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Biogeography of marine giant viruses reveals their interplay with eukaryotes and ecological functions

AUTHOR(S):

Endo, Hisashi; Blanc-Mathieu, Romain; Li, Yanze; Salazar, Guillem; Henry, Nicolas; Labadie, Karine; de Vargas, Colomban; ... Karp-Boss, Lee; Sunagawa, Shinichi; Ogata, Hiroyuki

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Biogeography of marine giant viruses reveals their interplay 1 with eukaryotes and ecological functions $\mathbf{2}$ 3 Hisashi Endo¹, Romain Blanc-Mathieu^{1,2}, Yanze Li¹, Guillem Salazar³, Nicolas Henry^{4,5}, 4 Karine Labadie⁶, Colomban de Vargas^{4,5}, Matthew B. Sullivan^{7,8}, Chris Bowler^{9,10}, $\mathbf{5}$ Patrick Wincker^{10,11}, Lee Karp-Boss¹², Shinichi Sunagawa³, Hiroyuki Ogata^{1,*} 6 7 8 **Affiliations:** 9 1. Bioinformatics Center, Institute for Chemical Research, Kyoto University, Gokasho, 10 Uji, Kyoto, 611-0011, Japan 11 2. Laboratoire de Physiologie Cellulaire & Végétale, CEA, Univ. Grenoble Alpes, 12CNRS, INRA, IRIG, Grenoble, France 133. Department of Biology, Institute of Microbiology and Swiss Institute of 14Bioinformatics, ETH Zürich, Zürich 8093, Switzerland 4. CNRS, UMR 7144, Station Biologique de Roscoff, Place Georges Teissier, 29680 1516Roscoff, France. 175. Sorbonne Universités, UPMC Université Paris 06, UMR 7144, Station Biologique 18 de Roscoff, Place Georges Teissier, 29680 Roscoff, France. 196. Genoscope, Institut de Biologie François-Jacob, Commissariat à l'Énergie Atomique 20(CEA), Université Paris-Saclay, Évry, France. 217. Department of Microbiology, The Ohio State University, Columbus, OH 43210, 22USA 238. Department of Civil, Environmental and Geodetic Engineering, The Ohio State 24University, Columbus, OH 43210, USA 259. Institut de Biologie de l'ENS (IBENS), Département de biologie, École normale 26supérieure, CNRS, INSERM, Université PSL, Paris 75005, France 2710. Research Federation for the study of Global Ocean Systems Ecology and Evolution, 28FR2022/Tara Oceans GOSEE, 3 rue Michel-Ange, 75016 Paris, France 2911. Génomique Métabolique, Genoscope, Institut de Biologie François Jacob, 30 Commissariat à l'Énergie Atomique (CEA), CNRS, Université Évry, Université Paris-Saclay, Évry, France. 31



- 32 12. School of Marine Sciences, University of Maine, Orono, ME, USA
- 33

34 **Corresponding author*:

- 35 H. Ogata, E-mail: ogata@kuicr. kyoto-u.ac.jp, Phone: +81-774-38-3270
- 36



37 Abstract

Nucleocytoplasmic large DNA viruses (NCLDVs) are ubiquitous in marine 3839 environments and infect diverse eukaryotes. However, little is known about their 40 biogeography and ecology in the ocean. By leveraging the Tara Oceans pole-to-pole metagenomic data set, we investigated the distribution of NCLDVs across size fractions, 41 42depths and biomes, as well as their associations with eukaryotic communities. Our 43analyses revealed a heterogeneous distribution of NCLDVs across oceans, with an elevated uniqueness in polar biomes. The community structures of NCLDV families were 44 45correlated with specific eukaryotic lineages including many photosynthetic groups. 46NCDLV communities were generally distinct between surface and mesopelagic zones, 47but at some locations, they exhibited a high similarity between the two depths. This 48vertical similarity was correlated to surface phytoplankton biomass but not to physical 49mixing processes, suggesting the potential role of vertical export in structuring 50mesopelagic NCLDV communities. These results underscore the importance of the coupling between NCLDVs and eukaryotes in biogeochemical processes in the ocean. 51

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54 Introduction

55The photic zone is the most productive layer of the ocean, containing a wide variety 56of microorganisms such as bacteria, autotrophic and heterotrophic protists and 57multicellular organisms. The population dynamics of these organisms determine the 58flows of energy and materials through marine food webs, playing a fundamental role in ecosystem functioning and biogeochemical cycles in the ocean^{1,2}. Viruses exert a top-5960 down control on marine organisms and release material to the pools of particulate and 61 dissolved organic matter³. This material and remineralized inorganic nutrients are utilized by autotrophic and mixotrophic phytoplankton⁴. The recycling of nutrients in the surface 6263 layer potentially reduces the transfer of fixed organic carbon to higher trophic levels and the deep sea^{5,6}. However, it is also possible that viruses enhance downward carbon flux 64 65 by facilitating cell aggregation and producing carbon-enriched materials from infected cells⁷⁻⁹. 66

67 Nucleocytoplasmic large DNA viruses (NCLDVs or so-called "giant viruses") 68 represent a monophyletic group of viruses that infect a variety of eukaryotic lineages¹⁰⁻¹². 69 Studies focusing on conserved marker genes such as family B DNA polymerase (*polB*) 70have revealed that NCLDVs are highly diverse and abundant in aquatic environments¹³⁻ 71¹⁶. The diversity of a family of NCLDVs, namely *Mimiviridae*, exceeds that of bacteria 72and archaea in the ocean¹⁷ and their richness in a few liters of seawater can reach more than 5,000 operational taxonomic units¹⁸. More recently, several thousand draft genomes 7374(i.e., metagenome-assembled genomes; MAGs) of NCLDVs were constructed from 75environmental sequences, thanks to the development of high-throughput sequencing and 76bioinformatics technologies^{19,20}. However, the global biogeography of marine NCLDVs 77still remains under-explored.

