Global Trends in Marine Plankton Diversity across Kingdoms of Life

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Highlights
- Most epipelagic planktonic groups exhibit a poleward decline of diversity
- No latitudinal diversity gradient was observed below the photic zone
- Temperature emerges as the best predictor of epipelagic plankton diversity
- Global warming may increase plankton diversity, particularly at high latitudes

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In Brief
The drivers of ocean plankton diversity across archaea, bacteria, eukaryotes, and major virus clades are inferred from both molecular and imaging data acquired by the Tara Oceans project and used to predict the effects of severe warming of the surface ocean on this critical ecosystem by the end of the 21st century.
Global Trends in Marine Plankton Diversity across Kingdoms of Life

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SUMMARY

The ocean is home to myriad small planktonic organisms that underpin the functioning of marine ecosystems. However, their spatial patterns of diversity and the underlying drivers remain poorly known, precluding projections of their responses to global changes. Here we investigate the latitudinal gradients and global predictors of plankton diversity across archaea, bacteria, eukaryotes, and major virus clades using both molecular and imaging data from Tara Oceans. We show a decline of diversity for most planktonic groups toward the poles, mainly driven by decreasing ocean temperatures. Projections into the future suggest that severe warming of the surface ocean by the end of the 21st century could lead to tropicalization of the diversity of most planktonic groups in temperate and polar regions. These changes may have multiple consequences for marine ecosystem functioning and services and are expected to be particularly significant in key areas for carbon sequestration, fisheries, and marine conservation.

INTRODUCTION

Our planet is dominated by interconnected oceans that harbor a tremendous diversity of microscopic planktonic organisms. They form complex ecological networks (Lima-Mendez et al., 2015) that sustain major biogeochemical cycles (Falkowski et al., 2008; Field et al., 1998) and provide a wide range of ecosystem services (Guidi et al., 2016; Ptacnik et al., 2008; Worm et al., 2006). The ongoing increase in atmospheric carbon dioxide concentrations is having knock-on effects on the ocean by altering, among others, temperature, salinity, circulation, oxygenation, and pH levels (Pachauri et al., 2014; Rhein et al., 2013). These changes have already left visible imprints on marine plankton, fish, mammals, and birds, with shifts in species phenology and distribution (Beaugrand et al., 2009; Boyce et al., 2010; Poloczanska et al., 2013; Richardson and Schoeman, 2004). Future increases in ocean temperatures are expected to modify phytoplankton diversity and distribution directly by altering metabolic rates and growth (Thomas et al., 2012; Toseland et al., 2013) or
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Indirectly through changes in ocean circulation and, consequently, the supply of nutrients to surface waters (Bopp et al., 2013), given that such modifications will most likely impair the functions, goods, and services provided by the ocean (Brun et al., 2019; Hutchins and Fu, 2017; Worm et al., 2006), predicting how plankton diversity will respond to climate change has become a pressing challenge (Cavicchioli et al., 2019).

Unraveling patterns of diversity across macroclimatic gradients, such as the latitudinal diversity gradient (LDG), is a way to anticipate the effects of climate change (Algar et al., 2009; Frenne et al., 2013). The LDG, historically studied principally in terrestrial macroorganisms, usually consists of a monotonic poleward decline of local diversity (known as alpha diversity; Whittaker, 1972) for both terrestrial and aquatic organisms (Hillebrand, 2004). The LDG is hypothesized to result from a range of non-exclusive ecological and evolutionary mechanisms that operate at multiple spatial and temporal scales (Clarke and Gaston, 2006; Pontarp et al., 2019; Willig et al., 2003). Among the mechanisms classically invoked, temperature is often thought to be one of the major drivers through two effects. The physiological tolerance hypothesis postulates that temperature structures the LDG by imposing abiotic constraints on species distribution range (Currie et al., 2004), with fewer species tolerating cold and conditions and tropical temperatures being generally below the upper thermal tolerance limit of most organisms. The “kinetic energy hypothesis” relates to the metabolic theory (reviewed in Brown, 2014), which posits that higher temperatures increase the rate of metabolic reactions, resulting in shorter generation times, faster ecological or physiological processes, and, ultimately, higher mutation and speciation rates, leading to higher local diversity. Beyond temperature, the “productivity/resources hypothesis” posits that greater resource availability and/or primary production in tropical terrestrial areas can support larger population sizes and limit local extinction, promoting species coexistence (reviewed in Clarke and Gaston, 2006). The “environmental stability hypothesis” asserts that short- to long-term environmental instability in extratropical latitudes should cause greater local extinction rates because life in such unstable environments requires particular and costly physiological adaptations, which would ultimately preclude speciation (Clarke and Gaston, 2006).
seriously hamper our ability to understand the drivers of these essential components of marine ecosystems and estimate their potential responses in a changing ocean.

Here we provide a unified view of plankton LDGs using systematically collected data from the Tara Oceans global expedition. We combine DNA sequencing of filtered seawater and imaging of net catches to study the diversity in molecular operational taxonomic units (MOTUs) and morphotype diversity of major groups from all domains of life as well as both small and large double-stranded DNA viruses (Karsenti et al., 2011). We then separately examine their respective LDGs while determining their best environmental correlates because they may be influenced by different drivers. Finally, to identify the regions that may experience the most drastic changes in plankton diversity in the future, we model the trends of plankton diversity at the global scale for the beginning (years 1996–2005) and end of the century (years 2090–2099, representative concentration pathway [RCP] 8.5).

RESULTS AND DISCUSSION

LDG across Marine Plankton Groups and Water Layers

In this study, we used a wide collection of uniformly collected datasets with broad latitudinal coverage to explore the local diversity trends of all organismal groups that make up plankton communities (Table S1A). Besides already publicly available resources from Tara Oceans, we included newly released metabarcoding data of the V9 region of the 18S rRNA gene and flow cytometry (FC) abundances from the Arctic Ocean. We further complemented these observations with new global-scale datasets obtained with amplicon sequencing, microscopy, and imaging techniques. Our datasets were derived from 189 sampling stations distributed worldwide (Figure S1; STAR Methods), where multiple water depths were sampled (surface, 5 m average depth; deep chlorophyll maximum, 17–188 m; mesopelagic, 200–1000 m). This extensive and standardized sampling of plankton diversity encompasses large gradients in temperature, resource/primary production, and environmental stability (Figure 1A). Using the taxonomic information retrieved from genomics and imaging data (Table S1A), we distinguished 12 marine plankton groups (MPGs; Figures 1B, S1A, and S2; Table S1B; STAR Methods) with different trophic modes (e.g., photosynthetic/mixotrophs versus non-photosynthetic/heterotrophs) and different life history strategies (e.g., parasitic protists, endophotosymbionts) or corresponding to highly dominant taxa having a significant contribution to the marine food web (e.g., copepods).

The LDG of each MPG was studied using the Shannon index, a diversity index that relates monotonically to species richness but differs in that it downweights rare species, whose numbers are highly sensitive to undersampling and molecular artifacts (Figure S3; STAR Methods). Focusing on surface waters first, we found that phyto-, bacterio-, and zooplankton MPGs all exhibited maximal MOTU diversity in tropical to subtropical regions that then decreased poleward (Figure 2A; see also Figure S4 for individualized curves for each MPG as well as specific taxonomic groups). Similar trends were found for parasites of eukaryotes (giruses and parasitic protists mainly composed of marine alveolates [MALVs]) and for eukaryotic photosynthetic intracellular symbionts (endophotosymbionts) as well as their eukaryotic hosts (photohosts). Different patterns emerged for two abundant families of prokaryotic viruses (Myoviridae and Podoviridae) which, unlike their hosts, did not exhibit a clear poleward decline in diversity. Because the diversity of hosts and their symbionts or parasites is often assumed to be linked through eco-evolutionary interactions (Morand, 2015), an explanation for this could be that these virus families have a broader spectrum of host species, which could potentially decouple certain eco-evolutionary constraints (de Jonge et al., 2019). However, other factors may be responsible for this trend as well, such as nutrient availability.

Figure 1. Latitudinal Trends of Oceanic Conditions and Marine Plankton Composition in Surface Waters

(A) In situ chl a concentrations and sea surface temperatures (SST) across latitude (Tara Oceans expedition), plus IAV of SST (STAR Methods). Solid lines represent the GAM smooth trends and gray ribbons the corresponding 95% confidence intervals of parameter latitudinal trends predicted by the GAMs. (B) Average relative abundances of MPGs as inferred from molecular datasets across latitude. Prokaryotes: 16S rRNA gene, 0.22–3 μm; eukaryotes: 18S rRNA gene, 0.8–2000 μm (STAR Methods). Dark gray represents other eukaryotic groups. P, photosynthetic/mixotrophic; H, non-photosynthetic/heterotrophic. The three viral groups are not represented here because of the absence of comparable abundance data. See also Figures S1 and S2 and Tables S1A and S1B.
or bacterial cell density (Gregory et al., 2019). Further data and analyses will be necessary to elucidate the underpinnings of this result.

