

# Giant virus with a remarkable complement of genes infects marine zooplankton

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As major consumers of heterotrophic bacteria and phytoplankton, microzooplankton are a critical link in aquatic foodwebs. Here, we show that a major marine microflagellate grazer is infected by a giant virus, *Cafeteria roenbergensis virus* (CroV), which has the largest genome of any described marine virus ( $\approx 730$  kb of double-stranded DNA). The central 618-kb coding part of this AT-rich genome contains 544 predicted protein-coding genes; putative early and late promoter motifs have been detected and assigned to 191 and 72 of them, respectively, and at least 274 genes were expressed during infection. The diverse coding potential of CroV includes predicted translation factors, DNA repair enzymes such as DNA mismatch repair protein MutS and two photolyases, multiple ubiquitin pathway components, four intein elements, and 22 tRNAs. Many genes including isoleucyl-tRNA synthetase, eIF-2 $\gamma$ , and an Elp3-like histone acetyltransferase are usually not found in viruses. We also discovered a 38-kb genomic region of putative bacterial origin, which encodes several predicted carbohydrate metabolizing enzymes, including an entire pathway for the biosynthesis of 3-deoxy-D-manno-octulosonate, a key component of the outer membrane in Gram-negative bacteria. Phylogenetic analysis indicates that CroV is a nucleocytoplasmic large DNA virus, with *Acanthamoeba polyphaga* mimivirus as its closest relative, although less than one-third of the genes of CroV have homologs in Mimivirus. CroV is a highly complex marine virus and the only virus studied in genetic detail that infects one of the major groups of predators in the oceans.

nucleocytoplasmic large DNA virus | horizontal gene transfer | viral evolution | DNA repair | 3-deoxy-D-manno-octulosonate

Predation by protistan grazers is a major pathway of carbon transfer and nutrient recycling in marine and freshwater systems (1); yet, viruses infecting phagotrophic protists in marine systems are largely unknown and completely unexplored genetically. The discovery of the giant *Acanthamoeba polyphaga* mimivirus in a freshwater amoeba, with its 1.2 million-base pair (bp) genome and 981 genes (2, 3), has sparked an intense debate about the biology and evolutionary origin of giant viruses. Whereas some researchers argue that giant viruses are “gene robbers” that have acquired their extensive gene collection by horizontal gene transfer (HGT) from cellular organisms (4–6), others favor the theory that these viruses date back to the emergence of eukaryotes and that most of their genes are viral in origin (7, 8). Recently, it has become evident that protists host the largest and most complex viruses known (9), that other giant viruses are likely widespread in oceans (10), and that some of these are pathogens of phytoplankton (11); yet, the only characterized giant viruses are those infecting species of *Acanthamoeba*. Ultimately, understanding the origin and evolution of giant viruses will be facilitated through the use of comparative genomics with other representative systems.

In this study, we used 454 pyrosequencing to sequence and de novo assemble the genome of a very large (300 nm capsid diameter) DNA virus, *Cafeteria roenbergensis virus* (CroV) strain BV-PW1, that was isolated from the coastal waters of Texas in the early 1990s (12). This lytic virus infects a marine heterotrophic flagellate, which is identical to *C. roenbergensis* strain VENT1 at the level of 18S rDNA. The host, which consumes bacteria and

viruses (13), was originally misidentified as *Bodo* sp. (12). It is a 2- $\mu\text{m}$ –to 6- $\mu\text{m}$ –long bicosoecid heterokont phagotrophic flagellate (Stramenopiles) that is widespread in marine environments and is found in various habitats such as surface waters, deep sea sediments, and hydrothermal vents (14, 15). Populations of *C. roenbergensis* may be regulated by viruses in nature (16).

