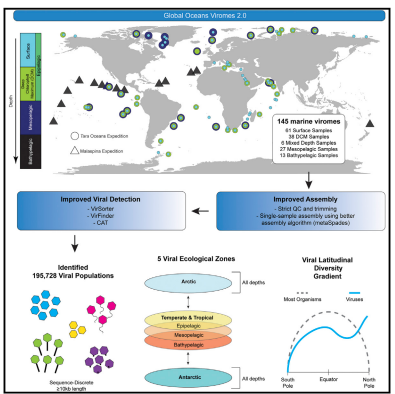
# Cell

# Marine DNA Viral Macro- and Microdiversity from Pole to Pole

## **Graphical Abstract**



### **Authors**

Ann C. Gregory, Ahmed A. Zayed, Nádia Conceição-Neto, ..., Shinichi Sunagawa, Patrick Wincker, Matthew B. Sullivan

## Correspondence

mbsulli@gmail.com

## In Brief

A global survey of ocean virus genomes vastly expands our understanding of this understudied community and reveals the Arctic as unexpected hotspot for viral biodiversity.

## **Highlights**

- Metagenomic assembly of 145 marine viromes uncovered 195,728 viral populations
- Read mapping revealed discrete sequence boundaries among >99% viral populations
- Viral communities separated into five distinct ecological zones in the global ocean
- Viral macro- and microdiversity did not follow the latitudinal diversity gradient





Article

# Marine DNA Viral Macro- and Microdiversity from Pole to Pole

Ann C. Gregory,<sup>1,24</sup> Ahmed A. Zayed,<sup>1,24</sup> Nádia Conceição-Neto,<sup>2,3</sup> Ben Temperton,<sup>4</sup> Ben Bolduc,<sup>1</sup> Adriana Alberti,<sup>5,17</sup> Mathieu Ardyna,<sup>6,25</sup> Ksenia Arkhipova,<sup>7</sup> Margaux Carmichael,<sup>8,17</sup> Corinne Cruaud,<sup>9,17</sup> Céline Dimier,<sup>6,10,17</sup> Guillermo Domínguez-Huerta,<sup>1</sup> Joannie Ferland,<sup>11</sup> Stefanie Kandels,<sup>12,13</sup> Yunxiao Liu,<sup>1</sup> Claudie Marec,<sup>11</sup> Stéphane Pesant,<sup>14,15</sup> Marc Picheral,<sup>6,17</sup> Sergey Pisarev,<sup>16</sup> Julie Poulain,<sup>5,17</sup> Jean-Éric Tremblay,<sup>11</sup> Dean Vik,<sup>1</sup> Tara Oceans Coordinators, Marcel Babin,<sup>11</sup> Chris Bowler,<sup>10,17</sup> Alexander I. Culley,<sup>18</sup> Colomban de Vargas,<sup>8,17</sup> Bas E. Dutilh,<sup>7,19</sup> Daniele Iudicone,<sup>20</sup> Lee Karp-Boss,<sup>21</sup> Simon Roux,<sup>1,26</sup> Shinichi Sunagawa,<sup>22</sup> Patrick Wincker,<sup>5,17</sup>

<sup>1</sup>Department of Microbiology, The Ohio State University, Columbus, OH 43210, USA

<sup>2</sup>Department of Microbiology and Immunology, Rega Institute for Medical Research, Laboratory of Viral Metagenomics, KU Leuven-University of Leuven, Leuven, Belgium

<sup>3</sup>Department of Microbiology and Immunology, Rega Institute for Medical Research, Laboratory for Clinical and Epidemiological Virology, KU Leuven-University of Leuven, Leuven, Belgium

<sup>4</sup>School of Biosciences, University of Exeter, Exeter, UK

<sup>5</sup>Génomique Métabolique, Genoscope, Institut François Jacob, CEA, CNRS, Univ Evry, Université Paris-Saclay, 91057 Evry, France

<sup>6</sup>Sorbonne Université, CNRS, Laboratoire d'Océanographie de Villefanche, LOV, 06230 Villefranche-sur-mer, France

<sup>7</sup>Theoretical Biology and Bioinformatics, Utrecht University, Utrecht, the Netherlands

<sup>8</sup>Sorbonne Université, CNRS, Station Biologique de Roscoff, AD2M ECOMAP, 29680 Roscoff, France

<sup>9</sup>CEA-Institut de Biologie François Jacob, Genoscope, Evry 91057, France

<sup>10</sup>Institut de Biologie de l'ENS (IBENS), Département de biologie, École normale supérieure, CNRS, INSERM, Université PSL, 75005 Paris, France

<sup>11</sup>Département de biologie, Québec Océan and Takuvik Joint International Laboratory (UMI 3376), Université Laval (Canada)-CNRS (France), Université Laval, Québec, QC G1V 0A6, Canada

<sup>12</sup>Structural and Computational Biology, European Molecular Biology Laboratory, 69117 Heidelberg, Germany

<sup>13</sup>Directors' Research, European Molecular Biology Laboratory, 69117 Heidelberg, Germany

<sup>14</sup>PANGAEA, Data Publisher for Earth and Environmental Science, University of Bremen, 28359 Bremen, Germany

<sup>15</sup>MARUM, Bremen University, 28359 Bremen, Germany

<sup>16</sup>Shirshov Institute of Oceanology of Russian Academy of Sciences, 36 Nakhimovsky prosp, 117997 Moscow, Russia

<sup>17</sup>Research Federation for the study of Global Ocean Systems Ecology and Evolution, FR2022/Tara Oceans GOSEE, 3 rue Michel-Ange, 75016 Paris, France

<sup>18</sup>Département de biochimie, microbiologie et bio-informatique, Université Laval, Québec, QC G1V 0A6, Canada

<sup>19</sup>Centre for Molecular and Biomolecular Informatics, Radboud University Medical Centre, Nijmegen, the Netherlands

<sup>20</sup>Stazione Zoologica Anton Dohrn, Villa Comunale, 80121 Naples, Italy

<sup>21</sup>School of Marine Sciences, University of Maine, Orono, ME, USA

<sup>22</sup>Department of Biology, Institute of Microbiology and Swiss Institute of Bioinformatics, ETH Zurich, 8093 Zurich, Switzerland

<sup>23</sup>Department of Civil, Environmental and Geodetic Engineering, The Ohio State University, Columbus, OH 43210, USA

<sup>24</sup>These authors contributed equally

<sup>25</sup>Present address: Department of Earth System Science, Stanford University, Stanford, CA 94305, USA

<sup>26</sup>Present address: Department of Energy Joint Genome Institute, Walnut Creek, CA 94598, USA

<sup>27</sup>Lead Contact

\*Correspondence: mbsulli@gmail.com

https://doi.org/10.1016/j.cell.2019.03.040

#### **SUMMARY**

Microbes drive most ecosystems and are modulated by viruses that impact their lifespan, gene flow, and metabolic outputs. However, ecosystem-level impacts of viral community diversity remain difficult to assess due to classification issues and few reference genomes. Here, we establish an  $\sim$ 12-fold expanded global ocean DNA virome dataset of 195,728 viral populations, now including the Arctic Ocean, and validate that these populations form discrete genotypic clusters. Meta-community analyses revealed five ecological zones throughout the global ocean, including two distinct Arctic regions. Across the zones, local and global patterns and drivers in viral community diversity were established for both macrodiversity (inter-population diversity) and microdiversity (intra-population genetic variation). These patterns sometimes, but not always, paralleled those from macro-organisms and revealed temperate and tropical surface waters and the Arctic as biodiversity hotspots and mechanistic hypotheses to explain them. Such further understanding of ocean viruses is critical for broader inclusion in ecosystem models.

#### INTRODUCTION

Biodiversity is essential for maintaining ecosystem functions and services (for review, see Tilman et al., 2014). In the oceans, the vast majority of biodiversity is contained within the microbial fraction containing prokaryotes and eukaryotic microbes, which represents ~60% of its biomass (Bar-On et al., 2018). Meta-analyses looking at changes in marine biodiversity show that biodiversity loss increasingly impairs the ocean's capacity to produce food, maintain water quality, and recover from perturbations (Worm et al., 2006). To date, marine conservation efforts have focused on specific organismal communities, such as fisheries or coral reefs, rather than conserving whole ecosystem biodiversity. However, emerging studies across diverse environments show that the stability and diversity of higher trophic level organisms rely upon diversity throughout the food web (Soliveres et al., 2016). Despite being the foundation of the food web, most marine microbial biodiversity numbers are based on a few well-studied locations (e.g., Hawaii Ocean Time Series, Bermuda Atlantic Time Series, and San Pedro Ocean Time Series). For ocean microbes and their viruses, global surveys that parallel century-old global terrestrial and decades-old marine macro-organismal global biodiversity surveys (Reiners et al., 2017) are only now emerging (de Vargas et al., 2015; Sunagawa et al., 2015; Brum et al., 2015; Roux et al., 2016; Ser-Giacomi et al., 2018) (Table S1). Key to assessing biodiversity changes across marine ecosystems is improving our understanding of current microbial biodiversity levels, distribution patterns, and their ecological drivers.

Despite their tiny size, viruses play a large role in marine ecosystems and food webs. For example, mortality due to viruses is credited with lysing  $\sim 20\%$ -40% of bacteria per day and releasing carbon and other nutrients that impact the food web (for review, see Suttle, 2007). Beyond mortality, viruses can alter evolutionary trajectories of microbial communities by transferring  $\sim 10^{29}$  genes per day globally (Paul, 1999) and biogeochemical cycling by metabolically reprogramming host photosynthesis, as well as central carbon metabolism and nitrogen and sulfur cycling (for review, see Hurwitz and U'Ren, 2016). Finally, as the oceans are estimated to capture half of human-caused carbon emissions (Le Quéré et al., 2018), it is notable that genes-to-ecosystems modeling has placed viruses as central players of the ocean "biological pump" (Guidi et al., 2016). Many of these discoveries are very recent as ocean viral genome sequence space is just now being explored at the level of viral macrodiversity (i.e., inter-population diversity) throughout the global oceans-at least for the most abundant double-stranded DNA viruses sampled (Table S2).

In spite of this progress in studying marine viral macrodiversity, virtually nothing is known about microdiversity (i.e., intra-population genetic variation). This is due to the controversy surrounding the existence of viral species (Gregory et al., 2016; Bobay and Ochman, 2018). In eukaryotic organisms, where species boundaries are more widely accepted, such microdiversity has been studied and is thought to drive adaptation and speciation to promote and maintain stability in ecosystems (Hughes et al., 2008; Larkin and Martiny, 2017). This is likely also true in viruses because even a few mutations can alter host interactions and ecological and evolutionary dynamics for the genotype (Marston et al., 2012; Petrie et al., 2018). In nature, viral microdiversity measurements have been limited to marker genes (e.g., genes encoding major capsid proteins), which capture neither community-wide variability (Sullivan, 2015) nor genome-wide evidence of selection (Achtman and Wagner, 2008). Recently, deeper metage-nomic sequencing and population genetic theory-grounded species delimitations (Shapiro et al., 2012; Cadillo-Quiroz et al., 2012) have begun to reveal such microdiversity in microbes, and this has elucidated unknown features of speciation, adaptation, pathogenicity, and transmission (Snitkin et al., 2011; Schloissnig et al., 2013; Rosen et al., 2015; Lee et al., 2017; Smillie et al., 2018). Although parallel species delimitations are now available for viruses (Gregory et al., 2016; Bobay and Ochman, 2018), no datasets are yet available to explore genome-wide microdiversity in viruses, particularly at the global scale.

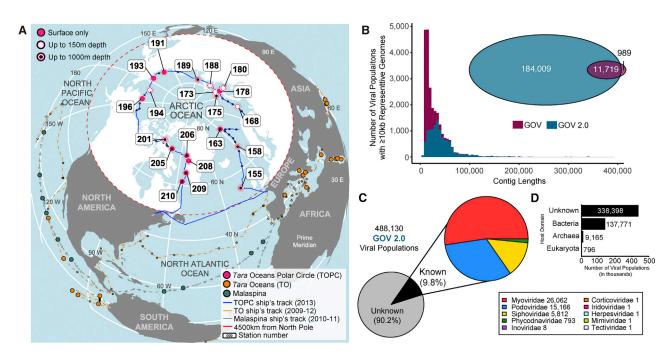
Here, we leverage the Tara Oceans global oceanographic research expedition sampling to establish a deeply sequenced, global-scale ocean virome dataset and use it to assess the validity of the current viral population definition and to establish and explore baseline macro- and micro-diversity patterns with their associated drivers across local to global scales. These data have been collected and analyzed in the context of the larger Tara Oceans Consortium systematically sampled, global-scale, viruses-to-fish-larvae datasets (de Vargas et al., 2015; Sunagawa et al., 2015; Brum et al., 2015; Lima-Mendez et al., 2015; Pesant et al., 2015; Roux et al., 2016) and help establish foundational ecological hypotheses for the field and a roadmap for the broader life sciences community to better study viruses in complex communities.

#### **RESULTS AND DISCUSSION**

#### **The Dataset**

The Global Ocean Viromes 2.0 (GOV 2.0) dataset is derived from 3.95 Tb of sequencing across 145 samples distributed throughout the world's oceans (Figure 1A; Table S3; STAR Methods). These data build on the prior GOV dataset (Roux et al., 2016) by increased sequencing for mesopelagic samples (defined in our dataset as waters between 150 m to 1,000 m) and upgrading assemblies, both of which drastically improved sampling of the ocean viruses in these samples (results below). Additionally, we added 41 new samples derived from the Tara Oceans Polar Circle (TOPC) expedition, which traveled 25,000 km around the Arctic Ocean in 2013. These 41 Arctic Ocean viromes were generated to represent the most significantly climate-impacted region of the ocean and an extreme environment. No such metagenome-based viral data exist for the Arctic region (Deming and Collins, 2017), and more generally, for many planktonic organisms, systematic sampling is uneven throughout the Arctic Ocean (Circumpolar Biodiversity Monitoring Program, 2017) due to geopolitical and physical challenges of sampling these regions.

The first step to studying viral biodiversity from the assembled GOV 2.0 dataset (Figure S1A; STAR Methods) was to identify contigs that likely derive from viruses using tools that collectively utilize homology to viral reference databases, probabilistic models on viral genomic features, and viral k-mer signatures (STAR Methods). These putative viral contigs were then



#### Figure 1. The Global Ocean Viromes 2.0

(A) Arctic projection of the global ocean highlighting the new sampling stations of viromes in the GOV 2.0 dataset. Datasets from non-arctic samples were previously published in Brum et al. (2015) and Roux et al. (2016).

(B) Histograms of the average assembled contig lengths for viral populations >10 kb shared between GOV and GOV 2.0. Inset: more than 92% of the unbinned GOV viral populations were reassembled and identified in GOV 2.0 >10 kb populations.

(C) Pie charts showing how many of the 488,130 total viral populations comprising GOV 2.0 can be annotated and, of those, their viral family level taxonomy. (D) Barplot showing the host affiliations for each viral population at the domain level.

