found in Mavirus and ALM (8, 10), whereas Sputnik encodes a putative tyrosine integrase (1). Among virophage genes, only PRO, ATPase, MCP, and mCP support the monophyly of virophages, whereas the remaining gene complement shows complex phylogenies suggestive of gene replacement (12).

Remarkably, phylogenetic analysis of the Mavirus rve integrase indicated that it is mostly related to homologs from eukaryotic mobile elements of the Maverick/polinton (MP) family (8). The polintons are widely distributed in diverse protists and animals and were initially classified as transposable elements (TEs) (13, 14). However, convincing arguments support the hypothesis that polintons encode capsid proteins and might be bona fide viruses (15). Because Mavirus was reported to display further synapomorphy with a putative MP from the slime mold *Polysphondylium pallidum*, it was hypothesized that MPs may have originated from ancient Mavirus relatives that would have acquired the capability of intragenomic transposition (8). However, this hypothesis was recently challenged by Yutin et al. (12). A critical prerequisite for such an evolutionary scenario is the integration of virophage DNA in the genome of a eukaryotic host that would permit vertical transmission and adaptation to an intracellular parasitic lifestyle. However, although Sputnik 2 was shown to integrate into the genome of its Mimivirus host (4), evidence of virophage insertions in eukaryotic genomes is lacking.

Significance

Virophages are viruses that hijack the replication machinery of giant viruses for their own replication. Virophages negatively impact giant virus replication and improve the survival chances of eukaryotic cells infected by giant viruses. In this study, we identified segments of the *Bigelowiella natans* genome that originate from virophages and giant viruses, revealing genomic footprints of battles between these viral entities that occurred in this unicellular alga. Interestingly, genes of virophage origin are transcribed, suggesting that they are functional. We hypothesize that virophage integration may be beneficial to both the virophage and *B. natans* by increasing the chances for the virophage to coinfect the cell with a giant virus prey and by defending the host cell against fatal giant virus infections.

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The coinfection of host cells by NCLDV and virophages has been shown to limit the production of NCLDV particles, accompanied by greater survival of the eukaryotic host (1, 5, 6, 8, 16). At the community level, these parasitic relationships result in complex interplays between virophages, NCLDVs, and eukaryotic hosts. As a result, virophages indirectly positively regulate the population size of eukaryotic hosts. At the global scale, these interactions may have significant impacts on biogeochemical cycles. For instance, in the marine environment, the co-occurrence of giant viruses and virophages in the context of algal blooms may influence the overall carbon flux as proposed for Antarctic lakes (7). Nevertheless, such tripartite community networks remain poorly explored except on theoretical grounds (17, 18).

The ecological prominence and diversity of virophages are largely unknown and wait for the isolation and sequencing of new specimens. Recently, we demonstrated the value of searching integrated viral DNA in the genomes of potential eukaryotic hosts to identify new members of the NCLDVs (19). Here we analyzed the nuclear genome assemblies of 1,153 fully sequenced eukaryotes and report the identification of integrated virophage elements in the genome of the Chlorarachniophyte *Bigelowiella natans* (supergroup Rhizaria). This discovery led to the prediction that this alga is also the host of viruses that are members of the NCLDVs. In support of this prediction, we also identified inserts of likely NCLDV origin. We investigated the transcriptional activity of the *B. natans* genome using RNA-sequencing (seq) data and show that virophage-like genes are actively transcribed in different *B. natans* strains whereas NCLDV-like genes tend to be silent—albeit with notable exceptions. Finally, we identified repeated genetic elements that have structural and genetic similarities to transpovirons, a distinct class of mobile genetic elements associated with giant viruses that were first discovered in members of the *Mimiviridae* (4). We discuss the biological relevance of integrated, actively transcribed virophages and propose a model for the mode of virophage–NCLDV coinfection. Altogether, our results contribute to the understanding of the genetic interactions occurring within microbial communities between eukaryotes, virophages, and NCLDVs.

Results

Integrated Virophage-Like Elements in the Algal Genome. A previous comparative analysis of fully sequenced virophage genomes revealed six core proteins or protein domains that are universally conserved, including S3 helicase, zinc-ribbon domain, major capsid protein, minor capsid protein, DNA-packaging ATPase, and a cysteine protease (12). The four latter proteins were shown to produce consistent monophyletic clades that contrasted virophages from polintons, a class of repeated elements related to virophages. We therefore used these proteins as markers of DNA inserts of putative virophage origin in eukaryotic genome sequences. In practice, we searched 1,153 predicted proteomes of protists, fungi, and basal metazoans for homologs of the four virophage markers using BLASTP. The proteome of *B. natans* was the only one to exhibit homologs for each of the four virophage core protein families. No homolog for any of the virophage markers was identified in the other proteomes using predefined family-specific score thresholds.

To better delineate the subset of *B. natans* proteins that have a potential virophage origin, we used a score plot approach. BLAST scores obtained between *B. natans* predicted proteins and their best virophage matches were plotted against the respective BLAST scores obtained between the *B. natans* predicted proteins and their best cellular matches (Fig. 1). Blue dots below the diagonal identify *B. natans* proteins that have higher similarity to a virophage protein than to a homolog in a cellular organism. Overall, 103 *B. natans* proteins had a match within a virophage proteome, of which 64 had a higher score with virophages than with cellular organisms. Furthermore, examination of the physical location of the virophage-like protein genes revealed that they tend to cluster in specific loci in the genome assembly, revealing large regions of possible virophage origin.