## Results and Discussion

**General Genome Features.** The genome of CroV is a linear double-stranded DNA molecule with a size of  $\approx 730$  kb, making this the second largest described viral genome. We sequenced and assembled the 618-kb central part of the viral chromosome, which is flanked on both ends by large and highly repetitive regions (Fig. 1). These terminal regions could potentially serve as protective caps for the protein-coding part of the genome, akin to telomeres in eukaryotes. The CroV genome is AT-rich (77% A+T), which is reflected in the distribution of codons and in the overall amino acid (aa) composition. AT-rich codons are consistently preferred over GC-rich ones, with the four most frequent aa (Lys, Ile, Asn, and Leu) each representing  $\approx 10\%$  of the overall aa (SI Appendix, Fig. S1).

Using conservative annotation criteria (SI Appendix), we identified 544 putative protein-coding sequences (CDSs) in the 618-kb central region of the CroV genome, which had a coding density of 90.1%. The average CDS was 1,025 nucleotides (nt) in length, and coding capacities ranged from 47 to 3,337 aa. Applying a BLASTP E-value cutoff of  $1e-05$ , 267 CDSs (49%) displayed similarity to sequences in GenBank and 134 CDSs (25%) could be assigned to one or more Clusters of Orthologous Groups of proteins (COGs,  $E < 0.001$ ) (SI Appendix, Fig. S2). CroV CDSs and their annotations are listed in Dataset S1. Based on the distribution of top BLASTP hits, approximately one-half of the CroV genes displayed similarities to proteins found in eukaryotes, bacteria, archaea, and other giant viruses (SI Appendix, Fig. S3). Twenty-two percent of CroV CDSs had their top BLASTP hit among eukaryotes, but in the absence of genomic information about *C. roenbergensis*, no statement can be made about potential gene transfer between CroV and its host. Although most CroV CDSs were of unknown function, 32% of CDSs could be assigned a putative function and they provide insights into the biology of this giant virus. Several of these enzymatic functions have not been reported to be encoded by any other virus (SI Appendix, Table S1).

**Translation Genes.** Viruses rely primarily on the protein translation apparatus of their hosts; it is therefore unusual to find viral genes

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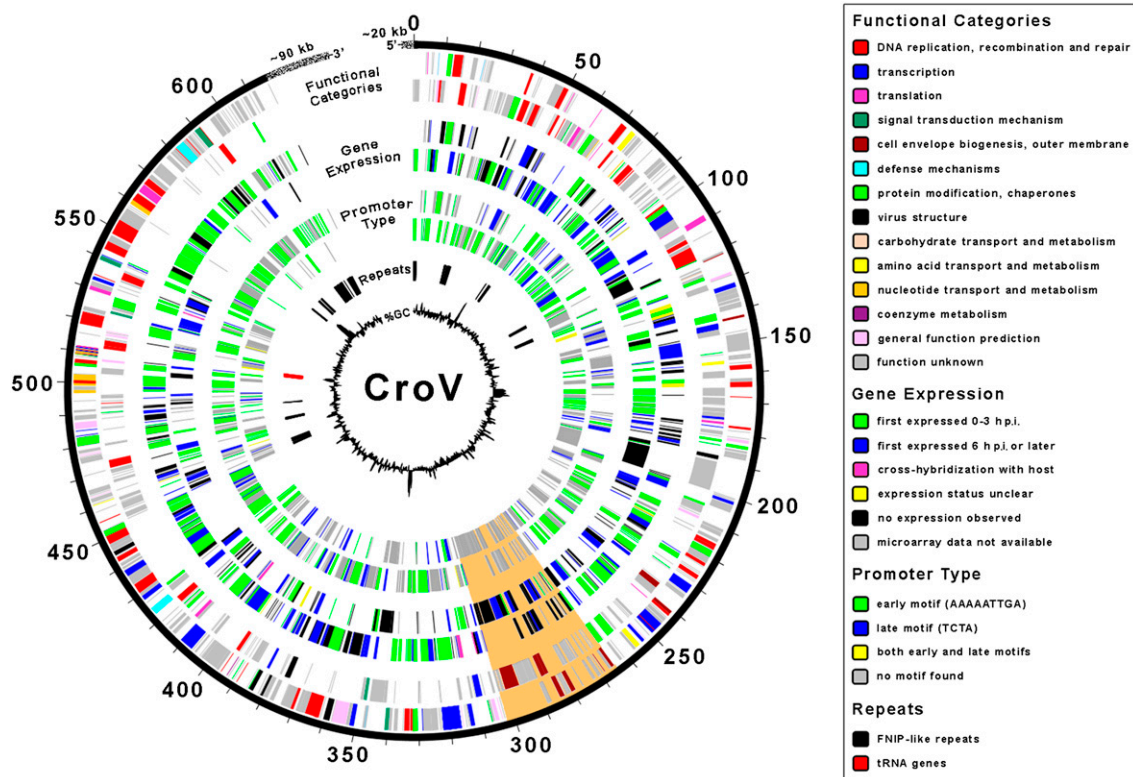
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Data deposition: The sequences reported in this paper have been deposited in the GenBank database [accession nos. GU244497 (CroV genome) and GU249597 (partial 18S sequence from *C. roenbergensis* strain E4-10)]. The microarray data reported in this paper have been deposited in the Gene Expression Omnibus (GEO) database, [www.ncbi.nlm.nih.gov/geo](http://www.ncbi.nlm.nih.gov/geo) (accession no. GSE19051).