See also Figures S1 and S7 and Tables S1, S2, and S3.

assigned to "populations," which are currently defined as viral contigs  $\geq 10$  kb where  $\geq 70\%$  of the shared genes have  $\geq$ 95% average nucleotide identity (ANI) across its members (Brum et al., 2015; Roux et al., 2016, 2018) (population definition also discussed below). This process identified 195,728 viral populations in the GOV 2.0 dataset, which is an  ${\sim}12$ -fold increase over the 15,280 identified in the original GOV dataset and assemblies (Roux et al., 2016) and augments prior marine viromic work (Table S2). Of these original GOV viral populations, 12,708 were represented by single contigs and, of these, most (92%) were recovered in GOV 2.0 (Figure 1B, inset), with average lengths increased 2.4-fold from 18 kbp to 44 kbp (Figure 1B). Outside these GOV-known and now improved viral populations, an additional 180,448 new GOV 2.0 viral populations were identified-derived mostly (58%) from improved assemblies and deeper sequencing of the original GOV samples and the rest (42%) from the 41 new Arctic Ocean viromes. Finally, new methods to identify shorter viral contigs (STAR Methods) were applied and these identified another 292,402 contigs as viral (5-10 kb length and/or circular), which, when added to the earlier data and clustered at  $\geq$  95% ANI, resulted in a total of 488,130 viral populations (N50 = 15,395; L50 = 105,286; mean read depth per population = 17x). Ninety percent of the populations could not be taxonomically classified to a known viral family, but the 10% that could were predominantly dsDNA viral families and bacteriophages (Figures 1C and 1D).

Although the focus of this study is DNA viruses, a remarkable diversity of RNA viruses has been described in nature, although largely outside of marine systems. For example, transcriptome sequencing from plants (Roossinck et al., 2010), arthropods (Shi et al., 2016), and birds and bats (for review, see Greninger, 2018) have shown a genomic and phylogenetic diversity of RNA viruses far beyond those in culture (Shi et al., 2018). In the oceans, however, RNA viral diversity and abundance remains largely unknown. The few estimates of marine RNA virus abundance are based on the relative quantification of RNA and DNA from purified viral particles and genome size extrapolations and suggest that up to half of the viral particles in seawater are RNA viruses (Steward et al., 2013; Miranda et al., 2016). Direct RNA virus counts are not yet available for any environment due to the lack of RNA-specific stains. To date, our understanding of marine RNA viral diversity is based on singlegene surveys that target subgroups of viruses (for review, see Culley, 2018) and a few viromes generated from extracellular viral particles (Culley and Steward, 2007; Culley et al., 2006; Miranda et al., 2016; Steward et al., 2013; Urayama et al., 2018; Zeigler Allen et al., 2017) or from RNA viral sequences identified in metatranscriptomes (Carradec et al., 2018; Moniruzzaman et al., 2017; Urayama et al., 2018; Zeigler Allen et al., 2017). Together, these studies suggest that the marine RNA virosphere is composed of a large diversity of positive-polarity single-stranded RNA (ssRNA) and double-stranded RNA (dsRNA) viruses diverge from established taxa, with an apparent predominance of viruses that infect eukaryotes (Culley, 2018). Due to current methodological limitations, comprehensive, systematic assessments of marine RNA viral diversity on the global scale are not yet available and are excluded from our analysis.

#### Validating Viral "Population" Boundaries

Defining species is controversial for eukaryotes and prokaryotes (Kunz, 2013; Cohan, 2002; Fraser et al., 2009) and even more so for viruses (Bobay and Ochman, 2018), largely because of the paradigm of rampant mosaicism stemming from rapidly evolving ssDNA and RNA viruses, whose evolutionary rates are much higher than dsDNA viruses (for review, see Duffy et al., 2008). The biological species concept, often referred to as the gold standard for defining species, defines species as interbreeding individuals that remain reproductively isolated from other such groups. To adapt this to prokaryotes and viruses, studies have explored patterns of gene flow to determine whether they might maintain discrete lineages as reproductive isolation does in eukaryotes. Indeed, gene flow and selection define clear boundaries between groups of bacteria, archaea, and viruses, although the required scale of data are only available for cyanophages and mycophages among viruses (Shapiro et al., 2012; Cadillo-Quiroz et al., 2012; Gregory et al., 2016; Bobay and Ochman, 2018).

Because measuring gene flow requires extensive datasets not yet available for many groups, the term "species" is rarely used for prokaryotes or viruses, and instead discrete lineages are described as "populations." Separate from these population genetic theory grounded observations, evidence of discrete lineages, or sequence-discrete populations, is to use metagenomic read-mapping to evaluate naturally occurring sequence variation across organisms. Sequence-discrete populations have now been observed for prokaryotes (Konstantinidis and Tiedje, 2005) and more recently for some dsDNA viruses (viral-tagged metagenomes and 142 isolate genomes for marine cvanophages) (Deng et al., 2014; Gregory et al., 2016) (Table S4). Buoyed by this and signatures of at least some double-stranded DNA (dsDNA) viruses obeying the biological species concept (Bobay and Ochman, 2018), viral ecologists have established the definition of viral populations described above (Brum et al., 2015; Roux et al., 2016, 2018). Notably, however, only deeply sequenced groups, cyano- and mycophages, have been evaluated to date (Gregory et al., 2016; Bobay and Ochman, 2018), and an emergent hypothesis suggests that phages evolve with different modes and tempos driven by differing temperate or obligately lytic lifestyles (Mavrich and Hatfull, 2017). Thus, there is a need to evaluate how generalizable this empirically derived  $\geq$ 95% ANI cut-off viral population definition is in nature.

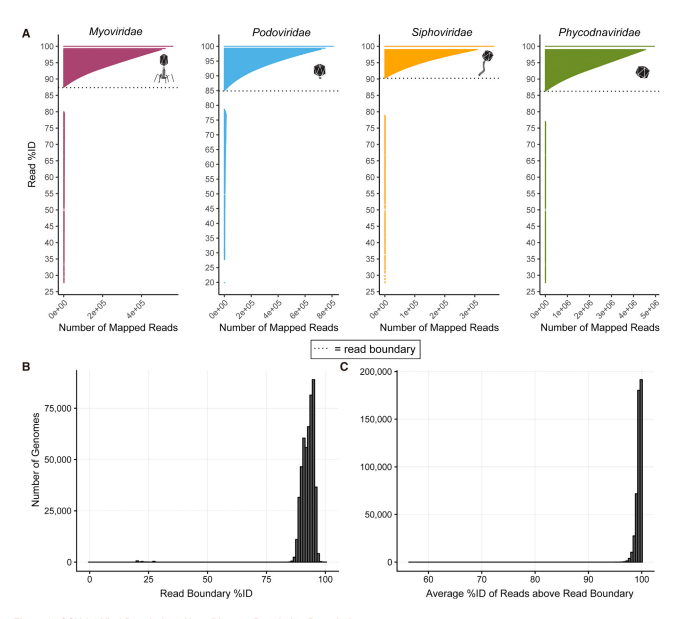
To test this, we permissively mapped metagenomic reads against our 488,130 GOV 2.0 viral populations by allowing "local" matching as low as 18% nucleotide identity and statistically identifying "breaks" in the resulting read frequency histograms (STAR Methods). This revealed that, on average, the break occurred such that reads <92% nucleotide identity failed to map (Figure 2C; Table S5 for full results), which resulted in a genome-wide signature of  $\geq$ 95% ANI for nearly all (99.9% or 487,875) of the GOV 2.0 viral populations, including the smaller

<10 kb viral populations (Figure 2D). This implies that the observed viral populations in the dataset are predominantly and detectably sequence-discrete. This result is consistent with data from viral-tagged metagenomes (Deng et al., 2014) and gene-sharing networks of prokaryotic virus genomes (Iranzo et al., 2016; Bolduc et al., 2017), which also showed that sampled viral genome sequence space is clustered at each "species" and "genus" levels, respectively. Thus, while ssDNA and RNA viruses have variable and elevated genome evolutionary rates that can erode species boundaries (for review, see Duffy et al., 2008), it appears that virtually all metagenome-assembled dsDNA viral populations form discrete genotypic clusters and can be appropriately delineated via a  $\geq 95\%$  genome-wide ANI cut-off.

#### Meta-Community Analysis Reveals Five Ecological Zones

Having organized this global sequence space into discrete and biologically meaningful populations, we next sought to use metagenome-derived abundance estimates to establish patterns and drivers of viral population diversity across the global ocean across multiple levels of ecological organization (Figure 3). This revealed that the 145 GOV 2.0 viral communities robustly assorted into just five meta-communities, denoted ecological zones, whether assessed using Bray-Curtis dissimilarity distances in principal coordinate analysis (Figure 4A), non-metric multidimensional scaling (Figure S2A), or hierarchical clustering (Figure S2B) and after accounting for variable sample sizes (see STAR Methods and Figure S3). We designated these five emergent ecological zones as the Arctic (ARC), Antarctic (ANT), bathypelagic (BATHY), temperate and tropical epipelagic (TT-EPI), and mesopelagic (TT-MES) and used these for further study. Depth ranges overlapped with those previously defined (Reygondeau et al., 2018), with epipelagic, mesopelagic, and bathypelagic being waters of depths 0-150 m, 150-1,000 m, and deeper than 2.000 m. respectively.

Comparison of our virome-inferred ecological zones to those inferred for the oceans in other ways was telling. Our zones differed from traditional oceanographic biogeographical biomes (e.g., Longhurst), where four biomes and  $\sim$ 50 provinces have been designated across surface ocean waters based on annual cycles of nutrient chlorophyll a (Longhurst et al., 1995; Longhurst, 2007), and from mesopelagic ecoregions and biogeochemical provinces based on biogeography and environmental climatology, respectively (Sutton et al., 2017; Reygondeau et al., 2018). However, they were similar to those observed for marine bacterial communities, which clustered by mid-latitude surface, high-latitude, and deep waters (Ghiglione et al., 2012). This implies that the physicochemical structuring of marine microbial communities is likely the most important factor in structuring marine viral communities, perhaps reflecting a relative stability in host range of viruses in the oceans (de Jonge et al., 2019). To evaluate this physicochemical structuring, we examined the universal predictors and drivers of viral ecological zones, across one (Figure 5A) and multiple ordination dimensions (Figure 5B; STAR Methods). This suggested that temperature was the major driver structuring these ecological zones, as previously shown from global microbial surveys (Sunagawa



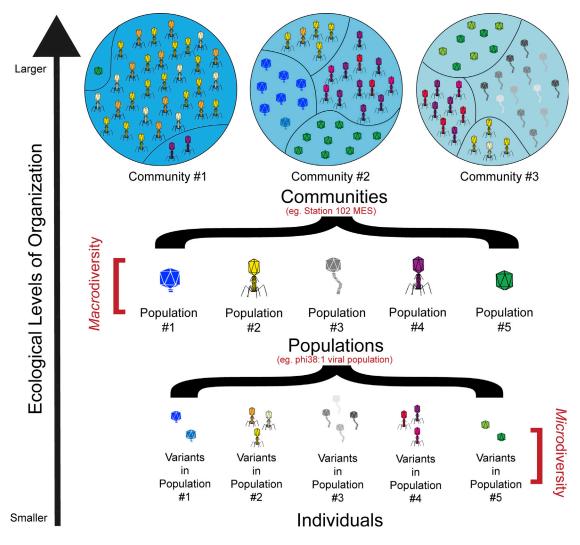


(A) Barplots showing the read mapping results for the most abundant viral population >10 kb in length for each of the top four viral families. Despite differences in read boundaries across the representative viral populations, there is no difference in the average read boundaries across the different viral families.(B) Histogram showing the read distribution frequency break (i.e., read boundary) between spuriously mapped reads and legitimate reads mapping to the genome.

(C) Histograms showing the average percent identity of reads mapped to each genome after removing spuriously mapped reads. See also Tables S4 and S5.

et al., 2015) and our own smaller ocean virome surveys, where we posited previously that temperature likely directly impacts microbial community structure, and indirectly viral community structure (Brum et al., 2015). Moreover, temperature has been shown to play an important role in virus-host interactions, especially in the Arctic (Maat et al., 2017).

To look for specific viral adaptations in each ecological zone, we identified genes under positive selection by evaluating the ratio of non-synonymous to synonymous mutations observed in gene sequences using the pN/pS equation (Schloissnig et al., 2013). Of 1,139,501 genes tested from populations with enough coverage ( $\geq$ 10× mean read depth; mean number of populations assessed per sample: 14,852 viral populations), 124,882 genes were identified as being under positive selection in at least one sample. Most (82%) of the positively selected genes were functionally unannotatable, with the remaining 18% annotatable as predominantly genes related to structure or DNA metabolism (Table S6). In model systems, such genes are often under strong selective pressures during adaptations to new hosts (Marston et al., 2012; Jian et al., 2012; Enav et al., 2018). Thus, we



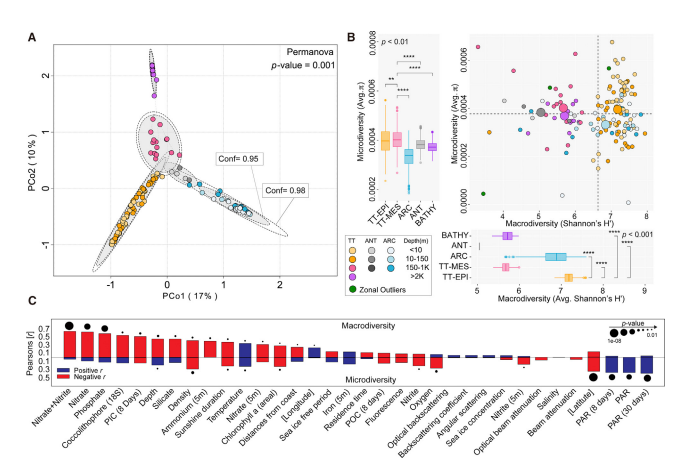
#### Figure 3. Ecological Levels of Organization

Schematic showing the different ecological levels of organization studied in this paper.

hypothesize that host availability in each ecological zone is a strong selective pressure on our marine viral populations. Given the lack of functional annotations for most of the genes, we clustered all translated GOV 2.0 viral genes into protein clusters (PCs) based on sequence homology (Sensu) (Holm and Sander, 1998) to identify positively selected zone-specific PCs. This resulted in 823,193 PCs, of which ~10% (79,588 PCs) appeared under positive selection, with a subset of these specific to a single zone (ARC = 80%; ANT = 33%; BATHY = 37%; TT-EPI = 75%; TT-MES = 69% of positively selected PCs per zone; see Table S6). These findings of many zone-specific positively selected PCs is indicative of niche-differentiation. However, functional stories from these data are challenging as 85% of these zone-specific PCs were of unknown function, with the remaining mostly being the structural and DNA metabolism genes described above. This suggests that we have a lot to learn about the function of genes that most likely drive niche-differentiation across the ecological zones.

# Viral Macro- and Microdiversity and Potential Drivers within and between Ecological Zones

To explore diversity patterns across ecological zones, we calculated per sample diversity using Shannon's H' for macrodiversity and a newly established method for community-wide microdiversity. This new method for community-wide microdiversity is limited in that it can only assess well-sampled, abundant populations because it estimates the average nucleotide diversity (or  $\pi$ ) from the mean of  $\pi$  from 100 randomly subsampled wellsequenced populations sampled 1,000 times (STAR Methods). These zone-normalized (STAR Methods) comparisons revealed that macrodiversity was highest in TT-EPI (p < 0.05), closely followed by the ARC, and lowest in TT-MES and ANT (Figure 4B, bottom), whereas microdiversity was highest in TT-MES (p < 0.05) and lowest in ARC (Figure 4B, left). At the zonal level, a negative trend between macro- and microdiversity emerges (Figure 4B, right), although we note that the small number of zonal points limits our statistical inferences, even in this global dataset.