Differences in the form of LDGs have been proposed previously to result from contrasting strategies in energy acquisition and processing (Hillebrand, 2004). To test this hypothesis, we compared LDG forms across MPGs (except for prokaryotic vi-

ruses) by conducting a segmented linear regression analysis and using the inferred parameters in a clustering analysis (absolute latitudinal breakpoints and slopes of the segment regressions; STAR Methods; Figure S5). Confirming our above assumption, parasites and endo-

photosymbionts did not cluster directly with their hosts. Endophotosymbionts have extensive free-living populations (Decelle et al., 2012), and parasitic pro-
tists might experience relatively long-lasting free-living stages under the form of resistant cysts waiting for host availability (Siano et al., 2010), which could explain this result. Rather, we found that MPGs with similar broad trophic modes (phototrophic versus heterotrophic/chemotrophic) tended to exhibit similar LDG forms. However, we noticed two particular exceptions: photosynthetic protists clustered with heterotrophs
(both prokaryotes and eukaryotes), whereas heterotrophic protists clustered with phototrophs. Whether these features result from the presence of still unknown mixotrophs in heterotrophic protists (i.e., photosymbioses) or a preferential heterotrophy of mixotrophic photosynthetic protists remains to be determined. Top-down or bottom-up controls by other trophic levels as well as interspecific competition could also contribute to these patterns. For example, most copepods preferentially feed on phototrophic protists (e.g., diatoms or dinoflagellates; Saiz and Calbet, 2011), which could explain why both groups exhibited similar LDGs (Figure S5).

We then examined the symmetry of the LDGs by performing separate linear regressions for each hemisphere. LDGs are commonly observed to be steeper in the northern hemisphere, supposedly because of stronger climate instability in this area of the globe (Chown et al., 2004). Our results contradict this expectation because we found that LDGs only tended to be asymmetric and steeper in the southern hemisphere for archaea and photosynthetic protists (Figure 2A; Table S1C). Although in agreement with another report on marine bacterioplankton (Sul et al., 2013), we suspect the absence or opposite trend observed here to arise from a significant undersampling at mid- to high latitudes in the southern hemisphere in our dataset (Figure S1). Another explanation could lie in differences in the timing of sampling between the two polar regions; i.e., from the end of spring to the beginning of autumn in the Arctic Ocean but only in a summer month in the Southern Ocean, when diversity is expected to be lowest because of intense blooms (Arrigo et al., 2008). In spite of some variations in the form of LDGs across MPGs or between hemispheres, our results nonetheless show that the poleward decline of diversity is a pervasive feature among marine plankton.

To ensure that the MOTU diversity trends observed with our molecular data (Figure 2A) were not biased by the DNA region studied or molecular approach used (i.e., DNA metabarcoding versus metagenomics for prokaryotes; Salazar et al., 2019), which may vary in taxonomic resolution or overrepresent certain taxa, we compared them against those obtained with other DNA markers (i.e., the V4 region of the 18S rRNA gene for eukaryotes and the V4–V5 regions of the 16S rRNA gene for prokaryotes) or with finer clustering thresholds (STAR Methods). All of these comparisons exhibited high correlation coefficient values, regardless of the clustering similarity thresholds used (Figures S6A–S6D). DNA-based measures of diversity can also be affected by organism size (multicellular organisms) or gene copy number (unicellular organisms), with organisms of smaller size or lower gene copy numbers likely to be more difficult to detect. To examine this potential issue, we compared the diversity trends of different planktonic groups observed with molecular versus optical data (Figures 2B and 2C; STAR Methods). Both zooplankton imaging data, which consist of morphological features (see Figure 2B for examples), as well as photosynthetic protist data obtained through confocal (Colin et al., 2017) or light microscopy (LM) were highly congruent with their corresponding molecularly based diversity trends (Figures 2C and S6E–S6H).

Although of much lower taxonomic resolution, FC-based diversity values, comprising mainly prokaryotes, also correlated well with molecularly based prokaryotic diversity (Figure 2C). These results, together with the high correlation of diversity trends between our data and those based on single-copy genes (Milanese et al., 2019) and the fact that relative abundances in DNA-based data correlate well with organism size/biovolume and even abundance at lower taxonomic resolution (de Vargas et al., 2015), suggest that the global diversity trends observed here are unlikely to be biased by differences in body size/gene copy number across taxa.

In contrast, we did not observe any LDGs, either in terms of MOTUs or morphological diversity, below the photic zone (>200 m depth, corresponding here to both mesopelagic [Tara Oceans] and bathypelagic waters; Salazar et al., 2016; Figures 2D and S7). These environments are isolated from sunlight and climatic gradients. Accordingly, although we did observe a weak latitudinal trend in temperature in our deep sea samples (linear regression on absolute latitudes: slope = −0.11, R² = 0.273, p < 0.001), the range of this parameter represented roughly half of the temperature range present in surface waters (0°C to 18°C versus −2°C to 31°C, respectively). Hence, reduced temperature variations could be one of the reasons for an absence of LDGs in the deep sea. In addition, there is overall more carbon export at high latitudes (Henson et al., 2012). This could compensate for the reduction of diversity potentially induced by low temperatures by increasing resource availability in polar deep waters (Pomeroy and Wiebe, 2001). Finally, migration of surface species to deep waters through passive or active vertical flux may also contribute to cancel out temperature effects and perhaps underpin the overall higher diversity values we observe in deep sea waters compared with the surface (Figures 2C and 2D; Mestre et al., 2018). Although the current sampling effort in these aphotic environments is insufficient to firmly support these hypotheses, our results are consistent with previous observations for brittle stars (Woolley et al., 2016) and bacteria (Ghioglio et al., 2012) in the deep sea, whose diversity did not follow LDG trends. Both sediments and water layers below the photic zone are populated by heterotrophic and chemolithoautotrophic organisms, whose diversity and abundance are strongly influenced by organic matter availability (Bergauer et al., 2018; Danovaro et al., 2016; Woolley et al., 2016). This supports the idea that life is sustained by different types of energy supply across water layers, from systems driven by solar energy or kinetic effects of temperature in epipelagic waters to chemically driven environments (i.e., carbon- or mineral-based) in the deep sea.

Global Drivers of MPG Diversity in the Surface Ocean

To further understand the mechanisms underlying the observed LDGs in the surface ocean, we considered contextual variables related to the most common LDG hypotheses. First, we used sea surface temperature (SST, in situ measurements) to assess the physiological tolerance and kinetic energy hypotheses. Second, we used chlorophyll a (chl a) concentrations (in situ measurements) and annual maximum of nitrate concentration (AM NO₃) to test the productivity/resources hypothesis. Chl a was considered a proxy for phytoplankton biomass. We acknowledge that the latter may be affected by the poleward increase of intracellular pigmentation in phytoplankton to compensate for limitations in light (Behrenfeld et al., 2015). Phytoplankton
carbon estimated via particulate backscattering has been proposed as a better proxy (Graff et al., 2015), but we lacked this parameter for many stations. Nevertheless, the available backscattering data exhibited a high correlation with chl $a$ ($r = 0.6, p < 0.001$; Figure S8). Regarding AM NO$_3$, the annual availability of this macronutrient is fundamental for primary production (Moore et al., 2013). We therefore considered this parameter to capture longer-term effects of primary production on the observed plankton diversity. Third, we considered intra-annual variation (IAV) of SST to test the environmental stability hypothesis. In addition, we included a set of other contextual parameters in our analysis to identify potential drivers of diversity patterns for MPGs that had not been resolved previously (i.e., for viruses and protists; see STAR Methods for more details; Figure S8). Among them, sunlight, which is the fundamental source of energy for photosynthetic groups, was accounted for as satellite-derived estimates of photosynthetically active radiation (PAR) and the median light in the mixed layer (STAR Methods). Their more scattered availability in our dataset indicated high correlations with SST and mixed layer depth, respectively (Figure S8), which, together with chl $a$ concentration, reflects well the light conditions at the different sampling stations.