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**Fig. 1.** Genome diagram of CroV. Genome coordinates are given in kbs. Nested circles from outermost to innermost correspond to (i) predicted CDSs on forward strand and (ii) reverse strand; (iii) expression data for CDSs on forward strand and (iv) reverse strand; (v) gene promoter type for CDSs on forward strand and (vi) reverse strand; (vii) location of repetitive DNA elements; (viii) GC content plotted relative to the genomic mean of 23.35% G+C. The speckled regions at the chromosome ends are not drawn to scale and indicate terminal repeats for which no sequence information is available. A 38-kb genomic segment of putative bacterial origin is shaded orange.

associated with protein synthesis. CroV encodes an isoleucyl-tRNA synthetase and putative homologs of eukaryotic translation initiation factors eIF-1, eIF-2 $\alpha$ , eIF-2 $\beta$ /eIF-5, eIF-2 $\gamma$ , eIF-4AIII, eIF-4E, and eIF-5B. Using the transfer RNA gene prediction software tRNAScan-SE, we identified 22 tRNA genes, clustered in a 2.8-kb region around position 510,000 (Fig. 1 and *SI Appendix, Table S2*). We also found two putative tRNA-modifying enzymes in CroV, tRNA pseudouridine 5S synthase and tRNA<sup>Leu</sup> lysidine synthetase. These genes add to a rapidly growing number of virus-encoded protein translation components. Some tRNA genes are scattered among bacteriophages and eukaryotic viruses such as the phycodnaviruses (17, 18), and four tRNA synthetases along with several putative translation factors are found in Mimivirus (2). These findings imply that CroV and similarly complex viruses encode genes to modify and regulate the host translation system to their own advantage, which results in a “lifestyle” that is less dependent on host cell components than that of smaller viruses.

**DNA Repair Genes.** The ability to repair various kinds of DNA damage is well documented among large DNA viruses (19, 20). Given that the AT-rich genome of CroV is exposed to high solar irradiance in surface waters of the ocean and is therefore likely to suffer from DNA lesions such as pyrimidine dimers, it is not surprising that CroV encodes multiple DNA repair proteins. We found putative components of several DNA repair mechanisms, including a presumably complete base excision repair pathway with formamidopyrimidine DNA glycosylase, a family 1 apurinic/aprimidinic (AP) endonuclease, a family X DNA polymerase, and an NAD-dependent DNA ligase. Further DNA repair proteins include DNA mismatch repair protein MutS, XPG endonuclease, a homolog of the alkylated DNA repair protein AlkB, and two DNA photolyases. Photolyases are classified into three

major groups: Cyclobutane pyrimidine dimer (CPD) photolyases, (6-4) photolyases, and single-stranded DNA photolyases (ref. 21 and references therein). The CPD photolyases are further subdivided into class I and class II enzymes, the former being more prevalent in bacteria and the latter more frequent in eukaryotes. The gene product of *croV115* is a predicted CPD class I photolyase and represents the first viral homolog in this class (*SI Appendix, Figs. S4 and S5*). The second CroV photolyase (*croV149*) does not belong to any of the established types of photolyases. Instead, it is related to a recently described group of photolyases/ cryptochromes that are present in several bacterial phyla and the euryarchaeotes (21) (*SI Appendix, Figs. S4 and S6*). The only eukaryotic member in this group (*Paramecium tetraurelia*) is also the closest homolog to the CroV and Mimivirus sequences and may have acquired this gene by HGT from a giant virus (*SI Appendix, Fig. S6*).