#### Figure 4. Viral Communities Partition into Five Ecological Zones with Different Macro- and Microdiversity Levels

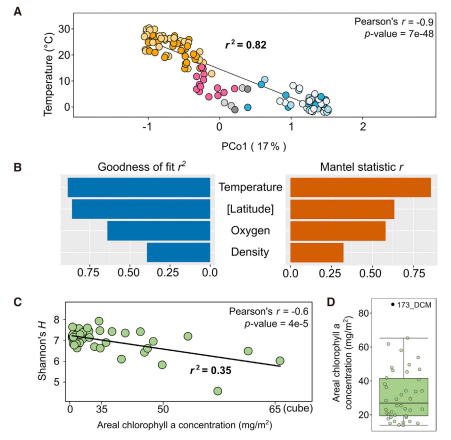
(A) Principal coordinate analysis (PCoA) of a Bray-Curtis dissimilarity matrix calculated from GOV 2.0. Analyses show that viromes significantly (Permanova p = 0.001) structure into five distinct global ecological zones: ARC, ANT, BATHY, TT-EPI, and TT-MES zones. Ellipses in the PCoA plot are drawn around the centroids of each group at 95% (inner) and 97.5% (outer) confidence intervals. Four outlier viromes that did not cluster with their ecological zones were removed (Figure S3A) and all the sequencing reads were used (see Figure S3B and STAR Methods).

(B) Right: scatterplots showing correlations between macrodiversity (Shannon's H') and microdiversity (average  $\pi$  for viral populations with  $\geq$  10× median read depth coverage; see STAR Methods) values for each sample across GOV 2.0. The larger circles represent the average per zone. Left: boxplots showing median and quartiles of average microdiversity per ecological zone. Bottom: boxplots showing median and quartiles of macrodiversity for each ecological zone. Zonal samples were randomly downsampled to n = 5 to account for zone sampling difference. All pairwise comparisons shown were statistically significant (\*\* p < 0.01 and \*\*\*\* p < 0.001) using two-tailed Mann-Whitney U tests.

(C) Positive (blue) and negative (red) Pearson's correlation results comparing macrodiversity (top) and microdiversity (bottom) with different biogeographical and biogeochemical parameters at the global scale (see Figure S3E; Table S3 for all abbreviations; STAR Methods). The significance of the correlations is indicated by the size of the black circles on top of the bars, and the variables on the x axis are ordered from the strongest to the weakest correlation with macrodiversity (except for the top four variables correlating with microdiversity for readability).

See also Figures S1, S2, S4, and S7 and Tables S6 and S7.

Recent work suggests that higher microdiversity can impede the maintenance of macrodiversity by promoting competitive exclusion (Hart et al., 2016). Thus, we posit that, if the zonal level negative macro-/microdiversity trends are real, this may result from increased intrapopulation niche variation that reduces interpopulation niche variation resulting in competitive exclusion by the superior competitors, which may occur slowly and may be why it only appears at this regional scale (Figure S4). Because estimates of microdiversity in our dataset and even currently available single virus genomics approaches (Martínez-Hernández et al., 2017) remain limited to only the most abundant populations, testing such a hypothesis awaits critically needed advances and scalability in single-virus genomics technologies. At the per-sample level, however, macro- and microdiversity were not correlated, even within each zone (Figure 4B, right). Although these are the first data available for viruses, for larger organisms, macro- and microdiversity are often correlated across habitats sharing similar species pools, presumably due to habitat characteristics altering immigration, drift, and selection (Vellend and Geber, 2005). These ecological correlations are generally positive and significantly stronger in discrete habitats (e.g., islands) in contrast to more connected communities like the ocean (for review, see Vellend et al., 2014). Thus, we posit that the lack of correlation between marine viral macro- and microdiversity at this per-sample level is driven by differences in local drivers (Figure 4C). Consistent with this, local potential



drivers differed as nutrients strongly (and negatively) correlated with viral macrodiversity, whereas photosynthetically active radiation (PAR; an indicator of productivity) best (and positively) correlated with viral microdiversity in the epipelagic waters (Figure 4C).

Mechanistically, these results suggest several possible hypotheses. We interpret that, at the viral macrodiversity level, decreased host diversity in algal blooms, which themselves rely on nutrient pulses (Faroog and Malfatti, 2007), could skew viral rank abundance curves toward dominance by increasing abundance of bloom-associated viral populations. Even though algal blooms were not targeted in the Tara Oceans expedition, we did find that viral macrodiversity negatively correlated with chlorophyll a (Figure 5C), and particulate inorganic carbon concentration (PIC) (Figure 4C), which is commonly used as a proxy for coccolithophore abundance (Groom and Holligan, 1987). Additionally, viral macrodiversity negatively correlated with the relative abundance of coccolithophores based on the V9 region of the 18S rRNA genes in the sequencing reads (Figure 4C). For viral microdiversity in epipelagic waters, we interpret that PAR is potentially the main driver (Figure 4C). PAR is known to impact host diversity, particularly in nutrient-poor surface waters, by inhibiting photoautotrophs through overwhelming their photosystems with too many electrons that can back up and even damage the photosystems (Feng et al., 2015). Further PAR can inhibit the growth of the dominant heterotroph, SAR11 (Ruiz-González et al., 2013), and

# Figure 5. Ecological Drivers of Global Viral Macrodiversity

(A) Regression analysis between the first coordinate of a PCoA (Figure 4A) and temperature showed that samples were separated by their local temperatures with an  $r^2$  of 0.82.

(B) Potential ecological drivers & predictors of betadiversity across GOV 2.0 for the first two dimensions (goodness of fit  $r^2$  using a generalized additive model) and across all dimensions (Mantel test based on Spearman's correlation). Temperature was uniformly reported as the best predictor of viral beta-diversity globally.

(C) Regression analysis between viral macrodiversity at the deep chlorophyll maximum (DCM) layer and areal chlorophyll *a* concentration (after cube transformation) showed that the negative correlation between viral macrodiversity and nutrients (Figure 4C) is mediated (at least partially) by primary productivity. The Shannon's H outlier 32\_DCM (Figure S3) and a chlorophyll *a* concentration outlier (173\_DCM; D) have been excluded from the regression analysis.

(D) Boxplot analysis of areal chlorophyll *a* concentrations showing a single outlier concentration that fell above the fourth quantile of the data points (function geom\_boxplot of ggplot).

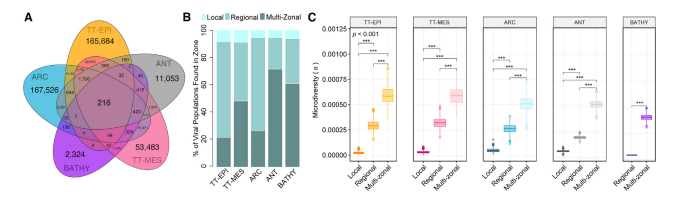
can stimulate other key microbes such as *Roseobacter, Gammaproteobacteria*, and NOR5 (Ruiz-González et al., 2013). We hypothesize that the shorter-term impacts of high PAR in the surface waters on host

communities may create new niches for viruses, whereby microdiversity increases to enable differentiation of existing viral populations. As above, advances in single-virus genomics would be invaluable for testing this hypothesis.

#### Viral Macro- and Microdiversity and Potential Drivers against Classical Ecological Gradients

Ecologists have long explored the relationship between diversity and geographic range, which in eukaryotes and bacteria are highly (and positively) correlated and thought to be due to the accumulation of niche-specific selective mutations across populations with large heterogeneous geographic ranges (i.e., the niche variation hypothesis) (Van Valen, 1965; Hedrick, 2006; Rosen et al., 2015). No parallel studies have looked at viruses. To explore this for viruses, we determined the geographic range of viral populations based on their distribution within and between ecological zones (Figure 6A) and then calculated their average  $\pi$  (STAR Methods) to assess patterns in macro- and microdiversity, respectively. Viral populations were designated as "multi-zonal" if they were observed in >1 ecological zone, "zone-specific regional" if they were observed in only one zone but  $\geq 2$  viral communities, or "zone-specific local" if they were observed in only 1 viral community within a single zone.

These analyses first revealed differences in the dominant viral geographic ranges across the different ecological zones. For example, multi-zonal viral populations dominated ANT and



#### Figure 6. Size of Geographic Range Positively Correlates with Microdiversity

(A) Venn diagram showing the number of viral populations found only in one zone (zone-specific) and those that are shared between and among the five ecological zones (multi-zonal).

(B) Stacked barplots showing the number of multi-zonal, regional, and local viral populations found within the species pool of each ecological zone.

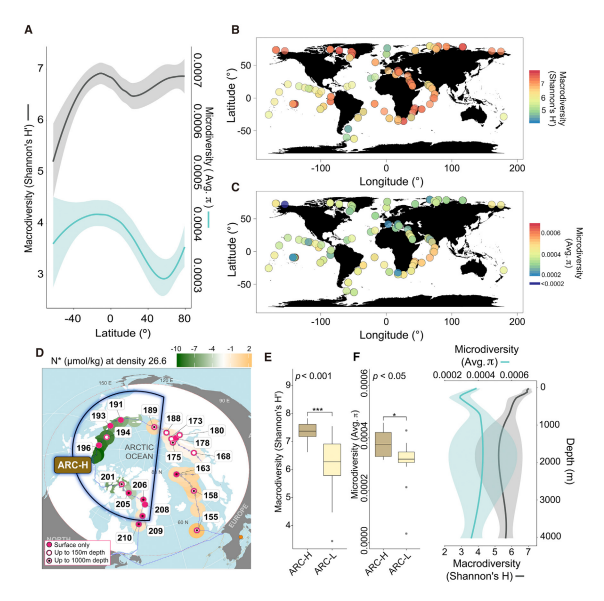
(C) Boxplots showing median and quartiles of microdiversity (average  $\pi$  for viral populations with  $\ge 10 \times$  median read depth coverage) per populations found within each zone defined as multi-zonal, regional, or local. Statistics were the same as in Figure 2. See also Figure S5.

BATHY (>60% of viral populations found within zone), both across the zone (Figure 6B) and within each station (Figure S5), whereas zone-specific regional viral populations dominated TT-EPI and ARC, and the multi-zonal and zone-specific viral populations were approximately equally represented in TT-MES (Figure 6B). The high levels of zone-specific viral populations in TT-EPI and ARC, as well as the high levels of viral macrodiversity (Figure 4B, bottom), are indicative of high endemism and suggest these regions may be biodiversity hotspots for marine viruses. In contrast, the ANT and BATHY are composed mostly of multi-zonal viral populations suggesting that they may be sink habitats that are more dependent on migration (Sensu) (Watkinson and Sutherland, 1995). However, across all ecological zones, viral population microdiversity increased with virus geographic range (Figure 6C; p < 0.05), presumably from varied ecologies providing differing selective niches for the single, widely distributed population that then drive differentiation through isolation-by-environment processes (Sensu) (Shapiro et al., 2012). Such findings are new for viruses, but parallel the results for eukaryotes (Hedrick, 2006) and bacteria (Rosen et al., 2015), and suggest a universality to isolation-by-environment processes across organismal kingdoms and viruses.

Ecologists have also long observed, across most flora and fauna, that there are latitudinal patterns in diversity across both terrestrial and marine environments. Briefly, the latitudinal diversity gradient suggests that both macro- and microdiversity are highest at mid-latitudes and decrease poleward (Pianka 1966; Hillebrand 2004; Mannion et al., 2014; Miraldo et al., 2016). We found that both viral macro- and microdiversity followed the latitudinal diversity gradient except in ARC, where both increased (Figure 7A). This high equatorial macro- and microdiversity was consistent across the Indian, Atlantic, and Pacific Oceans as expected (Figures 7B and 7C). The Arctic Ocean, however, was not only unexpectedly elevated in diversity, but it also displayed a unique pattern. Specifically, two distinct zones—definable by climatology-derived water mass nutrient stoichiometry (N\*) (Figure 7D; see "Comparing ARC-H and ARC-L" in STAR Methods)—emerged as high (ARC-H) and low (ARC-L) diversity regions that were significantly differentiable at both macro- and microdiversity levels (Figure 7E). Further, ARC-H was characterized by low nutrient ratios (N\*; >9× lower in ARC-H than ARC-L on average; p < 5E-04) and drove the divergence from the latitude diversity gradient (Figure S6A).

Mechanistically, we interpret these observations as follows. Prior work in this region has shown (1) strong denitrification in the Bering Strait (Devol et al., 1997), which explains the low N\* in the west, and (2) increasing oligotrophy in the Beaufort Gyre due to increasing vertical stratification, which selects against larger algae and for smaller algae and bacteria in the ARC-H (Li et al., 2009). As above, we hypothesize that shorter-term increased host diversity results in increased viral macro- and microdiversity in ARC-H. Although our GOV 2.0 dataset is confounded by seasonality of sampling, we posit that this elevated summertime macro- and microdiversity in ARC may fuel viral ecological differentiation and represent an unrecognized "cradle" of viral biodiversity beyond the tropics. Although this elevated diversity in the Arctic was surprising, together with a similar deviation seen in mollusks (Valdovinos et al., 2003) and recently reported in ray-finned fish (Rabosky et al., 2018), these results call into question whether this decades-old paradigm needs revisiting and suggests that polar regions may be important biodiversity hotspots for viruses, as well as larger organisms.

Finally, as ocean exploration accelerates (see Figure S7), patterns in diversity through the vertical layers of the ocean have become a focus. An emergent depth diversity gradient hypothesis suggests that macrodiversity decreases with depth (Costello and Chaudhary, 2017), which has been explored across the World Register of Marine Species that includes some microbes and viruses (http://www.marinespecies.org/), but microdiversity has not yet been explored for any organism. Overall, our virome-inferred diversity patterns were less obviously consistent with the depth diversity gradient, although deep water ocean



#### Figure 7. Viral Macro- and Microdiversity Global Biodiversity Trends

(A) Locally estimated scatterplot smoothing (LOESS) plots showing the latitudinal distributions of macro- and microdiversity.

(B) Equirectangular projections of the globe showing macrodiversity.

(C) Equirectangular projection of the globe showing microsdiversity levels within each sample across the global ocean. Samples collected at different depths from the same latitude and longitude are overlaid and the colors representing their macro- and microdiversity values are merged.

(D) Arctic projection of the global ocean showing the geographical division between ARC-H and ARC-L stations. The patterns are largely concordant with the Arctic division by climatology-derived N\*. While we did sample across different seasons, the calculated N\* values are not dependent on the season (see "Impact of the coast, depth, and seasons" in STAR Methods).

(E) Boxplots showing median and quartiles of macrodiversity (left) and microdiversity (right) of the ARC-H and ARC-L regions. Statistics were the same as in Figure 2.

(F) LOESS smooth plots showing the depth distributions of macro- and micropopulation diversity. On all the smooth plots, the line represents the LOESS best fit, while the lighter band corresponds to the 95% confidence window of the fit. Abbreviations: N\*, the departure from dissolved N:P stoichiometry in the Redfield ratio and a geochemical tracer of Pacific and Atlantic water mass (STAR Methods). See also Figure S6.

data were limited (Figure 7F). Briefly, viral macrodiversity largely followed the depth diversity gradient with high diversity in the surface waters and decreased diversity with depth, whereas viral microdiversity did not as it decreased until 200 m depth, but then

sharply increased (Figure 7F). This deep water increase coincided with an increase in bacterial macrodiversity in the mesopelagic region (Figures S6B and S6C), and in TT-MES, this bacterial macrodiversity correlated with viral microdiversity (Figure S6D).

