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Restructuring of genomic provinces of surface ocean plankton under climate change

Paul Frémont^{1,2*}, Marion Gehlen^{3*}, Mathieu Vrac³, Jade Leconte^{1,2}, Tom O. Delmont^{1,2}, Patrick Wincker^{1,2}, Daniele Iudicone⁴, Olivier Jaillon^{1,2*}

¹Génomique Métabolique, Genoscope, Institut François Jacob, CEA, CNRS, Université d'Evry, Université Paris-Saclay, 91057 Evry, France. ²Research Federation for the study of Global Ocean Systems Ecology and Evolution, FR2022/Tara Oceans, Paris, France. ³LSCE-IPSL, CEA/CNRS/Université Paris-Saclay, Gif-sur-Yvette, France. ⁴Stazione Zoologica Anton Dorn. Villa Comunale, 80121, Naples, Italy.

*Corresponding authors: pfremond@genoscope.cns.fr; marion.gehlen@lsce.ipsl.fr; ojaillon@genoscope.cns.fr

The impact of climate change on diversity, functioning and biogeography of marine plankton is a major unresolved issue. Here, niche theory is applied to plankton metagenomes of 6 size fractions, from viruses to meso-zooplankton, sampled during the *Tara* Oceans expedition. Niches are used to derive plankton size-dependent structuring of the oceans south of 60°N in *climato-genomic* provinces characterized by signature genomes. By 2090, assuming the RCP8.5 high warming scenario, provinces would be reorganized over half of the considered oceans and quasi-systematically displaced poleward. Particularly, tropical provinces would expand at the expense of temperate ones. Compositional shifts among planktonic grazers and nitrogen-fixing bacteria suggest impacts on the nitrogen and carbon cycles. Sea surface temperature is identified as the main driver of the changes (~51%) followed by phosphate (11%) and salinity (10%). These results demonstrate the potential of integration of genomics with physico-chemical data for higher scale modeling and understanding of ocean ecosystem dynamics.

Planktonic communities are composed of complex and heterogeneous assemblages of small animals, single-celled eukaryotes (protists), bacteria, archaea and viruses - that drift with currents. They contribute to the regulation of the Earth system through primary production via photosynthesis¹, carbon export to the deep oceans^{2,3} and form the base of the food webs that sustain the whole trophic chain in the oceans and beyond⁴.

The composition of communities varies over time at a given site with daily⁵ to seasonal fluctuations⁶ following environmental variability⁷. Overlying these relatively short scale

spatio-temporal variations, a more macroscale partitioning of the ocean has been revealed by different combinations of biological and physico-chemical data^{8–10}, and recently documented at the resolution of community genomics¹¹. The basin scale biogeographical structure has been proposed to result from a combination of multiple bio-physico-chemical processes named the seascape⁷. These processes include both abiotic and biotic interactions¹², neutral genetic drift¹³, natural selection^{14,15}, temperature variations, nutrient supply but also advection and mixing along currents^{11,13}.

Today, knowledge of global scale plankton biogeography at the DNA level is in its infancy. We lack understanding and theoretical explanations for the emergence and maintenance of biogeographical patterns at genomic resolution. Omics data (*i.e.* the DNA/RNA sequences representative of the variety of coding and non-coding sequences of organisms) provide the appropriate resolution to track and record global biogeographical features¹¹, modulation of the repertoire of expressed genes in a community in response to environmental conditions^{2,16,17} and eco-evolutionary processes^{13–15}. Moreover, metagenomic sequencing can be consistently analyzed across plankton organisms as recently demonstrated by global expeditions^{18–21}. The strong links between plankton and environmental conditions suggest potentially major consequences of climate change on community composition and biogeography^{22,23}. Time series observations have highlighted recent changes in the planktonic ecosystem attributed to anthropogenic pressures, such as changes in community composition^{24,25} or poleward shifts of some species²⁶. These changes are expected to intensify with ongoing climate warming²⁷ and could lead to major reorganization of plankton community composition²², with a potential decline in diversity^{28–30}. Another major consequence of global reorganization of the seascape on biological systems would be a decrease of primary production at mid-latitudes and an increase at higher latitudes²⁷.

Here we report the global structure of plankton biogeography south of 60°N based on metagenomic data using niche models and its putative modifications under climate change. First, we define environmental niches³¹, *i.e.* the envelope of environmental parameters suitable for an organism or a population, at the scale of genomic provinces across 6 organism size fractions representing major plankton groups from nano- (viruses) to meso-zooplankton (small metazoans). Then, we spatially extrapolate their

niches into *climato-genomic* provinces to derive the structure of plankton biogeography for each size fraction individually and for all combined. Next, considering the same niches, we assess the spatial reorganization of these provinces under climate change at the end of the century, with a focus on associated compositional shifts among copepods (planktonic grazers important for the carbon cycle) and nitrogen-fixing bacteria (important for both nitrogen and carbon cycles). Finally, we quantify the relative importance of the environmental drivers explaining projected changes.

Niche models and signature genomes from genomic provinces

We define and validate environmental niches using 4 machine learning techniques for 27 previously defined genomic provinces¹¹; they correspond to 529 metagenomes for 6 size fractions (ranging from 0 to 2000 μm) sampled at 95 sites from all oceans except the Arctic (Supplementary information 1, Supplementary Figs. 1-2). Predictor variables of the niches are sea surface temperature (SST), salinity, dissolved silica, nitrate, phosphate and iron, plus a seasonality index of nitrate.

The signal of ocean partitioning is likely due to abundant and compact genomes whose geographical distributions closely match provinces. Within a collection of 1778 bacterial, 110 archaeal and 713 eukaryotic environmental genomes^{32,33} characterized from *Tara* Oceans samples without cultivation, we find a total of 324 signature genomes covering all but 4 provinces, and displaying taxonomic signal coherent with the size fractions (Fig. 1 for eukaryotes and Supplementary Fig. 3 for Bacteria and Archaea). Some of the signature genomes correspond to unexplored lineages with no cultured representatives, highlighting the knowledge gap for organisms that structure plankton biogeography and the strength of a rationale devoid of any *a priori* on reference genomes or species.

Structure of present day biogeography of plankton

To extrapolate the niches to a global ocean biogeography for each size fraction, we define the most probable provinces, named hereafter as *dominant* and assigned to a climatic annotation (Supplementary Table 1), on each $1^\circ \times 1^\circ$ resolution grid point using 2006-13 WOA13 climatology³⁴ (Supplementary Fig. 4 and Fig. 2).

In agreement with previous observations¹¹, provinces of large size fractions ($>20 \mu\text{m}$) are wider and partially decoupled from those of smaller size fractions, probably due to differential responses to oceanic circulation and environmental variations, different life

cycle constraints, lifestyles^{7,11} and trophic network positions³⁵. Biogeographies of small metazoans that enrich the largest size fractions (180-2000 and 20-180 μm) are broadly aligned with latitudinal bands (tropico-equatorial, temperate and (sub)-polar) dominated by a single province (Fig. 2ab). A more complex oceanic structuring emerges for the smaller size fractions (<20 μm) (Fig. 2c-f) with several provinces per large geographical region. For size fraction 0.8-5 μm enriched in small protists (Fig. 2d), distinct provinces are identified for tropical oligotrophic gyres and for the nutrient-rich equatorial upwelling region. A complex pattern of provinces, mostly latitudinal, is found for the bacteria (Fig. 2e, 0.22-3 μm) and the virus enriched size classes (Fig. 2f, 0-0.2 μm) although less clearly linked to large-scale oceanographic regions. A single province extending from temperate to polar regions emerges for size fraction 5-20 μm enriched in protists (Fig. 2c), for which fewer samples were available (Supplementary Fig. 2b-c), which probably biases this result. A consensus map combining all size fractions, built using the PHATE algorithm³⁶, summarizes the main characteristics of the biogeographies described above (Supplementary information 2 and Supplementary Fig. 5).

Finally, we compare genomic biogeographies and existing ocean partitionings⁸⁻¹⁰. Though each of them is unique (Supplementary Fig. 8-10), common borders highlight a global latitudinal partitioning independent of the type of data (Supplementary information 3).

Future changes in plankton biogeography structure

We assess the impacts of climate change on plankton biogeography at the end of the century following the Representative Concentration Pathway 8.5 (RCP8.5)³⁷ greenhouse gas concentration trajectory. To consistently compare projections of present and future biogeographies, we use bias-adjusted mean of 6 Earth System Model (ESM) climatologies (Supplementary Table 2, Supplementary Fig. 11). The highest warming (7.2°C) is located off the east coast of Canada in the North Atlantic while complex patterns of salinity and nutrient variations are projected in all oceans (Supplementary Fig. 12). Following this trajectory, future temperature at most sampling sites will be higher than the mean and maximum contemporary temperature within their current province (Supplementary Fig. 13).

Our projections indicate multiple large-scale changes in biogeographical structure including expansions, shrinkages and shifts in plankton organism size-dependent provinces (Fig. 2a-d, Supplementary Fig. 14-16). A change in the *dominant* province in at least one size fraction would occur over 60.1% of the ocean surface, ranging from 12% (20-180 μm) to 31% (0.8-5 μm) (Fig. 2, Table 1).

Centroids of provinces with *dominance* areas larger than 10^6 km^2 within a basin would be moved at least 200 km away for 77% of them, 96% of which move poleward (Supplementary Figs. 15 and 16). While a few longitudinal shifts larger than 1000 km are projected, the distribution of latitudinal shifts is largely concentrated around the mean (290 km) with no shifts superior to 1000 km (Supplementary Fig. 16b). These important longitudinal shifts corroborate existing projections^{22,38,39} and differ from trivial poleward shifts due to temperature increase, reflecting more complex spatial rearrangements of the other environmental drivers (Supplementary Fig. 12). The average displacement speed of the provinces' centroids is $76 \pm 79 \text{ km.dec}^{-1}$ (latitudinally mean of $34 \pm 82 \text{ km.dec}^{-1}$, longitudinally $59 \pm 82 \text{ km.dec}^{-1}$).

Projected shifts in phytoplankton enriched provinces corroborate previously published shifts of North Pacific phytoplankton biomes: provinces C4 and C9 are projected to shift respectively at speeds of 118 km.dec^{-1} and 195 km.dec^{-1} comparable to 100 km.dec^{-1} and 200 km.dec^{-1} for the subtropical and tropical biomes of Polovina et al.³⁹.

For all size fractions, climate change would lead to a poleward expansion of tropical and equatorial provinces at the expense of temperate provinces (Supplementary information 4, Supplementary Table 2, Supplementary Figs. 14 and 17). This is illustrated by the temperate province F5 of size fraction 180-2000 (Supplementary Fig. 16), which is projected to shrink in the five major basins. In the North Atlantic, its centroid would move approximately 800 km to the northeast (Supplementary Fig. 16c). Similar trends are found comparing present day and end of the century consensus maps (Supplementary information 2, Supplementary Fig. 17).

We calculate a dissimilarity index (equation (3)) at each grid point between probabilities of future and present *dominant* provinces for all size fractions combined (Fig. 4a). Large dissimilarities are obtained over northern (25° to 60°) and symmetrically southern (-25 to -60°) temperate regions (mean of 0.29 and 0.24 respectively) mostly reflecting

the poleward retraction of temperate provinces (red arrows, Fig. 4a). In austral and equatorial regions, despite important environmental changes (Supplementary Fig. 12) and previously projected changes in diversity^{28–30} and biomass⁴⁰, the contemporary provinces remain the most probable at the end of the century (mean dissimilarities of 0.18 and 0.02 respectively).

To further study the future decoupling between provinces of different size fractions, we analyze the assemblages of *dominant* provinces of each size fraction. By using two differently stringent criteria, from 45.3 to 57.1% of ocean surface, mainly located in temperate regions, would be inhabited in 2090–99 by assemblages that exist elsewhere in 2006–15 (Fig. 4b versus Fig. 4c). Contemporary assemblages would disappear on 3.5 to 3.8% of the surface, and, conversely, novel assemblages, not encountered today, would cover 2.9 to 3.0% of the surface. While these changes are limited to a relatively modest area, they include important economic zones (Fig. 4b, Supplementary Fig. 19). Over 41.8% to 51.8% of the surface of the main fisheries and 41.2% to 54.2% of Exclusive Economic Zones, future assemblages would differ from those present today (Supplementary Fig. 19).

Future changes in the distribution of grazers and nitrogen-fixing bacteria

In order to elucidate the potential biogeochemical impact of biogeographical restructuring, we focus on compositional changes among copepods and nitrogen-fixing bacteria (a.k.a, diazotrophs), two groups considered important for the carbon and nitrogen cycles^{41,42} that are well represented among the Tara Oceans environmental genomes. Focusing on marine areas where *dominant* provinces are projected to be replaced, we compare the present and future distribution of environmental genomes corresponding to 198 copepods and 27 diazotrophs^{32,33}.

Copepods are cosmopolitan small crustaceans. These abundant grazers contribute to the biological pump and their body size is considered to be a key trait for carbon export⁴¹. They feed on smaller plankton⁴¹ and are prey to higher trophic levels. We characterize the composition of provinces using 198 environmental genomes detected in at least 5 samples and annotated as Marine Hexanauplia of clade A and B. They are further divided into five subgroups based on their differential abundances across the large size classes (>5 μ m): 8 mesozooplankton clade A, 19 unclassified clade A, 71

microzooplankton clade A, 10 unclassified clade B and 90 microzooplankton clade B. Therefore, large copepods are preferentially from clade A in our data.

The relative abundances of unclassified clade B in size class 20-180 μm (Fig. 3e) and of mesozooplankton clade A in size class 180-2000 μm , (Supplementary Fig. 20a) are projected to increase in regions where the temperate province is replaced by the tropico-equatorial province. In areas where the polar province is replaced by the temperate province, a greater relative abundance in microzooplankton clade A (1% to 10%) (Fig. 3e) is projected. No significant compositional differences in the provinces of size fraction 5-20 μm are found (Supplementary Fig. 20b). These results highlight potential significant compositional shifts by the end of the century in the different clades and sizes of main grazers.

Diazotrophy, the biotic fixation of atmospheric nitrogen, is an important process for both nitrogen and carbon cycles. It supports biological productivity in the nitrogen-limited tropical oceans⁴². Marine diazotrophs include cyanobacteria (e.g. *Trichodesmium*) described as the principal nitrogen fixers⁴³ and various heterotrophic bacterial diazotrophs (HBDs) that lack cultured representatives or *in situ* imaging^{33,44}. Forty-eight of the bacterial environmental genomes are diazotrophs (8 cyanobacteria and 40 heterotrophic bacterial diazotrophs (HBDs)), encapsulating 92% of the metagenomic signal for known *nifH* genes (a reference marker for nitrogen fixation⁴⁵) at the surface of the oceans. Twenty-seven are found in at least 5 samples and have been used for the analysis.

In size fraction 0.8-5 μm , we project significantly higher relative abundances in cyanobacteria in the tropico-equatorial regions of the Pacific ocean (C9 to C11 and C4, Fig. 3f), which might point towards an increase in nitrogen fixation in this region as previously suggested by other models⁴². Supporting this result, we find similar significant compositional changes towards an increase in cyanobacteria in size fraction 5-20 μm (Supplementary Fig. 21c). We also project significant compositional changes for some clades of HBDs (e.g. increase in gammaproteobacteria: C8 to C3, Fig. 3f) though no global trend can be identified here. In the other size classes (20-180, 180-2000 and 0.22-3 μm) compositional changes are not significant (Supplementary Fig. 22). To summarize, although we cannot estimate nitrogen fixation rates using genomic

data alone, genomic measurements of nitrogen-fixing cyanobacteria are in agreement with an increase in nitrogen fixation in the tropics, echoing results from other models⁴².

Drivers of plankton biogeography reorganization

We quantify the relative importance of environmental predictors (sea surface temperature, salinity, dissolved silica, phosphate, nitrate, iron and seasonality of nitrate) into niche definition and in driving future changes of the structure of plankton biogeography (equation (4)). Among these environmental properties, temperature is the first influential parameter (for 19 niches out of 27) but only at 22.6% on average (Supplementary Fig. 22a).