**Transcription Genes.** Large DNA viruses typically carry hundreds of genes, including several that regulate gene expression. Among the predicted transcriptional genes in CroV are eight DNA-dependent RNA polymerase II subunits, at least six transcription factors involved in transcription initiation, elongation, and termination, a tri-functional mRNA capping enzyme, a poly(A) polymerase, and several helicases. The complex transcriptional machinery encoded by CroV suggests that viral gene transcription does not depend on host enzymes and likely occurs in the cytoplasm. Interestingly, CroV contains a CDS with high similarity to an ELP3-like histone acetyltransferase (HAT, COG1243, 2e-46), a gene previously not seen in viruses. In combination with other unidentified viral gene products, the CroV HAT may enable the virus to directly modulate the genome condensation state of the host and, thus, exert control over its transcriptional activity. Alternatively, this enzyme may be involved



in replication and packaging of the virus genome itself. Another unusual characteristic of CroV is the presence of three DNA topoisomerase (Topo) genes of types IA, IB, and IIA. TopoIA and TopoIIA are very similar to their counterparts in Mimivirus, and HGT events from bacteria (eventually via a eukaryotic phagotrophic host) have been proposed for these genes (22). CroV TopoIB is the first viral homolog of the eukaryotic subfamily, whereas the TopoIB encoded by Mimivirus falls within the bacterial group (*SI Appendix, Fig. S7*) and is functionally more similar to the poxvirus enzymes (23). Despite apparently different evolutionary trajectories, the presence of three Topo genes in CroV and Mimivirus suggests a crucial role for these enzymes in transcription, replication, or packaging of giant virus genomes.

**Repetitive DNA and Ubiquitin Components.** Approximately 5% of the genome (excluding the terminal regions) consisted of repetitive elements. The most prevalent was a 22-aa-long leucine-rich repeat similar to the FNIP/IP22 repeat (Pfam entry PF05725) that had >400 copies in the CroV genome and was present in at least 28 CDSs (Fig. 1 and *Dataset S1*). This repeat also occurs in Mimivirus and *Dictyostelium discoideum* (24). Whereas leucine-rich repeats are known to mediate protein–protein interactions in a variety of proteins with diverse functions (25), the role of these repeats in CroV is unknown. In Mimivirus, FNIP/IP22 repeat-containing genes also possess an N-terminal F-box domain, which mediates interaction with the ubiquitin (Ub) pathway (26). Ub signaling appears to be a general strategy used by nucleocytoplasmic large DNA viruses (NCLDVs) to counter host defenses, because multiple Ub-conjugating and Ub-hydrolyzing enzymes have been found in these viruses (26). Furthermore, it has been shown that orthopoxvirus replication requires a functional Ub-proteasome system (27). In CroV, we identified a small arsenal of genes encoding proteins predicted to function in the Ub pathway, including an E1 Ub-activating enzyme, six E2 Ub-conjugating enzymes, two deubiquitinating enzymes, and one Ub gene. The specific means of how CroV and other giant viruses use Ub signaling to interact with their hosts remain to be determined.