A growing number of marine eukaryotes have been reported as host organisms of NCLDVs, particularly phytoplankton groups such as haptophytes, chlorophytes and dinoflagellates²¹⁻²³. Other eukaryotic lineages, including non-photosynthetic organisms such as bicosoecids and choanoflagellates, have also been reported as host organisms of



NCLDVs in marine environments^{24,25}. These studies collectively suggest the ecological
importance of NCLDVs in the ocean via top-down effects on eukaryotic communities.
However, our knowledge of NCLDV-host relationships is highly limited, given the large
phylogenetic diversities of NCLDVs and microeukaryotes.

Here we reveal patterns in the global biogeography of NCLDVs using the 86 87 metagenomic data from the Tara Oceans project. The metagenomic data cover varying 88 geographic regions including polar and deep-sea ecosystems, in which NCLDVs are under-researched²⁶⁻²⁸. We constructed NCLDV taxonomic abundance profiles for 283 89 90 samples, representing two viral size fractions, three ocean depth ranges (surface, deep 91 chlorophyll maximum and mesopelagic), and four biomes (coastal, trades, westerlies and 92polar). The global biogeography of NCLDVs derived from these data reveals strong 93 associations between NCLDVs and eukaryotic microorganisms. Furthermore, vertical connectivity of NCLDV communities indicates a possible mechanism for how 94 95mesopelagic NCLDV communities are structured with respect to ocean biogeochemical 96 processes.

97

98 **Results**

99 NCLDV phylotypes detected in *Tara* Oceans metagenomes

100 We detected 6,818 PolBs affiliated with NCLDVs in the second version of the Ocean Microbial Reference Gene Catalog (OM-RGC.v2)²⁸ using the pplacer phylogenetic 101 placement method²⁹ (see methods for details). The OM-RGC.v2 was built based on 370 102 103 Tara Oceans metagenomes from femto- (<0.2 µm; 151 samples), pico- (0.22–1.6 or 0.22– 1043.0 µm; 180 samples) and other (39 samples) size fractions. After removing 32 samples 105with a low NCLDV frequency and 55 samples from non-target size fractions and depths, 106 the remaining 283 samples contained 6,783 NCLDV PolB sequences. The pplacer 107 classified these PolBs into nine NCLDV families/lineages. The number of phylotypes 108 (distinct *polB* at 95% nucleotide sequence identity) was the largest in *Mimiviridae* (5,091 109 phylotypes), followed by *Phycodnaviridae* (981 phylotypes). The number of phylotypes



110 taxonomically assigned to Iridoviridae, Medusavirus and Asfarviridae, were 239, 120 111 and 109, respectively. We also detected PolBs assigned to *Pithoviridae* (93), Ascoviridae 112(78), Poxviridae (51) and Marseilleviridae (21). However, Poxviridae was omitted from 113 our discussion as the environmental gene sequences were distantly related to known 114 Poxviridae. Rarefaction analysis showed that, at the end of sampling, the number of 115NCLDV phylotypes increased by less than 0.01% per sample for all samples, and ranged 116 from 0.02% to 0.32% when samples were divided into different size fractions, depths and 117biomes (Extended Data Fig. 1).

118 To examine detailed phylogenetic affiliation and to visualize the dispersal 119 characteristics of each NCLDV phylotypes detected by pplacer, we constructed a 120 phylogenetic tree using selected PolB sequences (Extended Data Figs. 2–4). Among the 121*Mimiviridae* family, genes closely related to the algal-infecting subfamily, recently proposed as "Mesomimivirinae" (e.g., AaV, CeV, pkV, PgV, PoV and TetV)³⁰, which 122123 infect pelagophytes (the genus Aureococcus), haptophytes (the genera Haptolina, 124Prymnesium and Phaeocystis), and chlorophytes (the genera Pyramimonas and 125Tetraselmis), were relatively abundant. On the other hand, only a few sequences were 126 affiliated with the subfamilies "Megamimivirinae" and "Klosneuvirinae" except the 127*Cafeteria roenbergensis virus* (CroV), which is the only member of "Megamimivirinae" isolated from the marine environment²⁴. Among *Phycodnaviridae*, the genus 128129 Prasinovirus (e.g., BpV, MpV, OtV and OlV), which infect chlorophyte genera such as 130 Bathycoccus, Micromonas and Ostreococcus, showed the highest richness.

131

132 Heterogeneity in NCLDV community structure across size, depth and biomes

The dominant NCLDV taxa detected from all sample locations and depths in the picosize fraction were *Mimiviridae* and *Phycodnaviridae*, with average contributions of 64.6% and 25.4%, respectively (Fig. 1A). The dominant groups of NCLDVs varied widely among sites and depths in samples from the femto-size fraction (Fig. 1B). In this fraction, *Phycodnaviridae* and *Asfarviridae* had relatively high contributions to the total



NCLDVs with the mean values of 29.7% and 19.9%, respectively. *Mimiviridae* and *Ascoviridae* were also important contributors with mean values of 12.2% and 11.1%,
respectively.

141 A non-metric multidimensional scaling (NMDS) analysis showed that NCLDV 142 assemblages clustered according to size fraction, depth and biome (Fig. 2A–2C). 143 Significant differences in NCLDV community composition were detected among all 144 categories (PERMANOVA, p < 0.01), and size fraction, depth and biome explained 5.5%, 145 4.3% and 10.9% of the total variance, respectively.

146 Taxonomic richness (i.e., number of phylotypes) and Shannon's diversity index were 147used to investigate variation in NCLDV community diversity. In this study, we analyzed 148 the samples from all depths and size fractions to compare diversity differences among 149 depth ranges, although latitudinal trend in Shannon's diversity for pico-sized communities from the surface was reported previously³¹. In the pico-size fraction, mean 150151values for NCLDV richness at the surface and in the DCM layer were about 1.7 times 152higher than that in the mesopelagic layer (Kruskal-Wallis and Dunn's post hoc test, p 153<0.01) (Extended Data Fig. 5A). In the femto-size fraction, NCLDV richness was 154significantly higher at the surface and MES layer than in the DCM layer (Dunn's test, p 155= 0.04 - 0.05), although the differences were small and not consistent with the pico-size 156fraction.