The relative impact of each environmental parameter is calculated²² for each site presenting a significant dissimilarity between 2006-15 and 2090-99 (Fig. 5a). Overall, SST would be responsible for the reorganization of the provinces at 50% followed by Phosphate (11%) and Salinity (10.3%) (Supplementary Fig. 23). Over the majority of the ocean, SST is the primary driver of the reorganization (Fig. 5a). In some regions, salinity (e.g. eastern North Atlantic) and Phosphate (e.g. equatorial region) dominate (Fig. 5a). When excluding the effect of SST, salinity and phosphate become the primary drivers of the reorganization of the provinces (Fig. 5b). The impact of SST varies across size classes with a significantly higher contribution in large size classes (>20 μm) compared to the small ones (mean of ~73% versus ~49%, t-test $p < 0.05$; Fig. 5c). Though the contribution of combined nutrients to niche definition is similar for small and large size classes (mean of ~56% versus ~61%, Supplementary Fig. 22, Supplementary Table 3), their future projected variations have a higher relative impact on the reorganization of biogeographies of small organisms (mean of ~39% versus ~20%, t-test $p < 0.05$, Supplementary Fig. 22, Supplementary Table 3). For instance, in the tropical zone, the shrinkage of the equatorial province C9 (size fraction 0.8-5 μm , Fig. 2b,d, Supplementary Fig. 24e) is driven at 24% by reduction of dissolved phosphate concentrations and at 25% by SST increase. In contrast, SST drives at 56% the shrinkage of the temperate province F5 (size fraction 180-2000 μm , Supplementary Fig. 24d). Finally, non-poleward shifts are found only within small size fractions (<20 μm) (Supplementary Figs. 14, 15) highlighting differential responses to nutrients and SST

changes between large and small organisms, the latter being enriched in phytoplankton that directly rely on nutrient supplies.

Discussion

We propose a novel partitioning of the ocean in plankton size dependent *climato-genomic* provinces, complementing previous efforts based on other bio-physico-chemical data^{9–11}. Though initially built at genomic scale, our biogeographies paradoxically reveal basin scale provinces that are larger than BGCP¹⁰ and Fay and McKingley biomes⁹. These provinces are probably relatively stable across seasons suggesting limited effects of seasonality on the position of frontiers of BGCPs provinces¹⁰. We propose that this apparent paradox emerges from the combination of the scale, nature and resolution of sampling. First, two proximal samples from the *Tara* Oceans expedition are separated by ~300 km on average sampled over three years. This relatively large spatio-temporal scale overlies shorter scale compositional variations previously observed^{5,6}. Second, our estimates of plankton community dissimilarities are highly resolute as they are computed at genomic scale with billions of small DNA fragments^{11,18} thus smoothing out the more discrete species level signal. Together, from these combinations of processes and patterns occurring at multiple scales emerge basin scale provinces associated with coherent environmental niches and signature genomes.

These *climato-genomic* provinces are structured in broad latitudinal bands with smaller organisms (<20 µm) displaying more complex patterns and partially decoupled from larger organisms. This decoupling is the result of distinct statistical links between provinces based on organism size fractions and environmental parameters and could reflect their respective trophic modes³⁵.

Complex changes of the parameters defining the niches are projected under climate change leading to the reorganization of size-dependent provinces. Assuming a constant relationship to environmental drivers that define the *climato-genomic* provinces, climate change is projected to restructure them over approximately 50% of surface oceans south of 60°N by the end of the century (Fig. 4). The largest reorganization is detected in subtropical and temperate regions in agreement with other studies^{29,39} and is accompanied by appearance and disappearance of size-fractionated provinces'

assemblages. Out of contemporary range and novel environmental conditions are projected for tropico-equatorial and austral regions. While some studies extrapolate important diversity and biomass changes in these zones^{28–30,40}, here we project shifts of their boundaries and maintenance of their climatic label. The present approach does not account for putative changes in community composition or the emergence of novel niches over these regions for which novel environmental selection pressure is expected. Despite these limitations, genomic data allow us to project compositional shifts due to the reorganization of the provinces at a high phylogenetic and functional resolution. Large (size class 180–2000 μm) and small (size class 20–180 μm) copepods are expected to be in greater relative abundance in subtropical and subpolar regions, respectively. Though the *Tara* Oceans environmental genomes collection might not reflect the full diversity in copepods, this projected restructuring of the communities in small and large copepods could modify carbon export as grazers' size is a key trait for this process⁴¹. Secondly, it might lead to novel prey-predator interactions e.g. in regions where new assemblages of communities are projected.

Important compositional shifts among most abundant marine nitrogen-fixing bacteria are also found. The relative share of nitrogen-fixing cyanobacteria is projected to increase in tropico-equatorial regions congruent with modeling studies⁴². Though their contribution to nitrogen fixation is not characterized, heterotrophic bacterial diazotrophs are abundant in genomic samples³³ and significant compositional shifts in certain clades are reported here. Ultimately, further studies associating omics and physico-chemical data should be informative for biogeochemical modeling of nitrate fixation rates and carbon export fluxes.

Overall, our projections for the end of the century do not take into account possible future changes of major bio-physico-chemical factors such as the dynamics of community mixing, trophic interactions through transport⁴⁶, the dynamics of the genomes^{13–15} (adaptation or acclimation) and biomass variations⁴⁰. New sampling in current and future expeditions⁴⁷, as well as ongoing technological improvements in bio-physico-chemical characterization of seawater samples^{32,47,48}, will soon refine functional^{16,49}, environmental (micronutrients⁵⁰) and phylogenetic³² characterization of plankton ecosystems for various biological entities (genotypes, species or communities)

and spatio-temporal scales⁴⁷. Ultimately, integrating this varied information will allow a better understanding of the conditions of emergence of ecological niches in the seascape and their response to a changing ocean.

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This article is contribution number XX of *Tara Oceans*.

Competing interests

The authors declare no competing interests.

Author contributions

PF, OJ and MG conceived the study. MV wrote the bias correction algorithm. PF computed the results, compiled and analyzed the data. PF wrote the initial draft of the paper. JL, OJ and MG conducted a preliminary study. PF, OJ, MG, MV, TD, DI, and PW discussed the results and contributed to write the paper.

Online content

All data and codes used are available at http://www.genoscope.cns.fr/tara/SourceCodes/NCLIM-20102618_codes.tar.gz

Materials and methods

Genomic provinces of plankton

Environmental niches are computed for trans-kingdom plankton genomic provinces from Richter et al.¹¹. They consist of the clustering of metagenomic dissimilarity matrices (based on the amount of DNA k-mers shared between pairs of samples) from 6 available size fractions with sufficient metagenomic data from the *Tara* Oceans dataset. The six size fractions (0-0.2, 0.22-3, 0.8-5, 5-20, 20-180 and 180-2000 μm) represent major plankton groups. Two large size classes (180-2000 μm and 20-180 μm) are enriched in zooplankton dominated by arthropods (mainly copepods) and cnidarians. They are expected to directly depend on smaller eukaryotes as they feed on them. Size classes 5-20 μm and 0.8-5 μm are enriched in smaller eukaryotic algae, such as dinophytes (5-20 μm), pelagophytes and haptophytes (0.8-5 μm). The distribution of these photoautotrophs presumably depends on nutrient availability. Finally, size classes 0.22-3 μm and 0-0.2 μm are respectively enriched in bacteria and viruses. Bacteria are characterized by a wide range of trophisms including autotrophy (cyanobacteria), mixotrophy and heterotrophy, while viruses are mainly parasites. Within each size fraction (from large to small), there are respectively 8, 8, 11, 6, 6 and 8 (48 in total) provinces defined in Richter et al.¹¹ formed by *Tara* Oceans metagenomes (644 metagenomes sampled either at the surface (SUR) or at the Deep Chlorophyll Maximum (DCM) across 102 sites). The clustering of individual size fractions is independent.

Genome signature of the provinces

We analyzed the distribution of 713 eukaryotic and 1888 prokaryotic genomes^{32,33} within the genomic provinces. These genomes are Metagenome-Assembled Genomes (MAGs) obtained from *Tara* Oceans metagenomes. For each size class, we select MAGs that are present (according to a criteria defined in Delmont et al.³²) in at least 5 samples. We computed an index of presence enrichment of MAGs within provinces as the Jaccard index⁵¹, defined as follows:

$$J = \frac{M_{11}}{M_{11} + M_{01} + M_{10}} \quad (1)$$

M_{11} is the number of samples where the MAG is present and match a sample of the province. M_{01} and M_{10} are respectively the number of samples where the MAG is not present in a sample of the province and inversely. A MAG is considered to be signature of a province if the Jaccard index is superior to 0.5 with this province and inferior to 0.1 for all other provinces of the given size class (Fig. 1 and Supplementary Fig. 3).

World Ocean Atlas data

Physicochemical parameters proposed to have an impact on plankton genomic provinces¹¹ are used to define environmental niches: sea surface temperature (SST), salinity (Sal); dissolved silica (Si), nitrate (NO_3), phosphate (PO_4), iron (Fe), and a seasonality index of nitrate (SI NO_3). With the exception of Fe and SI NO_3 , these parameters are extracted from the gridded World Ocean Atlas 2013 (WOA13)³⁴. Climatological Fe fields are provided by the biogeochemical model PISCES-v2⁵². The seasonality index of nitrate is defined as the range of nitrate concentration in one grid cell divided by the maximum range encountered in WOA13 at the *Tara* Oceans sampling stations. All parameters are co-located with the corresponding stations and extracted at the month corresponding to the *Tara* Oceans sampling. To compensate for missing physicochemical samples in the *Tara* Oceans *in situ* data set, climatological data (WOA) are preferred. The correlation between *in situ* samples and corresponding values extracted from WOA are high (r^2 : SST: 0.96, Sal: 0.83, Si: 0.97, NO_3 : 0.83, PO_4 : 0.89). In the absence of corresponding WOA data, a search is done within 2° around the sampling location and values found within this square are averaged.

Nutrients, such as NO_3 and PO_4 , display a strong collinearity when averaged over the global ocean (correlation of 0.95 in WOA13) which could complicate disentangling their respective contributions to niche definition. However, observations and experimental data allow identification of limiting nutrients at regional scale characterized by specific plankton communities⁵³. The projection of niches into future climate would yield spurious results when the present-day collinearity is not maintained⁵⁴ but there is up to now no evidence for large scale changes in global nutrient stoichiometry⁵⁵.

Earth System Models and bias correction

Outputs from 6 Earth System Models (ESM) (Supplementary Table 2) are used to project environmental niches under greenhouse gas concentration trajectory RCP8.5³⁷. Environmental drivers are extracted south of 60° north for present day (2006-2015) and end of century (2090-2099) conditions for each model and the multi-model mean is computed. A bias correction method, the Cumulative Distribution Function transform, CDFt⁵⁶, is applied to adjust the distributions of SST, Sal, Si, NO₃ and PO₄ of the multi-model mean to the WOA database. CDFt is based on a quantile mapping (QM) approach to reduce the bias between modeled and observed data, while accounting for climate change. Therefore, CDFt does not rely on the stationarity hypothesis and present and future distributions can be different. CDFt is applied on the global fields of the mean model simulations. By construction, CDFt preserves the ranks of the simulations to be corrected. Thus, the spatial structures of the model fields are preserved.

Environmental niche models: training, validation and projections

Provinces with similar metagenomic content are retrieved from Richter et al.¹¹. From a total of 48 initial provinces, 10 provinces are removed either because they are represented by too few samples (7 out of 10) or they are found in environments not resolved by ESMs (e.g. lagoons of Pacific Ocean islands, 3 out of 10). This narrows down the number of samples from 644 to 595 metagenomes. Four machine learning methods are applied to compute environmental niches for each of the 38 provinces: Gradient Boosting Machine (gbm)⁵⁷, Random Forest (rf)⁵⁸, fully connected Neural Networks (nn)⁵⁹ and Generalized Additive Models (gam)⁶⁰. Hyper parameters of each technique (except gam) are optimized. These are (1) for gbm, the interaction depth (1, 3 and 5), learning rate (0.01, 0.001) and the minimum number of observations in a tree node (1 to 10); (2) for rf, the number of trees (100 to 900 with step 200 and 1000 to 9000 with step 2000) and the number of parameters used for each tree (1 to 7); (3) for nn, the number of layers of the network (1 to 10) and the decay (1.10^{-4} to 9.10^{-4} and 1.10^{-5} to 9.10^{-5}). For gam the number of splines is set to 3, respectively 2 only when not enough points are available (for fraction 0-0.2, 65 points). R packages gbm (2.1.3),

randomForest (4.6.14), mgcv (1.8.16) and nnet (7.3.12) are used for gbm, rf, nn and gam models.

To define the best combination of hyper parameters for each model, we perform random cross-validation by training the model on 85% of the dataset randomly sampled and by calculating the Area Under the Curve⁶¹ (AUC) on the 15% remaining points of the dataset. This process is repeated over 30 random subsets of the entire training set for each combination of hyperparameters. We work in a presence/absence framework *i.e.* for each province separately the dataset consists of the variable to predict (“presence” or not of the province in the sample) and the predictors consisting of the environmental variables for each sample. A fixed probability threshold of 0.5 for presence/absence detection is used to calculate the AUC, *i.e.* samples with P, the probability of presence calculated by the model, superior to 0.5 are assigned to 1 and inversely samples with P inferior to 0.5 are assigned to 0. Fixing the probability threshold allows optimization of all models according to this threshold so that within a size fraction the *dominant* province has a reasonably high probability of presence (at least in regions with similar environmental parameters to the training dataset) and for the four types of models we use (gbm, nn, rf and gam). The best combination of hyper parameters is the one for which the mean AUC over the cross-validation is the highest. A model is considered valid if at least 3 out of the 4 techniques have a mean AUC superior to 0.65, which is the case for 27 out of the 38 provinces (Supplementary Fig. 2a). A climatic annotation is given to the 27 validated niches (Supplementary Table 2). Final models are trained on the full dataset and only the techniques that have a mean AUC higher than 0.65 are considered to make the projections. The vast majority (23) of the 27 validated niches is validated by all four models and 4 by only 3 models. Relative influences of each parameter in defining environmental niches are calculated using the feature_importance function from the DALEX R package⁶² for all four statistical methods (Supplementary Fig. 22a). To evaluate the consistency and coherence of environmental niche models, we first make global projections on the 2006-13 WOA2013 climatology. Projections are consistent with sampling regions for provinces encompassing vast oceanic areas. For example, the genomic province sampled in temperate Atlantic regions of size fraction 180-2000 μm is projected to be present in the north and south

temperate Atlantic but also other temperate regions (Supplementary Fig. 4). For model training and projections, physicochemical variables are scaled to have a mean of 0 and a variance of 1. For this scaling, the mean and standard deviation of each WOA13 variable (+ PISCES-v2 Fe) co-localized with *Tara* Oceans stations with a value available are used. This standardization procedure allows for better performance of nn models. Finally, as statistical models often disagree on projection sets whereas they give similar predictions on the training set (Supplementary Fig. 6, 7), we use the ensemble model approach for global-scale projections of provinces⁶³ *i.e.* the mean projections of the validated machine learning techniques.