If more extensive deep water sampling confirms these patterns, we see several scenarios that could explain these data. First, we hypothesize that viral microdiversity may, in part, be driven by an increase in macrodiversity of zone-specific bacterial populations in TT-MES, which we interpret as an expansion of host 'niches' available for infection that could drive diversification in viruses (Elena et al., 2009). Second, we hypothesize that the decrease in viral macrodiversity may be driven by increased viral microdiversity of some viral populations in the mesopelagic region that can promote competitive exclusion (Sensu) (Hart et al., 2016) as discussed above. Alternatively, lower cell density in the mesopelagic layer (Sunagawa et al., 2015) may result in less encounters between "predator" and "prey," reducing viral speciation (as a function of reduced number of viral generations), but selecting for viruses with broader host range. Again, testing these hypotheses will require technological advances to measure in situ host ranges and sensitivities of viruses and cells, respectively, at scales relevant to the diversity in nature.

#### Conclusions

This study provides a systematic and global-scale view of patterns and drivers of marine viral macro- and microdiversity that reveals three overarching advances. First, five ecological zones emerge for the global ocean, which contrasts known Longhurst biogeographic patterning in other organisms, but is consistent with observations from the largely co-sampled ocean microbiome (Sunagawa et al., 2015). Second, patterns and drivers of viral macro- and microdiversity differ per sample and positively correlate to geographic range. These findings offer hints at underlying mechanisms that impact these two levels of diversity that will guide researchers from discovery to hypothesis-testing as technologies, such as scalable single virus genomics and in situ host range assays, advance toward sampling scales relevant to those in nature. Third, epipelagic waters and the Arctic Ocean emerge from our work as biodiversity hotspots for viruses. While this is surprising given the latitudinal diversity gradient paradigm that the tropics rather than the poles are the cradles of diversity, it is in line with other observations in larger organisms (Valdovinos et al., 2003; Rabosky et al., 2018) and emphasizes the importance of these drastically climateimpacted Arctic regions for global biodiversity. Together, these advances, along with the parallel global-scale ecosystem-wide measurements of Tara Oceans (de Vargas et al., 2015; Sunagawa et al., 2015; Brum et al., 2015; Lima-Mendez et al., 2015; Roux et al., 2016) provide the foundation for incorporating viruses into emerging genes-to-ecosystems models (Guidi et al., 2016; Garza et al., 2018) that guide ocean ecosystem management decisions that are likely needed if humans and the Earth System are to survive the current epoch of the planet-altering Anthropocene.

#### **STAR**\***METHODS**

Detailed methods are provided in the online version of this paper and include the following:

- KEY RESOURCES TABLE
- CONTACT FOR REAGENT AND RESOURCE SHARING

#### EXPERIMENTAL MODEL AND SUBJECT DETAILS

- Tara Oceans Polar Circle (TOPC) expedition sample collection and virome creation
- METHODS DETAILS
  - Tara Oceans Polar Circle (TOPC) expedition sample processing and sequencing analyses
- QUANTIFICATION AND STATISTICAL ANALYSIS
  - $\,\circ\,$  Viral contig assembly, identification, and dereplication
  - Viral taxonomy
  - $\odot$  Viral population boundaries
  - Calculating viral population relative abundances, average read depths, and population ranks
  - Subsampling reads
  - Macrodiversity calculations
  - Microdiversity calculations
  - O Annotating Genes & Making Protein Clusters
  - Selection Analyses
  - Drivers of Macro- and Micro-diversity
  - Subsampling macro- and micro- diversity
  - Classifying multi-zonal, regional, and local viral populations
  - Comparing ARC-H and ARC-L
  - Comparing GOV to GOV 2.0
  - Calculating 16S OTU Macrodiversity
- IMPACT OF THE COAST, DEPTH, AND SEASONS
  Assessment of microbial contamination
- DATA AND SOFTWARE AVAILABILITY
  - Code availability
  - Data availability

#### SUPPLEMENTAL INFORMATION

Supplemental Information can be found online at https://doi.org/10.1016/j. cell.2019.03.040.

#### CONSORTIA

The members of Tara Oceans coordinators are Silvia G. Acinas, Marcel Babin, Peer Bork, Emmanuel Boss, Chris Bowler, Guy Cochrane, Colomban de Vargas, Michael Follows, Gabriel Gorsky, Nigel Grimsley, Lionel Guidi, Pascal Hingamp, Daniele Iudicone, Olivier Jaillon, Stefanie Kandels-Lewis, Lee Karp-Boss, Eric Karsenti, Fabrice Not, Hiroyuki Ogata, Stéphane Pesant, Nicole Poulton, Jeroen Raes, Christian Sardet, Sabrina Speich, Lars Stemmann, Matthew B. Sullivan, Shinichi Sunagawa, and Patrick Wincker. Affiliations for Tara Oceans coordinators can be found in Document S1.

#### ACKNOWLEDGMENTS

Tara Oceans (that includes both the Tara Oceans and Tara Oceans Polar Circle expeditions) would not exist without the leadership of the Tara Expeditions Foundation and the continuous support of 23 institutes (https://oceans. taraexpeditions.org). We further thank the commitment of the following sponsors: CNRS (in particular Groupement de Recherche GDR3280 and the Research Federation for the study of Global Ocean Systems Ecology and Evolution, FR2022/Tara Oceans-GOSEE), European Molecular Biology Laboratory (EMBL), Genoscope/CEA, The French Ministry of Research, and the French Government "Investissements d'Avenir" programmes OCEANOMICS (ANR-11-BTBR-0008), FRANCE GENOMIQUE (ANR-10-INBS-09-08), MEMO LIFE (ANR-10-LABX-54), and PSL<sup>\*</sup> Research University (ANR-11-IDEX-0001-02). We also thank the support and commitment of Agnès b. and Etienne Bourgois, the Prince Albert II de Monaco Foundation, the Veolia Foundation, Region Bretagne, Lorient Agglomeration, Serge Ferrari, Worldcourier, and KAUST.

The global sampling effort was enabled by countless scientists and crew who sampled aboard the Tara from 2009-2013, and we thank MERCATOR-CORIOLIS and ACRI-ST for providing daily satellite data during the expeditions. We are also grateful to the countries who graciously granted sampling permissions. The authors declare that all data reported herein are fully and freely available from the date of publication, with no restrictions, and that all of the analyses, publications, and ownership of data are free from legal entanglement or restriction by the various nations whose waters the Tara Oceans expeditions sampled in. This article is contribution number 86 of Tara Oceans. Computational support was provided by an award from the Ohio Supercomputer Center (OSC) to M.B.S. Study design and manuscript comments from Bonnie T. Poulos, Ho Bin Jang, M. Consuelo Gazitúa, Olivier Zablocki, Janaina Rigonato, Damien Eveillard, Frédéric Mahé, Federico Ibarbalz, and Hisashi Endo are gratefully acknowledged. Funding was provided by the Gordon and Betty Moore Foundation (3790 to M.B.S.), NSF (OCE 1536989 and OCE 1829831 to M.B.S.), Oceanomics (ANR-11-BTBR-0008) and France Genomique (ANR-10-INBS-09) to Genoscope, ETH and Helmut Horten Foundation (to S.S.), a Netherlands Organization for Scientific Research (NWO) Vidi grant (864.14.004 to B.E.D.), and an NIH T32 training grant fellowship (AI112542 to A.C.G.).

#### **AUTHOR CONTRIBUTIONS**

M.B., C.B. and L.K.-B. directed the Tara Oceans Polar Circle expedition. M.C., C.D., J.F., S.K., C.M., S. Pesant, M.P., S. Pisarev, J.P., and Tara Oceans coordinators conceptualized and organized sampling efforts for the Tara Oceans Polar Circle expedition. S. Pesant annotated, curated, and managed all biogeochemical data. A.A., C.C., and P.W. coordinated all sequencing efforts. A.C.G., A.A.Z., N.C.-N., B.T., B.B., K.A., G.D.-H., Y.L., D.V., J.-E.T., M.B., C.B., C.d.V., A.I.C., B.E.D., D.I., L.K.-B., S.R., S.S., P.W., and M.B.S. created the study design, analyzed the data, and wrote the manuscript. All authors approved the final manuscript.

#### **DECLARATION OF INTERESTS**

The authors declare no competing interests.

Received: October 31, 2018 Revised: January 5, 2019 Accepted: March 20, 2019 Published: April 25, 2019

#### SUPPORTING CITATIONS

The following references appear in the Supplemental Information: Angly et al., (2006); Marston and Amrich (2009); Marston and Martiny (2016); Sul et al. (2013); Zinger et al. (2011).

#### REFERENCES

Abecasis, G.R., Auton, A., Brooks, L.D., DePristo, M.A., Durbin, R.M., Handsaker, R.E., Kang, H.M., Marth, G.T., and McVean, G.A.; 1000 Genomes Project Consortium (2012). An integrated map of genetic variation from 1,092 human genomes. Nature *491*, 56–65.

Achtman, M., and Wagner, M. (2008). Microbial diversity and the genetic nature of microbial species. Nat. Rev. Microbiol. *6*, 431–440.

Alberti, A., Poulain, J., Engelen, S., Labadie, K., Romac, S., Ferrera, I., Albini, G., Aury, J.M., Belser, C., Bertrand, A., et al.; Genoscope Technical Team; Tara Oceans Consortium Coordinators (2017). Viral to metazoan marine plankton nucleotide sequences from the *Tara* Oceans expedition. Sci. Data *4*, 170093.

Angly, F.E., Felts, B., Breitbart, M., Salamon, P., Edwards, R.A., Carlson, C., Chan, A.M., Haynes, M., Kelley, S., Liu, H., et al. (2006). The marine viromes of four oceanic regions. PLOS Biol. *4*, e368.

Bar-On, Y.M., Phillips, R., and Milo, R. (2018). The biomass distribution on Earth. Proc. Natl. Acad. Sci. USA *115*, 6506–6511.

Bateman, A., Coin, L., Durbin, R., Finn, R.D., Hollich, V., Griffiths-Jones, S., Khanna, A., Marshall, M., Moxon, S., Sonnhammer, E.L., et al. (2004). The Pfam protein families database. Nucleic Acids Res. *32*, D138–D141.

Bobay, L.M., and Ochman, H. (2018). Biological species in the viral world. Proc. Natl. Acad. Sci. USA *115*, 6040–6045.

Bolduc, B., Jang, H.B., Doulcier, G., You, Z.Q., Roux, S., and Sullivan, M.B. (2017). vConTACT: an iVirus tool to classify double-stranded DNA viruses that infect *Archaea* and *Bacteria*. PeerJ *5*, e3243.

Brum, J.R., Ignacio-Espinoza, J.C., Roux, S., Doulcier, G., Acinas, S.G., Alberti, A., Chaffron, S., Cruaud, C., de Vargas, C., Gasol, J.M., et al.; Tara Oceans Coordinators (2015). Ocean plankton. Patterns and ecological drivers of ocean viral communities. Science *348*, 1261498.

Buchfink, B., Xie, C., and Huson, D.H. (2015). Fast and sensitive protein alignment using DIAMOND. Nat. Methods *12*, 59–60.

Cadillo-Quiroz, H., Didelot, X., Held, N.L., Herrera, A., Darling, A., Reno, M.L., Krause, D.J., and Whitaker, R.J. (2012). Patterns of gene flow define species of thermophilic Archaea. PLoS Biol. *10*, e1001265.

Cambuy, D.D., Coutinho, F.H., and Dutilh, B.E. (2016). Contig annotation tool CAT robustly classifies assembled metagenomic contigs and long sequences. bioRxiv. https://doi.org/10.1101/072868.

Carradec, Q., Pelletier, E., Da Silva, C., Alberti, A., Seeleuthner, Y., Blanc-Mathieu, R., Lima-Mendez, G., Rocha, F., Tirichine, L., Labadie, K., et al.; Tara Oceans Coordinators (2018). A global ocean atlas of eukaryotic genes. Nat. Commun. 9, 373.

Cohan, F.M. (2002). What are bacterial species? Annu. Rev. Microbiol. 56, 457-487.

Circumpolar Biodiversity Monitoring Program (2017). State of the Arctic Marine Biodiversity Report (Conservation of Arctic Flora and Fauna).

Costello, M.J., and Chaudhary, C. (2017). Marine biodiversity, biogeography, deep-Sea gradients, and conservation. Curr. Biol. *27*, R511–R527.

Culley, A. (2018). New insight into the RNA aquatic virosphere via viromics. Virus Res. 244, 84–89.

Culley, A.I., and Steward, G.F. (2007). New genera of RNA viruses in subtropical seawater, inferred from polymerase gene sequences. Appl. Environ. Microbiol. 73, 5937–5944.

Culley, A.I., Lang, A.S., and Suttle, C.A. (2006). Metagenomic analysis of coastal RNA virus communities. Science *312*, 1795–1798.

de Jonge, P.A., Nobrega, F.L., Brouns, S.J.J., and Dutilh, B.E. (2019). Molecular and evolutionary determinants of bacteriophage host range. Trends Microbiol. *27*, 51–63.

de Vargas, C., Audic, S., Henry, N., Decelle, J., Mahé, F., Logares, R., Lara, E., Berney, C., Le Bescot, N., Probert, I., et al.; Tara Oceans Coordinators (2015). Ocean plankton. Eukaryotic plankton diversity in the sunlit ocean. Science *348*, 1261605.

Deming, J.W., and Collins, E. (2017). Sea ice as a habitat for Bacteria, Archaea and Viruses. In Sea Ice, Third Edition, D.N. Thomas, ed. (John Wiley and Sons), pp. 327–351.

Deng, L., Ignacio-Espinoza, J.C., Gregory, A.C., Poulos, B.T., Weitz, J.S., Hugenholtz, P., and Sullivan, M.B. (2014). Viral tagging reveals discrete populations in Synechococcus viral genome sequence space. Nature *513*, 242–245.

Devol, A.H., Codispoti, L.A., and Christensen, J.P. (1997). Summer and winter denitrification rates in western Arctic shelf sediments. Cont. Shelf Res. *17*, 1029–1033.

Dixon, P. (2003). VEGAN, a package of R functions for community ecology. J. Veg. Sci. *14*, 927–930.

Duffy, S., Shackelton, L.A., and Holmes, E.C. (2008). Rates of evolutionary change in viruses: patterns and determinants. Nat. Rev. Genet. 9, 267–276.

Elena, S.F., Agudelo-Romero, P., and Lalić, J. (2009). The evolution of viruses in multi-host fitness landscapes. Open Virol. J. 3, 1-6.

Enav, H., Kirzner, S., Lindell, D., Mandel-Gutfreund, Y., and Béjà, O. (2018). Adaptation to sub-optimal hosts is a driver of viral diversification in the ocean. Nat. Commun. 9, 4698. Enright, A.J., Van Dongen, S., and Ouzounis, C.A. (2002). An efficient algorithm for large-scale detection of protein families. Nucleic Acids Res. 30, 1575–1584.

Farooq, A., and Malfatti, F. (2007). Microbial structuring of marine ecosystems. Nat. Rev. Microbiol. *5*, 782–791.

Feng, J., Durant, J.M., Stige, L.C., Hessen, D.O., Hjermann, D.Ø., Zhu, L., Llope, M., and Stenseth, N.C. (2015). Contrasting correlation patterns between environmental factors and chlorophyll levels in the global ocean. Global Biogeochem. Cycles *29*, 2095–2107.

Fraser, C., Alm, E.J., Polz, M.F., Spratt, B.G., and Hanage, W.P. (2009). The bacterial species challenge: making sense of genetic and ecological diversity. Science *323*, 741–746.