**CroV Harbors Four Inteins.** No introns were detected in the genome, but four CDSs contained an intein, i.e., a self-splicing protein sequence inserted in highly conserved regions of a host protein (28). All four CroV inteins are part of NCLDV core genes that are thought to play a key role in DNA replication and transcription: DNA-dependent DNA polymerase B (PolB), TopoIIA, DNA-dependent RNA polymerase II subunit 2 (RPB2), and the large subunit of ribonucleotide reductase (RNR). Ten other inteins have been found in viruses infecting eukaryotes (29), including PolB inteins in Mimivirus (30), *Heterosigma akashiwo* virus (31), and *Chrysochromulina ericina* virus (32) as well as RNR inteins in four iridoviruses and the chlorella virus NY-2A (33). With the exception of a gene fragment from *Emiliana huxleyi* virus 163 (34), the CroV RPB2 intein constitutes the only viral report of an intein in RPB2. Finally, the CroV TopoIIA intein is a unique case of an intein in a DNA topoisomerase II gene, thus extending the known range of intein-containing genes. All four CroV inteins possess the conserved nucleophilic residues that are required for the standard splicing reaction [C/S at the N-terminal splice junction and N(C/S/T) at the C-terminal splice junction] (28) and are therefore probably capable of autocatalytic excision.

**Microarray Analysis.** A microarray experiment was undertaken to determine which CroV genes were unambiguously transcribed in infected cells and if there was a clear temporal pattern in the transcription of those genes. We detected viral transcripts in infected *C. roenbergensis* cells by fluorescently labeling mRNA isolated at different time points during the infection cycle, which lasted 12–18 h in *C. roenbergensis* strain E4-10. We then hybridized the labeled transcripts to glass slides spotted with oligonucleotide probes for 438 of the 544 predicted CroV genes (*SI Appendix*). Detectable levels of expression were found for 274 genes (63%), 152 genes (35%) were below the detection limit, 4

(1%) cross-hybridized with host mRNA isolated from uninfected cells, and 8 (2%) could not be assigned a clear on/off status (Fig. 1 and *Dataset S1*). Therefore, approximately one-half of the predicted genes and 63% of the genes we tested were expressed during infection under our laboratory conditions. This percentage is comparable with the observed expression of 65% of viral genes during infection of the marine phytoplankter *Emiliana huxleyi* by EhV-86 (35). However, recent gene expression studies in PBCV-1 and Mimivirus validated transcription for nearly all of their predicted genes (3, 36). It seems therefore likely that our microarray data underestimated the true extent of transcriptional activity in CroV. All of the previously mentioned translation-related genes in CroV, as well as most of the “virus-atypical” genes were expressed (*Dataset S1*), suggesting that these genes are functional. Although the microarray experiment was designed primarily to validate CroV gene predictions and cannot be exploited quantitatively, the data allowed us to recognize some general trends of CroV gene expression. Based on the time points at which transcripts were first detected, we could distinguish between an early and a late phase of CroV gene expression. The early phase lasted from 0 h to 3 h after infection (h p.i.) and affected 150 genes. The majority of DNA replication and transcription genes belonged to the early class. The late phase was characterized by genes that were first detected in the microarray at 6 h p.i. or later. The 124 genes in this class included all of the predicted structural components, such as the major and minor capsid proteins. Further and more extensive analysis of the CroV transcriptome may be able to refine this preliminary temporal classification.

**Promoter Analysis.** The intergenic regions had an average size of  $71 \pm 64$  bp. We examined the 100-nt region upstream of the predicted start codons for possible promoter motifs by using MEME software (37). A perfectly conserved “AAAAATTGA” motif, flanked by AT-rich sequences, was found to precede 127 CroV CDSs (23%) (Fig. 24). The MEME E-value for this motif was  $9e-170$ . Allowing one mismatch per sequence at the less strongly conserved positions one to six of the AAAAATTGA motif increased the number of positive CDSs to 191 (35%). The majority of CDSs that displayed this motif in their immediate upstream region belonged to the “early” temporal category (Fig. 24). We therefore classified this motif as an early gene promoter in CroV. Our results are in agreement with findings from Mimivirus, where a nearly identical early promoter motif (AAAATTGA) is associated with 45% of Mimivirus genes (3, 38). But, in contrast to Mimivirus, where the motif is found preferentially in the –50 to –110 region, the early promoter motif in CroV displayed a much narrower distribution, with a peak at position –40 relative to the predicted start codon (Fig. 2B).