Combined size class provinces and ocean partitioning comparisons

To combine all size classes' provinces, we use the PHATE algorithm^{36,64} from the R package phateR. This algorithm allows visualization of high dimensional data in the requested number of dimensions while best preserving the global data structure⁶⁴. We choose to train PHATE separately on WOA13 projections and present day and end of century projections including presence probabilities of *non dominant* provinces. We use 3 dimensions and set hyper parameter k-nearest neighbors (knn) and decay respectively to 1000 for WOA13 and 2000 for model data as in this case there are twice as many points. The hyper parameter knn reflects the degree to which the mapping of PHATE from high to low dimensionality should respect the global features of the data. We argue that 1000 and 2000 are good choices as it will be sufficient to have a highly connected graph, conserve global structure, allow visualization of structures of the size of the provinces (mean number of points in a province: 4867) and have a reasonable computational time. Decay is set to 20 in both cases. Then we cluster the resulting distance matrix using the k-medoids algorithm⁶⁵ and the silhouette average width criteria⁶⁶ is used as an indicator of good fit. The silhouette criterion is maximal for 2, 3 and 4 clusters and 2 peaks are found at 7 and 14 clusters (the peak at 7 is slightly less high than the one at 14, data not shown). We choose to present the 4 and 7 cluster geographical patterns as they seem more relevant with respect to the resolutions of our environmental datasets (WOA13 and climate models). We compare the three polar clusters of the 7 cluster geographical patterns with Antarctic Circumpolar Currents fronts⁶⁷ by overlying them on the map (black lines Supplementary Fig. 5b).

To visualize the global biogeography structure, the resulting 3 vectors of PHATE are plotted using an RGB color code. Each coordinate of each vector is respectively assigned to a given degree of color component between 0 and 255 (8 bits red, green or blue) using the following formula (Supplementary Figs. 5, 17):

$$C_{col}(i) = \frac{C(i) - \min(C(1), C(2), C(3))}{\max(C(1), C(2), C(3)) - \min(C(1), C(2), C(3))} * 255 \quad (2)$$

$C(i)$ is the i th component of the PHATE axes. Respectively, components 1, 2 and 3 are assigned to red, green and blue.

To compare the six size fraction provinces, the combined size class with existing biogeochemical partitions of the oceans^{9,10} and with each other, we use the adjusted rand index⁶⁸ (Supplementary Fig. 8-10) and overlay their masks above our partitions. In this case, presence probabilities of *dominant* provinces are not used anymore. Instead, each ocean grid point is assigned to the *dominant* provinces or the phate clusters.

Centroids and migration shifts

The centroid of each province is defined as the average latitude and longitude for which the probability of presence is superior to 0.5 and weighted by both the probability of presence at each grid point and the grid cell area. It is calculated for both present day conditions and end of the century conditions. The migration shift is calculated as the distance between the present day and the end of the century centroids considering the earth as a perfect sphere of radius 6371 km. For consistency (*i.e.* avoid long distance aberrant shifts), it is only calculated for provinces with an area of dominance larger than 10^6 km^2 in the given basin.

Bray-Curtis dissimilarity index

Climate change impact on global projections is calculated at each grid point as the Bray-Curtis dissimilarity index^{69,70} defined as follows:

$$BC = \frac{\sum_n |P_n^{future} - P_n^{present}|}{\sum_n |P_n^{future} + P_n^{present}|} \quad (3)$$

Where ($P_n^{present}$ and P_n^{future}) are respectively the probability of presence of the province n in present day and at the end of the century. Only the probabilities of *dominant* provinces are non-null and all others are set to zero. The mask of main fisheries⁷¹

(chosen as the first 4 deciles) and Exclusive Economical Zones⁷² is overlaid on the Bray-Curtis map.

Change in province assemblages

A province assemblage is defined as the assemblage of *dominant* provinces of each size fraction at a given grid point of the considered ocean. We consider two criteria of change in province assemblage between present day and end of the century conditions. The first one, more straightforward and less stringent, considers that a province assemblage occurs when a change of *dominant* province is found in at least one size fraction. In a more stringent way, a change of assemblage is considered significant for $BC > \frac{1}{6}$ (previous section). This threshold corresponds to an idealized case where each *dominant* province has a probability of one and a change of *dominant* province is found in only one size fraction. For example, the *dominant* province assemblage goes from vector (F5,E6,D3,C8,B7,A7) (with the size fractions in decreasing order) corresponding to all temperate provinces to vector (F8,E6,D3,C8,B7,A7). This example corresponds to the replacement of the temperate province of size fraction 180-2000 μm (F5) by the tropico-equatorial province (F8). This criterion allows us to discard assemblage changes for which the changes in probability of presence of *dominant* provinces are very low. With this criterion, only a small oceanic area is found to have no changes of assemblage (Fig. 4c light blue zones).

Composition of provinces in bacterial diazotrophs and marine copepods

We characterize the composition in Marine Hexanauplia (copepods) and marine diazotrophs of provinces by considering the mean relative abundances of groups of MAGs^{32,33} characterized taxonomically (for both copepods and diazotrophs) and by size (for copepods) in each province for all size fractions except the viral one (for diazotrophs) and for size fractions $>5 \mu\text{m}$ (for Marine Hexanauplia). Marine Hexanauplia are annotated taxonomically (either as belonging to clade A (109 MAGs) or clade B (105 MAGs)) from which 198 are found in at least 5 samples that we use (98 clade A and 100 clade B). To attribute a preferential size class to these MAGs, mean relative abundances over all sites of each of them are compared across size fraction 180-2000, 20-180 and 5-20 μm using Welch ANOVA⁷³ (p-value<0.05). When the Welch ANOVA

test is significant the MAG is either annotated as mesozooplankton (when most abundant in size class 180-2000 μm) or microzooplankton (when most abundant in size class 20-180 or 5-20 μm). When the Welch ANOVA test is not significant, the MAG is annotated as unclassified. Then for each group of MAG (defined by the preferential size class plus the clade), the mean of the sum over the MAGs from this group is calculated in each province to characterize the province. The same procedure is applied for 27 out of 48 bacterial diazotrophs (present in at least 5 samples) from the prokaryotic MAG collection³³ distinguishing groups at the phylum level (Gammaproteobacteria $n=8$, Cyanobacteria $n=8$, Deltaproteobacteria $n=2$, Alphaproteobacteria $n=4$, Planctomycetes $n=3$, Verrucomicrobiota $n=2$). Finally, significant differential composition between provinces are annotated using Holm corrected⁷⁴ pairwise Mann-Whitney U test⁷⁵ ($p<0.05$ for significance) comparing the abundance distributions of each group of either diazotrophs or copepods between each pair of provinces from a same size class.

Driver analysis

To assess the relative importance of each driver in province changes, the methodology from Barton et al.²⁴ is adopted. For a set N of n provinces (individual provinces or all provinces together), the probability of presence of each province is recalculated for present day conditions except for driver d (from the set of drivers D) for which the end of the century condition is used ($P_n^{\text{future for } d\text{th driver only}}$). The set of driver D can be either all drivers (Fig. 5a,c) or all drivers except SST (Fig. 5b). The relative importance of driver d at a given grid point for the set of N of provinces is computed as follows:

$$RI(d) = \frac{\sum_{n \in N} |P_n^{\text{future for } d\text{th driver only}} - P_n^{\text{present}}|}{\sum_{d \in D} \sum_{n \in N} |P_n^{\text{future for } d\text{th driver only}} - P_n^{\text{present}}|} \quad (4)$$

$RI(d)$ is computed at grid cells where $BC > \frac{1}{6}$ and calculated with either the set of all drivers (Fig. 5a,c) or all drivers except SST (Fig. 5b). When $RI(d)$ is calculated for individual provinces (Fig. 5c and Supplementary Fig. 24d,e), it is computed only at grid cells where $BC > \frac{1}{6}$ and the concerned province is either *dominant* in present day and/or end of century conditions.

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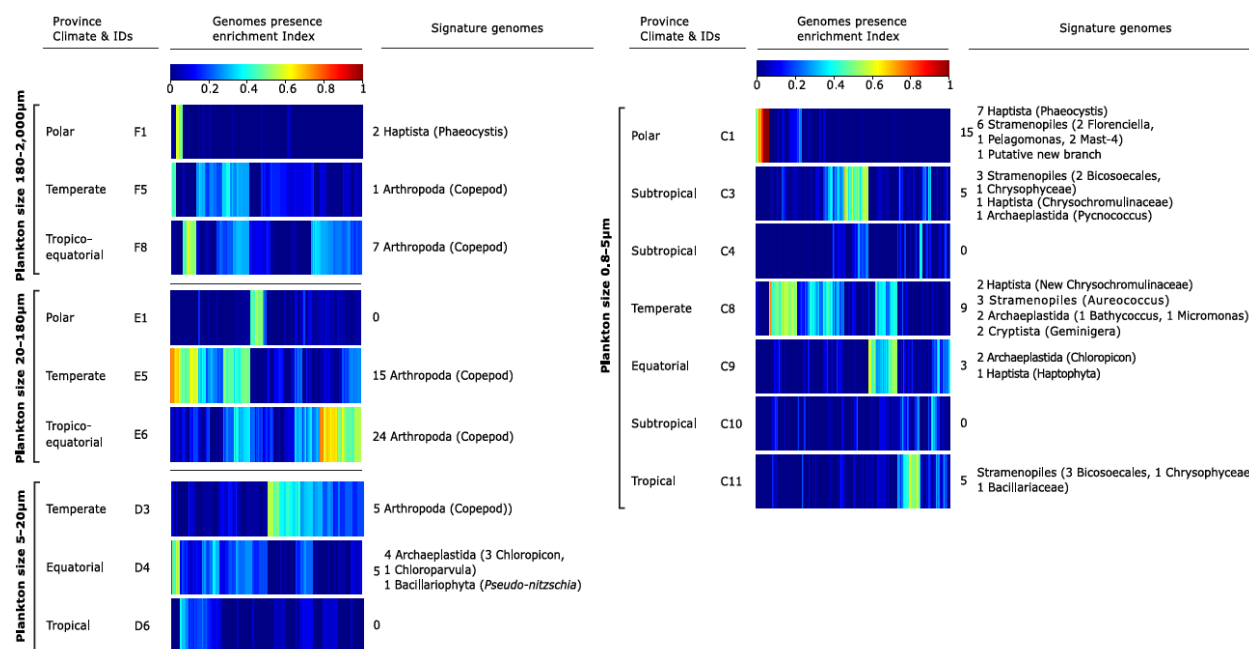


Fig. 1 | Eukaryotic signature genomes of provinces of eukaryote enriched size classes.

For each plankton size class, indexes of presence enrichment (equation (1)) for 713 genomes of eukaryotic plankton³⁸ in corresponding provinces are clustered and represented in a color scale. Signature genomes (see *Methods*) are found for almost all provinces, their number and taxonomies are summarized (detailed list in Supplementary Table 6).

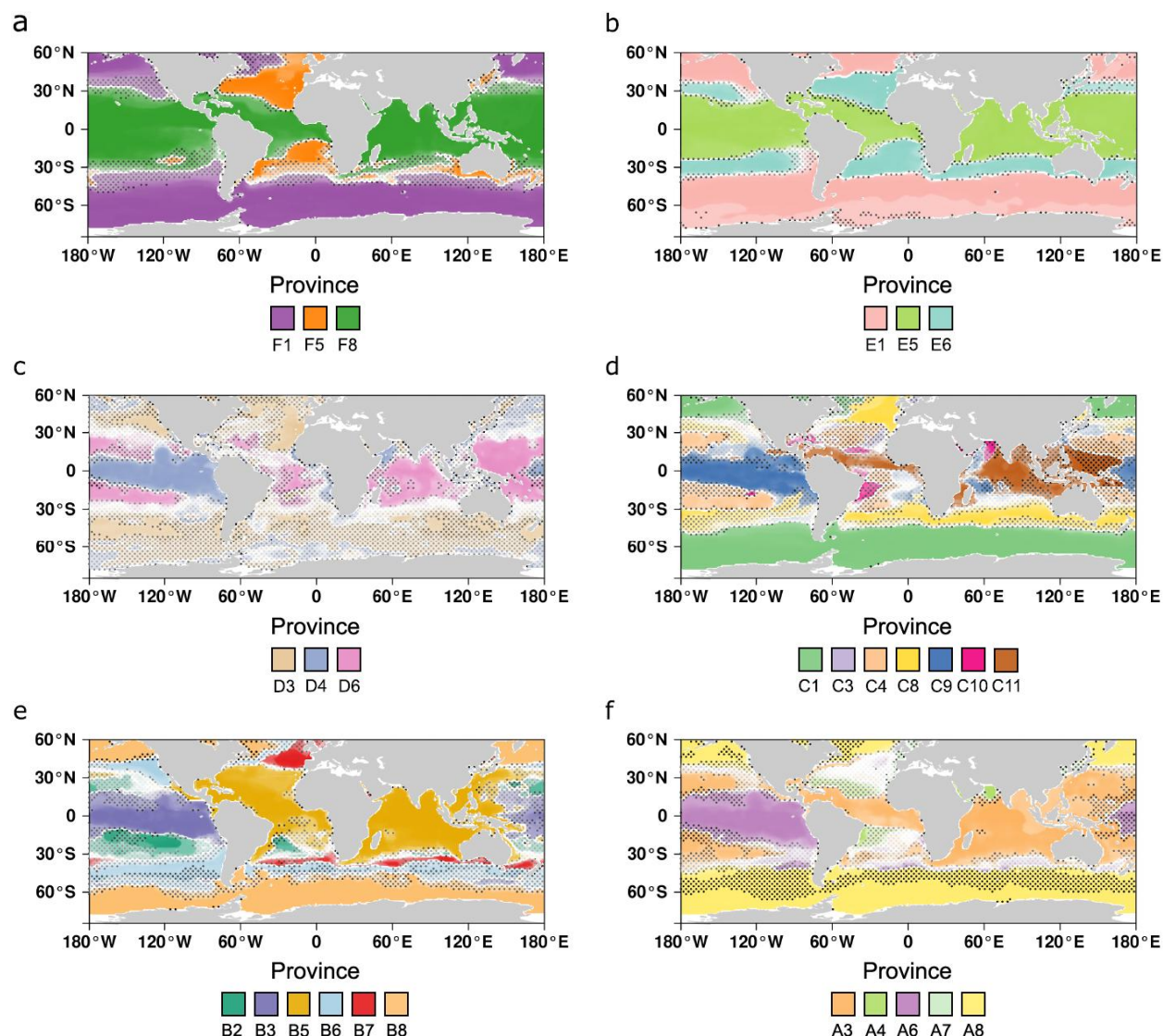


Fig. 2 | Global biogeographies of size-structured plankton provinces projected on WOA2013 dataset. (a) Metazoans enriched (180-2000 μm) (b) Small metazoans enriched (20-180 μm) (c) protist enriched (5-20 μm) (d) protist enriched (0.8-5 μm) (e) Bacteria enriched (0.22-3 μm) (f) Viruses enriched (0-0.2 μm). (a-f) Dotted areas represent uncertainty areas where the delta of presence probability of the dominant province and an other (from the same size fraction) is inferior to 0.5. Simple biogeographies are observed in large size fractions (>20 μm) with a partitioning in three major oceanic areas: tropico-equatorial, temperate and polar. More complex geographic patterns and patchiness are observed in smaller size fractions with the distinction of pacific equatorial provinces and provinces associated with oligotrophic tropical gyres