157

158 High uniqueness of NCLDV phylotypes in the Arctic Ocean

We analyzed the overlap and uniqueness of NCLDV phylotypes across different ecological zones (i.e., size fraction, depth and biome) to evaluate their ability to disperse across different environments. Each ecological category was divided into two major groups (i.e., pico- and femto-sizes, euphotic and mesopelagic zones, and polar and nonpolar biomes), because the NCLDV community in mesopelagic zone or polar biome was separated most significantly from other depths or biomes (Fig. 2). We found 4,003 (59.0% to the total NCLDVs) shared NCLDV phylotypes across size fractions, 4,737 (69.8%)



shared phylotypes across depth ranges, and 1,950 (28.7%) shared phylotypes across biomes (Fig. 3A). Only twelve unique phylotypes were detected in the femto-size fraction, whereas 2,768 unique phylotypes were identified in the pico-size fraction. The euphotic zone (surface and DCM) harbored 1,986 unique phylotypes, whereas the aphotic mesopelagic zone had only 60 unique phylotypes. The polar biome (the Arctic and the Southern Ocean) included 620 unique NCLDV phylotypes, whereas 4,213 unique NCLDVs were detected in non-polar biomes (i.e., trades, westerlies and coastal).

173 To further characterize regional differences in the NCLDV community, we 174investigated the total and unique NCLDV phylotypes observed in nine geographic regions 175and the phylotypes shared among regions. The total number of phylotypes was relatively 176 high in the Atlantic, Pacific and Indian Oceans and in the Mediterranean Sea, with values 177of between 3,665 and 4,685 (Fig. 3B). Lower numbers of NCLDV phylotypes were 178identified from the Red Sea (2,653) and the Arctic Ocean (2,467). The Southern Ocean 179presented the lowest number of NCLDV phylotypes (561), although this was based on 180 only 5 samples. The Arctic Ocean samples displayed a high number of unique NCLDV 181 phylotypes (551), which corresponded to 22.3% of the total phylotypes detected in this 182region. In contrast, the number of unique phylotypes from other regions ranged from 0 to 183 134 (0.0% to 3.4%).

There was no linear or saturation trend in the number of total or unique NCLDV phylotypes with increasing sample size (Fig. 3C). The high proportion of unique phylotypes in the Arctic Ocean was not a function of sample size, although the number of total phylotypes detected in the Southern Ocean may be limited by the low number of samples. The phylogenetic positions of unique NCLDVs from the polar biome were dispersed across most of the NCLDV families (Fig. 4)

190

191 NCLDV distributions correlate with eukaryotic communities

A partial Mantel test was conducted to assess community associations among the
 NCLDV families/lineages and major eukaryotic lineages. The pairwise partial correlation



194 coefficients (Spearman's ρ) varied from -0.17 to 0.76 (Fig. 5A), and 93.6% of the 195examined pairs (225 out of 234 for the pico-size fraction and 213 out of 234 for the femto-196 size fraction) showed statistically significant correlations (p < 0.01, permutation test) after 197 false discovery rate (FDR) correction. Pairs from pico-sized NCLDV communities with 198 a correlation coefficient ≥ 0.53 were considered to represent strong positive associations, 199 because 8 out of 9 known marine virus-host lineage associations were recovered by this 200 criterion (Figs. 5A and 5B). Using this threshold, 30 out of 234 NCLDV-eukaryote 201lineage pairs were found to have strong linkages (Fig. 5C). The NCLDV families/lineages 202were generally highly correlated with the known host groups among autotrophic and 203 mixotrophic microalgae (haptophytes, chlorophytes, dinophytes, pelagophytes and raphidophytes) ($\rho = 0.54-0.67$). Interestingly, *Mimiviridae* was strongly correlated with 204 205chrysophyte microalgae ($\rho = 0.65$), which are not currently known as NCLDV hosts. 206 Other than algal lineages, a strong positive correlation was found between *Mimiviridae* 207and heterotrophic eukaryote choanoflagellates ($\rho = 0.76$), which are a known lineage of 208Mimiviridae. A group of non-photosynthetic heterokonts bicosoecids are also a known 209host of the Mimiviridae species CroV in marine environments, but this group was not 210 highly correlated with *Mimiviridae* ($\rho = 0.30$).

211

212 Potential chrysophyte viruses constitute novel clades of *Mimiviridae*

213 To explore possible associations between NCLDVs and chrysophytes as indicated by 214 the Mantel's regression analysis (Fig. 5C), we tested for chrysophyte-derived genes in 215the metagenome-assembled genomes (MAGs) of NCLDVs generated by Schultz et al. $(2020)^{19}$ and Moniruzzaman et al. $(2020)^{20}$. The results showed that 89 (82 after removing 216217 redundancy) out of 2,263 MAGs contained genes closely related to the transcripts of the 218chrysophytes (Supplementary Data 1). Comparisons between PolB sequences revealed 219 27 PolBs from the OM-RGC.v2 that were closely related to the NCLDV MAGs with 220 chrysophyte homologs. Most of these PolBs constituted novel clades within the branches 221of Mimiviridae (Fig. 4; Extended Data Fig. 4). We confirmed that other genes in the



222 contigs that contained chrysophyte homologs are highly similar to the Mimiviridae or

223 *Phycodnaviridae* sequences in many cases (Extended Data Fig. 6).