We then searched for a possible late promoter motif starting with a representative set of six CDSs, all predicted to encode capsid components (*SI Appendix*). Five of the six genes exhibited the conserved tetramer “TCTA,” flanked by AT-rich regions on either side, in their –11 to –20 region (Fig. 2). Based on this profile, we expanded the search to all CroV CDSs and identified 72 that were positive for the TCTA motif signature. A MEME search on the 30-nt upstream region of the 124 genes classified as “late” yielded a very similar motif (MEME E-value  $5e-04$ ; *SI Appendix, Fig. S8*). As shown in Fig. 24, most CDSs with the TCTA promoter motif were first expressed at 6 h p.i. or later, supporting our conclusion that this sequence motif represents a promoter element for genes transcribed during the late phase of CroV infection. The CroV late promoter motif is unrelated to the putative late promoter motif identified in Mimivirus (3).

**A Thirty-Eight-Kilobase Genomic Fragment Involved in Carbohydrate Metabolism.** Upon examination of the CroV promoter distribution, we noticed that neither early nor late promoter motifs were associated with CDSs located between the genomic positions 264,800 and 302,500 (Fig. 1). Of these 34 CDSs (croV242–croV275), 14 were most similar to bacterial proteins (*SI Appendix, Table S3*; BLASTP E-value  $<1e-05$ ) and 7 of them are predicted to function in carbo-



may have been acquired from a bacterium, considering that CroV frequently encounters phagocytosed bacteria inside the host cytoplasm and encodes several enzymes that might catalyze an integration of foreign DNA (e.g., transposase *croV356*). Genomic islands of putative bacterial origin have been identified in other giant viruses such as phycodnaviruses and Mimivirus, but in contrast to CroV, these bacterial gene clusters tend to be located toward the ends of the linear viral chromosomes (40). However, given that the GC content of the 38-kb region is even lower than that of the rest of the CroV genome (19.4% vs. 23.6% G+C) and that some of these proteins occupy a phylogenetic position between bacterial and eukaryotic homologs (e.g., KDOPS and KDOPase; *SI Appendix, Figs. S10 and S11*), we cannot rule out alternative scenarios for the origin of this region.

**Phylogenetic Relationship.** Based on the presence and phylogenetic analysis of a set of core genes (*SI Appendix, Fig. S13*), CroV is an addition to the presumably monophyletic group of NCLDVs (2, 26, 41), which includes the families *Ascoviridae*, *Asfarviridae*, *Iridoviridae*, *Mimiviridae*, *Phycodnaviridae*, *Poxviridae*, and the newly discovered *Marseillevirus* (42). In a recent study by Yutin et al. (43), genes encoded by NCLDVs were categorized into groups that presumably evolved from a common ancestor and subsequently diversified in the various NCLDV families. Using this dataset of Nucleo-Cytoplasmic Virus Orthologous Genes (NCVOGs), we found that at least 172 CroV CDSs belonged to an existing NCVOG (*Dataset S1*). Thirty-two percent of CroV CDSs were significantly similar to a Mimivirus gene (any Mimivirus hit with a BLASTP E-value  $>1e-05$ ) and 22 CroV CDSs had their only detectable GenBank homolog in Mimivirus. CroV therefore appears to be the closest known relative to Mimivirus, despite large differences in genome (730 kb vs. 1,181 kb) and capsid size (300 nm vs. 500 nm). The CroV–Mimivirus relationship was further corroborated by phylogenetic analysis of PolB, a commonly used marker gene to infer phylogenetic relationships among NCLDVs. Bayesian Inference analysis of PolB resulted in a strongly supported clade comprising the largest known viruses: Mimivirus, CroV, and three partially sequenced viruses infecting the marine microalgae *Phaeocystis pouchetii* (PpV), *Chrysochromulina ericina* (CeV), and *Pyramimonas orientalis* (PoV) (*Fig. 4*). These three algal viruses, for which only PolB and major capsid protein (MCP) sequences are available, also possess very large