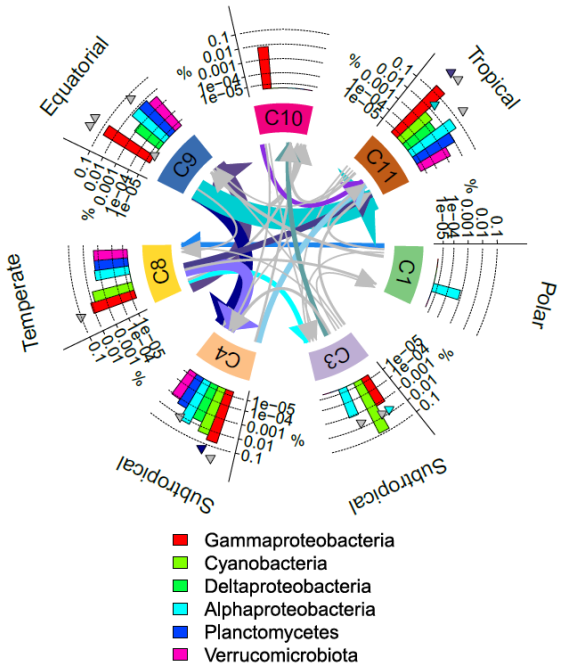
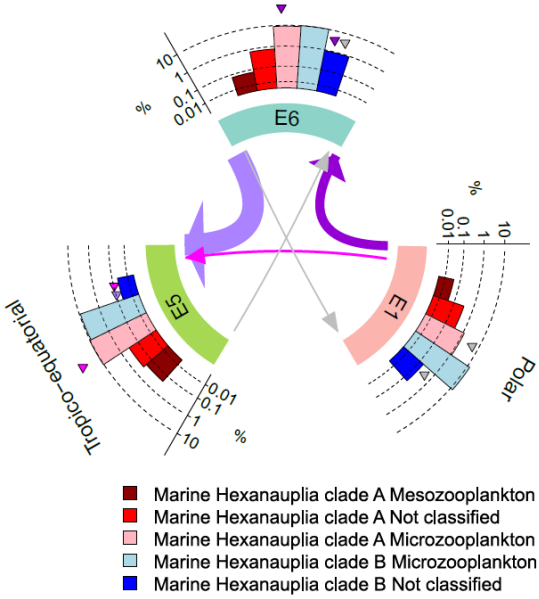
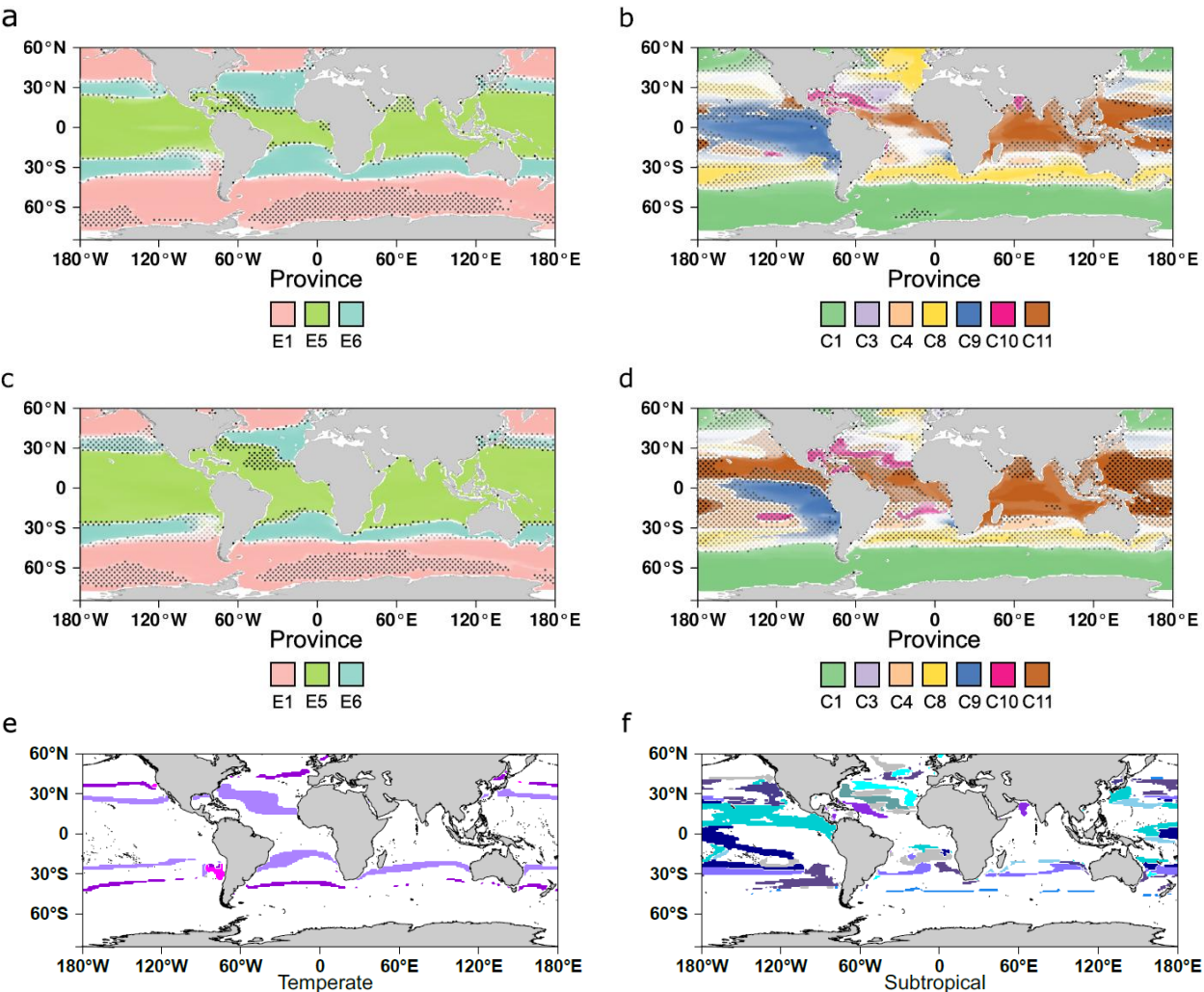


Fig. 3 | Global biogeographies of the small metazoan enriched size fraction (20-180 μm), the protist enriched size fraction (0.8-5 μm) in (a, b) modeled present day and (c, d) end of the century. Compositional shifts between provinces in (e) marine hexanauplia (180-2000 μm) and (f) diazotrophs (0.8-5 μm). (a-d) The dominant niche *i.e.* the one predicted to have the highest probability of presence is represented at each grid point of the map. The color transparency is the probability of presence of the dominant niche. A simple biogeography is observed for size fraction 180-2000 μm (a, b) with a polar niche, a temperate and a tropico-equatorial niche. Biogeography of size fraction 0.8-5 μm is more complex and patchy with several temperate and tropical niches (c, d). Biogeography of large size plankton and small size plankton are therefore decoupled. Climate change impacts tropical niches which are expanding towards the poles in both size fractions with a more complex behavior in size fraction 0.8-5 μm (b and d). Changes in community composition might also happen in areas with out of range projected conditions (*e.g.* equator, poles, Supplementary Fig. 12), but here the contemporary province is still the most probable and the climatic label is maintained. (e-f) Top: Maps highlighting areas of dominant community shifts in size fraction 20-180 μm (e) and 0.8-5 μm (f) Bottom: circular plots summarizing significant compositional shifts either in Marine Hexanauplia (copepods) (f) or bacterial diazotrophs (e): each type of transition is colored differently and accordingly to the one on the map or in grey if they represent less than 2% of the transitions. Barplots represent mean relative abundances in 5 types of Marine Hexanauplia (copepods) (f) or main clades of marine diazotrophs (e) based on genome abundance in the given province. Arrows represent dominant community shifts pointing towards the end of the century projected province and their widths are proportional to the area of change. Significant compositional changes in a type of genome (*e.g.* cyanobacteria) are represented by triangles of the associated transition color above the corresponding bar (*e.g.* yellowgreen for cyanobacteria) of the barplot of the province projected at the end of the century (which is at the tip of the arrow).

809

Size fraction (μm)	Present day covered area (%)	End of century covered area (%)	% area with a change of dominant province	Most frequent transition	%	2 nd most frequent transition	%
180-2000	74	74	13	temperate->tropico-equatorial (F5->F8)	67	polar->temperate (F1->F5)	14
20-180	78	77	12	temperate->tropico-equatorial (E6->E5)	67	polar->temperate (E1->E6)	29
5-20	45	49	22	temperate -> equatorial (D3->D4)	47	equatorial -> tropical (D4->D6)	25
0.8-5	56	59	31	equatorial -> tropical (C9->C11)	22	temperate -> equatorial (C8->C9)	15
0.22-3	60	61	15	temperate-> tropical (B7->B5)	22	polar -> temperate (B8->B6)	16
0-0.2	64	66	16	equatorial -> tropical (A6->A3)	32	temperate -> equatorial (A8->A6)	31
Total			60				

810 **Table 1 | Global statistics of covered areas and province changes and transitions.** From
811 12% to 31% of the total covered area is estimated to be replaced by a different province at the
812 end of the century compared to present day depending on the size fraction. In total, considering
813 all size fractions this represents 60% of the total covered area with at least one predicted
814 change of dominant province across the six size fractions.

815

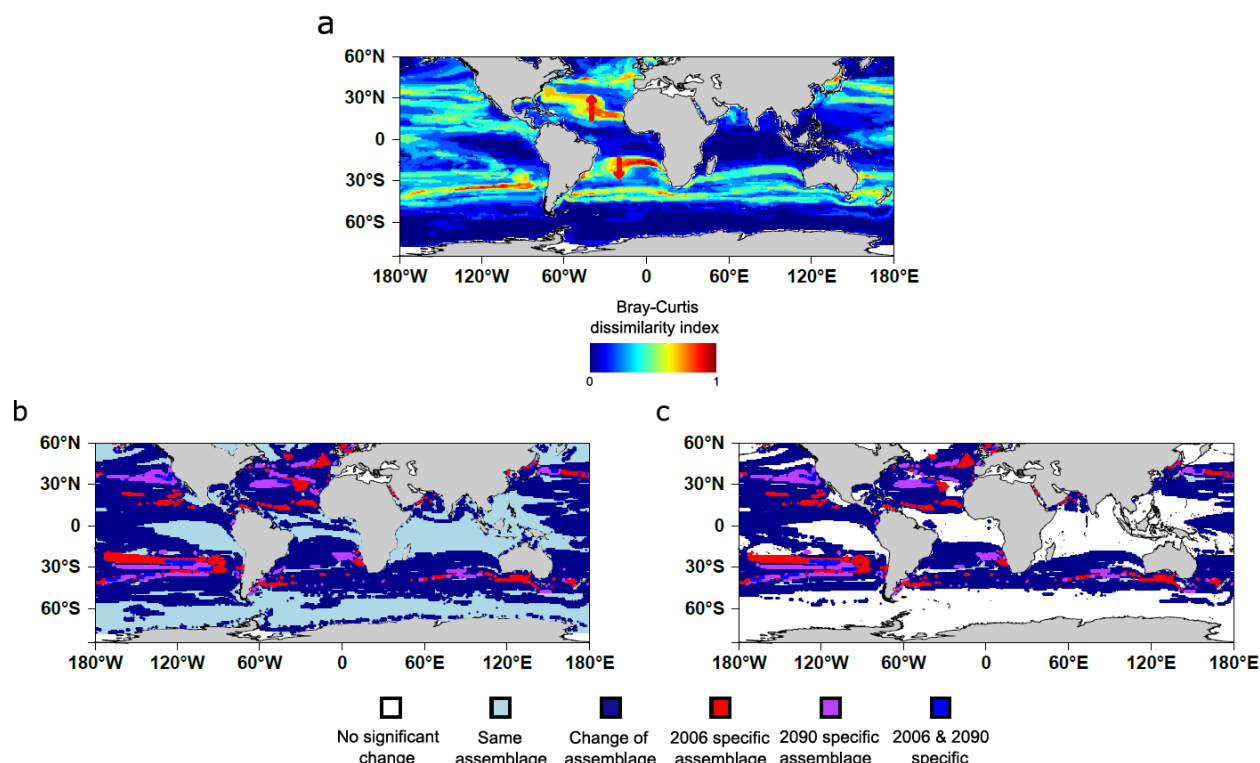


Fig. 4 | (a) Bray-Curtis dissimilarity index map comparing present day with end of the century projections of dominant provinces. Maps of trans-kingdom assemblage reorganization of dominant provinces (b) and with a criterion of significance (c). (a) Bray-Curtis dissimilarity index (equation (3)) is calculated by integrating all the dominant provinces presence probabilities over the six size fraction. Most important changes appear in subtropical, temperate and subpolar regions. These changes are due to the displacement of tropical and temperate provinces towards the pole but also the geographical decoupling between large and small size plankton. The mean change in niche dissimilarity index is 0.25. **(b)** An assemblage is the combined projected presence of the dominant province of each size class. Assemblage reorganization (present day versus end of the century) is either mapped on all considered oceans or with a criterion on the Bray-Curtis dissimilarity index ($BC > 1/6$, see *Materials and Methods*) **(c)**. Depending on the criterion from 60.1% **(b, dark blue)** to 48.7% **(c, dark blue)** of the oceanic area is projected to change of assemblage. New assemblages are expected to appear in 2090 (purple+blue) whereas some 2006 specific assemblages are projected to disappear (red+blue). New assemblages as well as lost assemblages are mostly found in temperate, subtropical and tropical regions where most of the rearrangements are projected.

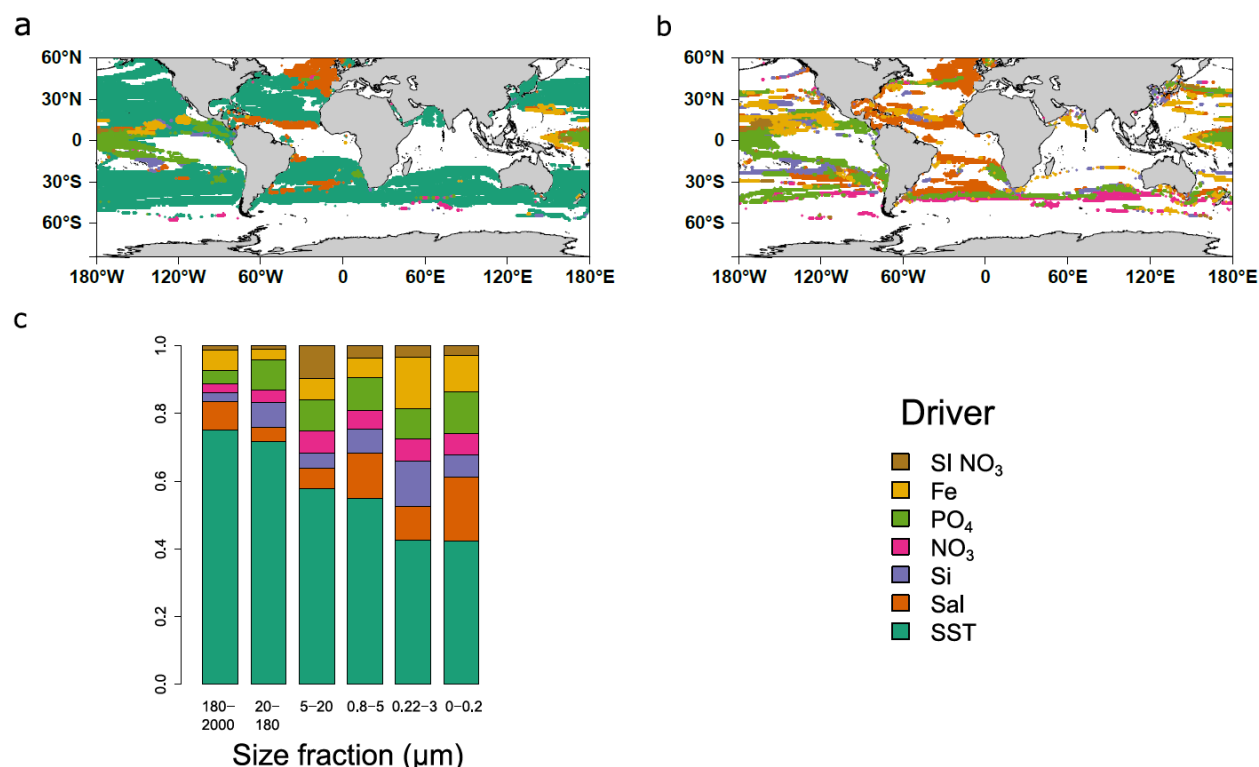


Fig. 5 | Map of most impacting drivers on dominant province changes (a), most impacting driver without considering temperature change (b) and relative importance of the drivers in the different size fractions (c). (a) Temperature appears as the top impacting driver on the majority of the projected ocean with a significant change of province (Fig. 4). **(b)** Salinity and dissolved phosphate are found to be the second and third drivers of province reorganization notably at tropical and subpolar latitudes. Note the importance of nitrate at temperate southern latitudes. **(c)** Temperature is found to be the most important driver for all size classes but has a more important impact in large size classes (>20 μm). Nutrients have on average a relatively more important impact in small size classes in driving province reorganization.