We conducted a combination of correlation analyses and generalized additive models (GAMs; Hastie, 2017), which allows us to deal with non-linear and/or non-monotonic relationships that could be found between diversity and environmental gradients (see below; STAR Methods). We restricted our analysis to surface planktonic communities because of their major contribution to oceanic biogeochemical cycles (Falkowski et al., 2008; Field et al., 1998) and their greater sensitivity to climate change (Bopp et al., 2013) and because of greater Tara Oceans data availability compared with the deep waters, allowing us to make more robust inferences and projections.

We found that SST was strongly and positively associated with the MOTU diversity patterns of most MPGs (Figures 3A, 3B, and S9A) and, therefore, was the best predictor of MPG diversity among the tested parameters (Figure 3C). Although new for most protist MPGs, these findings are consistent with previous observations for bacterioplankton (Fuhrman et al., 2008), copepods (Rombouts et al., 2009), and larger marine organisms (Tittensor et al., 2010; Woolley et al., 2016). SST and species thermal tolerance limits have been suggested to impose strong constraints on the distribution/abundance of marine ectotherms, including copepods (Beaugrand et al., 2009; Sunday et al., 2012). Our results extend this explanation to unicellular organisms as well; we observed a decline of phototrophic bacteria (mainly cyanobacteria) relative abundance at cooler higher latitudes (Figure 1B), whereas the relative abundance of phototrophic eukaryotes (mainly diatoms) increased. Such differences...
may result from contrasting thermal niches because diatoms generally have larger thermal breadths and lower minimal thermal growth than cyanobacteria (Chen, 2015). Similarly, the temperature-diversity relationship of several MPG categories increased until it reached a plateau, in particular for heterotrophic bacteria and archaea (Figure S8A). This may suggest that these groups have larger ranges of temperature optima, corresponding roughly to those encountered in tropical/subtropical waters, and should be less affected by climate change (Hutchins and Fu, 2017). Greater SST should also increase both speciation and extinction rates, according to the metabolic theory (reviewed in Brown, 2014). This assumption has been proposed for marine foraminifera (Allen et al., 2006) and diatoms (Lewitus et al., 2018), suggesting that temperature-dependent evolutionary processes are likely important in generating patterns of diversity across MPGs. However, our current approach remains correlative, and future phylogenetic studies will be critical to estimate speciation and diversification rates in relation to temperature.

MPG diversity also decreased noticeably and monotonically with increasing standing stocks of chl a and to a lesser extent, of AM NO3 (Figures 3A, 3B, S9B, and S9C). These negative relationships are counterintuitive to the productivity/resources hypothesis, which asserts that greater resource availability should promote species coexistence through niche partitioning. They also contrast with the unimodal biomass-diversity relationship often reported for phytoplankton (Irigoin et al., 2004; Li, 2002; Vallina et al., 2014). As explained above, it is very unlikely that this difference arises from the diversity indices used (i.e., richness versus Shannon index). Rather, we explain this difference by our broader sampling of plankton size classes and the increased detection and taxonomic resolution of our DNA-based identification methods. Accordingly, our results are in agreement with taxon-focused or DNA-based surveys, which have reported a higher diversity of copepods (Rombouts et al., 2008), bacteria (Smith, 2007), and microplankton (Raes et al., 2018) at sites of low primary production. Although our data preclude us to infer the exact mechanisms behind this negative relationship, we propose several potential explanations. First, this observation may be related to the “paradox of the plankton” (Hutchinson, 1961); i.e., the observation that a limited number of resources support unexpectedly highly diverse communities. Non-equilibrium and chaotic environmental and/or population dynamics in aquatic systems can occur at very small temporal and spatial scales, and this, together with the existence of dormant stages in plankton organisms, is usually thought to underlie this feature by preventing local extinction (Roy and Chattopadhyay, 2007; Scheffer et al., 2003; Ser-Giacomi et al., 2018). Also, recent genomic studies in prokaryotes suggest that adaptive gene loss and subsequent microbial feeding interdependencies are selectively favored in aquatic, nutrient-poor environments. These dependencies would constitute additional but currently unmeasurable niche axes, supporting more species (Giovannoni et al., 2014; the “black queen hypothesis”; Morris et al., 2012).

More generally, such trophic interdependencies probably do exist in the planktonic network without necessarily involving genome streamlining. On the other hand, high-nutrient or -chl a environments can correspond to areas with punctual/ mid-term strong physical forcing, such as winds (Demarco, 2009) or changes in light availability, which we did not measure on site. These environments are usually found to promote the growth of a few species at the expense of others through competitive or trophic interactions (Beihenfeld and Boss, 2014; Huisman et al., 1999; Irigoien et al., 2004; Li, 2002).

IAV of SST (Figure S8D) as well as other abiotic parameters, such as mixed layer depth together with its IAV or silicate or phosphate concentrations, exhibited comparatively weak or no correlation with MPG diversity (Figures 3A–3C; Table S1D). Finally, the GAMs, including the four focus parameters (i.e., SST, chl a, IAV SST, and AM NO3), did not exhibit latitudinal trends in their residuals (Table S1E), suggesting that the LDG was fully explained by these models. SST, followed by chl a concentration, thus appears to be the prominent driver of plankton diversity. This conclusion is further supported by additional GAMs where only SSTs, chl a concentrations, and their interaction were used as explanatory variables (see STAR Methods and their further use below). These latter models had exceptionally high explanatory power for most MPGs (42% to 81% of deviance explained) and also successfully explained the LDGs (Tables S1F and S1G). In addition, SSTs strongly correlated with microbial and photosynthetic abundances, and chl a strongly correlated with abundances of larger metazoans (Figure 3A; Figures S10A and S10B). This supports the idea that these two parameters regulate MPG diversity by controlling their population size and, therefore, also their extinction rates (Clarke and Gaston, 2006). Thus, our results lend support to the interplay of physiological tolerance, kinetic energy, and, to a lesser extent, productivity/resources effects in regulating MPG diversity in planktonic communities and causing latitudinal gradients of diversity in epipelagic waters.

Although our overall conclusions concur with those reported for marine macroorganisms (Hillebrand, 2004; Tittensor et al., 2010; Woolley et al., 2016), they partially contrast with recent findings regarding planktonic communities in the South Pacific Ocean (Raes et al., 2018), where primary productivity has been found to override temperature effects. This difference could be a consequence of the sampling extent of our study, which covers both the northern and southern hemispheres as well as multiple oceanic provinces that may differ in their diversity gradient (Chown et al., 2004; Sul et al., 2013). In contrast, Raes et al. (2018) characterized the latitudinal trends along a transect exhibiting marked environmental transitions caused by subtropical and subpolar frontiers. This should be confirmed by analyzing a larger number of sampling points in each basin than those available here. Another possible explanation lies in the different diversity measures used in the two studies. The Shannon index used here, although co-varying with species richness, down-weighs the influence of rarest species. The carrying capacity of a given environment strongly relies on resource availability and primary productivity, which control local extinction, in particular of rare species (Vallina et al., 2014). Such processes can solely be detected with species richness, which we did not assess here because of its strong sensitivity to sampling and technical biases (STAR Methods). More generally, several factors are also more confounding at the global scale, and their effects are more difficult to tease apart. For example, SST partially
correlates with chl concentration, PAR, and annual averages and IAVs in solar radiation as well as with the length of the productive season at the global scale (Figure S8; Clarke and Gaston, 2006). Hence, it is possible that both resource/sunlight-energy availability and stability effects partially contribute to the observed temperature effects. This feature may also explain why we could not find clear clustering of MPGs based on the drivers of their diversity according to their broad trophic modes, as found for their LDG patterns (Figure S5). Finally, we acknowledge that we considered environmental stability over short time-scales. Past glaciation cycles and associated sea level changes most likely contribute to current MPG diversity, as suggested for marine diatoms (Lewitus et al., 2018) and foraminifera (Yasuhara et al., 2012). Notwithstanding, fossil records do suggest that the poleward decline of zooplankton diversity and its temperature dependence are remarkably stable features through geological times (Yasuhara et al., 2012), albeit with variations in the overall levels of diversity. We therefore believe that paleoclimate effects are unlikely to alter our conclusions.