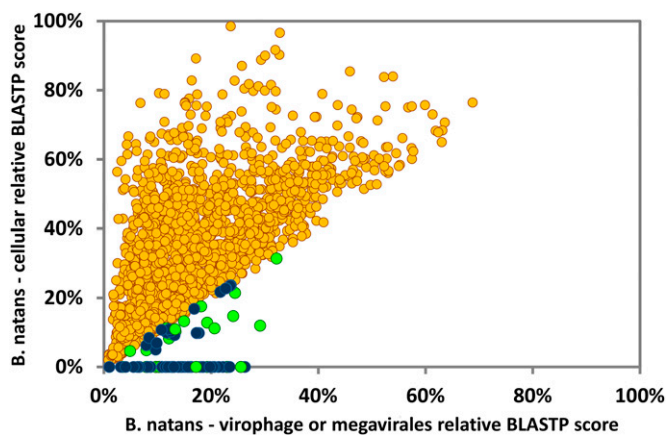


Fig. 1. Similarity plot of *B. natans* proteins against virophage/NCLDV and cellular best hits. Circles represent relative BLASTP scores of *B. natans* proteins aligned against their best cellular hits in the NR database (y axis) and their best viral hits among NCLDVs or virophages. When no cellular hit was recorded whereas a viral hit was obtained, the cellular score was set to zero. BLAST scores were normalized by dividing them by the score of the alignment of the query sequence against itself. Circles are colored according to the origin of the best overall scoring hit (yellow, cellular organisms; blue, virophages; green, NCLDVs).

A total of 38 virophage-like elements (VLEs) ranging from 100 base pairs (bp) to 33.3 kb were detected by nucleotide alignments in the *B. natans* genome assembly. Many VLEs correspond to truncated copies of larger elements, suggesting recurrent insertions followed by degradation. However, the number of VLEs may be misestimated because some of them lie at the end of contigs, suggesting that VLEs are difficult to assemble. None of the VLEs were located on the nucleomorph chromosomes, mitochondrial genome, or chloroplast genome. The cumulated size of VLE sequences reaches 327 kb. The VLEs were highly similar between each other—that is, nucleotide identities averaged 91.3%—indicating that they belong to the same family of closely related elements. However, sequence conservation was occasionally interrupted by unique sequences containing one or more genes as shown in Fig. S1, revealing insertion or deletion events that occurred subsequent to their divergence. Some VLEs may be unable to produce viable virophages (i.e., unable to complete a full replication cycle) because important genes may have been lost following integration. Alternatively, the difference in gene content between VLEs may reflect the genetic diversity of virophages before their integration. Six of the identified VLEs contained terminal inverted repeats (TIRs) of 2.0–2.6 kb at their two extremities. TIRs were also described at the extremities of the PgVV virophage-like element [associated with the virus PgV-16T infecting *P. globosa* (9)] and polintons, and are common among poxviruses, chloroviruses, and asfarviruses (7). In *B. natans*, each TIR contained at least two putative ORFs. Other VLEs contained a single TIR copy at one of their extremities, most probably because these elements were truncated.

As shown in Fig. 2A, the G+C content of VLEs (36.4% on average) was markedly lower than the background G+C content of the host genome (44.9% on average). Such a difference in G+C content suggests that the VLEs have been acquired horizontally in the relatively recent past.

Virophage-Like Element Genes. Overall, 298 ORFs (>90 codons) were predicted out of the 38 VLEs and organized into 54 gene families (Dataset S1). The largest element of 33.3 kb was identified on scaffold 2 (positions 1,655,224–1,688,550; Fig. 2B) and contained 27 predicted ORFs representing 25 distinct gene families listed in Table 1. Functional annotation could be predicted for only 14 of the pan-VLE gene families. Furthermore, 39 gene

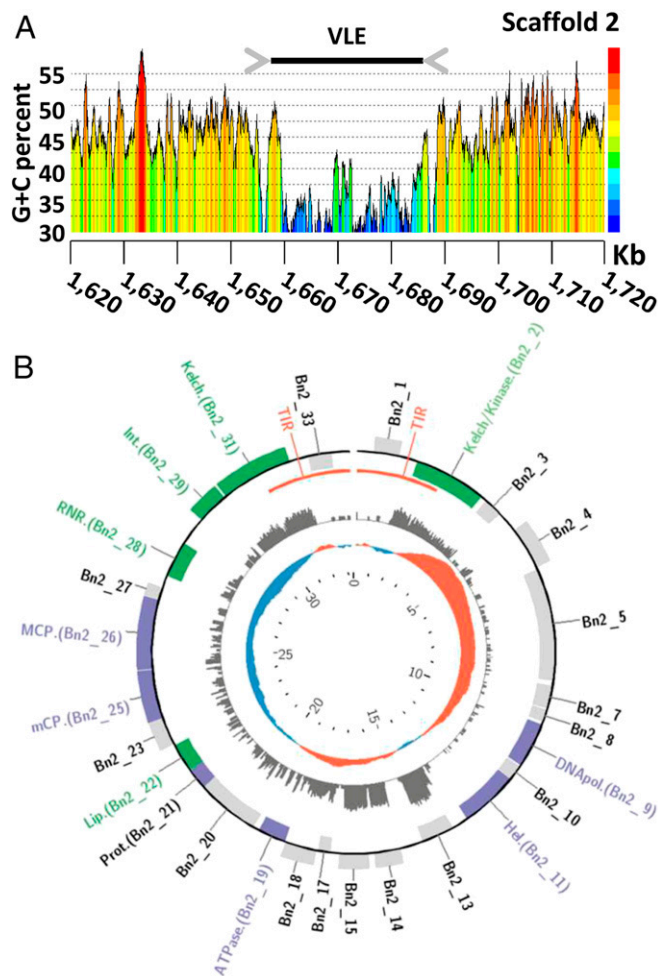


Fig. 2. Schematic representation of the largest viroplage-like element of *B. natans*. (A) G+C content curve in the region of the viroplage-like element on scaffold 2. (B) Gene map of the linear viroplage-like element. Genes displayed on the outer and inner rims of the outermost track are coded on the direct (Watson) and reverse (Crick) strand, respectively. Purple and green genes correspond to core viroplage genes and genes with annotated functions, respectively. The middle track indicates the number of RNA-seq reads mapped to the regions (logarithmic scale). The inner track represents the G+C skew, with positive values in red and negative values in blue. The genome coordinates (in kb) are indicated in the innermost track of the map.

families did not have a significant match in sequence databases (BLASTP E value < 0.01). Such a small fraction of predicted function is a trademark of viroplages that generally contain a majority of orphan genes (1, 7, 8). Also, the functions encoded by the *B. natans* VLEs are reminiscent of bona fide viroplages. Homologs could be identified for each of the six core genes that are ubiquitously conserved in complete viroplage genomes (12). Importantly, these include the major and minor capsid protein genes that are among the viral hallmark proteins distinguishing viruses from other types of mobile elements. In addition, the VLEs encode two key proteins involved in virion maturation, namely the protease and the packaging ATPase. The core viroplage zinc-ribbon domain is fused with a GIY-YIG nuclease domain like in Mavirus and OLV; noticeably, the corresponding protein (Bn119_7) is encoded by only one of the VLEs on scaffold 119, suggesting that the other VLEs have lost the gene at some point in their evolution. Furthermore, the VLEs were found to encode an rve family integrase closely related to the one encoded in Mavirus and ALM. This protein might have catalyzed the in-

tegration of the original viroplage-like DNA into the algal genome. The VLEs also contain five gene families homologous to sequences that have a more patchy distribution among viroplages, including genes for lipase, ribonucleotide reductase small subunit, B family DNA polymerase, and two ORFs with unknown function.