Supplementary information 1 | Niche models of genomic provinces

To compute and test the validity of realized environmental niches, we train four machine learning techniques to probabilistically associate genomic provinces with environmental data: sea surface temperature, salinity, three macronutrients (dissolved silica, nitrate and phosphate), one micronutrient (dissolved iron) plus a seasonality index of nitrate. The four machine learning techniques are Gradient Boosting Machine (gbm)¹, Random Forest (rf)², fully connected Neural Networks (nn)³ and Generalized Additive Models (gam)⁴. Following a cross-validation framework, a valid environmental niche is obtained for 27 out of 38 initial provinces (71%) comforting their definition and covering 529 samples out of 595 (89%, Supplementary Fig. 2). Rejected provinces contain relatively few stations (mean of 6 ± 2.6 versus 19 ± 15.3 for valid provinces, $p\text{-value} < 10^{-3}$ Wilcoxon test⁵). For spatial and temporal extrapolations of the provinces presented below, we use the ensemble model approach⁶ that considers mean predictions of machine learning techniques.

Supplementary information 2 | Integrated biogeographies using the PHATE algorithm

We use the PHATE dimension reduction algorithm⁷ to combine all provinces for all size classes into a single consensus biogeography revealing 4 or 7 robust clusters (Supplementary Fig. 5, *methods*). The 4 cluster consensus biogeography is mainly latitudinally organized distinguishing polar, subpolar, temperate and tropico-equatorial regions. The 7 cluster consensus biogeography distinguishes the equatorial pacific upwelling biome and three subpolar biomes that most likely reflect the chemico-physical structuring of the Southern Ocean and known polar fronts⁸ (red lines Fig. 2h). However, learning data are scarcer south of 60°S so these extrapolations need to be taken with caution.

PHATE is also used for modeled projections into two comparable consensus maps, one for present day and one for the end of the century (Supplementary Fig. 17). Some particularly visible patterns of geographical reorganization are common to several or even all size fractions and visible when comparing the two consensus maps (Supplementary Fig. 17 compared to Fig. 3a-d and Supplementary Fig. 14). For

example, the tropico-equatorial and tropical provinces expand in all size fractions and the provinces including the pacific equatorial upwelling shrink for size fractions smaller than 20 μm .

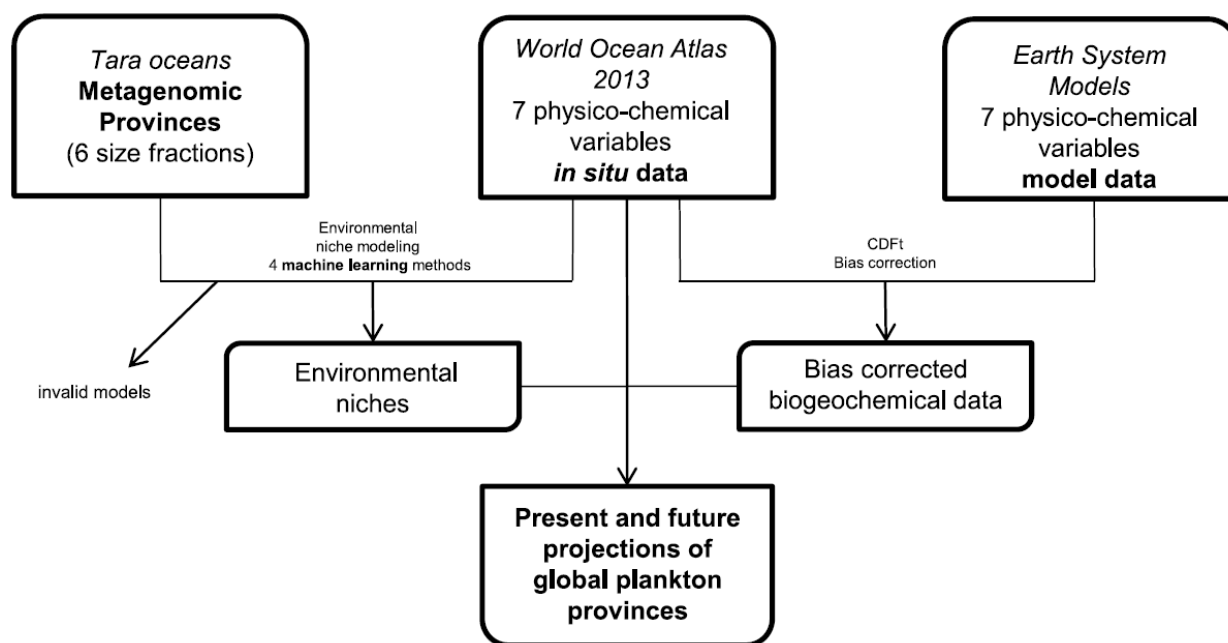
Supplementary information 3 | Comparison of the biogeographies with existing partitions of the oceans

Previous ocean partitioning either in biomes⁹⁻¹¹ or biogeochemical provinces (BGCPs)^{9,10} are based on physico-biogeochemical characteristics including SST⁹⁻¹¹, chlorophyll *a*⁹⁻¹¹, salinity⁹⁻¹¹, bathymetry^{9,10}, mixed layer depth¹¹ or ice fraction¹¹. Considering three of these partitions as examples we notice differences with our partitions (Supplementary Fig. 8-9) for example in terms of the number of regions in the considered oceans (56 for 2013 Reygondeau et al. BGCPs¹⁰, 17 for Fay and McKingley¹¹) and their structure (the coastal biome for 2013 Reygondeau et al. biomes¹⁰). Numerical comparison of our partitions with others (*Methods*) reveals low similarity between them, the highest being with Reygondeau biomes (Supplementary Figs. 8-10). However, biomes and BGCP frontiers closely match our province frontiers in many cases. Near the frontiers, *dominant* provinces have smaller probabilities in agreement with smooth transitions instead of sharp boundaries as already proposed⁹ and with a seasonal variability of the frontiers¹⁰ (Supplementary Fig. 8). Some of these transitions are very large and match entire BGCPs, for example in subtropical North Atlantic and subpolar areas where high annual variations are well known¹⁰.

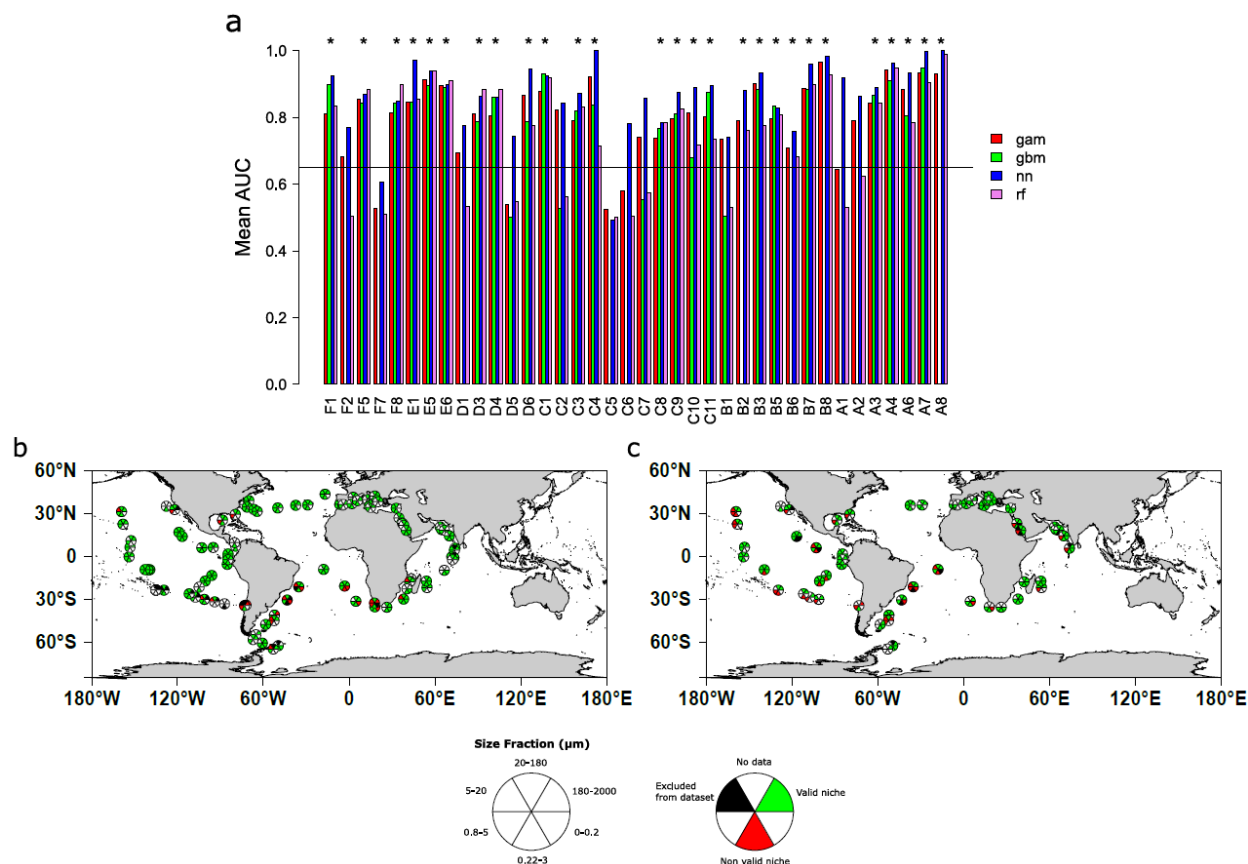
Supplementary information 4 | Expansion and shrinkage of provinces in response to climate change

To quantify patterns of expansion or shrinkage of the provinces, we calculate the surface covered by the *dominant* provinces weighted by probabilities of presence (Supplementary Fig. 18, Supplementary Table 2). In this way, *dominant* provinces are defined on 100% of the surface ocean (327 million km^2) but their presence probabilities correspond to the equivalent of 45 to 74% (due to sampling variability and niche overlaps) of the surface ocean depending on the plankton size fraction (Table 1). Overall, our results indicate expansions of the surface of tropical and tropico-equatorial

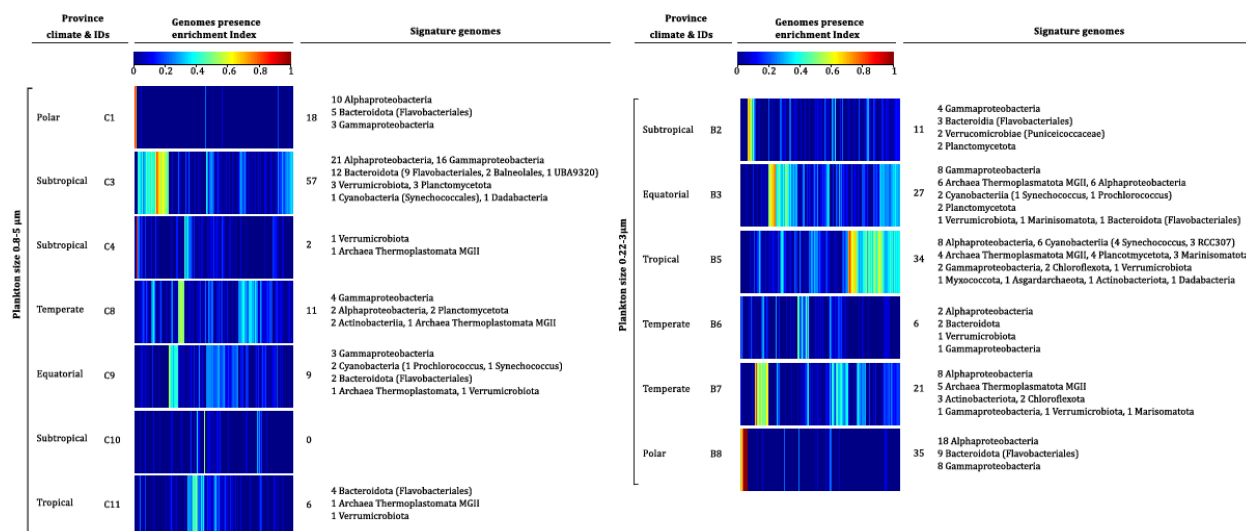
905 provinces but in very different ways depending on the size fractions of organisms. The
 906 surface area of temperate provinces is ~22 million km² on average (from 10 Mkm² for
 907 0-0.2 µm to 49 Mkm² for 20-180 µm) and should decrease by 36% on average (from -20
 908 % for 5-20 µm up to -54% for 0.8-5 µm, -12 million km² on average, +6% for 0.22-3
 909 µm). Tropical provinces cover ~118 million km² on average (from 86 Mkm² for 0.8-5 µm
 910 up to 169 Mkm² for 180-2000 µm) and their coverage should increase by 32% on
 911 average (from +13% for 0-0.2 µm up to +75% for 0.8-5 µm, +25 million km² on average)
 912 (Supplementary Fig. 18 and Supplementary Table 2).



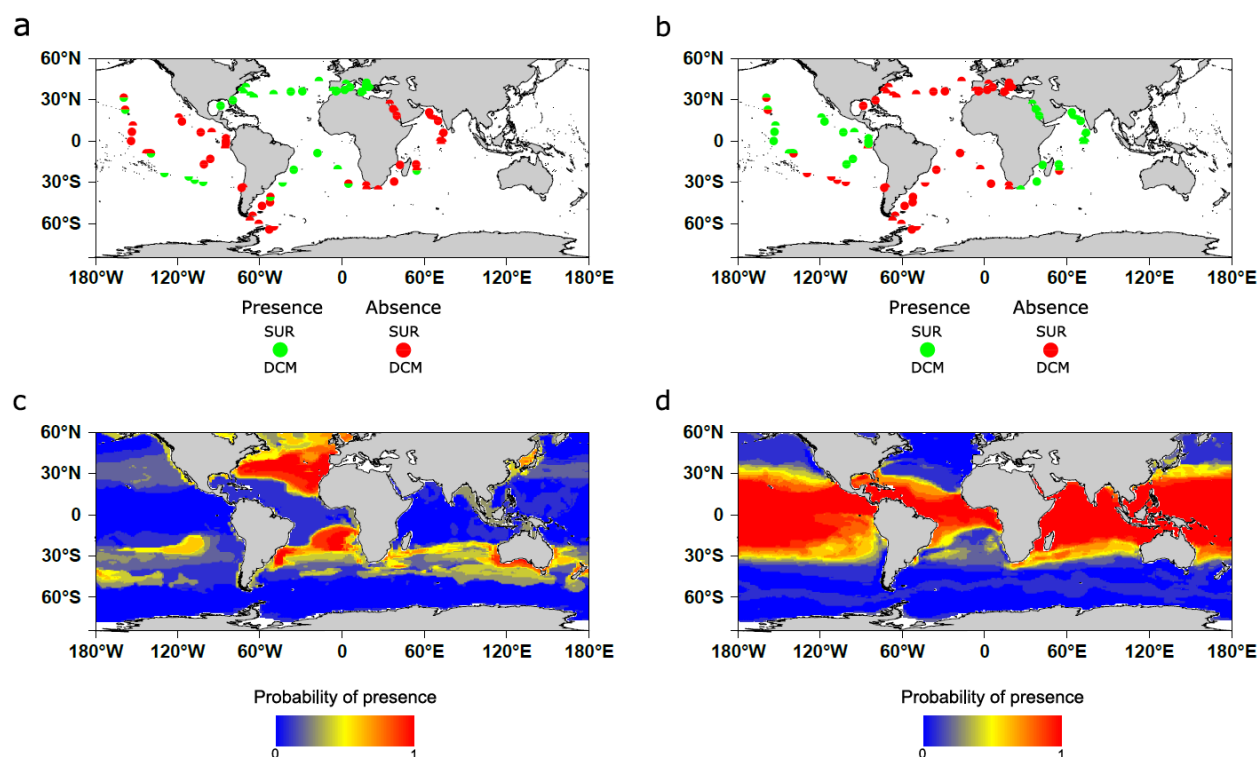
Supplementary Fig. 1 | Study pipeline. Metagenomic data from the 2009-2013 *Tara Oceans* expedition and *in situ* measurements of physicochemical variables (*World Ocean Atlas 2013*, WOA13)¹² are combined to define environmental niches at the plankton community level across 6 size fractions of the plankton realm. Bias corrected outputs from a mean model of 6 Earth System Models^{13–18} and WOA13 data are then used to project global plankton provinces for present day conditions and end of the century conditions under a high warming scenario (RCP8.5)¹⁹. Physico-chemical variables are Sea Surface Temperature (SST), Salinity (Sal), Dissolved silica (Si), Nitrate (NO₃), Phosphate (PO₄), Iron (Fe) and a seasonality index of nitrate (SI NO₃).