224

225 Vertical connectivity of NCLDV communities

226 The vertical connectivity of NCLDV communities was investigated using Bray-Curtis 227 community similarity measures to compare between epipelagic (surface or DCM) and 228mesopelagic samples at individual sampling locations. The Bray-Curtis similarities were 229less than 0.10 for about half of the tested locations (20 out of 36 surface sites and 13 out 230of 26 DCM sites; Fig. 6A; Extended Data Fig. 7A). All sites in the Arctic Ocean and 231several sites in tropical and subtropical regions showed relatively high similarities 232between the two depth (0.15 to 0.60). The NCLDV community similarity value was 233positively correlated with the chlorophyll a concentration in the epipelagic layer 234(Spearman's $\rho = 0.52$, p < 0.01, asymptotic t approximation, n = 36 for surface; $\rho = 0.44$, 235p = 0.02, n = 25 for DCM) and NCLDV richness in the mesopelagic layer ($\rho = 0.82$, p 236<0.01, n = 36 for surface; $\rho = 0.70$, p < 0.01, n = 26 for DCM) (Figs. 6B and 6C; Extended 237Data Figs. 7B and 7C). We also evaluated relationships between NCLDV vertical 238similarity and physical environmental factors including: the sampling depth of mesopelagic water, the mixed layer depth, and the temperature difference between 239240epipelagic and mesopelagic waters. No significant correlations were detected among 241these parameters (p > 0.05, n = 32-36 for surface samples and n = 25-26 for DCM 242samples) (Figs. 6D–F; Extended Data Figs. 7D–F).

We plotted correlations among the relative contributions of NCLDV phylotypes between the euphotic and aphotic zones at all sampling locations (Extended Data Figs. 8 and 9). Where there was a strong similarity in the NCLDV community found at different depths, *Phycodnaviridae* generally contributed highly to samples from the Arctic Ocean (e.g., TARA stations 158, 201 and 209), and both *Mimiviridae* and *Phycodnaviridae* contributed strongly in tropical and subtropical regions (e.g., stations 72, 110 and 122).



250 **Discussion**

251We investigated the diversity and community structure of NCLDVs based on 252metagenomic PolB sequences collected from the world oceans. NCLDV communities 253differed substantially between pico- and femto- size fractions (Fig. 1). NCLDV 254communities in the pico-size fractions were dominated by Mimiviridae and 255Phycodnaviridae, regardless of sampling location or depth (Fig. 1A). In marine 256environments, species from the haptophytes (the genera Prymnesium, Haptolina, and 257*Phaeocystis*), chlorophytes (*Pyramimonas*), pelagophytes (*Aureococcus*), bicosoecids 258(Cafeteria) and choanoflagellates (Bicosta) are known hosts of Mimiviridae, while 259species of haptophytes (Emiliania), chlorophytes (Ostreococcus, Micromonas and 260Bathycoccus) and raphidophytes (*Heterosigma*) have been reported as *Phycodnaviridae* 261hosts (Virus-Host DB)³². Although the dominance of Mimiviridae and Phycodnaviridae have been reported in previous studies, mainly from coastal seawater^{13,14}, our results 262263demonstrate the ubiquitous nature of these protist-infecting viruses across world ocean 264biomes. It is worth noting that most of the NCLDVs (99.7%) detected from the femto-265size fraction were also present in the pico-size fraction (Fig. 3A), despite the large 266differences in relative abundance between two size fractions at each location. Therefore, 267the abundance information can be important for characterizing the differences of NCLDV 268communities. A proportion of the NCLDVs in the pico-size fraction were present within 269infected cells, because cell sizes of some host species such as Aureococcus 270anophagefferens and Micromonas pusilla are less than 3 µm. Thus, the abundance of 271these lineages in the pico-size fraction may be partly enriched by the viruses replicating 272inside their hosts.

In addition to *Phycodnaviridae* and *Mimiviridae*, *Asfarviridae* also contribute an important proportion of NCLDVs in the femto-size fraction of most euphotic zones (Fig. 1B). Although very limited information is available regarding the natural hosts for this group, a representative *Asfarviridae*-like species in marine environments is *Heterocapsa circularisquama* DNA virus (HcDNAV), which infects the red-tide-forming



278 dinoflagellate *H. circularisquama*³³. In the terrestrial ecosystem, this viral family is 279 known to infect a wide variety of organisms such as amoebozoa, arthropods and 280 mammals^{32,34}. Given the broad range of host species for this viral lineage, there may be 281 an unknown but wide-spread host taxa for *Asfarviridae* in the ocean.

282Our study revealed a heterogeneous pattern in the distribution of NCLDVs across the 283oceans of the world (Fig. 2C). Although there are limited studies available on the factors 284controlling the large-scale distribution of viruses, it is widely accepted that both 285deterministic (environmental factors and inter-specific interactions) and stochastic 286processes (e.g., immigration and speciation) are important in making up microbial assemblages³⁵⁻³⁷. The distribution and diversity of viruses would not be directly affected 287288by environmental variables such as temperature and nutrient availability, but is directly 289 influenced by the geographic ranges of their host species^{3,38}. Recent work with 290cyanophages demonstrated that a significant number of free-living viruses are locally 291produced through active infection rather than from migration³⁹. Therefore, we expect that 292viral community structure will reflect host distribution as well as infectious activity.

293Despite significant differences in community composition across oceanic biomes, we 294found that most NCLDV phylotypes are dispersed throughout tropical and temperate 295regions (Figs. 3A and 3B), presumably following their host community composition, which is primarily determined by temperature⁴⁰. However, the polar biome (mainly the 296 297 Arctic Ocean) constitutes a "hotspot" of unique NCLDV phylotypes from a wide variety 298of families, despite having a low total richness in comparison to other regions (Figs. 3B 299 and 3C). We revealed that NCLDVs unique to non-polar biome were also abundant (Fig. 300 4), indicating a strong separation of NCLDV communities between polar and non-polar 301 biomes. A geographical barrier and steep environmental gradients may underlie this 302 distinct ecosystem structure (i.e., different host communities and their productivity) in the Arctic Ocean^{27,28,31}. Moreover, the Arctic Ocean is characterized by high amounts of river 303 304 discharge, contributing more than 10% to global runoff flux⁴¹. Consequently, biological 305 processes in the Arctic may be influenced by river inputs from terrestrial ecosystems.