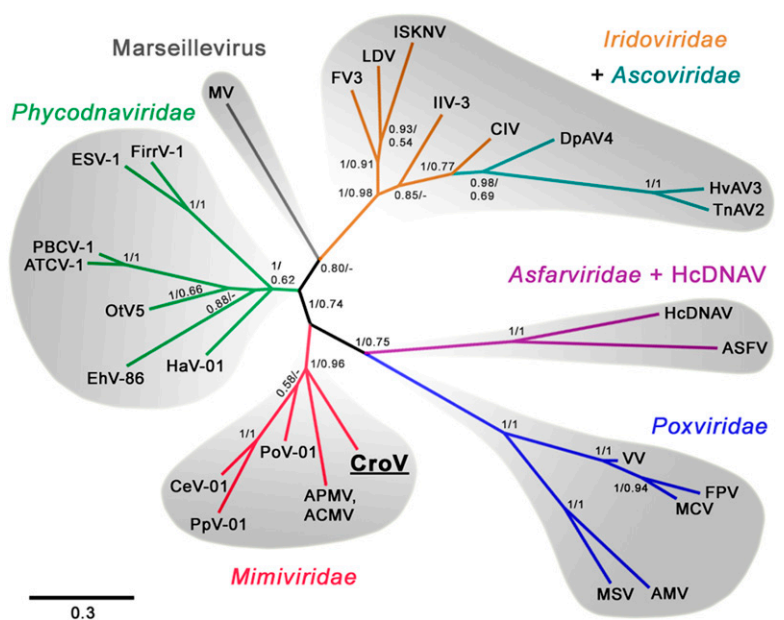
DNA genomes (485 kb, 510 kb, and 560 kb, respectively) and are proposed members of the family *Phycodnaviridae*, although a taxonomic revision of this tentative assignment has been proposed (10). Similarly, when the MCP was used to reconstruct the NCLDV phylogeny (*SI Appendix, Fig. S14*), these five viruses formed a monophyletic group that also included *Heterosigma akashiwo* virus, another large DNA virus that is assigned to the *Phycodnaviridae*. The topology of the NCLDV tree strongly suggests that the five largest viral genomes are more closely related to each other than to other NCLDV families and that they may have originated from a relatively recent ancestral virus that must have already been a bona fide NCLDV with a very large genome, probably encoding  $>150$  proteins.

## Conclusions

We present here the genetic analysis of a virus infecting a marine phagotroph. With a genome size larger than that of some cellular organisms, CroV is an example of an extraordinarily complex virus. It possesses a large number of predicted genes involved in DNA replication, transcription, translation, protein modification, and carbohydrate metabolism, indicating that CroV has a highly autonomous propagation strategy during infection.

The mechanisms by which such enormous virus genomes evolved have been much discussed (40, 44, 45). Most studies have focused on Mimivirus, because it represents the most extreme case of a giant virus and is the largest dataset available. The majority of Mimivirus genes have no cellular homologs and are presumably very ancient (46), up to one-third of its genes arose through gene and genome duplication (45), and  $<15\%$  of Mimivirus genes may have been horizontally transferred from eukaryotes and bacteria (6). Our analysis of the CroV genome is consistent with this general picture of giant virus genome evolution. Gene duplication and lineage-specific expansion of the FNIP/IP22 repeat are two factors that clearly contributed to the enormous size of the CroV genome. Examples of duplicated genes are the paralogous groups of CDSs *croV027–croV031* (contain FNIP/IP22 repeats), *croV420–croV422* (unknown function), and some of the tRNA genes. A potential case of large-scale HGT from a bacterium is represented by the 38-kb genomic segment that differs in coding content and promoter regions from the rest of the viral genome. The remaining CDSs with cellular homologs are more difficult to categorize, because genes can be transferred from cells to viruses and vice