Garza, D.R., van Verk, M.C., Huynen, M.A., and Dutilh, B.E. (2018). Towards predicting the environmental metabolome from metagenomics with a mechanistic model. Nat. Microbiol. 3, 456–460.

Ghiglione, J.F., Galand, P.E., Pommier, T., Pedrós-Alió, C., Maas, E.W., Bakker, K., Bertilson, S., Kirchmanj, D.L., Lovejoy, C., Yager, P.L., and Murray, A.E. (2012). Pole-to-pole biogeography of surface and deep marine bacterial communities. Proc. Natl. Acad. Sci. USA *109*, 17633–17638.

Gregory, A.C., Solonenko, S.A., Ignacio-Espinoza, J.C., LaButti, K., Copeland, A., Sudek, S., Maitland, A., Chittick, L., Dos Santos, F., Weitz, J.S., et al. (2016). Genomic differentiation among wild cyanophages despite widespread horizontal gene transfer. BMC Genomics *17*, 930.

Greninger, A.L. (2018). A decade of RNA virus metagenomics is (not) enough. Virus Res. 244, 218–229.

Groom, S.B., and Holligan, P.M. (1987). Remote sensing of coccolithophore blooms. Adv. Space Res. 7, 73–78.

Guidi, L., Chaffron, S., Bittner, L., Eveillard, D., Larhlimi, A., Roux, S., Darzi, Y., Audic, S., Berline, L., Brum, J., et al.; Tara Oceans coordinators (2016). Plankton networks driving carbon export in the oligotrophic ocean. Nature *532*, 465–470.

Hart, S.P., Schreiber, S.J., and Levine, J.M. (2016). How variation between individuals affects species coexistence. Ecol. Lett. *19*, 825–838.

Hedrick, P.W. (2006). Genetic Polymorphism in Heterogeneous Environments: The Age of Genomics. Annu. Rev. Ecol. Evol. Syst. *37*, 67–93.

Hillebrand, H. (2004). On the generality of the latitudinal diversity gradient. Am. Nat. *163*, 192–211.

Holm, L., and Sander, C. (1998). Removing near-neighbour redundancy from large protein sequence collections. Bioinformatics *14*, 423–429.

Hughes, A.R., Inouye, B.D., Johnson, M.T.J., Underwood, N., and Vellend, M. (2008). Ecological consequences of genetic diversity. Ecol. Lett. *11*, 609–623.

Hurwitz, B.L., and Sullivan, M.B. (2013). The Pacific Ocean virome (POV): a marine viral metagenomic dataset and associated protein clusters for quantitative viral ecology. PLOS One 8, e57355.

Hurwitz, B.L., and U'Ren, J.M. (2016). Viral metabolic reprogramming in marine ecosystems. Curr. Opin. Microbiol. *31*, 161–168.

Hyatt, D., Chen, G.L., Locascio, P.F., Land, M.L., Larimer, F.W., and Hauser, L.J. (2010). Prodigal: prokaryotic gene recognition and translation initiation site identification. BMC Bioinformatics *11*, 119.

Iranzo, J., Koonin, E.V., Prangishvili, D., and Krupovic, M. (2016). Bipartite network analysis of the archaeal virosphere: evolutionary connections between viruses and capsid-less mobile elements. J. Virol. *90*, 11043–11055.

Jang, H.-B., Bolduc, B., Zablocki, O., Kuhn, J.H., Adriaenssens, E.M., Krupovic, M., Brister, R., Kropinski, A.M., Koonin, E.V., Turner, D., et al. (2019). Gene sharing networks to automate genome-based prokaryotic viral taxonomy. BioRxiv. https://doi.org/10.1101/533240.

Jian, H., Xu, J., Xiao, X., and Wang, F. (2012). Dynamic modulation of DNA replication and gene transcription in deep-sea filamentous phage SW1 in response to changes of host growth and temperature. PLoS One 7, e41578.

Kanehisa, M., Goto, S., Kawashima, S., and Nakaya, A. (2002). The KEGG databases at GenomeNet. Nucleic Acids Res. *30*, 42–46. Konstantinidis, K.T., and Tiedje, J.M. (2005). Genomic insights that advance the species definition for prokaryotes. Proc. Natl. Acad. Sci. USA *102*, 2567–2572.

Kunz, W. (2013). Do species exist?: Principles of taxonomic classification (John Wiley & Sons).

Kurtz, S., Phillippy, A., Delcher, A.L., Smoot, M., Shumway, M., Antonescu, C., and Salzberg, S.L. (2004). Versatile and open software for comparing large genomes. Genome Biol. *5*, R12.

Langmead, B., and Salzberg, S.L. (2012). Fast gapped-read alignment with Bowtie 2. Nat. Methods 9, 357–359.

Larkin, A.A., and Martiny, A.C. (2017). Microdiversity shapes the traits, niche space, and biogeography of microbial taxa. Environ. Microbiol. Rep. 9, 55–70. Le Quéré, C., Andrew, R.M., Friedlingstein, P., Sitch, S., Pongratz, J., Manning,

A.C., Korsbakken, J.I., Peters, G.P., Canadell, J.G., Jackson, R., et al. (2018). Global carbon budget 2017. Earth Syst. Sci. Data *10*, 405–448.

Lee, S.T.M., Kahn, S.A., Delmont, T.O., Shaiber, A., Esen, Ö.C., Hubert, N.A., Morrison, H.G., Antonopoulos, D.A., Rubin, D.T., and Eren, A.M. (2017). Tracking microbial colonization in fecal microbiota transplantation experiments via genome-resolved metagenomics. Microbiome *5*, 50.

Lemos, L.N., Fulthorpe, R.R., Triplett, E.W., and Roesch, L.F. (2011). Rethinking microbial diversity analysis in the high throughput sequencing era. J. Microbiol. Methods *86*, 42–51.

Li, W.K.W., McLaughlin, F.A., Lovejoy, C., and Carmack, E.C. (2009). Smallest algae thrive as the Arctic Ocean freshens. Science *326*, 539.

Lima-Mendez, G., Faust, K., Henry, N., Decelle, J., Colin, S., Carcillo, F., Chaffron, S., Ignacio-Espinosa, J.C., Roux, S., Vincent, F., et al.; Tara Oceans coordinators (2015). Ocean plankton. Determinants of community structure in the global plankton interactome. Science *348*, 1262073.

Logares, R., Sunagawa, S., Salazar, G., Cornejo-Castillo, F.M., Ferrera, I., Sarmento, H., Hingamp, P., Ogata, H., de Vargas, C., Lima-Mendez, G., et al. (2014). Metagenomic 16S rDNA Illumina tags are a powerful alternative to amplicon sequencing to explore diversity and structure of microbial communities. Environ. Microbiol. *16*, 2659–2671.

Longhurst, A.R. (2007). Ecological geography of the sea (Academic Press).

Longhurst, A., Sathyendranath, S., Platt, T., and Caverhill, C. (1995). An estimate of global primary production in the ocean from satellite radiometer data. J. Plankton Res. *17*, 1245–1271.

Maat, D.S., Biggs, T., Evans, C., van Bleijswijk, J.D.L., van der Wel, N.N., Dutilh, B.E., and Brussaard, C.P.D. (2017). Characterization and temperature dependence of Arctic Micromonas polaris viruses. Viruses 9, E134.

Mannion, P.D., Upchurch, P., Benson, R.B.J., and Goswami, A. (2014). The latitudinal biodiversity gradient through deep time. Trends Ecol. Evol. 29, 42–50.

Marston, M.F., and Amrich, C.G. (2009). Recombination and microdiversity in coastal marine cyanophages. Environ. Microbiol. *11*, 2893–2903.

Marston, M.F., and Martiny, J.B. (2016). Genomic diversification of marine cyanophages into stable ecotypes. Environ. Microbiol. *18*, 4240–4253.

Marston, M.F., Pierciey, F.J., Jr., Shepard, A., Gearin, G., Qi, J., Yandava, C., Schuster, S.C., Henn, M.R., and Martiny, J.B.H. (2012). Rapid diversification of coevolving marine *Synechococcus* and a virus. Proc. Natl. Acad. Sci. USA *109*, 4544–4549.

Martínez-Hernández, F., Fornas, O., Lluesma Gomez, M., Bolduc, B., de la Cruz Peña, M.J., Martínez, J.M., Antón, J., Gasol, J.M., Rosselli, R., Rodríguez-Valera, F., et al. (2017). Single-virus genomics reveals hidden cosmopolitan and abundant viruses. Nat. Commun. *8*, 15892.

Mavrich, T.N., and Hatfull, G.F. (2017). Bacteriophage evolution differs by host, lifestyle and genome. Nat. Microbiol. 2, 17112.

Miraldo, A., Li, S., Borregaard, M.K., Flórez-Rodríguez, A., Gopalakrishnan, S., Rizvanovic, M., Wang, Z., Rahbek, C., Marske, K.A., and Nogués-Bravo, D. (2016). An Anthropocene map of genetic diversity. Science *353*, 1532–1535.

Miranda, J.A., Culley, A.I., Schvarcz, C.R., and Steward, G.F. (2016). RNA viruses as major contributors to Antarctic virioplankton. Environ. Microbiol. *18*, 3714–3727.

Moniruzzaman, M., Wurch, L.L., Alexander, H., Dyhrman, S.T., Gobler, C.J., and Wilhelm, S.W. (2017). Virus-host relationships of marine single-celled eukaryotes resolved from metatranscriptomics. Nat. Commun. *8*, 16054.

Nurk, S., Meleshko, D., Korobeynikov, A., and Pevzner, P.A. (2017). meta-SPAdes: a new versatile metagenomic assembler. Genome Res. *27*, 824–834.

Paul, J.H. (1999). Microbial gene transfer: an ecological perspective. J. Mol. Microbiol. Biotechnol. 1, 45–50.

Pesant, S., Not, F., Picheral, M., Kandels-Lewis, S., Le Bescot, N., Gorsky, G., ludicone, D., Karsenti, E., Speich, S., Troublé, R., et al.; Tara Oceans Consortium Coordinators (2015). Open science resources for the discovery and analysis of Tara Oceans data. Sci. Data 2, 150023.

Petrie, K.L., Palmer, N.D., Johnson, D.T., Medina, S.J., Yan, S.J., Li, V., Burmeister, A.R., and Meyer, J.R. (2018). Destabilizing mutations encode nongenetic variation that drives evolutionary innovation. Science *359*, 1542–1545.

Pianka, E.R. (1966). Latitudinal Gradients in Species diversity: A Review of Concepts. Am. Nat. *100*, 33–46.

Quinlan, A.R., and Hall, I.M. (2010). BEDTools: a flexible suite of utilities for comparing genomic features. Bioinformatics *26*, 841–842.

Rabosky, D.L., Chang, J., Title, P.O., Cowman, P.F., Sallan, L., Friedman, M., Kaschner, K., Garilao, C., Near, T.J., Coll, M., and Alfaro, M.E. (2018). An inverse latitudinal gradient in speciation rate for marine fishes. Nature *559*, 392–395.

Reiners, W.A., Lockwood, J.A., Reiners, D.S., and Prager, S.D. (2017). 100 years of ecology: what are our concepts and are they useful? Ecol. Monogr. 87, 260–277.

Ren, J., Ahlgren, N.A., Lu, Y.Y., Fuhrman, J.A., and Sun, F. (2017). VirFinder: a novel *k*-mer based tool for identifying viral sequences from assembled meta-genomic data. Microbiome *5*, 69.

Reygondeau, G., Guidi, L., Beaugrand, G., Henson, S.A., Koubbi, P., MacKenzie, B.R., Sutton, T.T., Fioroni, M., and Maury, O. (2018). Global biogeochemical provinces of the mesopelagic zone. J. Biogeogr. *45*, 500–514.

Roossinck, M.J., Saha, P., Wiley, G.B., Quan, J., White, J.D., Lai, H., Chavarría, F., Shen, G., and Roe, B.A. (2010). Ecogenomics: using massively parallel pyrosequencing to understand virus ecology. Mol. Ecol. *19* (*Suppl 1*), 81–88.

Rosen, M.J., Davison, M., Bhaya, D., and Fisher, D.S. (2015). Microbial diversity. Fine-scale diversity and extensive recombination in a quasisexual bacterial population occupying a broad niche. Science *348*, 1019–1023.

Roux, S., Adriaenssens, E.M., Dutilh, B.E., Koonin, E.V., Kropinski, A.M., Krupovic, M., Kuhn, J.H., Lavigne, R., Brister, R., Varsani, A., et al. (2018). Minimum Information about an Uncultivated Virus Genome (MIUViG). Nat. Biotechnol. *2018*. Published online December 17. https://doi.org/10.1038/nbt. 4306nbt.4306.

Roux, S., Krupovic, M., Debroas, D., Forterre, P., and Enault, F. (2013). Assessment of viral community functional potential from viral metagenomes may be hampered by contamination with cellular sequences. Open Biol. *3*, 130160.

Roux, S., Enault, F., Hurwitz, B.L., and Sullivan, M.B. (2015). VirSorter: mining viral signal from microbial genomic data. PeerJ *3*, e985.

Roux, S., Brum, J.R., Dutilh, B.E., Sunagawa, S., Duhaime, M.B., Loy, A., Poulos, B.T., Solonenko, N., Lara, E., Poulain, J., et al.; Tara Oceans Coordinators (2016). Ecogenomics and potential biogeochemical impacts of globally abundant ocean viruses. Nature *537*, 689–693.

Roux, S., Emerson, J.B., Eloe-Fadrosh, E.A., and Sullivan, M.B. (2017). Benchmarking viromics: an *in silico* evaluation of metagenome-enabled estimates of viral community composition and diversity. PeerJ *5*, e3817.

Ruiz-González, C., Simó, R., Sommaruga, R., and Gasol, J.M. (2013). Away from darkness: a review on the effects of solar radiation on heterotrophic bacterioplankton activity. Front. Microbiol. *4*, 131.

Schloissnig, S., Arumugam, M., Sunagawa, S., Mitreva, M., Tap, J., Zhu, A., Waller, A., Mende, D.R., Kultima, J.R., Martin, J., et al. (2013). Genomic variation landscape of the human gut microbiome. Nature *493*, 45–50.

Ser-Giacomi, E., Zinger, L., Malviya, S., De Vargas, C., Karsenti, E., Bowler, C., and De Monte, S. (2018). Ubiquitous abundance distribution of non-dominant plankton across the global ocean. Nat. Ecol. Evol. *2*, 1243–1249.

Shapiro, B.J., Friedman, J., Cordero, O.X., Preheim, S.P., Timberlake, S.C., Szabó, G., Polz, M.F., and Alm, E.J. (2012). Population genomics of early events in the ecological differentiation of bacteria. Science 336, 48–51.

Shi, M., Lin, X.D., Tian, J.H., Chen, L.J., Chen, X., Li, C.X., Qin, X.C., Li, J., Cao, J.P., Eden, J.S., et al. (2016). Redefining the invertebrate RNA virosphere. Nature *540*, 539–543.

Shi, M., Zhang, Y.Z., and Holmes, E.C. (2018). Meta-transcriptomics and the evolutionary biology of RNA viruses. Virus Res. *243*, 83–90.

Smillie, C.S., Sauk, J., Gevers, D., Friedman, J., Sung, J., Youngster, I., Hohmann, E.L., Staley, C., Khoruts, A., Sadowsky, M.J., et al. (2018). Strain tracking reveals the determinants of bacterial engraftment in the human gut following fecal microbiota transplantation. Cell Host Microbe *23*, 229–240.