These factors may collectively contribute to the remarkable number of unique NCLDV phylotypes found in the Arctic, that were undetectable in other regions. The biogeography of NCLDVs on a global scale implies a tight link between the NCLDVs and the distribution of their hosts, which is strongly influenced by physicochemical and biological factors.

311 Tight coupling between NCLDVs and their hosts was further corroborated by our 312partial Mantel statistics, which described both known virus-host interactions and 313 additional but currently unrecognized associations between viruses and eukaryotic 314 lineages at the community level. Using the pico-sized NCLDV community, we detected 315almost all known virus-host interactions, except for those involving Bicoecea (Fig. 5C). 316 This demonstrates that distance-based correlation analysis using global ocean samples is 317 useful for detecting virus-host interplay in natural environments, although the validations 318 of the previously unknown associations remain to be further explored. Strong positive 319 relationships between NCLDVs and eukaryotes involved many phytoplankton lineages 320 including haptophytes, chlorophytes, dinophytes, pelagophytes and raphidophytes, all of 321 which include known host lineages of NCLDVs (Fig. 5C). Strong correlations were also 322 detected with heterotrophic choanoflagellates, which have recently been identified as a novel host of *Mimiviridae*²⁵. Some NCLDVs, especially *Mimiviridae*, had strong 323 324 correlations with chrysophytes, although no host species have yet been reported for this 325lineage. Many environmental NCLDV genomes were found to encode genes that are 326 likely to be derived from marine chrysophytes (Supplementary Data 1–3). Taxonomic 327 analyses based on PolB phylogeny and homology search revealed that most of these 328 phylotypes represent previously unknown clades of the Mimiviridae tree (Extended Data 329 4 and 6; Supplementary Data 4), suggesting that chrysophytes may be an important host 330 lineage of Mimiviridae in the ocean.

The global distribution of NCLDVs are determined by the geographic ranges of their host organisms. Therefore, the virus-eukaryote associations that we detected likely arose under these constraints. On the other hand, it is expected that NCLDVs influence the



334 abundance of eukaryotes at a local scale. Previous studies show that bacterial viruses have 335 an important role in determining bacterial mortality, because they substantially 336 outnumber their hosts and have highly specific infection mechanisms⁴². Similarly, 337 NCLDVs are reported to be more abundant than their host cells and have high infection specificity^{11,14,43}. For example, *Emiliania huxlevi* viruses (EhVs) of the *Phycodnaviridae* 338 family are responsible for almost all of the mortality of the haptophyte E. huxleyi during 339 blooms^{22,44,45}. Another field study suggests that viral lysis can explain a greater proportion 340 341 of phytoplankton mortality than grazing by zooplankton⁶. These studies, combined with 342 the global associations that were detected in this study, emphasize the potential 343 importance of NCLDVs in structuring eukaryotic communities.

344 Our results indicate that marine phytoplankton lineages could represent one of the 345most important host groups of NCLDVs. Therefore, NCLDVs could be involved in the 346 regulation of biogeochemical processes mediated by phytoplankton. We investigated this 347 by assessing the vertical connectivity of viral communities. The NMDS analysis showed 348clear differences between the NCLDV community composition of epipelagic (euphotic) 349 and mesopelagic (aphotic) zones at most sampling sites (Fig. 2B). Similar results were also reported for phage communities in the Pacific Ocean⁴⁶. The vertical separation of 350351viral communities may be caused by the stable stratification below the mixed layers 352 (typically above 200 m depth), which severely inhibits vertical water exchange. Despite 353 this limitation, mesopelagic ecosystems shared a significant number (98.7%) of NCLDV 354phylotypes with the upper epipelagic layers (Fig. 3A), suggesting the vertical connectivity 355 of NCLDVs and their local adaptation. Indeed, some mesopelagic NCLDV communities 356were very similar to surface communities (Fig. 6A and Extended Data Fig. 7A). This 357 implies that the surface and mesopelagic NCLDV communities may be connected at some 358 locations. The major source of energy and materials in the mesopelagic layer is the gravitational export of organic particles from the surface layer (i.e., the biological carbon 359 360 pump)⁴⁷⁻⁴⁹. Therefore, some surface viruses may be exported to mesopelagic layers with sinking aggregated phytoplankton cells⁵⁰⁻⁵². 361



362 A significant positive correlation existed between surface phytoplankton biomass and 363 NCLDV community similarity across depths (Fig. 6B and Extended Data Fig. 7B). Since 364 highly productive areas are likely to have a greater flux of settling particles to the deep 365 layers, this result supports the idea that NCLDVs are transported with the sinking particles. 366 High vertical connectivity was consistently associated with an increase in NCLDV 367 richness in the mesopelagic zone (Fig. 6C and Extended Data Fig. 7C). Previous studies 368 showed that sinking particles can transfer bacterial and phage populations to the deep layer^{52,53}. Mestre et al.⁵² demonstrated that particle-attached prokaryotes had higher 369 370 capacity for immigration than free-living ones. Based on the particle-driven vertical 371 dispersion model, we can expect that NCLDVs, inside or attached to their host cells or 372 cell debris, might be preferentially exported into the deep sea. Numerous studies based 373 on sediment trap measurement have shown that larger phytoplankton, such as diatoms, contribute strongly to vertical flux because of their high sinking velocities^{54,55}. However, 374 375 recent studies show that smaller phytoplankton including haptophytes and chlorophytes, 376known hosts of marine NCLDVs, also contribute greatly to downward carbon export^{8,9,56}. 377 The high vertical connectivity of NCLDVs was not affected by the extent of the depth 378 range nor by proxies for vertical mixing (Figs. 6D-F and Extended Data Figs. 7D-F), 379 indicating that the migration of NCLDVs occurred regardless of physical processes such 380 as upwelling, turbulent mixing, and convection. This result suggests that sinking export 381 is a major source of a variety of NCLDVs to deeper waters, where NCLDV diversity is 382 relatively low without this effect. A recent study revealed that some Phycodnaviridae and 383 Miniviridae potentially accelerate biological carbon export from the productive surface 384layer to deep layers, presumably by promoting cell death and aggregation of their host 385species⁵⁷. *Phycodnaviridae* and *Mimiviridae* also contributed strongly to high vertical 386 connectivity in our study (Extended Data Figs. 8 and 9). The infection of the 387 coccolithophore by the Phycodnaviridae EhV was observed to facilitate the sinking of 388 host cells, likely by enhancing the production of transparent exopolymer particles and subsequent aggregation⁹. Therefore, the high vertical connectivity of NCLDVs detected 389



in our analysis may be partly associated with enhanced vertical export of their infectedhosts.