Each TIR region encompasses two ORFs, resulting in two pairs of duplicated genes in complete VLEs. The outward duplicated ORFs within TIRs lack functional annotation. On the other hand, the inward duplicated ORFs encode Kelch-repeat domains most similar to eukaryotic and poxvirus homologs (Fig. S24) that may be involved in protein-protein interaction. Interestingly, these ORFs terminate outside the TIR regions, so that the N termini differ between the two protein copies. One of the N termini (Bn2_31) encodes a domain with no match in sequence databases, whereas the N-terminal region of the other protein (Bn2_2) encodes a protein kinase (PK) domain that is indirectly related to those encoded in various giant viruses (Fig. S2B). Sequenced viroplages have been shown to contain genes that are closely related to homologs from their respective giant virus hosts (1, 7, 8, 12). These apparent host-derived genes encode different repetitive proteins (distinct forms of collagen-like repeats in Sputnik and OLV, and FNIP repeats in Mavirus) that could be implicated in the interaction of the viroplages with their giant virus hosts (12). The VLEs encode two gene families that are most similar to giant virus homologs (i.e., Bn2_18 and Bn2_28), but not one encodes repetitive proteins. However, the Kelch domain proteins (i.e., Bn2_2 and Bn2_31), ankyrin domain protein (Bn161_7), and adhesion-like protein (Bn2_5) are proteins with repetitive domains likely implicated in the interaction of the viroplages with their giant virus hosts; however, they have no significant match in sequenced giant viruses.

Origin of Viroplage-Like Elements. We performed a phylogenetic analysis of the viroplage-like proteins that had sufficient evolutionary conservation with homologs in sequence databases. For each protein family, the *B. natans* paralogs aggregated in a single clade containing short branches, further confirming the very close relationships between the different VLEs. The packaging ATPase and protease were previously shown to support the monophyly of viroplages (12). The corresponding *B. natans* proteins cluster within the viroplage clade (Fig. 3A and B). In both these trees, the *B. natans* proteins group with Sputnik, OLV, and YSLVs, whereas Mavirus and ALM cluster in a sister clade. The phylogenetic tree of the major capsid proteins is compatible with the scenario of a closer evolutionary proximity of the *B. natans* VLEs to OLV, YSLVs, and Sputnik than to the ALM-Mavirus clade (Fig. 3C). The same conclusion can be drawn from the phylogenetic reconstruction of the viroplage lipase (Fig. S2C). Moreover, the fast-evolving minor capsid protein was conserved between Sputnik, OLV, Zamilon, YSLVs, and the *B. natans* elements, whereas the Mavirus and ALM homologs were comparatively too diverged to allow accurate phylogenetic reconstruction, further pointing to a more distant relationship with the latter (Fig. 3D). Altogether, these features suggest that the *B. natans* viroplage elements share a common ancestor with Sputnik-OLV-YSLVs. However, the exact timing of speciation events within this subtree could not be established, owing to incongruences between the phylogenetic reconstructions.

The phylogenetic trees for the other *B. natans* viroplage-like proteins disclosed more complex evolutionary scenarios. It was suggested that gene replacements and horizontal gene transfers might explain the entangled phylogenetic relationships between viroplages, large DNA viruses, and certain classes of mobile elements related to viroplages such as polintons (12). An elegant example is the ORFs encoding a superfamily 3 helicase domain that exists in all fully sequenced viroplages (core gene) but fails to support monophyly of the latter, revealing multiple origins of the viroplage genes (Fig. S2D). Nevertheless, the corresponding *B. natans* proteins cluster with the YSLV3 homolog, suggesting that

Table 1. Virophage-like genes

ORF no.	Putative function	RPKM	Percentile expression,* %	Best hit [†]
Scaffold 2 largest elements				
Bn2_1	Orphan protein	0.2	3.9	
Bn2_2	Kelch and kinase domain protein	16.7	74.5	<i>Dendroctonus ponderosae</i> 478256302 (2e-32)
Bn2_3	Orphan protein	6.3	55.1	
Bn2_4	Unknown virophage protein	1.8	27.5	Zamilon 563399747 (2e-04)
Bn2_5	Adhesin-like protein	1.0	19.0	<i>Escherichia coli</i> 693111543 (9e-13)
Bn2_7	Orphan protein	0.9	16.7	
Bn2_8	Orphan protein	0.0	0.0	
Bn2_9	DNA polymerase B	0.9	17.2	Mavirus 326439151 (2e-09)
Bn2_10	Orphan protein	0.5	10.4	
Bn2_11	S3 helicase	0.8	15.1	YSL5 701905635 (2e-28)
Bn2_13	Orphan protein	604.1	99.2	
Bn2_14	Orphan protein	125.7	95.6	
Bn2_15	Orphan protein	194.4	97.2	
Bn2_17	Orphan protein	21.8	78.7	
Bn2_18	DnaJ domain protein	24.4	80.4	<i>Ostreococcus lucimarinus</i> virus 313843979 (2e-39)
Bn2_19	DNA-packaging ATPase	14.2	71.8	OLV 322510450 (7e-25)
Bn2_20	Orphan protein	9.4	63.8	
Bn2_21	Cysteine protease	1.5	24.5	OLV 322510453 (5e-08)
Bn2_22	Lipase	5.1	50.5	Mavirus 326439161 (6e-04)
Bn2_23	Orphan protein	4.2	46.1	
Bn2_25	Minor capsid protein	4.2	45.8	OLV 322510454 (2e-11)
Bn2_26	Major capsid protein	3.9	44.0	OLV 322510455 (3e-16)
Bn2_27	Unknown protein	1.4	24.0	<i>Guillardia theta</i> 551643434 (4e-16)
Bn2_28	Ribonucleotide reductase small subunit	1.6	25.9	<i>Cafeteria roenbergensis</i> virus 310831442 (3e-80)
Bn2_29	rve integrase	2.6	35.3	<i>Dictyostelium fasciculatum</i> 470248944 (3e-39)
Bn2_31	Kelch domain protein	33.4	84.5	<i>Strongylocentrotus purpuratus</i> 390342441 (6e-35)
Bn2_33	Orphan protein	0.2	5.2	
Other remarkable ORFs found in smaller elements				
Bn119_7	ZnR and GIY-YIG domains	3.6	42	<i>Phytophthora sojae</i> 695382398 (3e-05)
Bn161_7	Ankyrin domain protein	93.4	94	<i>Amoebophilus asiaticus</i> 501449850 (3e-34)
Bn92_2	Unknown phage protein	0.0	1	<i>Synechococcus</i> phage 472343273 (1e-05)
Bn92_3	Ankyrin domain protein	0.1	4	<i>Pseudogymnoascus pannorum</i> 682412062 (3e-08)
Bn187_9	Unknown phage protein	20.4	78	<i>Vibrio</i> phage 510792797 (5e-06)
Bn84_33	Unknown virophage protein	3.0	39	YSLV5 701905611 (2e-07)