**Future Global Trends of MPG Diversity**

Climate change scenarios predict a general increase of SST, with major changes in the Arctic Ocean (Figure S11; Pachauri et al., 2014). Future ocean primary production is expected to decrease in the northern hemisphere and increase in the southern Ocean, although these projections are more uncertain (Figure S11; Bopp et al., 2013). To search for trends in diversity variation in response to these changes, we mapped the MOTU diversity of several MPGs at the beginning (years 1996–2005; Figure S12) and end of the 21st century (2090–2099) under a scenario of severe climate warming (RCP 8.5; STAR Methods). We used SST and chl concentration values simulated by Earth System Models of the Coupled Model Intercomparison Project Phase 5 (CMIP5; Bopp et al., 2013; Table S1H) and GAM models from epipelagic plankton that explained 60% or more of diversity (6 MPGs of 12; Table S1F). After ensuring that SST and chl concentration values were within the range of values used to train our models, we projected current and future MPG diversity at the global scale and calculated diversity anomalies (i.e., percentage of diversity change) between contemporary and future climates to identify areas where plankton diversity is most likely to be affected by the environmental changes in the ocean (Figures 4 and S12). To ensure reliability of our predictions, we generated 13,000 models for each MPG and each projection time to account for the uncertainties in the parameters of the GAMs and the output from different CMIP5 models used in this study (STAR Methods). Here we report averaged predictions from these models (see Figure S12 for prediction uncertainties). We also cross-validated our GAM models with independent datasets from other studies (STAR Methods; Figure S13) and obtained predictions congruent with the observed diversity.

Our projections suggest a general increase in MPG diversity, in particular in the northern hemisphere and at latitudes that encompass the limits of the subtropical gyres (25°–50°) (Figures 4A and S12). These results support tropicalization of temperate planktonic diversity or biomass, as suggested previously for bacterioplankton (Morán et al., 2010, 2017), zooplankton (Beaugrand et al., 2015), and also fish (Cheung et al., 2013; Vergès et al., 2016). Following SST trends, the most dramatic changes in diversity across most MPGs are expected to occur in the Arctic Ocean (more than 100% average increase over latitude; Figure 4B). In this biome, copepods and photosynthetic bacteria should experience the most dramatic increases in diversity, mostly because these communities are currently poorly diverse. The low values for endophyto-symbionts resulted in them exhibiting a large relative increase in diversity as well, especially at high latitudes (for absolute anomalies values, see Figure S12). All of these observations are in line with the poleward range expansion predicted for phytoplankton (Barton et al., 2016), in particular the cyanobacterial group *Synechococcus* (Flombaum et al., 2013) as well as for boreal fish species (Frairer et al., 2017), as a short-term response to poleward shifts in thermal niches (Thomas et al., 2012).

Hence, epipelagic planktonic communities are predicted to be strongly affected in the future, primarily by rising temperatures. Changes in chl (a higher uncertainty), either bound to primary production or photo-acclimation (Behrenfeld et al., 2015), should have more secondary effects, except in restricted areas for heterotrophic bacteria and marine copepods, where their effects seem to override those of SST (Figure S12D). In any case, the changes in MPG diversity predicted to occur by the end of the century will most probably induce cascading changes over the entire marine food web; e.g., by causing trophic mismatches or altering host-parasite/symbiont interactions (Doney et al., 2012; Edwards and Richardson, 2004; Gilg et al., 2012). For example, the increase in diversity and abundance of phototrophic bacteria suggested by our results and others (Hutchinson and Fu, 2017) would reduce upward energy flow in marine food webs because these taxa are usually less palatable for higher trophic levels (Ulhas et al., 2018). Likewise, increased temperature and diversity may also lead to reduced organism body size (Sommers et al., 2017). If the diversity and abundance of small-bodied organisms are to increase, then this may again reduce the energy transfer to higher trophic levels (Beaugrand et al., 2008).

Finally, we assessed the current ocean socio-economic and conservation status of the most affected latitudes in terms of MPG diversity (i.e., the 25% of latitudes with the highest mean absolute diversity anomalies). We did so by quantifying the current particulate carbon export, the maximum marine fisheries catch, and the number of marine protected areas at each latitude relative to global average expectations. We found that the most affected latitudes in the future currently exhibit a higher fisheries catch (32%–70% above average), carbon export (23%–70% above average), and fraction of marine protected areas (up to 100% above average; Figure 4B; Table S1I). This raises the question of how changes in diversity under the most severe climate warming scenario will affect global biogeochemical processes such as carbon export and sequestration, which are believed to have already been affected by climate change (Brun et al., 2019), and what would be the consequences for marine life in general, from already vulnerable marine animals and fish landings to life in the deep sea.

**Concluding Remarks**

The present findings and projections need to be interpreted carefully. Although being the largest systematic sampling effort of...
oceanic plankton to date, our sampling is limited by its punctual nature in space and time. Our models are also correlative and do not directly account for the effects of other abiotic parameters, such as in situ solar irradiance and their seasonal variations as well as biotic interactions and their dynamics, which should all influence plankton diversity. Regarding our projections, there are strong uncertainties about the potential lag between environmental changes and the response of plankton diversity as well as the adaptation potential of planktonic species to climate change, which can be relatively rapid, as proposed recently for zooplankton (Peijnenburg and Goetze, 2013) and diatoms (Schaum et al., 2018). Further studies better accounting for these different and intertwined mechanisms that operate at multiple spatial and temporal scales will be instrumental to improve our understanding of the drivers underlying ocean plankton ecosystems and their feedback with global change. Nevertheless, our approach is a first attempt to embrace this biological complexity at the global scale, and our results broadly agree with other statistical or theoretical projections (Barton et al., 2010, 2016; Righetti et al., 2019; Rombouts et al., 2009; Thomas et al., 2012; Tittensor et al., 2010). Our results should therefore be seen as a baseline and a framework for testing new hypotheses about changes in diversity within the whole plankton community across the global ocean, identifying the most vulnerable areas, and to better appreciate and anticipate functional and socio-economic consequences (Cavicchioli et al., 2019). These results will be helpful for guiding future broad and macroscale strategies to mitigate the effects of climate change on marine diversity and ecosystem services.

**STAR METHODS**

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

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DATA AND CODE AVAILABILITY

SUPPLEMENTAL INFORMATION

Supplemental Information can be found online at https://doi.org/10.1016/j.cell.2019.10.008.
A video abstract is available at https://doi.org/10.1016/j.cell.2019.10.008#mmc3.

CONSORTIA
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AUTHOR CONTRIBUTIONS

DECLARATION OF INTERESTS
The authors declare no competing interests.

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STAR METHODS

KEY RESOURCES TABLE

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LEAD CONTACT AND MATERIALS AVAILABILITY

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Chris Bowler (cbowler@biologie.ens.fr).

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Samples were derived from 189 stations over the 210 surveyed during the Tara Oceans expedition (2009-2013; Figure S1). They were collected across all major oceanic provinces using the sampling strategy and methodology described in Pesant et al. (2015). Briefly, the sampling was conducted at different water depths, i.e., at the sea surface (<5 m), the deep chlorophyll maximum (17-188 m) and the mesopelagic realm (200-1000 m). Sampling of the full, trans-kingdom planktonic diversity was performed with different protocols depending on their post processing, i.e., either for DNA-based analyses or for imaging analyses.

For samples dedicated to DNA analyses, we maximized the taxonomic breadth of our diversity assessment by fractionating planktonic communities from pumped seawater with filters of different mesh size. We considered samples collected with filters of 0.22-1.6/3 µm (hereafter 0.22-3 µm) for viruses and bacteria (prokaryotic viruses from the filtrate, giruses and bacteria from the 0.22 µm filter), and 0.8-2000 µm (0.8-3 µm for non-Arctic mesopelagic samples), 5-200 m, 20–1800 µm and 180–2,000 µm for eukaryotes. Prokaryotic viruses were flocculated using iron chloride (John et al., 2011). Preliminary analyses showed that the samples obtained at 0.8-2000 µm mesh size were representative of the whole structure and diversity of protists and even of copepods, whose high dominance allowed a straightforward detection by this protocol too, probably due to the presence of small life-stages or individuals, pieces of large ones, extracellular DNA from cell turnover, or fecal pellets. We hence restricted our analysis of the main eukaryotic planktonic groups on this particular subset of samples, but also provide diversity estimates at the scale of the whole eukaryotic planktonic community for each filter size to support the robustness of the results.
Imaging data acquisition followed different protocols depending on the organisms targeted. First, microphytoplankton were sampled at the sea surface with nets of 20-180 μm mesh size for microscopy analyses (see below), as described in Malviya et al. (2016). Large protists and metazoans were collected with four different nets: WP2, bongo and regent, with mesh sizes of 200, 300 and 680 μm, respectively, which were towed vertically from 500 m to the surface. We also used a muntinet, with mesh size of 300 μm, from which only the deepest level matching the mesopelagic realm was analyzed (Pesant et al., 2015). Picoplankton samples were prepared for flow cytometry from three aliquots of 1 ml of seawater (pre-filtered through 200-μm mesh), as described in Hingamp et al. (2013) and Sunagawa et al. (2015). Finally, we made use of data derived from the Malaspina expedition (Salazar et al., 2016) to account for diversity patterns of free-living prokaryotes in the bathypelagic realm (Figures 2D and S7). It should be noted that the numbers of stations and samples examined varied according to the combination of protocol and size fraction being analyzed (Figure S1; Table S1A).