392 The present study expands our knowledge of marine NCLDV biogeography. Most 393 NCLDV phylotypes are ubiquitously distributed over the oceans of the globe, although a 394 high proportion of unique NCLDVs was detected in the Arctic Ocean. Our comparison 395 of community distribution patterns highlighted the tight interplay between NCLDVs and 396 microeukaryotes. As marine ecological and biogeochemical processes are governed 397 primarily by microbes, NCLDVs would have an important influence on the dynamics of 398 marine systems. We also identified unexpected similarity of NCLDV communities 399 between surface and deep waters at some locations. This supports the idea that viral 400 activity may be related to the strength of the biological carbon pump, because the 401 efficiency and sinking rate of export production depends largely on surface phytoplankton composition and their infection status^{8,9,55,58}. Our findings underscore the importance of 402 403 NCLDVs as a component of marine microbial communities, and contribute to refine our 404 knowledge of marine ecosystems, a key regulator of the Earth's climate.

405

406 Methods

407 Sample collection

408 Metagenomic datasets were generated from samples collected by the *Tara* Oceans 409 expeditions from 2009 to $2013^{26-28,31,59}$. The second version of the Ocean Microbial 410 Reference Gene Catalog (OM-RGC.v2) is a non-redundant gene catalog constructed from 411 370 metagenomic samples from the *Tara* Oceans project²⁸ (https://www.ocean-412 microbiome.org). The catalog includes 46,775,154 genes in total, and the gene abundance 413 profiles are expressed as the sum of within-reads aligned base pairs normalized by gene 414 length, in *Tara* Oceans samples²⁸.

415

416 **Recruitment of NCLDV marker genes from the OM-RGC.v2**

417 To assess the community composition of NCLDVs, we used family B DNA



418 polymerase (polB) as a marker gene of NCLDVs. Initially, amino acid sequences of the 419 OM-RGC.v2 were searched against an in-house profile hidden Markov model (HMM) of 420 NCLDV PolB sequences using the software HMMER, hmmsearch (version 3.1)⁶⁰ with a 421threshold E-value $<1\times10^{-5}$. Consequently, 29,315 PolB sequences were obtained from the 422 OM-RGC.v2, although this collection included sequences other than NCLDVs. To 423remove the sequences not derived from NCLDVs and classify the taxonomic identity of 424each NCLDV sequence, phylogenetic mapping was performed within known PolB sequences. A maximum-likelihood (ML) reference phylogenetic tree was built based on 425426 211 PolB reference protein sequences from eukaryotes, bacteria, archaea, phages and 427NCLDVs. These sequences were aligned using the default settings of the multiple sequence alignment program MAFFT-linsi (version 7)⁶¹ and ML tree was constructed 428 429 with the use of randomized axelerated maximum likelihood (RAxML) program (version $(7.2.8)^{62}$. In the reference trees, we included sequences from eight proposed families of 430 431 NCLDVs⁶³: *Mimiviridae* (synonymous with Megaviridae), Phycodnaviridae, 432Pithoviridae, Marseilleviridae, Ascoviridae, Iridoviridae, Asfarviridae, and Poxviridae 433 (Extended Data Figs. 2-4). A sequence from a novel NCLDV clade Medusavirus was also included as a reference⁶⁴. Query sequences were aligned against the reference 434435alignment using the MAFFT 'addfragments' option, and then mapped onto the reference 436 tree using the software program $pplacer^{29}$.

437

438 Abundance profiling of NCLDVs

We used the abundance profile of NCLDV genes from the OM-RGC.v2 to evaluate the relative frequency and diversity of NCLDVs. In the abundance matrix, we only included samples from the pico-size (0.22–1.6 or 0.22–3.0 μ m) and femto-size (<0.22 μ m) fractions. Samples used in the analysis were from three depth ranges: the surface (2– 9 m), the deep chlorophyll maximum (DCM, 15–180 m) and the mesopelagic (MES, 250– 1,000 m). The sum of length-normalized PolB abundances ranged from 5.3 to 22,847.5 across samples. The samples containing low PolB abundances tended to yield lower



446 diversity estimates (i.e., number of phylotypes and Shannon's entropy) (Extended Data 447 Fig. 10). To avoid bias due to the low sequencing effort, samples for which the sum of 448 length-normalized PolB abundance was less than 50 (set as a proxy for low NCLDV 449 frequency) were removed from the analysis. The abundance matrix was then standardized 450by the sample with the lowest sum of length-normalized PolB abundance value. The 451minimum value of PolB abundance among NCLDV phylotypes in the sample having the 452lowest sum of length normalized PolB was set as the cutoff threshold. For each sample, 453NCLDV phylotypes with a length-normalized abundance of less than this threshold were 454treated as absent. A sample of a femto-size fraction of surface water from station 155 was 455also removed, because it contained only one NCLDV PolB after standardization. 456Consequently, our dataset was comprised of 283 samples (172 pico-fraction samples and 457111 femto-fraction samples), covering 88 sampling sites. These sites were categorized 458into four biomes (coastal, trades, westerlies and polar biomes) according to latitude or 459distance from the shore, and nine oceanic regions, as defined by Longhurst⁶⁵ 460 (Supplementary Table 1).