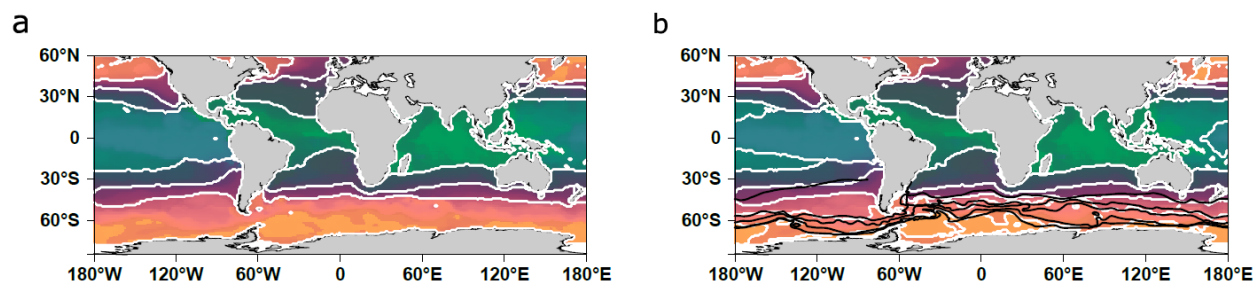


Supplementary Fig. 2 | (a) Barplot of mean AUC over a random 30-fold cross validation process of the 38 initial metagenomic provinces. (b,c) Map of validated and non validated stations across the six size fractions in surface samples (b) and DCM samples (c). (a) Mean AUC (Area Under the receiver operating Characteristic)²⁰ is plotted for the best hyperparameter combination of the 4 machine learning techniques used for each of the 38 metagenomic provinces. General Additive Models⁴ (gam) are shown in red (no optimization), Gradient Boosting Machines¹ (gbm) in green, fully connected Neural Networks³ (nn) in blue and Random Forest² models in purple (rf). A star for valid models is drawn at the top of each considered valid model. A model is validated when at least 3 out of the four models have a mean AUC superior to 0.65. A valid model is found for 27 out of 38 initial provinces. Out of the 27 validated models 4 are not valid for the gbm method. (b-c) For each Tara sample present in the dataset at surface (b) or Deep Chlorophyll Maximum (DCM) (c) and for each size fraction, the filter (one size fraction at one location and one depth) belongs either to a validated niche (green), a non validated niche (red), an excluded filter (see *Materials & Methods*) (black) or no data is available (white). Non validated niches represent only 11% of filters present in the dataset (66 out of 595).

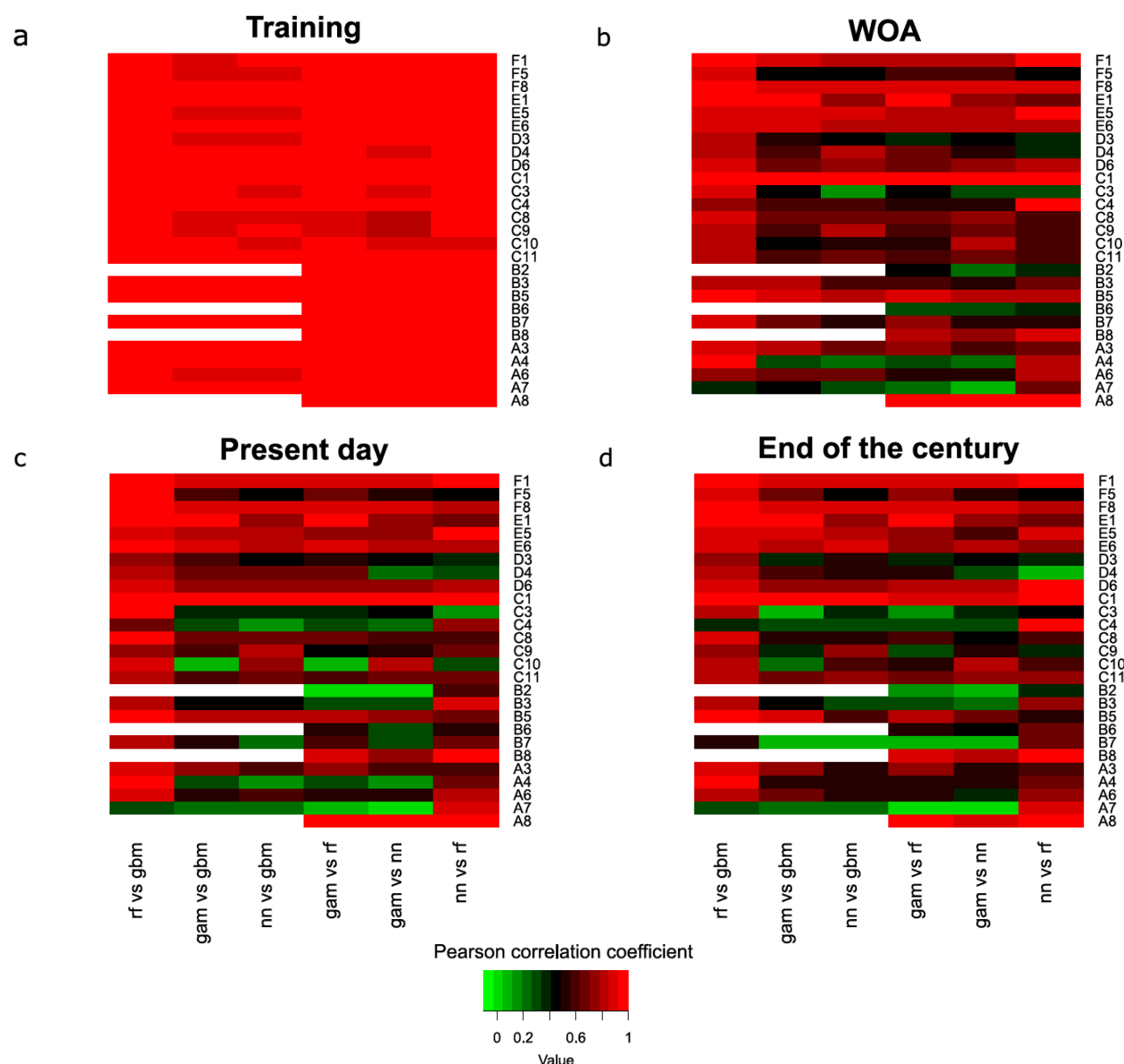


Supplementary Fig. 3 | Prokaryotic signature genomes of provinces of the prokaryote (0.22-3 μm) and protist (0.8-5 μm) enriched size classes. Indexes of presence enrichment²¹ for 1888 genomes of prokaryotic plankton²² in corresponding provinces are clustered and represented in a color scale. Signature genomes (see *Methods*) are found for almost all provinces, their number and taxonomies are summarized (detailed list in Supplementary Table 6).



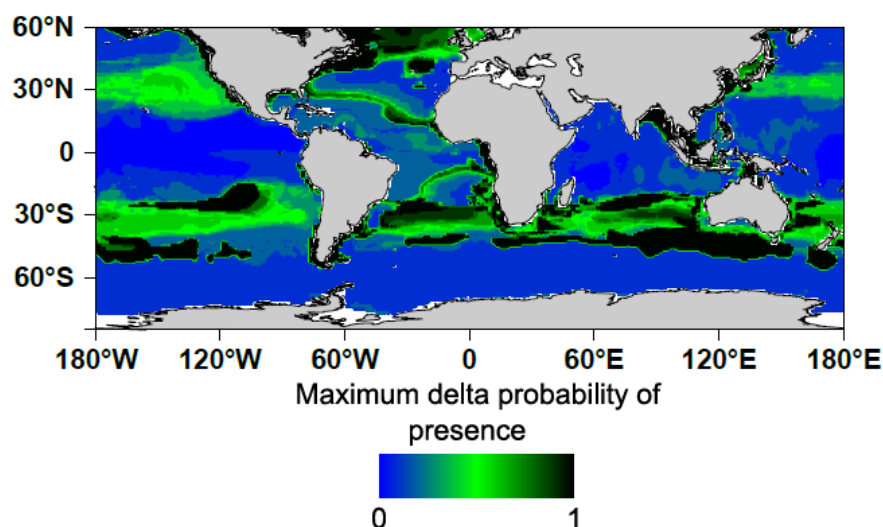


Supplementary Fig. 5 | Present day combined size class biogeographies using the PHATE algorithm on WOA13 dataset. (a) Combined size class biogeography using PHATE algorithm⁷ partitioned in 4 clusters. Each grid point is associated to a color depending on its PHATE coordinates (using three axes). Each coordinate is respectively assigned to a given degree of color between 0 and 255 according to equation 2 (either red green or blue depending on the axis). **(b)** Combined size class biogeography using PHATE algorithm partitioned in 7 clusters overlaid with Antarctic Circumpolar Current boundaries (black). Colors are the same as in **(a)**.

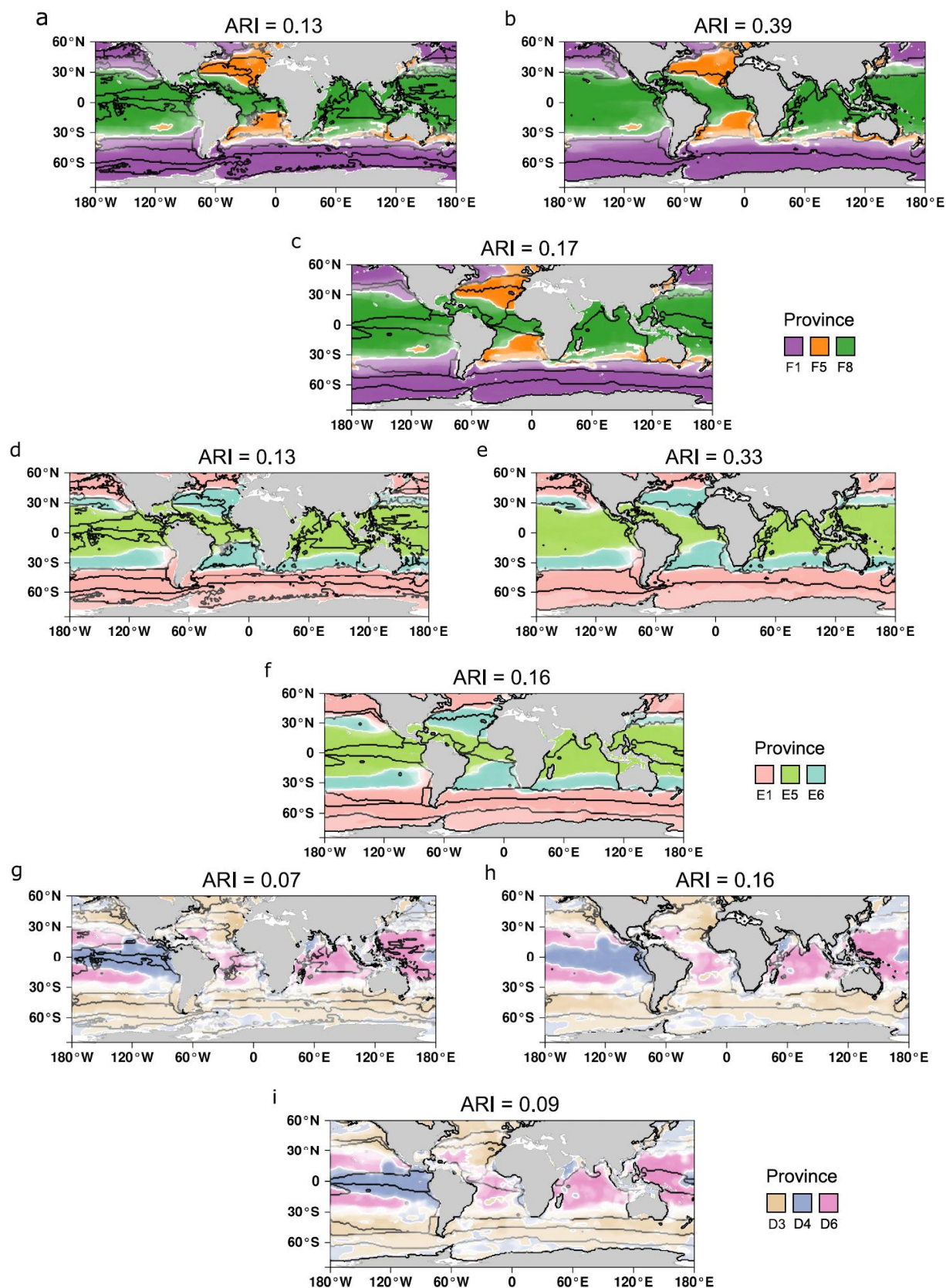


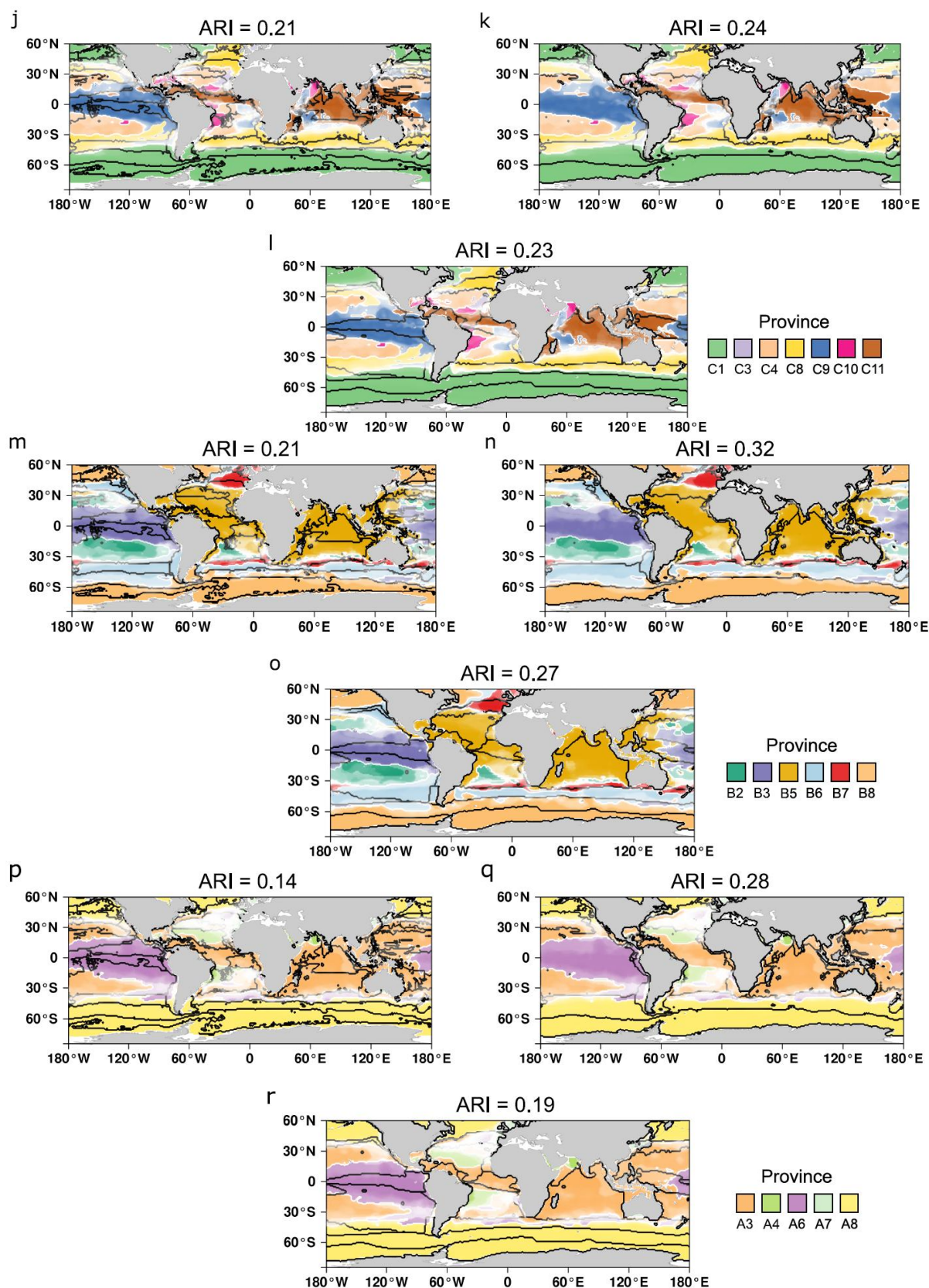
Supplementary Fig. 6 | Pairwise Pearson correlation coefficient heat maps between outputs of the 4 machine learning models. rf: Random Forest, gbm: gradient boosting machines, gam: general additive models, nn: neural networks. The rows are the provinces of the different size fractions. The columns are the pairwise comparisons of each machine learning technique. (a) Training set outputs. (b) WOA13 average data projections outputs (except for Iron, PISCES-v2²³ is used). (c) Present day data projection outputs (bias corrected mean model of 6 Earth System Models). (d) End of the century projection outputs. On the training set, models are in agreement with most of the correlation coefficients superior to 0.9. A drop in correlation is observed for modeled data (c, d) especially in small size fractions. This shows modeled data are more distant from the training set than WOA data. Random Forest and