**METHOD DETAILS**

**Physical and environmental measurements**

Measurements of temperature, conductivity, salinity, depth, pressure, and oxygen were carried out at each station with a vertical profile sampling system (CTD-rosette) and Niskin bottles following the sampling package described in Picheral et al. (2014). Chlorophyll a (chl a) concentrations were measured using high-performance liquid chromatography (Ras et al., 2008; Van Heukelem and Thomas, 2001). Phosphate and silicate concentrations were determined using segmented flow analysis (Aminot et al., 2009). The contribution of three pigment size classes (micro-, nano-, and picoplankton) to the total phytoplankton biomass was estimated based on high pressure liquid chromatography (HPLC) analyses (Ulitz et al., 2006). A full description of the performed measurements is described in (Pesant et al., 2015). Finally, we complemented these in situ measurements with (i) the average intra-annual variation of sea surface temperature (IAV SST) between years 1997-2017, which we obtained from the Extended Reconstructed Sea Surface Temperature v5 (Huang et al., 2017), (ii) the annual maximum of nitrate concentration (AM NO₃) retrieved from the World Ocean Atlas 2009 (Garcia et al., 2010), (iii) the intra-annual variation of the mixed layer depth (IAV MLD), which was derived from a monthly climatology (Holte et al., 2017), (iv) iron levels, which were derived from a global circulation model (Menemenlis et al., 2008), and (v) median sunlight in the mixed layer, which was estimated as in Behrenfeld et al. (2015).

**Plankton classification, diversity, and abundance estimates**

A combination of molecular and optical methods were used to describe the planktonic diversity of the ocean. A full description of the molecular data production is available in Alberti et al. (2017). Viral and prokaryotic metagenomes were obtained by shotgun metagenomics, for which sequencing, assembly and/or annotation are described in Gregory et al. (2019) for bacterial and archaeal DNA viruses, Hingamp et al. (2013) for nucleo-cytoplasmic large DNA viruses (i.e., giruses, also referred to as NCLDVs or giant viruses in the literature), and Sunagawa et al. (2015) for bacteria and archaea.

Two families of prokaryotic viruses, Myo- and Podoviridae, were studied on the basis of a capsid protein gene (gp23) and a DNA polymerase (polA), respectively (Adriaenssens and Cowan, 2014). We kept populations with a gp23/polA match either via annotation (Pfam, InterProScan and KEGG) or a set of in silico primers to increase the sensitivity (Adriaenssens and Cowan, 2014), and their abundance corresponded to the normalized number of reads that mapped against these genes. Analogously, the diversity of giruses was based on another DNA polymerase gene, polB, specifically recruited with pplacer (Matsen et al., 2010) from a non-redundant gene catalog (OM-RGCv2; Salazar et al., 2019). The corresponding frequency data was obtained by mapping the raw reads to the gene catalog. Note here that ssDNA and RNA viruses were not analyzed.

Prokaryotic taxa were defined on the basis of metagenomic reads that contained signatures of the 16S rRNA genes (referred to as mTags; Logares et al., 2014; Salazar et al., 2019; data accessible at https://www.ocean-microbiome.org/). Briefly, mTags were mapped to cluster centroids of taxonomically annotated 16S/18S reference sequences from the SILVA database (Pruesse et al., 2007) (release 128; SSU Ref NR 99) that had been clustered at 97% sequence identity beforehand using USEARCH v9.2.64 (Edgar, 2010). Mapping of mTags to a unique reference sequence were used to compute the abundances of MOTUs. A MOTU abundance table was built by counting the number of mTags assigned to each reference sequence in each sample. The abundance table was normalized by the total sum for each sample after excluding MOTUs that corresponded to eukaryotes and chloroplasts.

Additionally, we analyzed data obtained by amplicon sequencing of the V4-V5 region of the 16S rRNA gene (primers 515F-Y and 926R; Parada et al., 2016), following the pipeline described in https://github.com/SushiLab/Amplicon_Recipes. Briefly, paired-end reads were merged at a minimum 90% identity, and those with ≤1 mismatches were selected. Primer matching was performed with CUTADAPT v.1.9.1. Dereplication, MOTU clustering at 97% (UPARSE algorithm) and zOTUs denoting 100% similarity (UNOISE algorithm) were performed with USEARCH v.10.0.240 (Edgar 2010). MOTUs and zOTUs were taxonomically annotated against the SILVA database v132 (Quast et al., 2013) with the Last Common Ancestor approach. Non-prokaryotic MOTUs (eukaryotes, chloroplast, and mitochondria) were removed, whereas singletons were maintained. This dataset was used to ensure that the diversity estimates obtained with metagenomics and amplicon sequencing approaches were consistent (see below; see Figure S6). For flow cytometry data we defined six different groups: low and high nucleic acid-content heterotrophic bacteria, Prochlorococcus, Synechococcus and two groups of picoeukaryotes (see Data and Code Availability). This latter set of samples was mainly used to determine the cell density of each bacterial/picoeukaryote group, which we considered as contextual data in downstream analyses.
Nevertheless, we also retrieved diversity values from these data in order to assess their congruence at broad scale with those obtained through molecular approaches. They were not used to standardize DNA-based taxa abundances.

The taxonomic composition of protists and small metazoan communities was characterized through DNA metabarcoding using mainly the V9 region of the 18S rRNA gene, and the V4 region was also used to assess the congruence of the MOTU diversity estimates between the two DNA markers (Figure S6). For both sequencing reads datasets, we obtained a list of MOTUs as defined by the “swarm” algorithm (Mahé et al., 2014). Each MOTU was represented by a number of sequencing reads, which we used as a proxy for abundance. A full description of the sequencing reads processing (i.e., data curation, clustering into MOTUs, taxonomic classification, etc.) is available at http://taraoceans.sb-roscorf.fr/EukDiv/. Microphytoplankton were also identified and quantified manually using an inverted light microscope, as described in Malviya et al. (2016). The identification was performed by experts and reached the genus level for most of the 440 morphotypes identified (Table S1A). About half of these taxa corresponded to diatoms. Smaller protists in 5-20 μm size fractions from surface and deep chlorophyll maximum layers were retrieved from Colin et al. (2017) and obtained by environmental high content fluorescence microscopy (eHCFM).

The taxonomic classification of mesozooplankton collected with nets was performed on formaldehyde fixed samples scanned with the ZooScan imaging system (Gorsky et al., 2010) and identified with the help of an automatic recognition algorithm to the deepest possible taxonomic level using Ecotaxa (Picheral et al., 2017). The resulting identifications were validated by specialists, and reached different taxonomic levels, mostly the family level (or genus in some cases, e.g., copepods from the WP2 net). All images are accessible within Ecotaxa (https://ecotaxa.obs-vlfr.fr/). Mesozooplankton absolute abundances were calculated by taking into account the volume of water filtered by the nets. Together with images, various morphological measurements were obtained (also accessible within Ecotaxa). Major and minor best ellipsoidal axis were used to calculate the ellipsoidal biovolume of each organism that was used as a proxy of biomass. All other morphological measurements (such as length, elongation, gray level values and distribution; except those related to position of organisms within the initial scan) were recovered, normalized and used in a t-SNE analysis (van der Maaten and Hinton, 2008) using MATLAB software using the default settings (Euclidean distances; Barnes-Hut algorithm; perplexity of 30; exaggeration of 4; learning rate of 500). Different combinations of parameters were tested without clear improvements to the final result shown in Figure 2B for the bongo net. t-SNE results were used to overlay taxonomic information on the morphological overview of the imaging datasets.