*Percentile rank calculated over all *B. natans* genes.

[†]Species name and GenBank identifier (BLAST *E* value).

they originate from a common virophage ancestor. *B. natans* genes encoding rve family integrase (Fig. S2E), an unknown protein family represented by Bn2_27 (Fig. S2F), DNA polymerase (Fig. S2G), and GIY-YIG nuclease domains (Fig. S2H) exhibit preferential phylogenetic affinities, albeit with moderate bootstrap support with eukaryotic homologs encoded by polinton-related elements. These mixed clades are nested within larger clades containing virophages and environmental sequences, suggesting that gene acquisitions or replacements most likely occurred in the polinton-like elements. In contrast, the phylogenetic trees of the unknown protein family represented by Bn2_18 (Fig. S2I), the ribonucleotide reductase small subunit (Fig. S2J), and the Kelch protein family (i.e., PK and Kelch domains have distinct origins; Fig. S2A and B) support scenarios of gene acquisition from different

sources (bacteria, eukaryotes, or dsDNA viruses). Thus, the VLE genes reveal a mosaic origin that is typical of bona fide virophages (1, 7, 8). Altogether, the structure, gene content, and phylogenetic affinities of VLEs provide substantial evidence that they represent remains of integrated virophage genomes.

VLE Genes Are Transcribed. To investigate the transcriptional activity of the *B. natans* virophage-like elements, we analyzed a previously published RNA-seq dataset generated from cultivated *B. natans* cells (20). A total of 45.3 million Illumina paired-end reads were aligned onto the *B. natans* genome assembly (Dataset S2), of which 116,671 mapped within one of the virophage-like regions (Dataset S1). Two hundred seventy-eight out of the 302 predicted virophage-like ORFs had at least one read mapped to

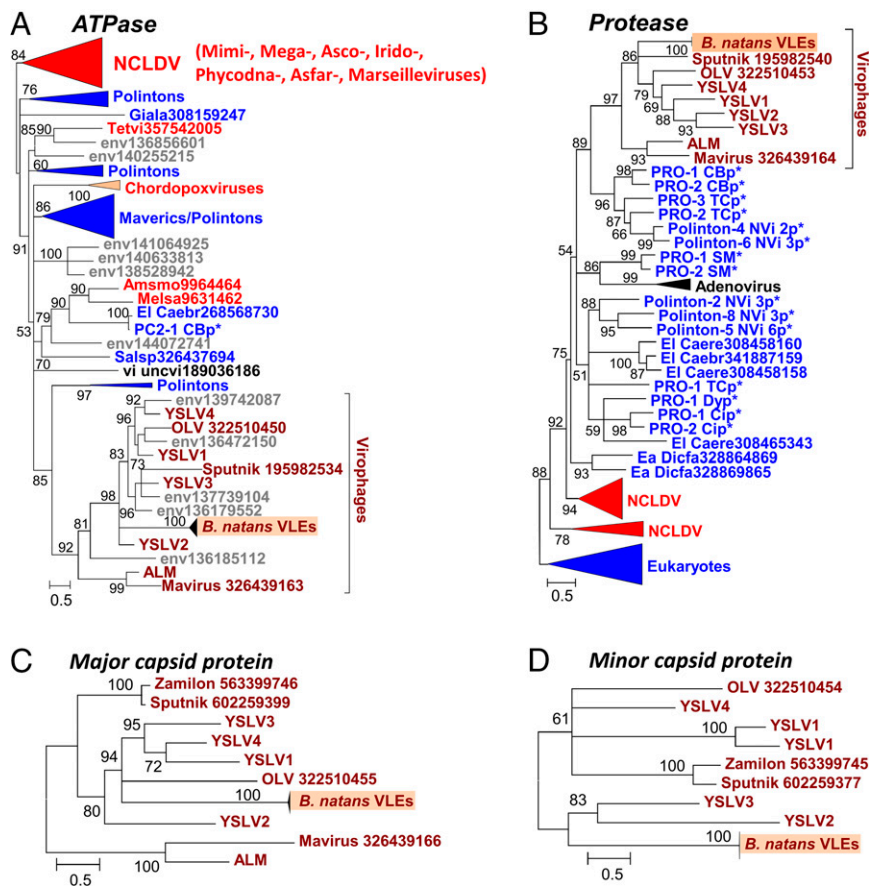


Fig. 3. Maximum-likelihood phylogenetic trees of conserved virophage proteins. (A) Packaging ATPase. (B) Maturation protease. (C) Major capsid protein. (D) Minor capsid protein. Branches with bootstrap support (expected-likelihood weights) less than 50% were collapsed. Sequences marked with an asterisk were taken from Repbase (38). For other sequences, the species name abbreviation and GenBank accession number are indicated; env, marine metagenome. Species: Amsmo, *Amsacta moorei* entomopoxvirus "L"; Caer, *Caenorhabditis brenneri*; Caere, *Caenorhabditis remanei*; Dicfa, *Dictyostelium fasciculatum*; Giala, *Giardia lamblia*; Melsa, *Melanoplus sanguinipes* entomopoxvirus; Salsp, *Salpingoeca rosetta*; Tetvi, *Tetraselmis viridis* virus; uncv, uncultured virus. Taxa: Ea, Amoebozoa; El, Opisthokonta; u2, Entomopoxvirinae. Dark red, virophages; blue, (predicted) polintons and related elements; light red, NCLDV; gray, unassigned environmental sequences. The numbers of validated amino acid positions in cleaned alignments are 210 (A), 166 (B), 548 (C), and 408 (D).