975 Gradient Boosting Machine are in very good agreement (first columns) which could be expected
976 as they are both based on multiple decision trees.
977



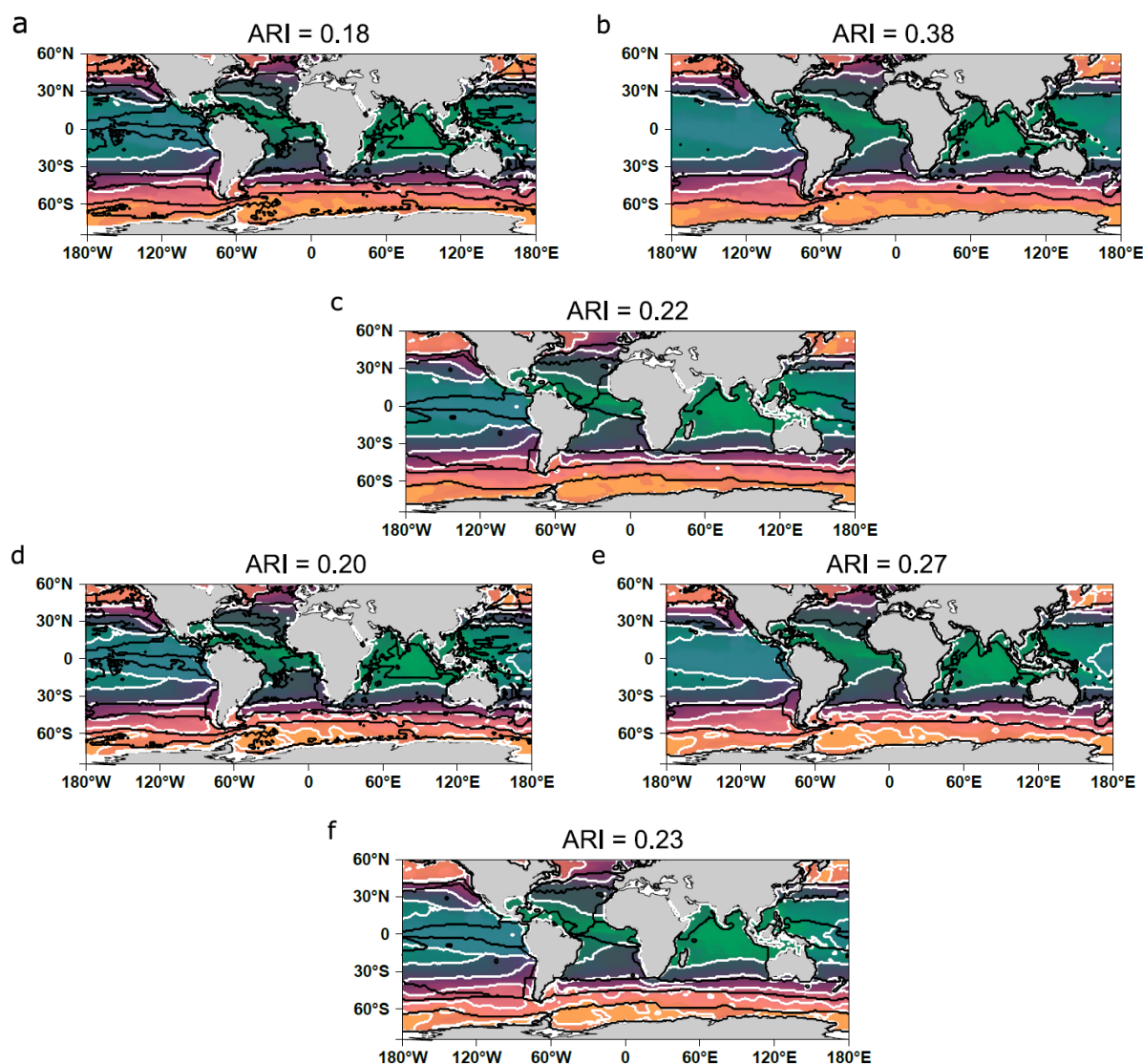
Supplementary Fig. 7 | Probability range map (WOA data) of province F5 of size fraction 180-2000 μm . At each point of the grid, the maximum delta probability of presence between the 4 machine learning projections is calculated. In black are the zones where two models completely disagree: one model predicts presence with certainty whereas the other predicts absence with certainty. Disagreement appears mostly in uncertain presence areas (Supplementary Fig. 4c) whereas models are generally in good agreement in absence areas (blue zones).



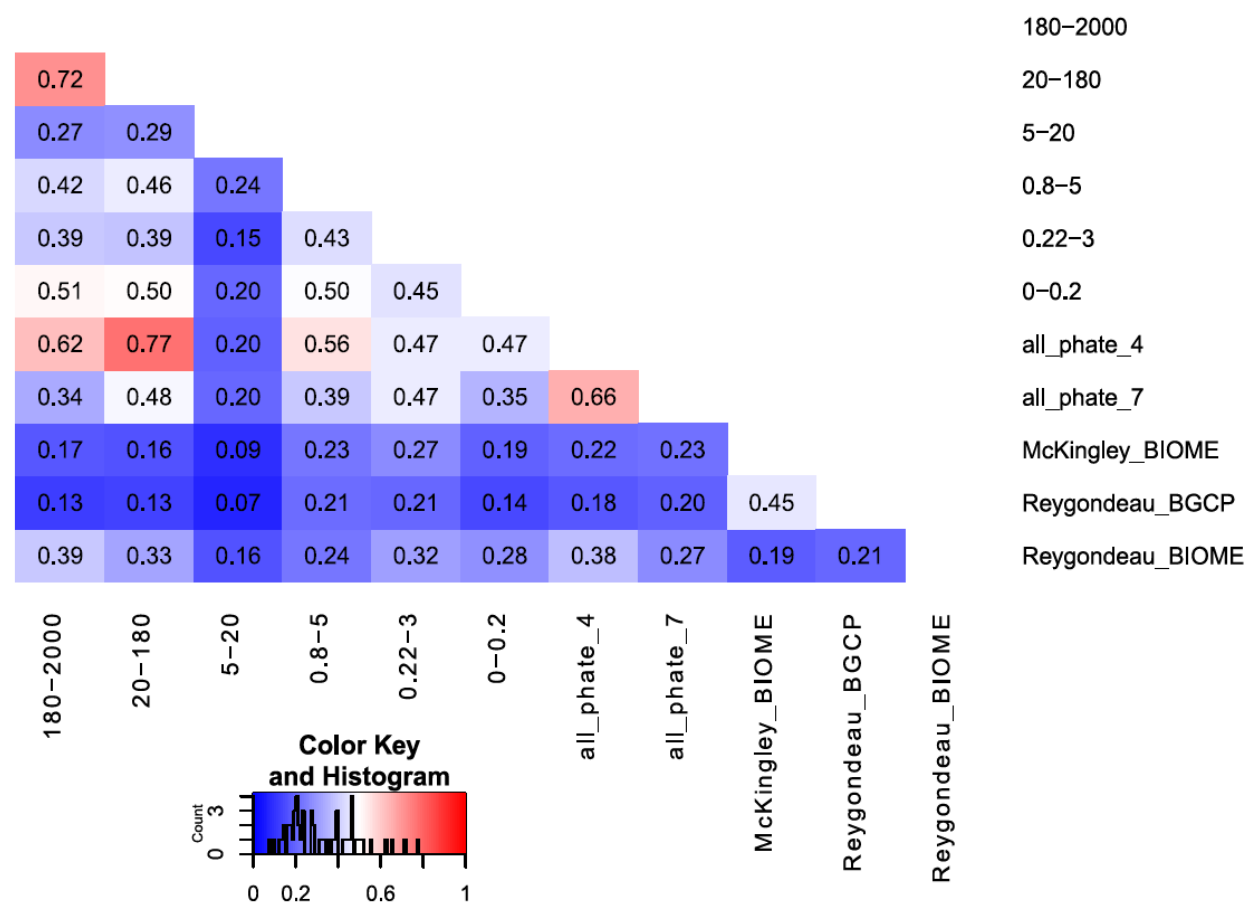


Supplementary Fig. 8 | Three existing oceanic partitions overlaid on top of plankton

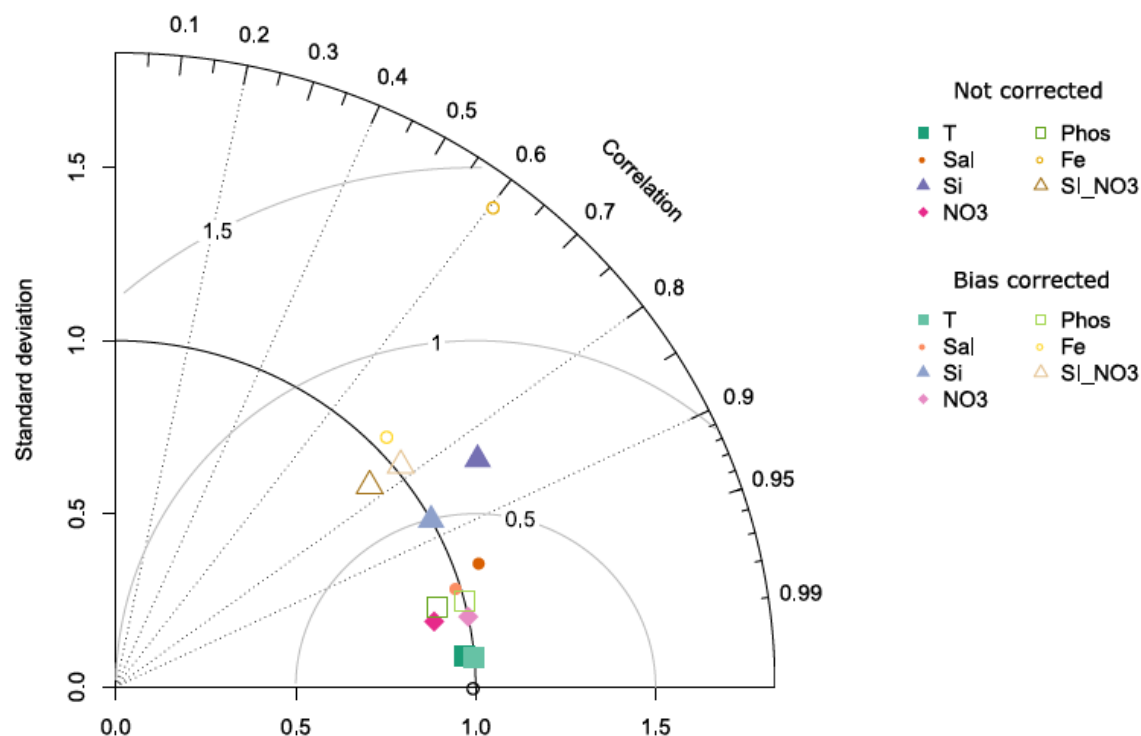
provinces for the six fractions and combined size fractions. The three oceanic partitions are Reygondeau et al. Biogeochemical Provinces (BGCP)¹⁰; Reygondeau et al. Biomes¹⁰; Fay and McKingley Biomes¹¹. Each partitioning mask is overlaid in the above order on top of plankton provinces for the six size fractions. **(a-c)** 180-2000 µm **(d-f)** 20-180 µm **(g-i)** 5-20 µm **(j-l)** 0.8-5 µm **(m-o)** 0.22-3 µm and **(p-r)** 0-0.2 µm. Above all maps, the adjusted rand index (ARI, an index used to compare partitions), for the comparison of mapped biogeographies with the black line mask is shown.