**Fig. 4.** Phylogenetic reconstruction of NCLDV members. The unrooted Bayesian Inference (BI) tree was generated from a 263-aa alignment of conserved regions of DNA polymerase B. Intein insertions were removed before alignment. Nodes are labeled with BI posterior probabilities and maximum likelihood bootstrap values (500 replicates). Abbreviations and accession numbers (GenBank unless stated otherwise) are as follows: ACMV, *Acanthamoeba castellanii* mamavirus, from ref. 43; AMV, *Amsacta moorei* entomopoxvirus, NP\_064832; APMV, *A. polyphaga* mimivirus, YP\_142676; ASFV, African swine fever virus, NP\_042783; ATCV-1, *Acanthocystis turfacea* chlorella virus 1, YP\_001427279; CeV-01, *C. ericina* virus 01, ABU23716; CIV, Chilo iridescent virus, NP\_149500; CroV, *C. roenbergensis* virus; DpAV4, *Diadromus pulchellus* ascovirus 4a, CAC19127; EhV-86, *E. huxleyi* virus 86, YP\_293784; ESV-1, *Ectocarpus siliculosus* virus 1, NP\_077578; FIRR-V-1, *Feldmannia irregularis* virus 1, AAR26842; FPV, Fowlpox virus, NP\_039057; FV3, Frog virus 3, YP\_031639; HaV-01, *H. akashiwo* virus 01, BAE06251; HcDNAV, *Heterocapsa circularisquama* DNA virus, DDBJ accession no. AB522601; HvAV3, *Heliothis virescens* ascovirus 3e, YP\_001110854; IIV-3, Invertebrate iridescent virus 3, YP\_654692; ISKNV, Infectious spleen and kidney necrosis virus, NP\_612241; LDV, Lymphocystis disease virus, YP\_073706; MCV, Molluscum contagiosum virus, AAL40129; MSV, *Melanoplus sanguinipes* entomopoxvirus, NP\_048107; MV, *Marseillevirus*, MAR\_ORF329, GU071086; OtV5, *Ostreococcus tauri* virus 5, YP\_001648316; PBCV-1, *P. bursarium* chlorella virus 1, NP\_048532; PoV-01, *P. orientalis* virus 01, ABU23717; PpV-01, *P. pouchetti* virus 01, ABU23718; TnAV2, *Trichoplusia ni* ascovirus 2c, YP\_803224; VV, Vaccinia virus, AAA98419.





versa. However, the majority of CroV CDSs show no significant similarity to any sequences in the public databases and their evolutionary origin remains hidden.

The array of “organismal” genes found in CroV further closes the overlap in metabolic coding capacity between large viruses and cellular life forms. This continued blurring of the distinction between what is considered living and nonliving adds to the ongoing debate about the puzzling evolutionary history of giant viruses (7, 8, 44). Moreover, the PolB gene of CroV has high similarity with those of other marine virus isolates, relatives of which appear to be widespread in the oceans (10), suggesting that CroV represents a major group of largely unknown but ecologically important marine viruses.

## Materials and Methods

**Flagellate Growth and Virus Purification.** *C. roenbergensis* strain E4-10 was isolated from coastal waters near Yaquina Bay, OR, as described (13).

Cultures of *C. roenbergensis* were grown in f/2-enriched seawater medium supplemented with 0.01% (wt/vol) yeast extract to stimulate bacterial growth. The mixed assembly of bacteria in the cultures served as the food source for *C. roenbergensis*. Cultures were kept at room temperature ( $\approx 22^\circ\text{C}$ ) in the dark. Typically, 1-L plastic Erlenmeyer flasks containing 250 mL of exponentially growing *C. roenbergensis* were infected at a cell density of  $5 \times 10^4$

cells per mL by adding 100  $\mu\text{L}$  (multiplicity of infection  $\approx 0.5$ ) of crude CroV-containing lysate. CroV purification is described in *SI Appendix*.

**Genome Sequencing and Assembly.** Phenol-chloroform extracted genomic DNA was sequenced by 454 pyrosequencing on GS 20 and GS FLX platforms. The two datasets were assembled individually and resulting contigs were analyzed with Sequencher (Gene Codes). Gap closing was achieved by a combination of multiplex PCR, bioinformatic prediction methods followed by PCR verification and sequencing, and a genomic shotgun library using the pSMART vector (Lucigen).

*SI Appendix* contains further details and experimental procedures on genome annotation and microarray analysis.

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