Snitkin, E.S., Zelazny, A.M., Montero, C.I., Stock, F., Mijares, L., Murray, P.R., and Segre, J.A.; NISC Comparative Sequence Program (2011). Genomewide recombination drives diversification of epidemic strains of *Acinetobacter baumannii*. Proc. Natl. Acad. Sci. USA *108*, 13758–13763.

Soliveres, S., van der Plas, F., Manning, P., Prati, D., Gossner, M.M., Renner, S.C., Alt, F., Arndt, H., Baumgartner, V., Binkenstein, J., et al. (2016). Biodiversity at multiple trophic levels is needed for ecosystem multifunctionality. Nature *536*, 456–459.

Steward, G.F., Culley, A.I., Mueller, J.A., Wood-Charlson, E.M., Belcaid, M., and Poisson, G. (2013). Are we missing half of the viruses in the ocean? ISME J. 7, 672–679.

Sul, W.J., Oliver, T.A., Ducklow, H.W., Amaral-Zettler, L.A., and Sogin, M.L. (2013). Marine bacteria exhibit a bipolar distribution. Proc. Natl. Acad. Sci. USA *110*, 2342–2347.

Sullivan, M.B. (2015). Viromes, not gene markers, for studying double-stranded DNA virus communities. J. Virol. *89*, 2459–2461.

Sunagawa, S., Coelho, L.P., Chaffron, S., Kultima, J.R., Labadie, K., Salazar, G., Djahanschiri, B., Zeller, G., Mende, D.R., Alberti, A., et al.; Tara Oceans coordinators (2015). Ocean plankton. Structure and function of the global ocean microbiome. Science *348*, 1261359.

Suttle, C.A. (2007). Marine viruses–major players in the global ecosystem. Nat. Rev. Microbiol. 5, 801–812.

Sutton, T.T., Clark, M.R., Dunn, D.C., Halpin, P.N., Rogers, A.D., Guinotte, J., Bograd, S.J., Angel, M.V., Perez, J.A.A., Wishner, K., et al. (2017). A global biogeographic classification of the mesopelagic zone. Deep Sea Res. Part I Oceanogr. Res. Pap. *126*, 85–102.

Suzek, B.E., Wang, Y., Huang, H., McGarvey, P.B., and Wu, C.H.; UniProt Consortium (2015). UniRef clusters: a comprehensive and scalable alternative for improving sequence similarity searches. Bioinformatics *31*, 926–932.

Tilman, D., Isbell, F., and Cowles, J.M. (2014). Biodiversity and ecosystem functioning. Annu. Rev. Ecol. Evol. Syst. 45, 471–493.

Tremblay, J.-É., Anderson, L.G., Matrai, P., Coupel, P., Bélanger, S., Michel, C., and Reigstad, M. (2015). Global and regional drivers of nutrient supply, primary production and  $CO_2$  drawdown in the changing Arctic Ocean. Prog. Oceanogr. *193*, 171–196.

Urayama, S.I., Takaki, Y., Nishi, S., Yoshida-Takashima, Y., Deguchi, S., Takai, K., and Nunoura, T. (2018). Unveiling the RNA virosphere associated with marine microorganisms. Mol. Ecol. Resour. *18*, 1444–1455.

Valdovinos, C., Navarrette, S.A., and Marquet, P.A. (2003). Mollusk species diversity in the Southeastern Pacific: Why are there more species towards the pole? Ecography *26*, 139–144.

Van Valen, L. (1965). Morphological variation and width of ecological niche. Am. Nat. *99*, 377–389.

Vellend, M., and Geber, M.A. (2005). Connections between species diversity and genetic diversity. Ecol. Lett. 8, 767–781.

Vellend, M., Lajoie, G., Bourret, A., Múrria, C., Kembel, S.W., and Garant, D. (2014). Drawing ecological inferences from coincident patterns of populationand community-level biodiversity. Mol. Ecol. *23*, 2890–2901.

Watkinson, A.R., and Sutherland, W.J. (1995). Sources, sinks, and pseudosinks. J. Anim. Ecol. 64, 126–130.

Worm, B., Barbier, E.B., Beaumont, N., Duffy, J.E., Folke, C., Halpern, B.S., Jackson, J.B., Lotze, H.K., Micheli, F., Palumbi, S.R., et al. (2006). Impacts of biodiversity loss on ocean ecosystem services. Science *314*, 787–790.

Zdobnov, E.M., and Apweiler, R. (2001). InterProScan–an integration platform for the signature-recognition methods in InterPro. Bioinformatics *17*, 847–848.

Zeigler Allen, L., McCrow, J.P., Ininbergs, K., Dupont, C.L., Badger, J.H., Hoffman, J.M., Ekman, M., Allen, A.E., Bergman, B., and Venter, J.C. (2017). The Baltic Sea virome: diversity and transcriptional activity of DNA and RNA viruses. mSystems 2, e00125-16.

Zinger, L., Amaral-Zettler, L.A., Fuhrman, J.A., Horner-Devine, M.C., Huse, S.M., Welch, D.B., Martiny, J.B., Sogin, M., Boetius, A., and Ramette, A. (2011). Global patterns of bacterial beta-diversity in seafloor and seawater ecosystems. PLOS One *6*, e24570.

#### **STAR\*METHODS**

#### **KEY RESOURCES TABLE**

REAGENT or RESOURCE	SOURCE	IDENTIFIER(S)
Sequencing Reagents and Kits		
NEBNext DNA Sample Prep Master Mix	New England Biolabs, Ipswich, MA	Cat n° E6040S
NEXTflex PCR free barcodes	Bioo Scientific, Austin, TX	Cat n° NOVA-514110
Kapa Hifi Hot Start Library Amplification kit	KAPA Biosystems, Wilmington, MA	Cat n° KK2611
DNA SMART ChIPSeq Kit	Takara Bio USA, Mountain View, CA	Cat N° 634865
Deposited Data		
Fara Oceans Viromes Raw Reads	Brum et al., 2015; Roux et al., 2016	European Nucleotide Archive (ENA) - see Table S3 for details
Fara Oceans Polar Circle Raw Reads	This paper	European Nucleotide Archive (ENA) - see Table S3 for details
<i>Malaspania</i> Viromes Raw Reads	Roux et al., 2016	Integrated Microbial Genomes (IMG) with Joint Genome Institute - see Table S3 for details
6S rRNA gene Tara Oceans data	Logares et al., 2014	Supplementary materials in Logares et al. (2014)
Biogeographical and Physicochemical data	Pesant et al., 2015	PANGAEA (Data Publisher for Earth & Environmental Science) - see Table S3 for details
N* Arctic Data	This paper	Table S3
Software and Algorithms		
nucmer (MUMmer3.23)	Kurtz et al., 2004	https://sourceforge.net/projects/mummer/
obmap 37.57	https://jgi.doe.gov/data-and-tools/bbtools/	https://jgi.doe.gov/data-and-tools/bbtools/ RRID:SCR_016965
netaSPAdes 3.11	Nurk et al., 2017	https://github.com/ablab/spades/releases; RRID:SCR_000131
prodigal 2.6.1	Hyatt et al., 2010	https://github.com/hyattpd/Prodigal; RRID:SCR_011936
liamond	Buchfink et al., 2015	https://github.com/bbuchfink/diamond; RRID:SCR_016071
/irSorter v1.0.4	Roux et al., 2015	https://github.com/simroux/VirSorter
/irFinder	Ren et al., 2017	https://github.com/jessieren/VirFinder
DAT	Cambuy et al., 2016	https://github.com/dutilh/CAT
blast 2.4.0+	ftp://ftp.ncbi.nlm.nih.gov/blast/ executables/blast+/	ftp://ftp.ncbi.nlm.nih.gov/blast/ executables/blast+/
/ConTACT2	Jang et al., 2019	https://bitbucket.org/MAVERICLab/ vcontact2
powtie2	Langmead and Salzberg, 2012	https://github.com/BenLangmead/bowtie2
3amM	https://github.com/Ecogenomics/BamM	https://github.com/Ecogenomics/BamM
Bedtools	Quinlan and Hall, 2010	https://github.com/arq5x/bedtools2/blob/ master/docs/content/overview.rst; RRID:SCR_006646
/egan (R package)	Dixon, 2003	https://cran.r-project.org/web/packages/ vegan/index.html; RRID:SCR_011950
	https://cran.r-project.org/web/packages/	https://cran.r-project.org/web/packages/

(Continued on next page)

Continued		
REAGENT or RESOURCE	SOURCE	IDENTIFIER(S)
heatmap3 (R package)	https://cran.r-project.org/web/packages/ heatmap3/index.html	https://cran.r-project.org/web/packages/ heatmap3/index.html
ggplot2 (R package)	https://cran.r-project.org/web/packages/ ggplot2/index.html	https://cran.r-project.org/web/packages/ ggplot2/index.html; RRID:SCR_014601
ggpubr (R package)	https://cran.r-project.org/web/packages/ ggpubr/index.html	https://cran.r-project.org/web/packages/ ggpubr/index.html
Analyses scripts (per Figure)	This paper	https://bitbucket.org/MAVERICLab/GOV2

#### **CONTACT FOR REAGENT AND RESOURCE SHARING**

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Matthew B. Sullivan (mbsulli@gmail.com).

#### **EXPERIMENTAL MODEL AND SUBJECT DETAILS**

#### Tara Oceans Polar Circle (TOPC) expedition sample collection and virome creation

Between June 2013 and December 2013, 41 samples were collected at different depths from 20 different sites near or within the Arctic Ocean (see full list of samples in Table S3). Physicochemical measurements, sample collection, and DNA extractions were performed using the methods described in Roux et al. (2016). Extracted DNA was prepared for sequencing using library preparation method described in Alberti et al. (2017) for viral samples collected during the *TOPC* campaign (section 4.2) and sequenced using the HiSeq 2000 system (101 bp, paired end reads). Importantly, our sample collection and library preparation methods have known bias toward < 0.2um dsDNA viruses (Roux et al., 2017). The *TOPC* samples were combined with the previously published viromes in (Brum et al., 2015; Roux et al., 2016). Of the previously published dataset, the mesopelagic samples at (*Tara* stations 37, 39, 56, 68, 70, 76, 78, 111, 122, 137, 138) and the Southern Ocean samples (*Tara* stations 82\_DCM, 84, 85) were sequenced deeper. These combined samples comprise the GOV 2.0 dataset. The number of reads found in each sample can be found in Table S3.

#### **METHODS DETAILS**

#### Tara Oceans Polar Circle (TOPC) expedition sample processing and sequencing analyses

Due to different library preparation for the *TOPC* samples than the original *Tara* Oceans samples, the previously sequenced mesopelagic samples (*Tara* stations 68, 78, 111, 137) were prepped using the *TOPC* library preparation to determine if it impacted our ability to assemble viral populations. We found no significant difference between library preparations in terms of the number of viral genomes assembled and the average genome length (Figures S7A and S7B). Additionally, to directly assess the impact of experimental variation between *Tara* Oceans and *TOPC* on our ecological interpretations, we applied hierarchical clustering on a Bray-Curtis dissimilarity matrix of our viromes and we found that all of the mesopelagic samples prepared using the *TOPC* protocols clustered with their respective samples prepared using the original *Tara* Ocean protocols, and the variation between them was far less than the ecological variation across our viromes (see distances in hierarchical clustering in Figure S7D). For two surface samples (*Tara* Stations 100 and 102), we also re-prepped the DNA using the DNA SMART ChIP-Seq kit which allows us to catch ssDNA in the library preparation (Takara) and further sequenced these two samples using the HiSeq 2000 system.

While the *Tara* Oceans and *Malaspina* expeditions used the same sampling and storage approaches (described in Roux et al., 2016), the sequencing reads were longer for the latter (101 bp for *Tara* and 151 bp for *Malaspina*). Given this, we have performed further analyses to evaluate whether the contribution of this experimental method variation surpasses the ecological variation presented in this study or not. These analyses, which are further described below, showed that ecological variation much better explained the data than experimental methods. To evaluate this, we compared the deep ocean samples collected from the *Tara* Oceans and *Malaspina* expeditions to assess their power to predict the correct ecological zone (mesopelagic or bathypelagic) based on the depth of collection (ecological variation) and the sequencing read length (experimental variation). Using three different metrics, namely the  $r^2$  value in a univariate regression analysis, the bayesian information criterion (BIC) of such constructed univariate model, and the *p*-value associated with different components in a multivariate regression analysis, we found that the depth of collection, rather than the experimental variation, best predicts the ecological zone (higher  $r^2$ ), with a better model fit (lower BIC), and lower *p*-value (Figure S7C). Additionally, we have one *Malaspina* sample from the mesopelagic ecological zone (the rest are *Tara* samples), and there is no significant difference between the *Malaspina* sample and Tara samples in the mesopelagic (Figures S3C and S3D). Together these findings demonstrate that the differences between the samples collected during the different expeditions are predominantly the result of ecology and community structure rather than experimental artifact.

All the remaining STAR Methods we used are quantifications and statistical analyses. All the details related to these STAR Methods are therefore provided in the following section, QUANTIFICATION AND STATISTICAL ANALYSES

#### **QUANTIFICATION AND STATISTICAL ANALYSIS**

#### Viral contig assembly, identification, and dereplication

All samples in the GOV 2.0 dataset (Roux et al., 2016) as well as the previously sequenced *TOPC* library-prepped mesopelagic samples and the DNA SMART ChIP-Seq kit surface samples were individually assembled using metaSPAdes 3.11.1 (Nurk et al., 2017). Prior to assembly, *Malaspina* samples from GOV 2.0 were further quality controlled. Briefly, adaptors and Phix174 reads were removed and reads were trimmed using bbduk.sh (https://jgi.doe.gov/data-and-tools/bbtools/; minlength = 30 qtrim = rl maq = 20 maxns = 0 trimq = 14 qtrim = rl). Following assembly, contigs  $\geq$  1.5kb were piped through VirSorter (Roux et al., 2015) and VirFinder (Ren et al., 2017) and those that mapped to the human, cat or dog genomes were removed. Contigs  $\geq$  5kb or  $\geq$  1.5kb and circular that were sorted as VirSorter categories 1-6 and/or VirFinder score  $\geq$  0.7 and p < 0.05 were pulled for further investigation. Of these contigs, those sorted as VirSorter categories 1 and 2, VirFinder score  $\geq$  0.9 and p < 0.05 or were identified as viral by both VirSorter (categories 1-6) and VirFinder (score  $\geq$  0.7 and p < 0.05) were classified as viral. The remaining contigs were run through CAT (Cambuy et al., 2016) and those with < 40% (based on an average gene size of 1000) of the genome classified as bacterial, archaeal, or eukaryotic were considered viral. In total, 848,507 viral contigs were identified. Viral contigs were grouped into populations if they shared  $\geq$  95% nucleotide identity across  $\geq$  80% of the genome (*sensu*) (Brum et al., 2015) using nucmer (Kurtz et al., 2004). This resulted in 488,130 total viral populations found in GOV 2.0 (see Table S5 for VirSorter, VirFinder, and CAT results), of which 195,728 were  $\geq$  10kb.

#### **Viral taxonomy**

For each viral population, ORFs were called using Prodigal (Hyatt et al., 2010) and the resulting protein sequences were used as input for vConTACT2 (Jang et al., 2019) and for blastp. Viral populations represented by contigs > 10kb were clustered with Viral RefSeq release 85 viral genomes using vConTACT2. Those that clustered with a virus from RefSeq based on amino acid homology based on diamond (Buchfink et al., 2015) alignments were able to be assigned to a known viral taxonomic genus and family. For GOV 2.0 viral populations that could not be assigned taxonomy or were < 10kb, family level taxonomy was assigned using a majority-rules approach, where if > 50% of a genome's proteins were assigned to the same viral family using a blastp bitscore  $\geq$  50 with a Viral RefSeq virus, it was considered part of that viral family.