it. The levels of transcription between *B. natans* genes were compared by the mean of the RPKM metric (reads per kilobase per million mapped reads). Ninety-three virophage-like ORFs had RPKM values >5 , which ranks them in the top half of the most-transcribed genes in *B. natans*, including 10 genes that figure in the top 10% (i.e., RPKM >50). Interestingly, these highly transcribed virophage-like ORFs encode one major and one minor capsid protein, one DNA-packaging ATPase, one *re* integrase, two Kelch domain proteins, and three families of orphan proteins. Other virophage core genes are generally transcribed at low to moderate levels (i.e., the majority of them have RPKM values ranking between the 20th and 36th percentiles), yet 6 of the 10 ATPase gene copies have substantial transcription levels, as indicated by RPKM values ranking between the 50th and 90th percentiles. Interestingly, we identified transcript sequences closely related to the VLEs in assembled RNA-seq datasets generated from various *B. natans* isolates that were sequenced as part of the Marine Microbial Eukaryote Transcriptome Sequencing Project (MMETSP) (21) (Dataset S3). This suggests that the same virophage elements are present and transcribed in other *B. natans* strains, possibly because they were inherited vertically from a common ancestor.

This observation questions the biological significance of the expression of virophage-like genes. One possibility is that the integrations of virophage genomes were accidental events, and

that the residual transcriptional activity reflects the fortuitous recognition of regulatory signals of the virophage genes by the cellular host transcription complex. According to this scenario, the expression of the virophage genes has no expected biological effect (neutral) and would disappear as the virophage sequences decay by accumulating random mutations. Alternatively, the observed transcriptional activity might reflect an adaptive strategy that benefits the cellular host population, the virophage, or both. Experimental evidence indicates that virophages have a positive effect on the host-cell population. Mavirus interferes with *Cafeteria roenbergensis* virus propagation and increases the survival of the host-cell population (8). Sputnik causes a 70% decrease in infective Mavirus particles and a threefold decrease in amoeba cell lysis (1); it also delays or abolishes replication of Marseillevirus (22). A model of population dynamics suggests that the presence of virophages reduces overall mortality of the host algal cell after a bloom (7). Hence, eukaryotes that are susceptible to infection by giant viruses will gain a selective advantage if they can stably associate themselves with virophages (8). An analogous hypothesis was briefly exposed by Katzourakis and Aswad (23) to explain the possible emergence of Maverick/polinton elements from hypothetical integrated viro-phage genomes in eukaryotes. Under this scenario, the hijacked viro-phage genes evolve under negative selection in the new eukaryotic genome environment.

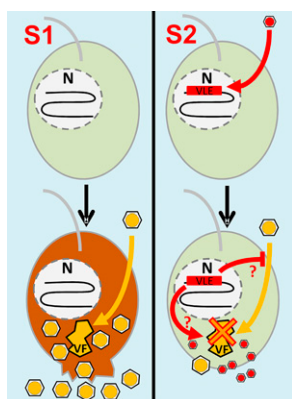


Fig. 4. Scenarios of infection of *B. natans* cells by NCLDVs and virophages. Scenario (S)1 starts with *B. natans* cells devoid of VLE insertions. An NCLDV particle (yellow hexagon) or its DNA enters the cell and establishes viral factories (VFs) that produce new NCLDV particles. The infection causes cell lysis and death accompanied by release of NCLDV particles. Scenario 2 begins with *B. natans* cells carrying VLEs in the form of functional provirophages integrated in the nuclear genome (N). VLEs may have been produced by independent entry of a virophage followed by active DNA integration in the host nuclear genome (delayed-entry mode). Upon NCLDV infection, expressed VLE proteins may inhibit virus penetration or trigger reactivation and excision of the provirophage, which in turn inhibits NCLDV replication and takes advantage of the viral factories for its own replication. As a result, a limited number of NCLDV particles are created compared with S1, leading to increased rates of cell survival. Potentially, new virophages and a limited number of NCLDV particles are released in the environment through exocytosis or another unknown mechanism that does not kill the host cell.

Another possible explanation, which is not mutually exclusive with the former, relates to the adaptive strategy of virophages to increase the frequency of coinfection with a host virus. Different modes of virophage entry into the cell have been evaluated on theoretical grounds (18). However, these scenarios exclusively consider the simultaneous entry of the virophage and the host virus, either independently or in a paired mode [i.e., the virophage adheres to the virus before infection; see Taylor et al. (18)]. Indirect evidence supports the paired-entry mode for Sputnik (1, 22), whereas an independent-entry mode has been observed for Mavirus (8). The transcribed *B. natans* virophage elements bring out a third hypothetical mode that we have dubbed the delayed-entry mode (Fig. 4). According to this scenario, the virophage is expected to enter the cell first by an unknown mechanism; its DNA reaches the cell nucleus, integrates into the cellular genome, and remains latent until superinfection by a virus reactivates and rescues the virophage, allowing its replication in the virus factory. This hypothetical scenario gets support from the observation that the *B. natans* virophage elements encode integrases, which suggests that genome integration is an active process rather than an incidental event. To go further into the delayed-entry scenario, it is possible that the transcription of virophage genes leads to the production of sentinel proteins that are able to detect infection of the host cell by another virus and transduce a signal triggering virophage reactivation. There is an obvious advantage of the delayed-entry mode over the independent-entry mode when the simultaneous independent entry of both the virus and virophage is a rare event due to, for example, high dilution of the virus particles in the environment. Furthermore, the integrated virophage can be passed on to the next generations of host cells, contributing to its spread and multiplication. In the form of an integrated provirophage, the virophage can potentially wait for virus superinfection during long periods of time, whose length depends on the rate at which random mutations inactivate the endogenous virophage element.