461

462 **Phylogenetic tree construction**

463 To construct a phylogenetic tree, the NCLDV-derived PolB sequences obtained from 464 the OM-RGC.v2 were filtered by length (≥700 amino acid sequences) because the 465 inclusion of short sequences yields unreliable phylogenies. Amino acid sequences from 466 the resulting 911 genes were aligned with known NCLDV sequences using the linsi 467 option from the MAFFT. The ML tree was constructed using RAxML with the use of a 468 known NCLDV sequence tree as a backbone constraint. We confirmed the validity of the 469 pplacer family assignment for 905 out of 911 selected sequences. The remaining six 470 sequences that were incorrectly placed within the phylogenetic tree were removed. The ML tree was visualized using the program iTOL⁶⁶. 471

472

473 Prediction of potential chrysophyte viruses using metagenomic assembled genomes



474To explore the genomic contents of environmental NCLDVs, we made use of two sets 475of metagenome-assembled genomes (MAGs) of NCLDVs (GVMAGs high and medium 476 quality¹⁹; MoMAGs²⁰), which were generated from environmental metagenomic datasets 477 collected on global scales. Gene prediction was made for all MAGs using the program GeneMarkS⁶⁷, then the predicted genes were searched using BLASTP against a database 478479that combines the NCBI Reference Sequence database (RefSeq release 90) and the marine 480 microbial eukaryote transcriptomes project (MMETSP) database⁶⁸. We identified MAGs 481 whose genes exhibited the best hit to transcripts of chrysophytes with >50% amino acid 482identity and >100 alignment length (Supplementary Data 1). For these MAGs, we 483 checked the redundancy between the MoMAG and GVMAG datasets using average nucleotide identity of \geq 95% and an alignment fraction of \geq 50% with FastANI (version 484 $1.3)^{69}$. Although seven MAGs were found to be overlapped between the two datasets 485486 (Supplementary Data 1), all of the MAGs were retained for downstream analyses as these 487 had different contig structures. The chrysophyte-related genes were considered potential 488 candidates for horizontal gene transfer between chrysophytes and NCLDVs, and were 489BLASTP searched against the RefSeq database for additional functional annotation 490 (Supplementary Data 2). We then extracted PolB sequences from the NCLDV MAGs 491 which had a chrysophyte-related gene using the HMMER hmmsearch program. These 492PolBs were BLASTP searched against the NCLDV PolBs from the OM-RGC.v2. MAG-493 derived PolBs aligned with over 700 amino acid sequences with >90% identity were 494 assigned to the PolB phylotypes derived from the OM-RGC.v2 (Supplementary Data 3). 495Phylogenetic affiliations of PolB from the chrysophyte-related MAGs were confirmed 496 using a phylogenetic tree. To further test the credibility of our analysis, we checked other 497 genes on the contigs that harbored the chrysophyte homologs using BLASTP against the 498 RefSeq database (Supplementary Data 4; Extended Data Fig. 6).

499

500 **Diversity analyses**

501 Diversity and multivariate analyses were performed using the statistical software R



502(version 3.6.2) (https://www.r-project.org/). To evaluate the diversity of each sample, the 503number of NCLDVs (richness) and Shannon's entropy were assessed by the package 504'vegan' (https://cran.r-project.org/web/packages/vegan). NCLDV richness among sizes 505and depths were compared using a Kruskal-Wallis test followed by Dunn's multiple 506 comparison. Compositional variation among samples was assessed with a non-metric 507multidimensional scaling (NMDS) ordination based on Bray-Curtis dissimilarity. 508Statistical significance of differences among the sample groups (size, depth and biomes) was tested using a permutational multivariate analysis of variance (PERMANOVA)⁷⁰ 509 510with 9,999 permutations.

511

512 Partial Mantel test

513A partial Mantel test was performed to assess the correlation between two multivariate 514matrices while controlling the potential effects of geographic distance (spatial 515autocorrelation) using the R package 'vegan'. Abundance matrices for the NCLDV and 516eukaryotic lineages were constructed from the integrated abundance tables, and the total 517abundance at each site was normalized to 1. The eukaryote abundance table was 518constructed based on 18S rRNA gene metabarcoding⁷¹. Data for NLCDVs were obtained 519from pico- $(0.22-1.6/3.0 \,\mu\text{m})$ or femto-size (<0.2 μ m) fractions and for the eukaryotic 520community from the pico- to meso-size fraction (0.8-2,000 µm). There were 84 521overlapping sampling events between pico-size NCLDVs and eukaryotic communities and 55 overlapping sampling events between femto-size NCLDVs and eukaryotic 522523communities. All overlapping samples were derived from the surface or DCM depth 524layers. Distance matrices for viruses and eukaryotes were calculated using the Bray-525Curtis measure. Geographic distances among sample sites were also measured using 526 Haversine distance and were used as a third distance matrix. Partial Mantel correlations 527were computed between all pairs of distance matrices of eukaryotic communities and 528NCLDVs with 9,999 permutations for each comparison. The false discovery rate (FDR) was computed using the Benjamini-Hochberg method⁷². 529



530	
531	Statistical test
532	Two-sided test was applied for all statistical tests.
533	
534	Data availability
535	The complete sequence data of the OM-RGC.v2 and the abundance profile can be
536	downloaded from https://www.ocean-microbiome.org. All sequences of 18S rRNA gene
537	metabarcoding have been deposited at European Nucleotide Archive (ENA) under the
538	BioProject ID PRJEB6610 and PRJEB9737. Environmental metadata are archived at
539	https://doi.pangaea.de/10.1594/PANGAEA.875582. Files used for recruiting NCLDV
540	PolB genes as well as processed abundance profiles of eukaryotes and NCLDVs with
541	corresponding environmental data are available at the GenomeNet FTP:
542	ftp://ftp.genome.jp/pub/db/community/tara/Biogeography/.
543	
544	Code availability

545 Custom scripts developed for this study are available at GitHub: 546 https://github.com/HisashiENDO/NCLDV_Biogeography.