#### Viral population boundaries

To determine if our viral populations had discrete sequence boundaries, all reads across the GOV 2.0 dataset (excluding the *Tara* stations 68, 78, 111, 137 prepped using the *TOPC* library preparation methods and the DNA SMART ChIP-Seq kit prepped libraries) were pooled and mapped non-deterministically to our viral populations using the 'very-sensitive-local' setting in bowtie2 (Langmead and Salzberg, 2012). The percent nucleotide identity (% ID) of each mapped read and the positions in the genome where the read mapped were determined. The frequency of reads mapping at a specific % IDs were weighted based on the length of each read mapped across the genomes. Frequencies of reads mapping at specific % IDs were smoothed using Loess smooth functions (span = 1 to be more permissive of lower % ID reads) to create read frequency histograms (% ID versus frequency). To determine break in the distribution of read frequencies between the different % IDs, Euclidean distances calculated were calculated between % ID frequencies and then hierarchically clustered in R.

#### Calculating viral population relative abundances, average read depths, and population ranks

To calculate the relative abundances of the different viral populations in each sample, reads from each GOV 2.0 virome were first nondeterministically mapped to the GOV 2.0 viral population genomes using bowtie2. BamM (https://github.com/ecogenomics/BamM) was used to remove reads that mapped at < 95% nucleotide identity to the contigs, bedtools genomecov (Quinlan and Hall, 2010) was used to determine how many positions across each genome were covered by reads, and custom Perl scripts were used to further filter out contigs without enough coverage across the length of the contig. For downstream *macro*diversity calculations, contigs  $\geq$  5kb in length that had < 5kb coverage or less than the total length of the contig covered for contigs < 5kb were removed. For downstream *micro*diversity calculations, all contigs with < 70% of the contig covered were removed. BamM was used to calculate the average read depth ('tpmean' -minus the top and bottom 10% depths) across each contig. For the *macro*diversity calculations, the average read depth was used as a proxy for abundance and normalized by total read number per metagenome to allow for sample-to-sample comparison. The rank abundance of all the viral populations was calculated using the normalized abundances and the 'rankabundance' in the BiodiversityR R package.

#### Subsampling reads

Unequal sequencing depth can have large impacts on diversity measurements, specifically α-diversity measurements (Lemos et al., 2011). Due to 5x more sequencing depth in *TOPC* samples and the deeply sequenced mesopelagic and Southern Ocean samples (Table S3), all viromes in the GOV 2.0 dataset were randomly subsampled without replacement to 20M reads for *Tara* or 10M reads for

*Malaspina* (as many *Malaspina* samples were < 20M reads and there was no significant difference between the 10M and 20M reads assemblies; p = 1) using reformat.sh from bbtools suite (https://sourceforge.net/projects/bbmap/). The subsampled read libraries were assembled using metaSPAdes 3.11.1. Contigs  $\geq$  1.5kb that shared  $\geq$  95% nucleotide identity across  $\geq$  80% of the genome with the 488,130 viral populations in GOV 2.0 were pulled out and grouped into populations to be used as the subsampled GOV 2.0 viral populations. In total, there were 46,699 viral populations. Relative abundances were calculated per sample as aforementioned for *macro*diversity calculations, but using the subsampled GOV 2.0 viral populations and the subsampled reads.

#### **Macrodiversity calculations**

The *macro*diversity  $\alpha$ - (Shannon's *H*) and  $\beta$ - (Bray-Curtis dissimilarity) diversity statistics were performed using vegan in R (Dixon, 2003). The  $\alpha$ -diversity calculations were based on the relative abundances produced from the subsampled reads. Loess smooth plots with 95% confidence windows in ggplot2 in R were used to look at changes in Shannon's *H* across latitude (Figure 7A) and depth (Figure 7F). For the  $\beta$ -diversity, both the subsampled and the total reads abundances were used to look at community structure (Figure S3). Principal Coordinate analysis (function capscale of vegan package with no constraints applied) and NMDS analysis (function metaMDS; K = 2 and trymax = 100) were used as the ordination methods on the Bray-Curtis dissimilarity matrices from both the subsampled and total reads calculated from GOV 2.0 (function vegdist; method "bray") after a cube root transformation (function nthroot; n = 3). The ecological zones that emerged were verified using a permanova test (function "adonis") and the confidence intervals were plotted using function "ordiellipse" at the specified confidence limits (95% and 97.5%) using the standard deviation method. There were no significant differences in clustering between the subsampled and all reads Bray-Curtis dissimilarity PCoA plots (Figure S3). Hierarchical clustering (function pvclust; method.dist = "cor" and method.hclust = "average") was conducted on the same Bray-Curtis dissimilarity matrices using 1000 bootstrap iterations and only the approximately unbiased (AU) bootstrap values were reported. The heatmaps were generated using the heatmap3 package with appropriate rotations of the branches in the dendrograms. Samples that did not cluster with their ecological zone (*Tara* mesopelagic stations 72, 85, and 102 and *Tara* surface station 155) were considered outliers and removed from further analyses (Figures S3A and S3C).

#### **Microdiversity calculations**

Viral populations with an average read depth of  $\geq 10x$  across 70% of their representative contig in at least one sample in the GOV 2.0 dataset were flagged for *micro*diversity analyses. We used 10x as the minimum coverage because population genetic statistics were found to be relatively consistent down to 10x based on previous downsampling coverage analyses (Schloissnig et al., 2013). BAM files containing reads mapping at  $\geq 95\%$  nucleotide identity were filtered for just the flagged viral populations. Samtools mpileup and bcftools were used to call single nucleotide variants (SNVs) across these populations. SNV calls with a quality call > 30 threshold were kept. Coverage for each allele for each SNV locus was summed across all the metagenomes. For each SNV locus, the consensus allele was re-verified and those with alternative alleles that had a frequency > 1% (Abecasis et al., 2012), the classical definition of a polymorphism, and supported by at least 4 reads were considered SNP loci (Schloissnig et al., 2013). Nucleotide diversity ( $\pi$ ) per genome were calculated using equation from Schloissnig et al. (2013). Due to the variable coverage across the genome, coverage was randomly downsampled to 10x coverage per locus in the genome. For the downsampling, if there was not the target 10x coverage for the locus, all of the alleles were sampled. Nucleotide diversity ( $\pi$ ) was calculated for each genome with an average read depth  $\geq 10x$  across 70% of their contig in each sample. For each sample,  $\pi$  values of 100 viral populations were randomly selected and averaged. This was repeated 1000x and the average of the all 1000 subsamplings was used as the final microdiversity value for each sample. Loess smooth plots with 95% confidence windows in ggplot2 in R were used to look at changes in average  $\pi$  across latitude (Figure 7A) and depth (Figure 7F).

#### **Annotating Genes & Making Protein Clusters**

Genes were annotated by translating the sequences into proteins and running a combination of reciprocal best blast hit analyses against the KEGG database (Kanehisa et al., 2002), and blast against the UniProt Reference Clusters database (Suzek et al., 2015), searching for matches against the InterPro protein signature database using InterProScan (Zdobnov and Apweiler, 2001), and running HMM searches against Pfams (Bateman et al., 2004). A diamond 'blastall' alignment search (Buchfink et al., 2015) of all the protein sequences was performed against all the protein sequence was performed and the protocol "Clustering similarity graphs encoded in BLAST results" with a granularity of I = 2 from the MCL website (https://micans.org/mcl/; Enright et al., 2002) was used to create protein clusters.

#### **Selection Analyses**

Natural selection (pN/pS) was calculated using the method from Schloissnig et al. (2013). The pN/pS method compares the expected ratio of non-synonymous and synonymous substitutions based on a uniform model of occurrence of mutations across the genome with the observed ratio of non-synonymous and synonymous substitutions. The original method treats each SNP locus as independent from each other. Thus, if two SNPs occur in the same codon, the alternate codon produced from each SNP would be considered in the pN/pS calculation. Thus, if two SNPs occur in one codon, the effect of the SNPs could potentially cancel each other out or amplify a non-synonymous signal leading to false positive selection calls. In order to minimize this bias, SNPs found within the

same codon in the same gene were tested for linkage in each metagenome. If SNP alleles from loci within the same codon had depth coverage within 15% of each other within each metagenome, they were considered linked in that sample.

For each codon with SNP loci in a gene, the minimum coverage was identified based on the lowest read depth coverage among the three base pair position. The initial number of the consensus codon was determined based on the lowest coverage of the consensus alleles at the SNP locus or loci if linked. The initial numbers of potential alternate codons was based on the coverage of the alternate allele at that position or the lowest coverage between two linked SNPs. The final coverage of the each codon per SNP locus was calculated by taking the rounded down number of the product of the initial number x (initial number/ minimum coverage for the codon). These codons then subsampled down to 10x. The number of observed non-synonymous and synonymous substitutions were counted and pN/pS was calculated. Genes were considered under positive selection if pN/pS was > 1.

#### **Drivers of Macro- and Micro-diversity**

Regression analysis between the first coordinate of the PCoA (Figure 5A) and available temperature measurements was conducted using the Im function in R. The environmental variables were fitted to the first two dimensions of the PCoA using a generalized additive model (function envfit; permutations = 9999 and na.rm = TRUE). Then, they were correlated with all the PCoA dimensions using a mantel test (function mantel; permutations = 9999 and method = "spear") after scaling (function scale) and calculating their distance matrices (function vegdist; method "euclid" and na.rm = TRUE). Finally, they were correlated with Shannon's *H* and  $\pi$  using Pearson's correlation (function cor; use = "pairwise.complete.obs") after removing Shannon's *H* outliers based on a boxplot analysis (Figure S4). Both Pearson's and Spearman's correlations are provided in (Table S7).

#### Subsampling macro- and micro- diversity

Due to unequal sampling across each ecological zone, we chose to normalize the number of samples between each ecological zone by subsampling the down to lowest zone sample size (ANT; n = 5). Shannon's *H* outliers were not included in the subsampling. Five samples within each zone were randomly subsampled without replacement and their *macro-* and *micro-* diversity values averaged, respectively. We subsampled 1000x and plotted the averages and assessed for significant differences using Mann-Whitney U-tests in ggboxplot from the R package ggpubr (Figure 4B).

#### Classifying multi-zonal, regional, and local viral populations

To determine geographic range, viral populations were evaluated for their distributions across the five ecological zones and plotted using the VennDiagram package in R (Figure 6A). If present in  $\geq$  1 sample in more than one ecological zone, it was considered multizonal (58% GOV 2.0 viral populations). If present only in samples found within a single zone, it was considered zone-specific (48% GOV 2.0 viral populations). Zone-specific viral populations were further divided into regional ( $\geq$ 2 samples within a zone) and local (only 1 sample within a zone). The proportion of multi-zonal, regional, and local viral populations found across each zone (Figure 6B) and across each station (Figure S6) were calculated by dividing the number of each type by the total number of viral populations found across a zone or station, respectively. To assess the impact of geographic range on *micro*diversity per zone, stations were randomly subsampled without replacement as described above. Within each sample,  $\pi$  values of 50, 100, and 20 viral populations of each geographic distribution (multi-zonal, regional, and local, respectively) were randomly selected and averaged. All the viral populations with a geographic range were sampled and averaged in samples that lacked enough deeply-sequenced viral populations with particular geographic range. This was repeated 1000x and the averages plotted and assessed for significant differences using Mann-Whitney U-tests in ggboxplot from the R package ggpubr (Figure 6C).

#### **Comparing ARC-H and ARC-L**

The ARC-H and ARC-L regions were defined based on their biogeography; the ARC-H stations were located in the Pacific Arctic region, the Arctic Archipelago, and the Davis-Baffin Bay, in addition to one station (Station 189) in the Kara-Laptev sea, which was separated by a land mass from the rest of the stations in the same area (Figure 7D). The ARC-L stations were located in the Kara-Laptev Sea (except Station 189), the Barents Sea, and subpolar areas (stations 155 and 210). The departure from the dissolved N:P stoichiometry in the Redfield ratio (N\*) was calculated as in Tremblay et al. (2015) to represent the deficit in dissolved inorganic nitrogen (DIN) in the ratio and as a geochemical tracer of pacific and atlantic water masses. *Macro-* and *micro-* diversity values for each station in ARC-H and ARC-L were plotted and assessed for significant differences using Mann-Whitney U-tests in ggboxplot from the R package ggpubr (Figure 7E).

#### **Comparing GOV to GOV 2.0**

Viral populations assembled in the GOV (Roux et al., 2016) were compared to the GOV 2.0 viral populations (Figure 1B) using blastn. Unbinned GOV viral populations with a nucleotide alignment to a GOV 2.0 viral populations with  $\geq$  95% nucleotide identity and an alignment length  $\geq$  50% the length were considered present in the GOV 2.0. These results were plotted in a venn diagram using the VennDiagram package in R. The frequency of contig lengths of viral populations that were shared across both samples were plotted using ggplot2 (function "geom\_histogram"; binwidth = 5000).

#### **Calculating 16S OTU Macrodiversity**

Previously published 16S OTU data were taken from Logares et al. (2014). The macrodiversity  $\alpha$ - (Shannon's H) statistics were performed using vegan in R (Dixon, 2003). Loess smooth plots with 95% confidence windows in ggplot2 in R were used to look at changes in bacterial Shannon's H down the depth gradient. Differences between surface, deep chlorophyll maximum, and mesopelagic bacterial samples were compared using Mann-Whitney U-tests and plotted in ggboxplot from the R package ggpubr. Finally, viral microdiversity was correlated with bacterial Shannon's H using Pearson's correlation (function cor; use = "pairwise.complete.obs") and a linear regression (Figure S6D).

#### IMPACT OF THE COAST, DEPTH, AND SEASONS

GOV 2.0 samples are largely open ocean samples. Even though the arctic samples were more coastal, we didn't observe any significant coastal impact on the global *macro*diversity (Pearson's r = -0.25; Bonferroni-corrected *p*-value = 0.15) and *micro*diversity (Pearson's r = 0.11; *p*-value = 0.23) levels (Figure 4C). Although nitrate and phosphate levels generally increase with depth, we observed higher negative correlations and significantly lower *p*-values for these nutrients with *macro*diversity levels than between depth and *macro*diversity (Figure 4C) which suggests an impact of nutrients on viral diversity via primary production (Figure 5C). Additionally, since the sampling was largely at discrete depth layers with different densities in the TT region (epipelagic, mesopelagic, and bathypelagic), rather than sampling gradients, we discerned a clearer signal for the separation between these ecological zones (Figure 4A). On the other hand, all the arctic epipelagic and mesopelagic samples fell within the same ecological zone due to the absence of a pycnocline in this area (Figure 4A). Finally, the circumnavigation of the Arctic Ocean spanned multiple seasons (spring, summer, and fall). Based on our previous observation from a time-series data in a sub-arctic system (Hurwitz and Sullivan, 2013), our viral *macro*diversity is expected to be lowest during the spring and summer and increase toward the winter season. However, our calculated N\* values are not dependant on the season and represent the largest magnitude of change among all of the environmental variables that correlated with *macro*diversity between the ARC-H and ARC-L regions.