Putative NCLDV Insertions. All characterized virophages so far have been shown to infect members of the *Mimiviridae* (NCLDVs). Thus, although no large DNA virus of *B. natans* has been identified so far, the discovery of virophage sequences in *B. natans* suggests that this alga is the prey of giant viruses that are themselves infected by virophages. DNA fragments that are relics of integrated NCLDV genomes have been discovered in various eukaryotic genomes (24–26), revealing footprints of past interaction between viruses and hosts (19). We therefore searched the *B. natans* genome for sequences related to NCLDVs using the BLAST score plot approach. As shown in Fig. 1, a total of 36 *B. natans* predicted proteins have better alignment scores with homologs in NCLDVs than in cellular organisms (Dataset S4). The closer relationship of these proteins with giant virus homologs was confirmed by phylogenetic reconstruction (for examples, see Fig. S3). This phylogenetic affinity may reflect horizontal gene transfer between viruses and eukaryotic hosts, the polarity of which (virus to host or host to virus) cannot always be determined with certainty. However, some proteins are homologous to typical NCLDV core genes, including major capsid protein and packaging ATPase (distantly related to the virophage ones), which have most likely been acquired by the *B. natans* host (Fig. S3A and B). Interestingly, some NCLDV-like genes are physically clustered in discrete regions of the *B. natans* genome assembly (Fig. S4). For example, we identified a 165-kb region of scaffold 10 (position ~340–505 kb) that is transcriptionally silent according to our analysis of the RNA-seq dataset (Dataset S4). Such a large untranscribed region is unique in the *B. natans* genome. This region contains 83 predicted genes, of which 9 have obvious NCLDV affinities, including genes for DNA-packaging ATPase, exonuclease, major capsid protein, and transcription factor. Of the remaining genes, only 7 have best matches in eukaryotes (albeit with low similarity) and 4 have best matches with homologs in bacteria and phages. The large majority of predicted genes ($63/83 = 76\%$) are orphans, a characteristic shared with giant virus genomes. Thus, it is likely that this large DNA stretch is the remnant of an integrated NCLDV genome similar to those previously observed in various eukaryotic genomes (19, 24–27). Inserts of likely NCLDV origin identified in the moss genome were also reported as transcriptionally silent (19). These hypothesized viral inserts probably behave like neutrally evolving nonfunctional DNA. The G+C content of the NCLDV-like region (45.5%) is similar to the background G+C content of the *B. natans* genome (44.8%), which prevents precisely identifying the insert boundaries.

Outside of the large NCLDV-like insert, some *B. natans* genes with viral phylogenetic affinity were found to be transcribed (Dataset S4). Six genes have transcription levels among the top 50% of *B. natans* genes. These genes may have been inherited from the *B. natans* ancestor and captured by large DNA viruses from eukaryotic hosts that are closely related to *B. natans*. This scenario can explain the preferential affinity between *B. natans* proteins and NCLDV homologs, whereas the corresponding genes actually have a eukaryotic origin. Alternatively, the corresponding genes may have been recruited in the metabolism of a Bigelowiella ancestor subsequent to their acquisition from viral donors, perhaps now fulfilling new cellular functions. For example, *B. natans* has one copy for each of the four regular B family eukaryotic DNA polymerase catalytic subunits alpha, delta, epsilon, and zeta. However, the alga possesses two extra delta DNA polymerases that group in phylogenetic positions compatible with distinct viral origins (Fig. S3C). The two viral-like DNA polymerase catalytic subunits are transcribed (i.e., 43th and 53th percentiles) at higher levels than those of the four regular eukaryotic isoforms (25th–40th percentiles, respectively). Furthermore, they have a large number of introns (i.e., 19 and 30 introns, respectively), whereas viral genes are generally devoid of introns or, in rare cases, only contain a small number of them.

The two extra DNA polymerases have no identifiable orthologs in other sequenced eukaryotes, except in the transcriptomes of other *Bigelowiella* isolates (Dataset S4). We can therefore not exclude a scenario in which the corresponding genes have been captured from viral donors and were progressively “eukaryogenized” through the accumulation of introns.

Putative Transposon Insertions. Four of the six NCLDV-like DNA-packaging ATPase genes are carried by four closely related repeated genetic elements (18.8–21.8 kb in length) flanked by TIRs 300–900 bp in length (Fig. 5). These repeated elements contain between 17 and 22 ORFs (Dataset S5), but some of the original genes seem to have accumulated internal stop codons and frameshifts, resulting in truncated translation. Very few proteins encoded by the repeated elements exhibit detectable similarity in public databases. They include a homolog of the *Aureococcus anophagefferens* virus protein AaV202, a homolog of the *P. globosa* virus virophage protein PgvV_00016, a homolog of the Mavirus virophage protein MV06, and a homolog of a hypothetical protein of the fungus *Batrachochytrium dendrobatidis*. Interestingly, two additional proteins are homologous to core transposon proteins, including the C-terminal superfamily I helicase domain protein and C2H2 Zn-finger protein. Transposons form a distinct class of mobile genetic elements (6.5–7.5 kb) associated with *Mimiviridae* (4). The genes for helicase and C2H2 protein are adjacent in the *B. natans* repeated elements and Mimivirus-associated transposons. Furthermore, Mimivirus-associated transposons are flanked by TIRs (~530 bp). Thus, the *B. natans* repeated elements and transposons share structural and genetic similarities, suggesting that the *B. natans* elements might belong to the transposon family. However, the putative integrated *B. natans* transposons are substantially bigger in size relative to their Mimivirus counterparts, possibly due to the incorporation of foreign genes of diverse origins, including NCLDVs and virophages. Some of the integrated transposon genes show evidence of transcription, including six genes that have transcription levels in the *B. natans* top 50%. In addition, homologous transcripts were identified in the transcriptome data of the other *Bigelowiella* isolates (Dataset S6).