548 **Figure legends**

- 549Figure 1Latitudinal patterns in NCLDV community composition.Relative550contributions of NCLDV families at each depth range of (A) pico- and (B)551femto-size fractions. The number of phylotypes detected in each sample is also552indicated with a white circle. Sampling stations were arranged in rows from553south to north, and color-coded based on biome (for a map of the sampling554stations, please see Salazar et al., 2019²⁸).
- 555Figure 2Community characteristics of NCLDVs. Non-metric multidimensional556scaling (NMDS) ordination based on the NCLDV community showing results557for all samples (A) and separately for pico- and femto-size fractions (B and C).558Sample groups are color-coded by size fraction (A), depth (B) and biome (C).559Ellipses represent 90% confidence levels for each group. All group categories560are significantly different from each other as analyzed using PERMANOVA (p561<0.01). Sample sizes for the test are noted in Supplementary Table 1.</td>
- 562Figure 3 Structural differentiation of NLCDV community across ecological zones. 563(A) Venn diagrams showing the numbers of shared or unique NLCDVs 564phylotypes across size fractions (left), depths (center) and biomes (right). (B) 565Map showing the number of total, unique and shared NCLDVs across nine 566 oceanic regions. The map was drawn using the R package 'maps' 567 (https://cran.r-project.org/web/packages/maps). (C) Relationships among 568sample size and total or unique NCLDVs detected in each region. 569Abbreviations: SO: Southern Ocean; RS: Red Sea; MS: Mediterranean Sea; 570NPO: North Pacific Ocean; NAO: North Atlantic Ocean; SAO: South Atlantic 571Ocean; SPO: South Pacific Ocean; IO: Indian Ocean; AO: Arctic Ocean.
- Figure 4 Phylogenetic affiliations of environmental NCLDVs and their dispersal 572573characteristics. Phylogenetic tree constructed from 905 long (≥700 amino 574acid) PolB sequences from the OM-RGC.v2 and 67 known NCLDV sequences (see also Extended Data Figs. 2-4 for details). The first six layers indicate the 575576occurrence of NCLDVs unique to each size fraction, depth and biome. The 577 outside layer denotes phylogenetic positions of known sequences (color code as in the legend) and the phylotypes closely related (>90% amino acid identity) 578to those of NCLDV MAGs having chrysophyte homologs (indicated in yellow). 579580Abbreviations: OLPV-2: Organic Lake phycodnavirus 2; OLPV-1: Organic Lake phycodnavirus 1; CeV: Chrysochromulina ericina virus 1; PgV: 581582Phaeocystis globosa virus 16T; HeV: Haptolina ericina virus RF02; PkV-2; 583Prymnesium kappa virus RF02; TetV-1: Tetraselmis virus 1; PoV: 584Pyramimonas orientalis virus 1: AaV: Aureococcus anophagefferens virus 585BtV-01; PkV-1; Prymnesium kappa virus RF01; ChoanoV: ChoanoVirus; 586 CroV: Cafeteria roenbergensis virus BV-PW1; MpV-1: Micromonas sp.

587 RCC1109 virus MpV1; OlV-1: Ostreococcus lucimarinus virus 1; Otv-1: 588Ostreococcus tauri virus 1; Otv-2: Ostreococcus tauri virus 2; MpV-12T: 589Micromonas pusilla virus 12T: BpV-1: Bathycoccus sp. RCC1105 virus; BCV-590FR483: Paramecium bursaria Chlorella virus FR-483: ACTV-1: 591Acanthocystis turfacea Chlorella virus 1; PBCV-1: Paramecium bursaria Chlorella virus 1; EhV-86: Emiliania huxleyi virus 86; FsV: Feldmannia 592593species virus; EsV-1: Ectocampus siliculou virus 1; P. salinus: Pandoravirus 594salinus; P. dulcis: Pandoravirus dulcis; HaV-1: Heterosigma akashiwo virus 5951.

596Figure 5 Associations between NCLDVs and eukaryotic communities. (A) Partial 597 Mantel correlation coefficients (Spearman's ρ) between NCLDVs and 598eukaryotic communities. Each plot shows the value of ρ computed based on pico- (x-axis) and femto-sized (y-axis) NCLDV communities. Known virus-599host associations are shown as red dots. (B) Histogram and density estimates 600 601 showing the distribution of ρ values in known (red) and unknown (gray) pairs. (C) Pairwise comparisons of the partial Mantel correlation coefficients between 602 NCLDV and eukaryotic lineages. Correlation coefficients $\rho > 0.53$ based on 603 pico-size NCLDV communities are drawn as edges. Known virus-host 604 605 associations are shown in red, whereas unknown associations are shown in gray.

606 Figure 6 Vertical linkage of NCLDV communities between the surface and 607 mesopelagic layers. (A) Latitudinal trend in NCLDV community similarity 608 between two depths (with the station numbers). Relationship between NCLDV 609 vertical similarity and (B) the surface chlorophyll a biomass, (C) NCLDV richness in the mesopelagic layer, (D) sampling depth of mesopelagic seawater, 610 611 (E) the mixed layer depth and (F) temperature difference between epipelagic and mesopelagic samples. All NCLDV data were generated based on the pico-612613 size fraction. Shaded areas represent 90% confidence intervals.

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837 Author contributions

HE and HO designed the study. HE performed most of the bioinformatics analysis.
RB-M and YL contributed to the bioinformatics analysis. GS, NH, KL, CdV, MBS, CB,
PW, LK-B, and SS contributed to the generation of primary data. CdV, MBS, CB, PW,





- 841 LK-B, SS, and HO coordinated Tara Oceans. All authors contributed to the writing of the
- 842 manuscript.
- 843

844 Materials & Correspondence

845 Correspondence and material requests should be addressed to HO (email: 846 ogata@kuicr.kyoto-u.ac.jp).

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848 **Competing financial interests**

- 849 The authors declare no competing financial interests.
- 850
- 851



detected Number of phylotypes









Mantel correlation ρ by Pico-sized NCLDVs





Depth of MES sample (m)



200

400

Mixed layer depth (m)

600

0.4

0.2 -

0.0





Temperature difference between two sample depths (°C)