Summary statistics of our datasets and their taxonomic resolution are provided in Table S1A. Based on their taxonomic affiliation, we classified all taxa into marine plankton groups (MPGs) following the criteria indicated in Table S1B. We did so not only to separate organisms of different broad functions, but also to minimize biases that could arise when comparing organisms with contrasting body size or marker gene copy number per organism (see below). For viruses we considered the three families mentioned above separately as they are the most abundant groups and have different ecologies (Brum et al., 2015; Hingamp et al., 2013; Roux et al., 2016). For prokaryotes, we distinguished photosynthetic bacteria (i.e., cyanobacteria) from heterotrophic/chemotrophic bacteria and archaea. For protists, we used an extended version of the functional database used in de Vargas et al. (2015), which encompasses a wide variety of protist taxa that are assigned to major functional groups: photosynthetic/mixotrophic protists, endophotysymbiotic hosts with endophotosymbionts (hereafter photohosts), parasitic protists, and free-living heterotrophs or phagotrophs (hereafter heterotrophic protists). Note that the endophotosymbiont group is probably the most incomplete due to the difficulties in currently being able to define comprehensively these organisms. For the mesozooplankton, the categories used corresponded to the most abundant taxonomic groups (such as copepods and chaetognaths) or feeding strategies (Figure S2). We here only considered MPGs for which the total relative abundance in the molecular dataset was > 1%, a threshold under which we considered that the detection level was too low to obtain reliable detection and diversity estimates. In total, we thus studied the diversity of 12 MPGs, of which a full list is provided in Table S1B.

**QUANTIFICATION AND STATISTICAL ANALYSIS**

**Diversity estimates calculation and validation**

Plankton diversity was estimated at each station with the Shannon diversity index, a robust measure of entropy. We chose this index rather than richness because, unlike richness, the Shannon index is insensitive to sampling effort, provided that the sampling is not too shallow (Jost, 2006). As such, the sampling effort in our datasets – albeit very deep – varied across samples but the rarefaction curves drawn with the Shannon index were largely saturating contrary to those based on richness (Figure S3). The Shannon index has also been shown to provide more reliable diversity estimates when using DNA-based data (Bálint et al., 2018; Haegeman et al., 2013). Finally, by construction, it also relates monotonically with species richness, and should therefore exhibit similar patterns (Jost, 2006). The Shannon index was calculated separately for each MPG using the samples filtered at < 0.22 m and 0.8-2000 m for prokaryotic viruses, bacteria/giruses (metagenomics) and eukaryotic plankton (DNA metabarcoding), respectively. We also calculated the Shannon index for the full local planktonic communities (i.e., not parsed into MPGs) for each sampling protocol (i.e., metagenomics, metabarcoding, and imaging for each sampling mesh size). To ensure that our Shannon values were robust, we computed their variation from 100 Monte Carlo simulated communities (function EntropyCI, R-package ‘entropart’ v1.6-1, https://cran.r-project.org/web/packages/entropart/; evaluated only for the eukaryotic plankton). Each variation range was very narrow and seldom overlap with the rest (difference between Shannon values from simulated communities, 0.003 ± 0.003; difference between Shannon values from samples; 0.969 ± 0.700), making this uncertainty negligible as compared to that generated by our downstream
analyses. Shannon diversity for each MPG retrieved from microscopy/imaging data was in general not assessed due to low taxonomic resolutions, with the exception of microphytoplankton/diatoms identified with light microscopy, and the copepods collected with the WP2 net due to their relatively high taxonomic resolution (Figure S6), as explained above.

Because every method to assess biodiversity has limitations, either due to technical issues (e.g., sampling difficulties, taxonomic resolution, lack of morphological/genetic differences) or more fundamentally due to the difficulties of classifying the diversity of life, we also assessed the congruence of diversity trends across methodologies to ensure the reliability of our conclusions. We therefore provide in Figure S6 several correlation analyses of the diversity patterns observed with a wide array of methodologies used in Tara Oceans, using previously published or newly released datasets. These comparisons include DNA-based diversity trends obtained with different markers, taxonomic resolutions, and size fractions, as well as diversity trends obtained with different optical methods.

More specifically, we compared the diversity trends for (a) different DNA markers (V9 versus V4 region of the 18S rRNA gene), (b) different taxonomic resolutions using different clustering similarity thresholds, and (c) molecular versus optical approaches (the latter based both on abundance or biovolume). More details about the datasets are available in the legend of Figure S6. Additionally, we evaluated the potential effect of marker gene copy number in prokaryotes by correcting for gene copy numbers of the 16S rRNA gene. The correction was performed using copy number estimates of references in the SILVA database (v136; Louca et al., 2018), from which we could assign a 16S rRNA gene copy number value for almost all miTags in our dataset (99%). After the correction, Shannon values remained essentially unchanged (Pearson’s r correlation between corrected and uncorrected Shannon values = 0.99, p < 0.001). Furthermore, we relied on the strong correlation we showed in a previous study between Shannon values for bacterial OTUs defined either by 16S rRNA gene or by single-copy genes (Milanese et al., 2019).

**Latitudinal diversity gradient**

Our first objective was to explore visually the LDG trend across all domains of life using both MOTU and morphotype diversity at each station. To this end, we used generalized additive models (GAMs; Hastie 2017) due to their ability to fit non-linear and/or non-monotonic functions which we expected between diversity and latitude. GAMs are further highly suitable for modeling large scale trends (Guisan et al., 2002). Additionally, we preferred the GAM smoothing approach over simple moving averages because priors are directly learned from the data and the sensitivity to extreme values is relatively low. For this particular analysis, GAMs were used only for visualizing the diversity trendline, and were not used in downstream analyses (see “Diversity modeling of MPGs”). Next, we analyzed the shape of the LDG of each marine planktonic group in two ways. First, a segmented regression analysis was conducted to describe the form of the latitudinal gradient on absolute latitude. More specifically, we aimed at detecting latitudinal breakpoints and changes in slopes across latitude. For this we used the R package ‘segmented’ 0.5-3.0 (https://cran.r-project.org/web/packages/segmented). In order to determine which MPG displayed similar LDG forms, we computed pairwise Euclidean distances between the obtained set of latitudinal breakpoints and slope values for each MPG, and subjected the resulting distance matrix to a hierarchical clustering analysis (Figure S5). In an additional analysis, we further determined whether the LDG of each MPG exhibited a North-South asymmetry. To this end, we performed separate linear regressions for each hemisphere (Table S1C).

**Diversity modeling of MPGs**

Our second objective was to find predictors for local diversity of marine planktonic groups. To this end, we also used GAMs (see “Latitudinal diversity gradient” for description and references). All GAMs were built using the R library ‘mgcv’ 1.8-24 (https://cran.r-project.org/web/packages/mgcv/), using only MOTU diversity of each MPG as response variable. Smooth terms (‘s’) were based on a thin plate regression splines and estimated by a Laplace approximation marginal likelihood criterion (Wood, 2010). The rest of the parameters were set on default mode.

In order to test the different hypotheses explaining plankton latitudinal diversity gradients and patterns of diversity in general, we first made a selection of variables from the Tara Oceans contextual data and public databases (see “Physical and environmental measurements”) that relate to these hypotheses and/or that were sufficiently available across our global sampling. We then evaluated their redundancy and link with MPG diversity by conducting multiple pairwise correlation analyses with the Spearman rank correlation test (Figures 3A and S8). We considered the contextual variables associations having |Spearman’s r| > 0.6 to be highly correlated and kept only the most representative and biologically meaningful variable among correlated ones to avoid colinearity in downstream analyses. We excluded null hypotheses for the LDG, such as the area (Willig et al., 2003) or mid-domain effect (Colwell and Lees, 2000) due to the high interconnectivity of the global ocean, which should limit the geometric constraints imposed on species distribution on lands.

The associations of plankton diversity with the four selected contextual variables (i.e., SST, chlorophyll a, annual maximum of nitrate concentration, and intra-annual variation of SST) were further analyzed using GAMs, as we expected them to be non-linear or non-monotonic functions based on previous studies (e.g., Tittensor et al., 2010; Vallina et al., 2014). Except for SST, the contextual variables were log10-transformed. For chl a, three low-concentration outlier values were excluded. “Individual” GAMs were built for each MPG and each contextual parameter, from which the explained deviance was used as an association measure and the approximate p value of the smooth term to account for effect significance (Figures 3B and S9). All p values obtained were corrected for multiple comparisons (Holm, 1979). To further test the different LDG hypotheses, we then built “full” GAMs for each MPG that included all four contextual variables, with the settings explained above. Their additive contribution was calculated by a sequential removal of the different parameters and a normalization with a null model (Figure S5).
From these analyses, we identified temperature and chlorophyll \( a \) to be the best correlates with most MPGs diversity. These two variables also capture relatively well other environmental gradients, such as cyanobacteria and mesozooplankton abundance (Figure S1A). Given their strong explanatory power and because their current and future state at the sea surface can be simulated with global ocean circulation models (Figure S11; Bopp et al., 2013), we used them to predict the current global-scale distribution of MPGs diversity, as well as its response to a severe scenario of climate and oceanic change. To this end, we built a set of “reduced” GAMs with surface diversity for each MPG as response variable, and SST, chlorophyll \( a \), as well as an interaction-like term (included as a tensor product, ‘\( t^2 \)’), as explanatory variables. SST and chlorophyll \( a \) (in situ measurements) were only partially anti-correlated, probably due to a decoupling in upwelling systems (Demarcq, 2009). Accordingly, the explained deviance of some of our GAMs was increased by 10% or more when considering these two parameters without affecting their parsimony.