Conclusion

The first discovered virophages have the form of small icosahedral virion particles and have been shown to infect giant viruses. However, integrated virophages have been found more recently in a Mimivirus genome (4). Here we show for the first time, to our knowledge, that virophage genomes can also integrate in a eukaryotic genome. This finding led us to predict that *B. natans* might be the prey for NCLDVs. Additional integrated DNA fragments that most probably originate from NCLDV genomes provide data showing that *B. natans* or its recent ancestor had physical contacts with NCLDV members. Furthermore, we also identified repeated genetic elements that resemble transposons associated with *Mimivirus*. Thus, the *B. natans* genome appears to have recorded genetic footprints of molecular “battles” between virophages, transposons, and giant viruses. *B. natans* belongs to the Chlorarachniophytes, a group of unicellular marine algae that acquired a plastid by secondary endosymbiosis involving engulfment of a green alga by a eukaryotic heterotroph host (28). They are typically mixotrophic, ingesting bacteria and smaller protists as well as conducting photosynthesis. *B. natans* is the only species of the supergroup Rhizaria for which a complete genome sequence is available (29). During endosymbiosis, hundreds of genes of green origin have been transferred toward the host genome in a process called endosymbiotic gene transfer (29). The acquisition of DNA from giant viruses and transposons as well as from virophages through horizontal gene transfer represents an additional component in the melting pot of genes composing the *B. natans* nuclear genome.

One of the most intriguing findings of our analysis is that integrated virophage genes are highly transcribed, suggesting that they are biologically functional. We speculated on three potential adaptive scenarios to explain this observation. Certainly the most interesting of them is the possibility that both the cellular host and virophage take advantage of the integration strategy, by providing the cell with a defense mechanism against giant viruses and providing the virophage with a mechanism to increase the rate of coinfection with a viral prey (i.e., delayed-entry mode S2, schematized in Fig. 4). From the perspective of the cellular host, it is tempting to speculate that integrated virophages can act as

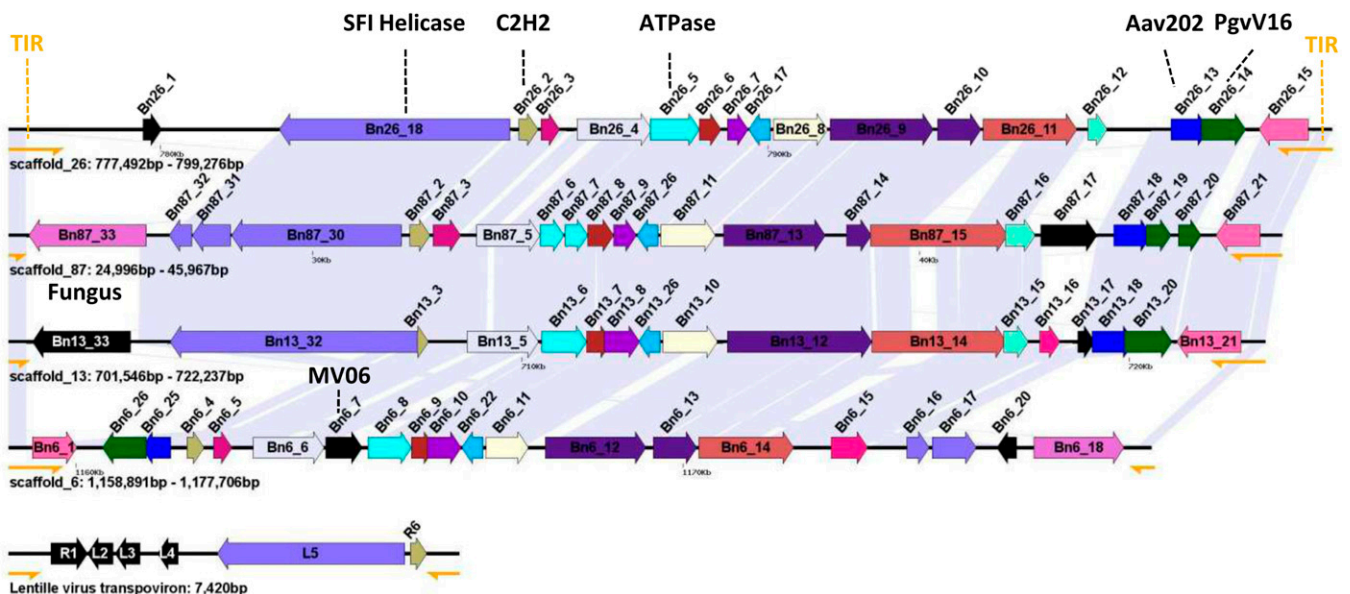


Fig. 5. Organization of putative inserted transposon genomes. The position and strand orientation of each ORF (>90 codons) are indicated by an arrow. Homologous ORFs across transposon-like elements are indicated with the same color. ORFs with no homologs in the other transposons are shown in black. TIRs are represented by yellow semiarrows. Shaded areas between elements indicate regions of high nucleotide similarity.

molecular weapons against viral pathogens, conferring a sort of immunity transmissible from generation to generation. Sputnik and Zamilon strains have been shown to have a broad host spectrum among the *Mimiviridae* (5, 16), and Sputnik even affects the replication of Marseillevirus, which is distantly related to the *Mimiviridae* (22). By extrapolation, we can further hypothesize that the putative defense function triggered by integrated virophages can be efficient over a diversity of viral pathogens.

It is currently difficult to estimate the prevalence and biological significance of virophage insertion in eukaryotes. At first glance, the fact that only *B. natans* showed virophage genes out of 1,153 screened eukaryotic genomes might suggest that genomic integration of virophages is highly unusual. However, there is a historical bias among sequenced eukaryotes, which include a majority of model organisms, crop plants, fungi, animals, and pathogens (30), all of which are apparently not infected by giant viruses and, hence, virophages. As a result, the organisms tested here included relatively few potential hosts of giant viruses and virophages [i.e., known hosts are amoebas and microalgae (1, 5, 8); Dataset S6]. This bias could explain the apparent low prevalence of virophage insertions detected in our study. In contrast, giant viruses and virophages are readily identified in environmental metagenomes (7, 10, 11, 31), suggesting that these viral entities are common in natural ecosystems. Thus, the question of the prevalence and biological significance of virophage insertions can only be reasonably addressed when more genomes of putative hosts are sequenced.