#### Assessment of microbial contamination

To quantifying microbial contamination across our samples, we screened our metagenomic reads using singleM (https://github.com/ wwood/singlem) for 16S sequences using the dedicated 16S SingleM package. We found that our viromes are exceptionally clean. Specifically, the number of 16S sequences in our samples ranged from 0-40 per million reads (Table S3), and hence the samples are considered to have "likely negligible bacterial contamination" according to the metric proposed by authors evaluating such signals in published viromes (threshold was 200 16S sequences per million; Roux et al., 2013). In spite of our viromes being exceptionally clean, we sought to evaluate the impact of any variation in 16S, and hence bacterial contamination, however small, on our findings. We found that even though microbial contamination increases with depth (most probably due to the decrease in cell size; linear regression  $r^2 = 0.89$ ), this increase was driven mainly by the bathypelagic samples. Briefly, the average contamination in BATHY was 28.7 per million reads (standard deviation = 6.8) as compared to the rest of the samples (average contamination = 1.7 per million reads and standard deviation = 2). These bathypelagic samples were not included in any of the ecological driver analyses due to the unavailability of the environmental data to us. Further, it is clear that our estimates of diversity were not influenced by the minor variations in the negligible contamination in our viroomes as a linear regression between Shannon's H and the number of 16S reads from deep ocean samples resulted in a negligible r<sup>2</sup> value (0.06). These data (used for conducting the regression analysis) represent a large range of diversity (3.3-7.8) and the full range of contamination (0-40), but avoid the convolution from the ecological difference between the surface and deep ocean layers. Thus, we conclude that the diversity observations we make in this study are driven by ecological variation far greater than microbial contamination.

#### DATA AND SOFTWARE AVAILABILITY

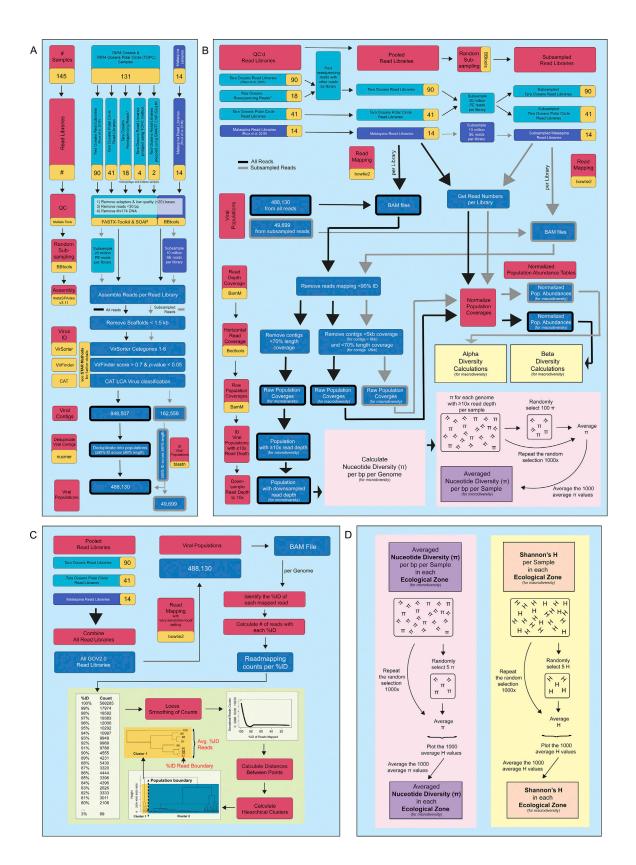
#### **Code availability**

Scripts used in this manuscript are available on the Sullivan laboratory bitbucket under GOV 2.0.

#### **Data availability**

All raw reads are available through ENA (*Tara* Oceans and *TOPC*) or IMG (Malapsina) using the identifiers listed in Table S3. Processed data are available through iVirus, including all assembled contigs, viral populations and genes.

# **Supplemental Figures**



Cell

Figure S1. Bioinformatic Workflow, Related to Figures 1 and 4

Flow diagrams showing the bioinformatic workflow for (A) the assembly and identification of viral populations, (B) the population coverages and abundances and how they were used to calculate *macro-* and *micro-*diversity calculations, (C) prediction of population boundaries, and (D) how average *macro-* and *micro*diversity calculations per ecological zone were calculated.

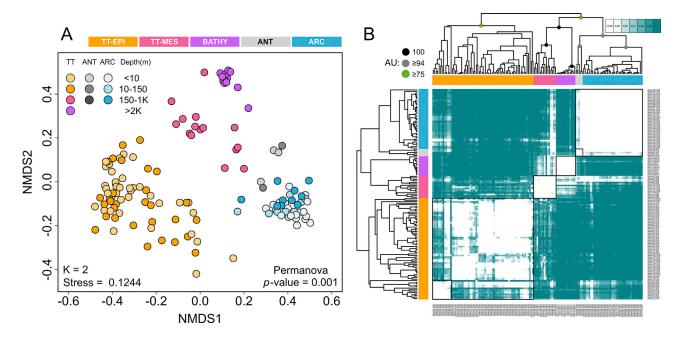
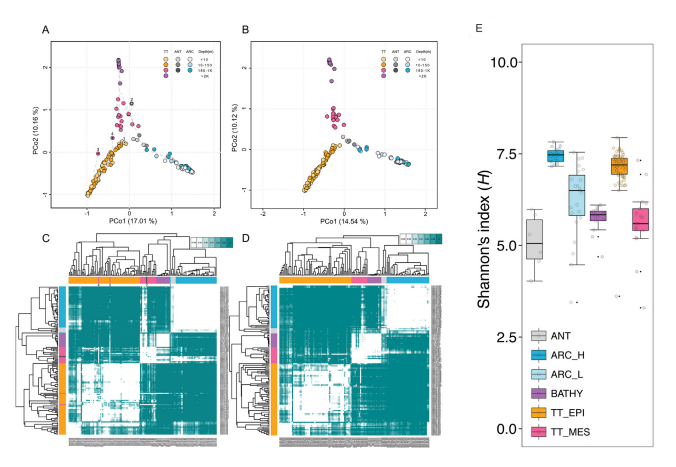


Figure S2. Non-metric Multidimensional Scaling and Hierarchical Clustering of GOV 2.0, Related to Figure 4 As observed with the Principal Coordinate analysis (Figure 4A), NMDS analysis (A) and correlation-based hierarchical clustering (B) of a Bray-Curtis dissimilarity matrix calculated from GOV 2.0 structured the viromes into five distinct global ecological zones with an approximately unbiased (AU) bootstrap value  $\geq$  77 in the hierarchical clustering. Four outlier viromes were removed and all the sequencing reads were used, with justification provided in Figures S3C and S3D, respectively. Abbreviations: ARC, Arctic; ANT, Antarctic; BATHY, bathypelagic; TT-EPI, temperate and tropical epipelagic; TT-MES, temperate and tropical mesopelagic.



#### Figure S3. Beta Diversity of the Total Reads and Subsampled Reads GOV 2.0 Dataset and Outlier Analyses, Related to Figure 4

PCoA of a Bray-Curtis dissimilarity matrix calculated from GOV 2.0 using all the sequencing reads (A) and after randomly subsampling the reads to the same sequencing depth (B). The dissimilarity matrices from (A) and (B) were used to conduct hierarchical clustering on the samples as shown in (C) and (D), respectively. The four viromes which were removed from Figures 4 and S2 are highlighted with asterisks; sample 1 (station 155\_SUR) is the only surface sample in the North Atlantic Drift Province and could have been influenced by the warm surface currents going northward due to the Atlantic Meridional Overturning Circulation; sample 2 (station 85\_MES) is the only mesopelagic sample from the Southern Ocean and could have been influenced by the upwelling of ancient deep ocean water (which is also congruent with the similarity observed between deep water bacterial communities of polar and lower latitude) (Ghiglione et al., 2012); sample 3 (station72\_MES) fell outside the 97.5% confidence intervals of all the ecological zones; sample 4 (station102\_MES) was located in El Niño-Southern Oscillation region and could have been influenced by the upwellings and downwellings in this area. Additionally, samples 1, 3, and 4 were among the Shannon's *H* outliers (E). Viral communities still partitioned into five ecological zones. Outliers that fell below the first quantile or above the fourth quantile (function geom\_boxplot of ggplot) of each ecological zone were removed before examining the predictors of viral *macro*diversity (Figure 4C). Outliers: 32\_SUR, 155\_SUR, 56\_MES, 70\_MES, 72\_MES, 102\_MES, MSP131, and MSP144.

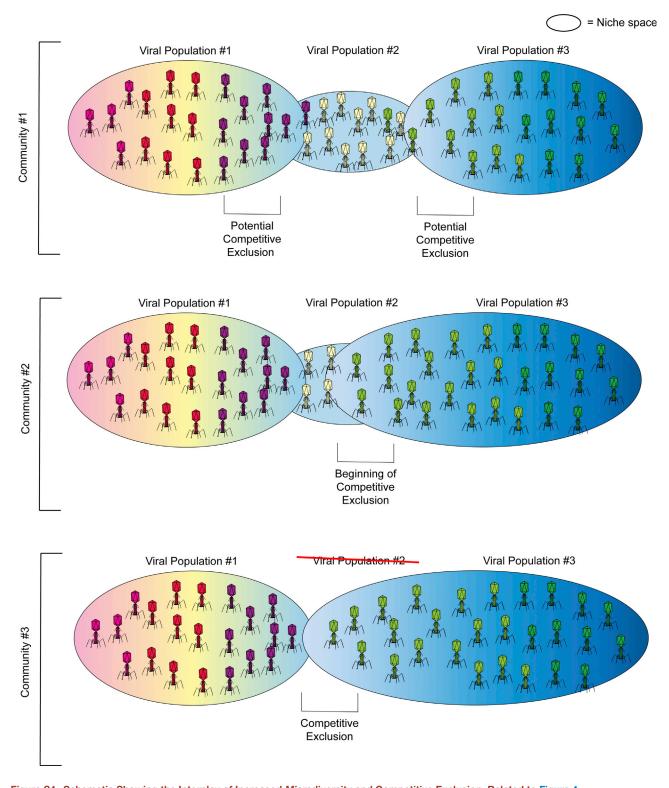


Figure S4. Schematic Showing the Interplay of Increased *Micro*diversity and Competitive Exclusion, Related to Figure 4 Viral populations with more *micro*diversity usually have larger niche sizes and therefore can outcompete viral populations with smaller overlapping niche sizes. This process of competitive exclusion may not be visible in each community as seen across the three communities. Thus, the average of communities such as across ecological zones can better show this relationship.

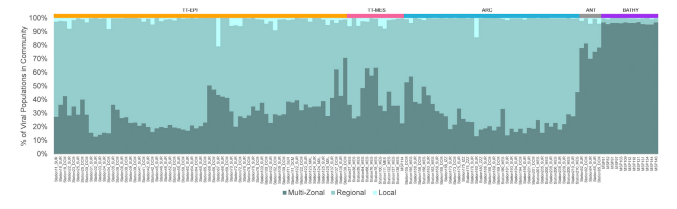


Figure S5. Stacked Barplots Showing the Number of Multi-Zonal, Regional, and Local Viral Populations Found within the Species Pool of Each Station, Related to Figure 6

Ecological zone outliers (see Figure S3) are excluded.

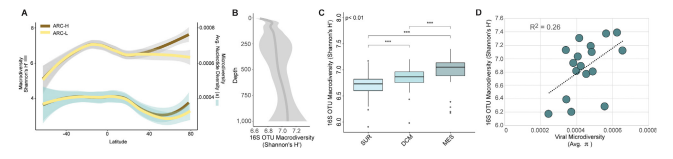
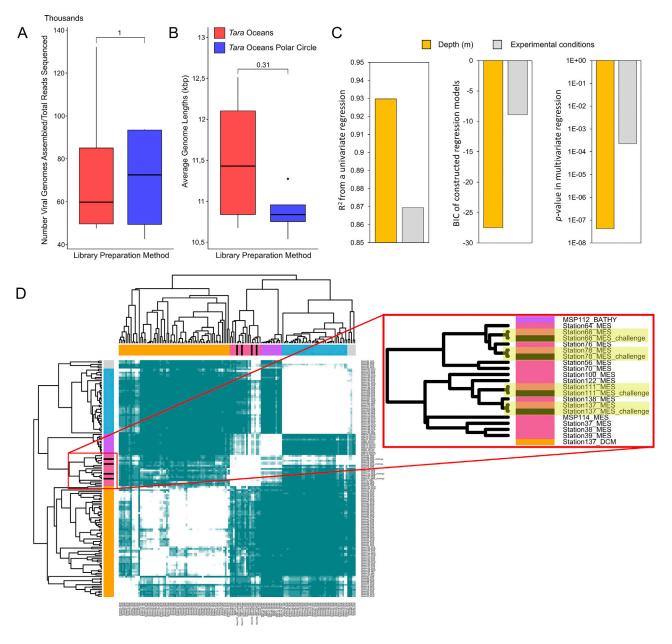


Figure S6. ARC-H Drives the Divergence from the Latitudinal Diversity Gradient and Microbial 16S OTUs Biodiversity Deviate from the Depth Diversity Gradient and Positively Correlates with Viral Microdiversity in the Mesopelagic, Related to Figure 7

(A) Loess smooth plots showing the latitudinal distributions of *macro*- and *micro*- population diversity with ARC-H and ARC-L regions. The line represents the loess best fit, while the lighter band corresponds to the 95% confidence window of the fit. (B) Loess smooth plots showing 16S OTUs (Logares et al., 2014) *macro*diversity distributions down the depth gradient. The line represents the loess best fit, while the lighter band corresponds to the 95% confidence window of the fit. (C) Boxplots showing median and quartiles of surface, deep chlorophyll maximum (DCM), and mesopelagic 16S OTU data taken from Logares et al. (2014). All pairwise comparisons shown were statistically significant (p < 0.05) using two-tailed Mann-Whitney U-tests. (D) Scatterplot showing the positive correlation (Pearson's correlation r = 0.51; *p-value* = 0.036) and linear regression ( $r^2 = 0.26$ ) between *Tara* Oceans mesopelagic samples shared between the 16S OTU samples in Logares et al. (2014) and our viral samples in GOV 2.0.



#### Figure S7. Library Preparation and Experimental Conditions Comparisons, Related to Figures 1 and 4

(A & B) Boxplots showing median and quartiles of the number of assembled viral genomes per total reads sequenced and the average genome lengths in *TO* and *TOPC* preparations of *Tara* mesopelagic stations 68, 78, 111, and 137, respectively. All pairwise comparisons shown were not statistically significant using twotailed Mann-Whitney U-tests. (C) Depth (as an ecological variable) predicts the ecological zone of the deep ocean (mesopelagic or bathypelagic) better than experimental variation between *Tara* and *Malaspina* expeditions, with a higher *r*<sup>2</sup> (left), lower BIC (middle), and lower *p-value* (right). The first two metrics were calculated from a univariate regression analysis (using depth alone or experimental variation alone as a predictor of the ecological zone), while the third metric was calculated from a multivariate multiple regression analysis that uses both depth and experimental variation as predictors. (D) Hierarchical clustering of a Bray-Curtis dissimilarity matrix calculated from GOV 2.0 viromes to which four additional viromes (black bars) have been added to control for the impact of experimental variation between the *Tara* Oceans and *Tara* Oceans Polar Circle expeditions. The four viromes prepared using the *Tara* Oceans Polar Circle protocols clustered with their respective original samples, which were prepared using the *Tara* Oceans protocols indicating that experimental variation was far less than ecological variation.