Both “full” and “reduced” GAMs were built with the same approaches as described above and were further validated with two additional analyses. First, we quantified the congruence between observed and GAM-modeled Shannon diversity values through a Pearson’s correlation analysis. Second, we ensured that the GAM residuals did not exhibit latitudinal or longitudinal trends, a way to control for spatial autocorrelation (Tables S1E–S1G). Checking the absence of latitudinal trends in the model residuals further indicates if our models successfully explained the latitudinal gradients of diversity. To further assess the performance of the “reduced” models used downstream for predicting current and future trends of diversity, we cross-validated them with other independently collected molecular datasets from open-ocean studies targeting either heterotrophic bacteria (Zinger et al., 2011) or the whole planktonic eukaryotic community (Raes et al., 2018) from different sampling dates and locations. Splitting this latter dataset into MPGs was not possible due to the unavailability of functional databases for the DNA marker used (V4 region of the 18S rRNA gene), so we conducted this cross-validation with a reduced GAM model built for the whole eukaryotic community. Figure S13 shows that the GAM models built with our datasets are able to predict correctly the diversity trends of plankton communities observed in these independent datasets (see complementary details in the caption of Figure S13 and below for CMIP5 models). Additionally, as both datasets include sampling points in the Southern Ocean, which was undersampled in ours, this agreement confirms the decrease in plankton diversity we observed toward the south.

Next, the MOTU diversity of the six MPGs for which “reduced” GAMs yielded a deviance explained \( \geq 60 \% \) was modeled for the beginning and the end of the 21st century. To do so, we first defined a coarse-grained arrange of 1° grid-cells. SST and chlorophyll \( a \) content across space and time were obtained from the Coupled Model Intercomparison Project Phase 5 (CMIP5), a multi-model simulation of the ocean (Bopp et al., 2013). Each model within CMIP5 is an Earth system simulation generated by different research groups (Table S1H), which allows us to account for different mechanistic weights. We extracted the two variables from each CMIP5 model for each grid cell for the beginning of the 21st century (averaged values for years 1996-2005) and the end of the 21st century (averaged values for years 2090-2099), the latter considering RCP 8.5 scenario, the most pessimistic IPCC trajectory for greenhouse gases concentration (radiative forcing level reaches 8.5 W/m\(^2\)). To obtain an average and assess the uncertainty in our predictions, we generated a combined calculation of the uncertainty of the GAM parameters and the multiple CMIP5 models. We did so by first obtaining posterior distributions of the fitted GAM parameters for the different plankton groups. We then sampled values from these distributions randomly to generate 1,000 models. For each of these models, we used each of the CMIP5 models \( (n = 13) \) as current and future temperatures and chl \( a \). Shannon diversity was then predicted with each of the 13,000 models we generated for each MPG and each time of projection (beginning and the end of the century), from which we assessed the uncertainty of our predictions.

Anomalies between future ocean projections and estimates for the beginning of the 21st century were calculated as the difference between the average diversity of each planktonic group projected for the time interval 2090-2099 and the one for 1996-2005 (Figures 4 and S12). In other words, a positive anomaly means that the predicted diversity will increase toward the end of the century. Confidence intervals of anomalies were based on the standard deviations of the average Shannon diversity estimates across the different CMIP5 models. Finally, to assess potential areas in the future ocean where the effect of primary production change on diversity could override the effect of temperature, we performed diversity projections holding either SST or chlorophyll \( a \) constant, and then comparing their output. That is, we assessed per grid-cell whether the effect of chlorophyll \( a \) on diversity was significantly larger than zero and larger than the one of SST (in absolute values; see Figure S12D). Manipulation and visualization of the CMIP5 spatial data and the corresponding diversity projections were performed combining R packages ‘ncdf4’ 1.16 (https://cran.r-project.org/web/packages/ncdf4), ‘raster’ 2.5-8 (https://cran.r-project.org/web/packages/raster) in R v.3.5.1 (https://www.r-project.org).

**Comparison of future trends with current areas of high socioeconomic and conservation value**

We identified the latitudes that are expected to experience the most dramatic changes in diversity (defined as the 25% of latitudes with the highest mean absolute diversity anomalies) and analyzed whether these areas overlap with current ecosystem services and reserves. To this end, we compared their corresponding current status in terms of (i) carbon export, using satellite-derived estimates at 100 m depth (Henson et al., 2012), (ii) maximum marine fisheries catch between years 2010-2014 (Watson, 2017), and (iii) number of marine protected areas (Bruno et al., 2018) relative to the global average. In all cases, the difference was expressed as the relative (%) increase or decrease in relation to the global average (Table S1I).
DATA AND CODE AVAILABILITY

Raw reads of Tara Oceans are deposited at the European Nucleotide Archive (ENA). In particular, newly released 18S rRNA gene metabarcoding reads are available under the number ENA: PRJEB9737. ENA references for the metagenomics reads corresponding to the size fraction \(< 0.22 \, \mu m\) (for prokaryotic viruses) analyzed in this study are included in Gregory et al. (2019); see their Table S3. ENA references for the metagenomics reads corresponding to the size fraction 0.22-1.6/3 \(\mu m\) (for prokaryotes and viruses) correspond to Salazar et al. (2019) (see https://zenodo.org/record/3473199). Imaging datasets from the nets are available through the collaborative web application and repository EcoTaxa (Picheral et al., 2017) under the address https://ecotaxa.obs-vlfr.fr/prj/412 for regent data, within the 3 projects https://ecotaxa.obs-vlfr.fr/prj/397, https://ecotaxa.obs-vlfr.fr/prj/398, https://ecotaxa.obs-vlfr.fr/prj/395 for bongo data, and within the 2 projects https://ecotaxa.obs-vlfr.fr/prj/377 and https://ecotaxa.obs-vlfr.fr/prj/378 for WP2 data. A table with Shannon values and multiple samples identifiers, plus a table with flow cytometry data split in six groups are available (https://doi.org/10.17632/p9r9wtjkm.1). Contextual data from the Tara Oceans expedition, including those that are newly released from the Arctic Ocean, are available at https://doi.org/10.1594/PANGAEA.875582.
Supplemental Figures

A

Myoviridae | n = 60
Podoviridae | n = 60
Giruses | n = 83

Bacteria (P) | n = 82
Bacteria (H) | n = 83
Archaea | n = 81

Protists (P) | n = 89
Endophotosymb. | n = 89
Photohosts | n = 89

Parasitic protists | n = 89
Protists (H) | n = 89
Copepods | n = 99

B

16S | 0.22-3 µm | n = 83
18S | 0.8-2000 µm | n = 89
18S | 20-180 µm | n = 134

18S | 180-2000 µm | n = 128
FC | <20 µm | n = 110
LM | 20-180 µm | n = 58

Imaging | 200 µm | n = 152
Imaging | 300 µm | n = 81
Imaging | 680 µm | n = 120

C

16S | 0.22-3 µm | n = 35
18S | 0.8-3 µm | n = 40
Imaging | 300 µm | n = 28

(legend on next page)
Figure S1. Tara Oceans Stations and Shannon Diversity Patterns, Related to Figures 1 and 2

(A) MPGs at the sea surface (< 5 m depth), (B) whole planktonic community using different sampling protocols at the sea surface (except for “Imaging,” integrative depth from 500 m depth to the surface) and (C) whole planktonic community of the mesopelagic realm (200–1000 m depth). Number of stations are specified in the inset titles. Color represents the Shannon index. For more details on the different size fractions and sampling protocols, please refer to the caption in Figure 1 and STAR Methods.
Figure S2. Average Abundances of a Broader List of Plankton Groups across Latitude, Related to Figures 1 and 2