Methods

Sequence Analysis. Annotations and sequences of 1,153 eukaryotic genome assemblies representing various protists, fungi, and basal metazoans were downloaded from GenBank. Dataset S6 lists the investigated species together with their GenBank accession numbers, including members of the Alveolate (103), Amoebozoa (35), Apusozoa (1), Chlorophyta (12), Choanoflagellida (2), Cryptophyta (1), Euglenozoa (49), Fonticula (1), Fornicata (6), Fungi (858), Haptophyta (1), Heterolobosea (2), Parabassalia (1), Metazoa (8), Opisthokonta (2), Rhizaria (3), Rhodophyta (4), Stramenopiles (63), and Streptophyta (2). We also downloaded the genome annotations of nine sequenced virophages, including Mavirus (GenBank accession no. GCF_000890715), Sputnik (GenBank accession no. EU606015), Zamilon (GenBank accession no. NC_022990), ALM (GenBank accession no. KC556923), OLV (GenBank accession no. HQ704801), YSLV1 (GenBank accession no. KC556924), YSLV2 (GenBank accession no. KC556925), YSLV3 (GenBank accession no. KC556926), and YSLV4 (GenBank accession no. KC556922).

For each virophage, the major capsid protein, minor capsid protein, DNA-packaging ATPase, and cysteine protease were aligned against the predicted eukaryotic proteomes using BLASTP. Experience showed that computational annotation of eukaryotic genomes can be inefficient in predicting genes of viral origin, because they have become pseudogenes, often resulting in truncation or in-frame stop codons and/or because they can have very distinct GC content relative to the host genome and no introns. Therefore, we also aligned the virophage markers against the translated products of ORFs (>90 codons) lying between predicted genes. Based on prior analysis of the distribution of BLASTP scores against the nonredundant (NR) database, we applied family-specific score thresholds to avoid false detection of remote homologs that are of cellular origin or nonhomologs with similar low-complexity sequences. The score threshold was set as the minimal BLASTP score between any two virophage proteins in the family. Scores obtained between any virophage proteins and cellular homologs were always lower. These score thresholds were 44.7 for proteases, 46.6 for ATPases, 41.2 for mCPs, and 45.1 for MCPs.

The genome assembly of *B. natans* CCMP2755 exhibited homologs for each of the four marker proteins with BLASTP *E* value >1E-5, except proteases, for which we used an *E*-value threshold of 1E-3, because this protein

is less conserved among virophages. To identify additional candidate viral-like protein genes in *B. natans*, we performed a BLAST-score plot analysis previously described in Maumus et al. (19). Briefly, the full complement of *B. natans* predicted proteins together with intergenic ORFs was used to probe the National Center for Biotechnology Information database using BLASTP (*E* value < 1E-5). For each protein query, the alignment scores with the best cellular hit and the best virophage or NCLDV hit were recorded. When no cellular hit was recorded whereas a viral hit was obtained, the cellular score was set to zero. BLAST scores were then normalized by the score of the alignment of the query sequence against itself (i.e., self-score), resulting in relative scores expressed in percent of self-score. Nonviral hit scores are plotted against viral hit scores in Fig. 1.

Identification and Delineation of Individual VLEs. The physical location of the virophage-like protein genes identified by BLASTP revealed that they tend to cluster in specific loci in the genome assembly, unveiling large regions of possible virophage origin. Six of these regions were bordered by long inverted repeats on each side, which coincide with sharp changes in GC content (e.g., Fig. 2). We made the assumption that the long inverted repeats mark the beginning and the end of VLEs. We extracted the nucleotide sequences of these putative complete VLEs from the genome assembly and used BLASTN to align the VLEs back to the genome assembly to identify additional truncated VLEs. Adjacent BLASTN matches that had an *E* value <1E-25 and a minimal length of 100 pb were assembled to identify a total of 38 VLEs up to 33.3 kb in length. Every candidate VLE was checked and validated manually.

Phylogenetic Analysis. Construction of adequate homologous protein sets for phylogenetic analysis was performed using the BLAST-EXPLORER website (32). Homologous proteins were aligned using MUSCLE (33), and amino acid positions in multiple alignments containing >30% gaps were removed. We used this criterion for alignment cleaning to keep coherence with the pioneering study of Yutin et al. (12), which produced a comprehensive phylogenetic study of virophage genes. Maximum-likelihood (ML) phylogenetic reconstruction was performed using the PhyML program (34). Before phylogenetic reconstruction, the best-fitting substitution model for each sequence dataset was determined using the ProtTest program (35). Sequences, alignments, ProtTest outputs, and phylogenetic trees are available in Dataset S7.

RNA-Seq Analysis. To analyze the transcriptional activity of the *B. natans* genome, we downloaded an RNA-seq dataset (ID MMETSP0045) from the CAMERA database (36). This dataset contained 61.6 million Illumina paired reads generated from polyadenylated RNA extracted from *B. natans* CCMP2755 cells grown in f/2-Si media for a month under a 12-h:12-h light:dark cycle at room temperature (20). Reads were aligned onto the reference genome sequence using Bowtie 2 (37) with default parameters. Due to the high nucleotide similarity between VLEs, 63% of the reads that mapped onto a VLE also had valid alignments in at least another VLE. For these cases the origin of the read is ambiguous, and we only picked one of the alignments at random for read-count purposes. In addition, we downloaded assembled RNA-seq datasets (contigs) of other *Bigelowiella* isolates publicly available in the CAMERA database: MMETSP1052 (*B. natans* CCMP623), MMETSP1054 (*B. natans* CCMP1259), MMETSP1055 (*B. natans* CCMP1258.1), MMETSP1358 (*B. natans* CCMP1242), and MMETSP1359 (*Bigelowiella longifila* CCMP242). These datasets were generated as part of the Marine Microbial Eukaryote Transcriptome Sequencing Project (21). Homologs of virophage-like encoded proteins were searched in the transcribed sequences using TBLASTN.

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