For 18S rDNA metabarcodes (relative abundances), imaging from net catches and flow cytometry (absolute abundances). Numbers refer to the filter mesh size (µm). H: non-photosynthetic/heterotrophic, P: photosynthetic/mixotrophic. See Figures 1 and S1 for further sampling details. The three viral groups are not represented here due to absence of comparable abundance data. Note that the differences between protocols relate, among other things, to resolution (e.g., potential photohosts from the nets are classified as Protists (H)), marker gene copy number (e.g., high in photohosts), lack of detection (many small copepods are lost when sampling with nets), or water column sampling differences (surface [SRF] or deep chlorophyll maximum [DCM] versus integrative [INT] for molecular/cytometry versus net catches, respectively).
Figure S3. Rarefaction Curves for the Plankton Community, Related to Figure 2 and STAR Methods
Based on richness (A, C) and Shannon (B, D), for prokaryotes (16S rRNA gene miTags, A, B), and eukaryotes (18S rRNA gene V9 metabarcoding, C, D). Each line corresponds to a surface water sample. Colors correspond to different latitudinal bands (absolute values).
Figure S4. Sea Surface LDG, Related to Figure 2A

(A) For viral, prokaryotic and eukaryotic MPGs, and (B) for well-known protist phyla and the dominant class of bacteria, Alphaproteobacteria, for which \( \sim 50\% \) corresponds to the SAR11 clade. Solid lines represent the GAM smooth trends and gray ribbons the corresponding 95% confidence intervals of the Shannon latitudinal trend predicted by the GAMs. The percentages provided below inset titles correspond to the deviance explained by GAMs when significant. Viruses and bacterial diversities are inferred from samples filtered at 0.22-3 \( \mu \text{m} \) and analyzed through marker genes derived from metagenomics. Eukaryote diversity shown here is inferred from 18S rDNA metabarcoding of samples filtered at 0.8-2000 \( \mu \text{m} \). H: non-photosynthetic/heterotrophic, P: photosynthetic/mixotrophic.
Figure S5. Classification of MPG Sea Surface LDG Analyzed by Segmented Regression, Related to Figure 2A

The estimated break is the absolute latitude between the two segments of slope $s_1$ and $s_2$, respectively. We used pink lines for $s_1 \leq 0$ (plateau or peak around the equator) and blue lines for $s_1 > 0$ (extra-equatorial peak). The dendrogram is the result of a hierarchical clustering based on the differences in break and slope values across MPGs (Euclidean distance on standardized values).
Figure S6. Correlation between the Shannon Values Derived from Multiple Datasets of Tara Oceans, Related to STAR Methods

(A) OTUs (as defined with “swarm”; Mahe et al., 2014) obtained with the V9 (x axis) and V4 (y axis) regions of the 18S rRNA gene using surface water samples (SRF); size fraction 0.8-2000 μm; (B) OTUs either as defined with swarm (x axis) or defined at 100% sequence identity from the V9 region of the 18S rRNA gene for SRF samples, size fraction 0.8-2000 μm; (C-D) 16S rRNA gene miTags (x axis) versus OTUs defined at 97% [C] and 100% sequence similarity [D] (y axis) obtained from the V4-V5 regions of the 16S rRNA gene for SRF samples, size fraction 0.22-3 μm; (E) OTUs of photosynthetic protists obtained with the V9 region of the 18S rRNA gene (x axis) versus protists (mostly photosynthetic) as identified with environmental High Content Fluorescence Microscopy (eHCFM; data from Colin et al., 2017) in SRF-DCM samples, size fraction 5-20 μm; (F) Diatom OTUs obtained with V9 region of the 18S rRNA gene (x axis) versus diatom species counted by light microscopy (y axis) in SRF samples; size fraction 20-180 μm; (G/H) Copepod OTUs obtained with the V9 region of the 18S rRNA gene, SRF samples, size fraction 180-2000 μm (x axis) versus abundances [G] and biovolumes [H] of copepods collected by the WP2 net, > 200 μm. Inset titles show the Pearson’s correlation coefficient and its associated p value. Note the differences in axes scales. Dashed line represents 1:1 relation. Refer to STAR Methods for details on each method.
Figure S7. LDG across Size and Depth, Related to Figures 2B–2D
For the whole prokaryotic (16S miTags, 0.22–3 μm, and 16S OTUs, 0.2 μm for bathypelagic (BAT)) and eukaryotic communities (18S OTUs, 0.8–2000, 20–180 and 180–2000 μm; imaging, > 300 and > 680 μm) at different depths (SRF: surface, < 5 m; DCM: deep chlorophyll maximum, 17–188 m, and MES: mesopelagic, > 200 m, BAT: bathypelagic, > 4000 m, INT: integrative, depth from 500 m depth to the surface). Non-significant GAMs are denoted with “NS.” See Figure S4 legend for more information on the plot. Note that the particular trend for the regent net, i.e., Imaging | 680 μm might be due to undersampling of small zooplankton.
Figure S8. Multiple Pairwise Spearman Correlation Analysis of the Full Matrix of Contextual Parameters for the Surface Ocean, Related to Figure 3A.

Rows and columns were clustered based on the absolute pairwise Spearman correlation turned into distance (1 - |ρ|). MLD: mixed layer depth. E_median ML: median light in the mixed layer. IAV: intra-annual variability. Part.backscat.coef: particle backscattering coefficient. For more information on parameters, see Figure 3 and STAR Methods.
Figure S9. Relationships between Diversity and 4 Contextual Variables for Viral, Prokaryotic, and Eukaryotic MPGs, Related to Figure 3B
(A) SST, (B) chl a, (C) AM NO3, and (D) IAV SST. Solid lines represent the GAM smooth trends and gray ribbons the corresponding 95% confidence intervals of the x-y relationship predicted by the GAMs. The percentages provided below inset titles correspond to the deviance explained by GAMs when significant (p value corrected for multiple comparisons). Non-significant GAMs are denoted with “NS.”
Figure S10. Relationship between Plankton Abundance and Two Contextual Variables, Related to Figure 3A
(A) SST, (B) chl a. Abundance values were obtained with flow cytometry (reported in [cells/ml]) or from counts of individuals captured with nets of different mesh size and identified by imaging (reported in [individuals/m^3]), respectively. Solid lines represent the GAM smooth trends and gray ribbons the corresponding 95% confidence intervals of the X-Y relationship predicted by the GAMs. The percentages provided below inset titles correspond to the deviance explained by GAMs when significant.
Figure S11. Projected Latitudinal Changes in SST and chl $a$, Related to Figures 4 and S12
Anomalies (%) in (A) SST and (B) chl $a$ at the end of the 21st century (2090-2099, RCP 8.5) relative to the beginning of the century (1996-2006). Data was obtained from 13 CMIP5 models (Table S1H). Grey ribbons represent the standard error.
Figure S12. Modeled Patterns of Diversity of MPGs in the Global Ocean, Related to Figure 4

(A) Shannon index modeled at the global scale for oceanic conditions at the beginning of the 21st century (1996-2006). Predicted Shannon values $\leq 0$ obtained at high latitudes, particularly for copepods and endophotosymbionts, were excluded. (B) Anomalies were calculated as the difference of their Shannon index at the end (2090-2099, RCP 8.5) and the beginning of the century (1996-2006). A positive value means that diversity will increase by the end of the century. Note that the scale is not symmetric and that white means zero change. (C) Uncertainty maps (standard deviation) for (B). (D) Areas where the effect of chl $\alpha$ on plankton diversity is likely to be higher than the one of SST. To determine this, either chl $\alpha$ or SST were held constant in the projections by the end of the century. Then, if the anomaly caused only by the change of chl $\alpha$ was different than zero and higher (absolute terms) than the one caused only by the change in SST, the pixel was colored. (E) Latitudinal diversity gradient at the beginning (solid line) and the end (dashed line) of the 21st century. Values represent averages over longitude for each latitudinal degree. Dots are observed values (Figure 2). 13 Earth system models from CMIP5 were used (Table S1H).
Figure S13. Correlation between the Shannon Values Observed in Independent Datasets and Those for the Same Locations that Are Predicted by GAM Models Built in This Work, Related to STAR Methods

(A) Heterotrophic bacteria from the surface, open ocean water sites of the International Census of Marine Microbes (iCoMM; Zinger et al., 2011). (B) Eukaryotic community retrieved from Raes et al. (2018). Note that for the latter the mesh size of the filters used were not exactly the same (Tara Oceans > 0.8 μm; Raes et al., 2018 > 0.22 μm). Note also that both datasets have different sampling dates and locations in relation to Tara Oceans. In both cases we used as predictors the temperature and chl a of each site predicted by 13 CMIP5 models at the month of sampling averaged over 1996-2006. Inset titles show the Pearson’s correlation coefficient and its associated p value.