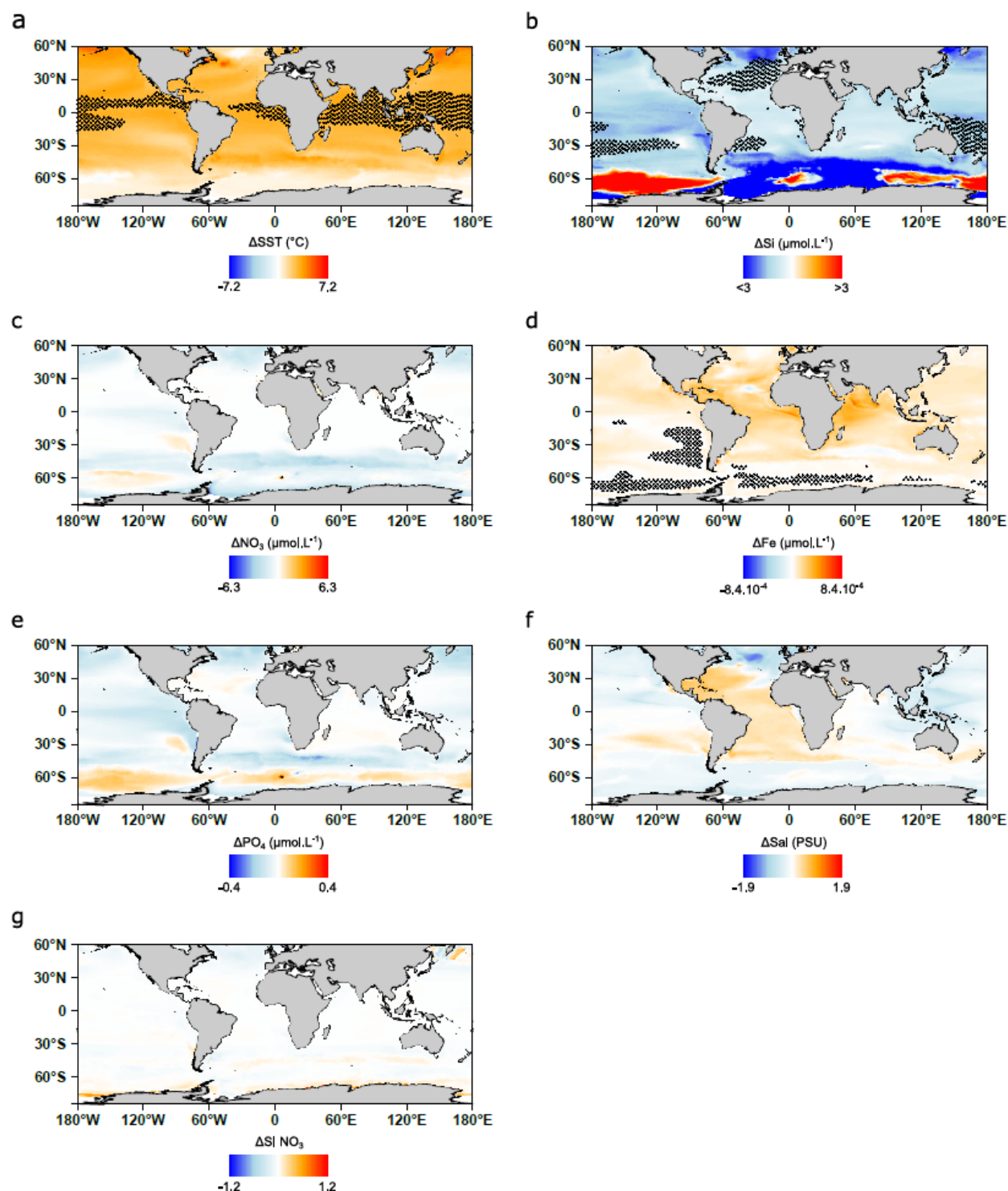
Supplementary Fig. 9 | Three existing oceanic partitions overlaid on top of plankton provinces for the combined size fractions. The three oceanic partitions are Reygondeau et al.¹⁰ Biogeochemical Provinces (BGCP); Reygondeau et al. Biomes¹⁰; Fay and McKingley Biomes¹¹. Each partitioning mask is overlaid in the above order on top of size fraction plankton provinces combined by the PHATE algorithm in (a-c) 4 clusters and (d-f) 7 clusters. Each grid point is associated with a color depending on its PHATE coordinates (using three axes). Each coordinate is respectively assigned to a given degree of color between 0 and 255 according to equation 2 (either red green or blue depending on the axis).



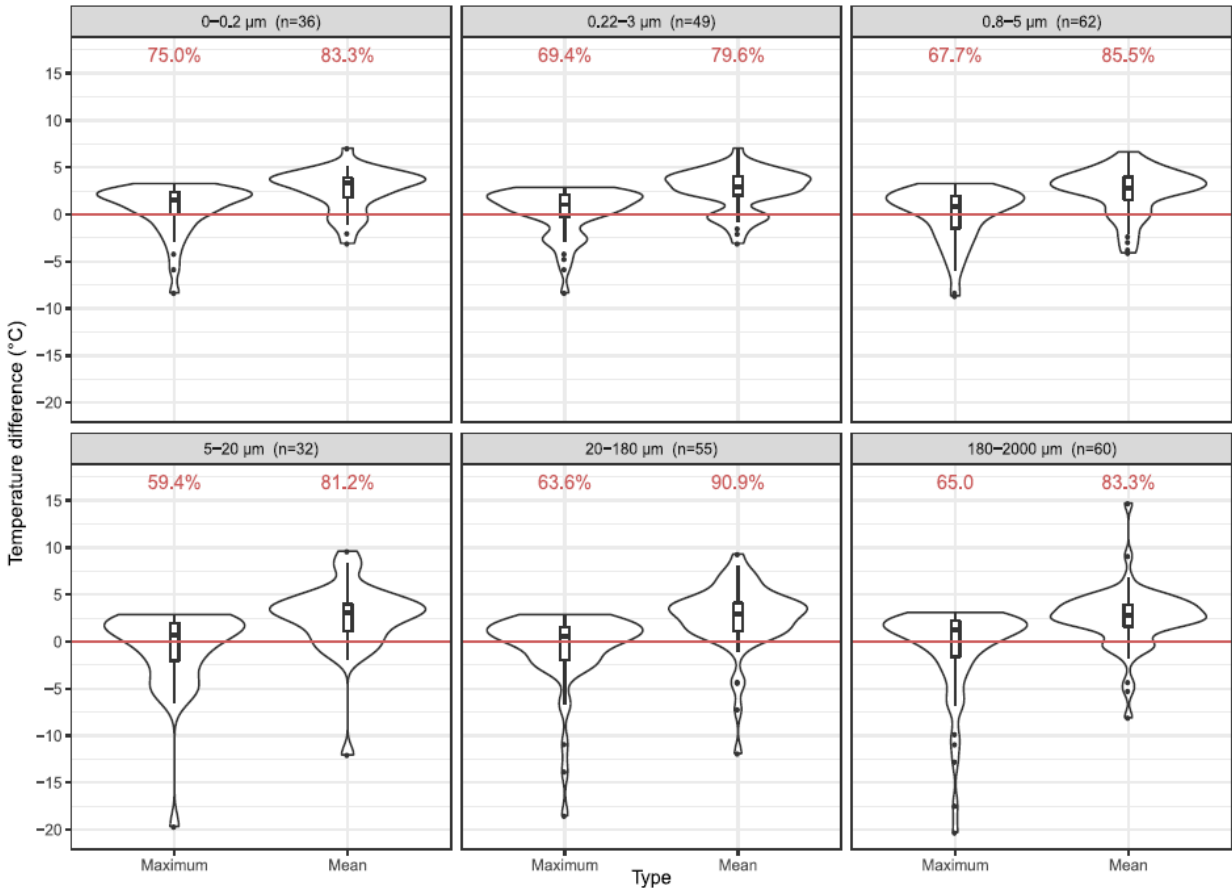
Supplementary Fig. 10 | Pairwise comparisons of ocean partitions based on plankton provinces and existing biogeochemical partitions. The three oceanic partitions are Reygondeau et al.¹⁰ Biogeochemical Provinces (BGCP); Reygondeau et al. Biomes¹⁰; Fay and McKingley Biomes¹¹.



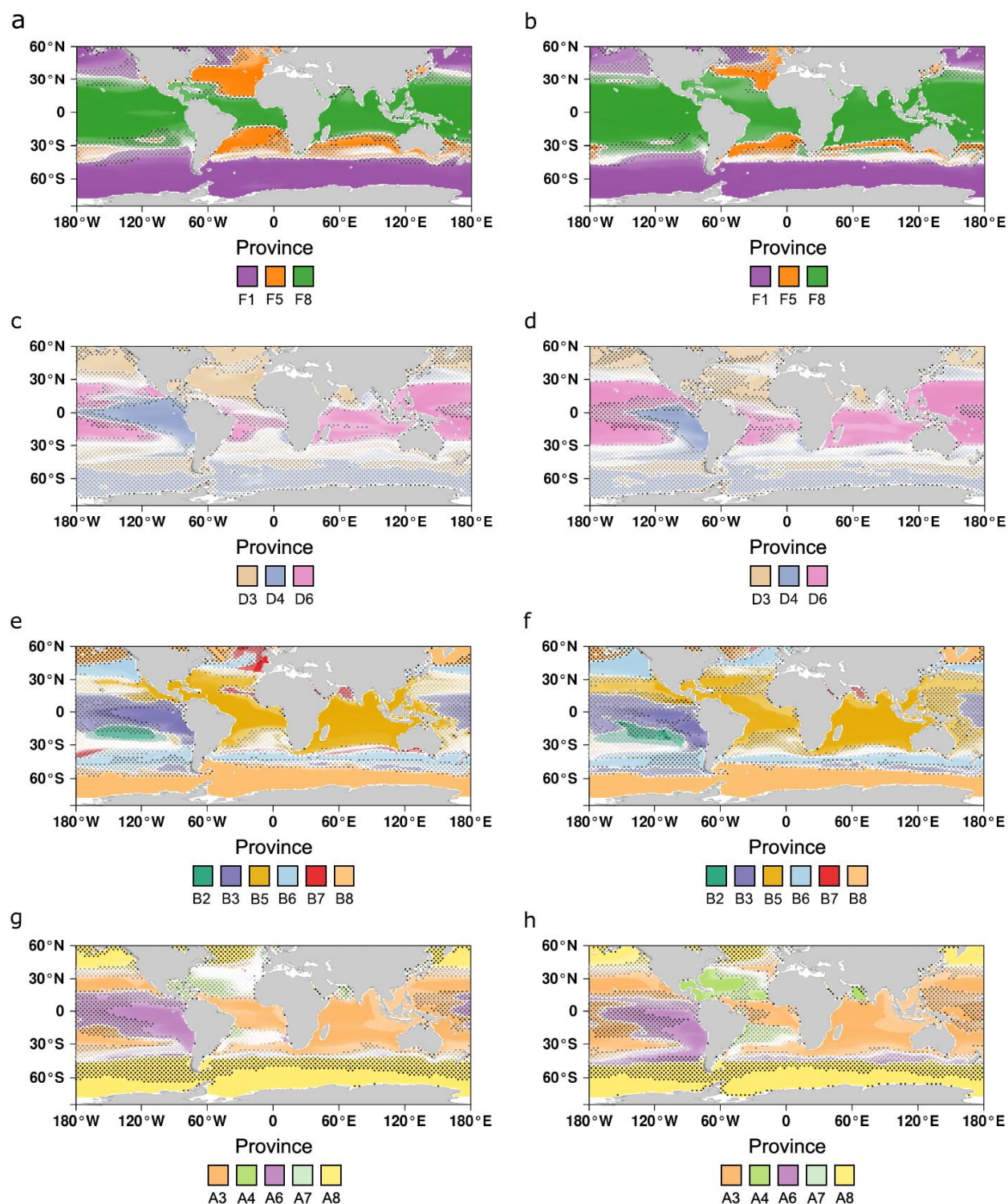
Supplementary Fig. 11 | Taylor diagram exhibiting statistical comparison of WOA 2013 observations and present day ESM model mean of the different drivers. Each variable is centered and scaled according to the mean and standard deviation of the observed variable (black circle point at standard deviation 1 on the x-axis). Dark color points are the ESM model mean without Cumulative Distribution Function transform (CDFt²⁴) bias correction. They are to be compared with light color points for which the bias correction is performed. Overall, good spatial correlations are found between the model mean and the observations. CDFt bias correction performs well by bringing standard deviations of the model to the observed standard deviations without decreasing correlations.



1033 maximum found in 2006-15) are reached at the end of the century are highlighted with small
1034 stars reflecting high uncertainty zones for machine learning approaches.
1035

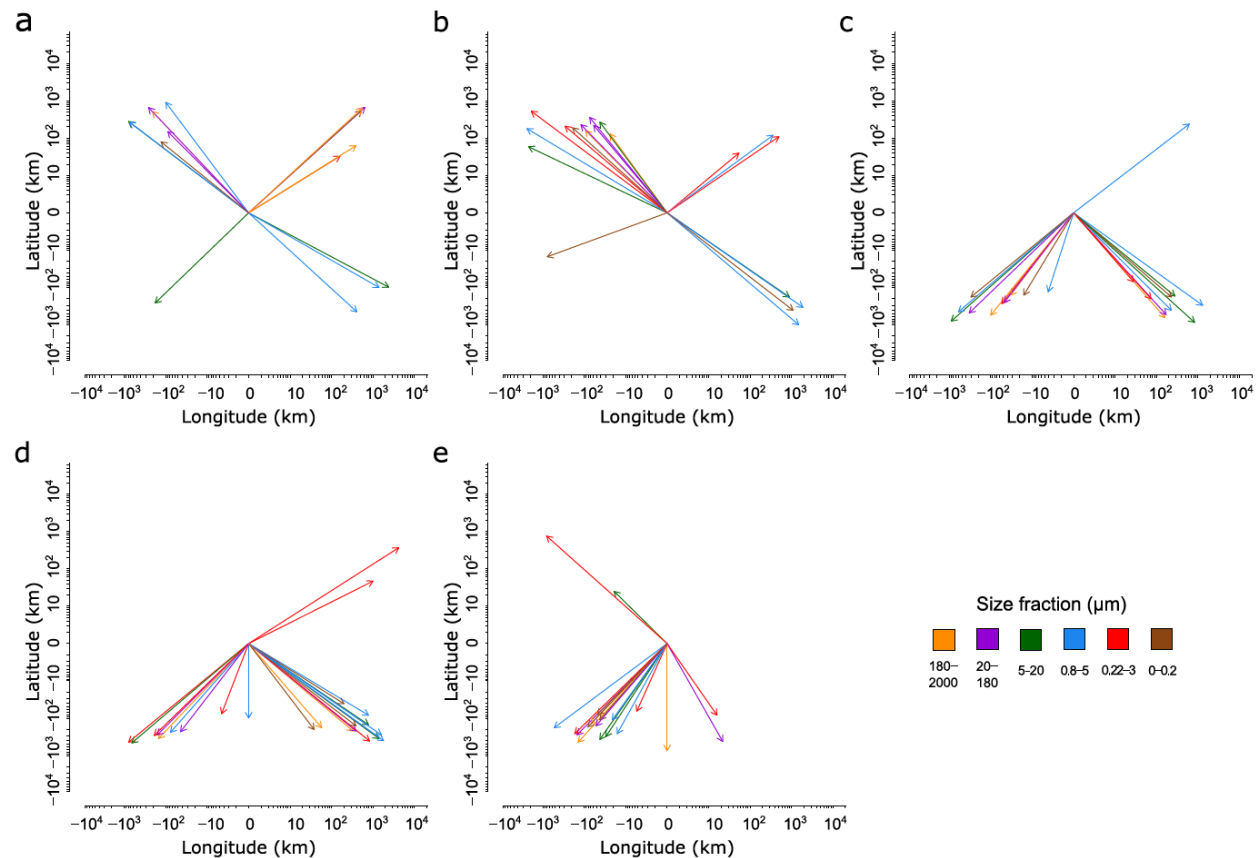


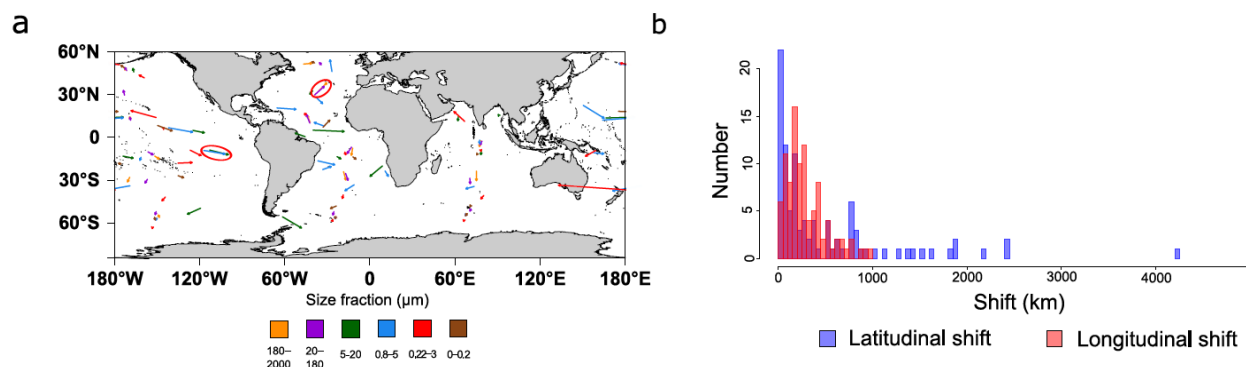
Supplementary Fig. 13 | Distribution of deltas between future temperature at each sampling site minus either the mean or maximum temperature within their contemporary genomic province. For most of the sites and across size fractions the future temperature projected by the bias adjusted ESM ensemble model is higher than both the maximum and mean contemporary temperature of their genomic province.



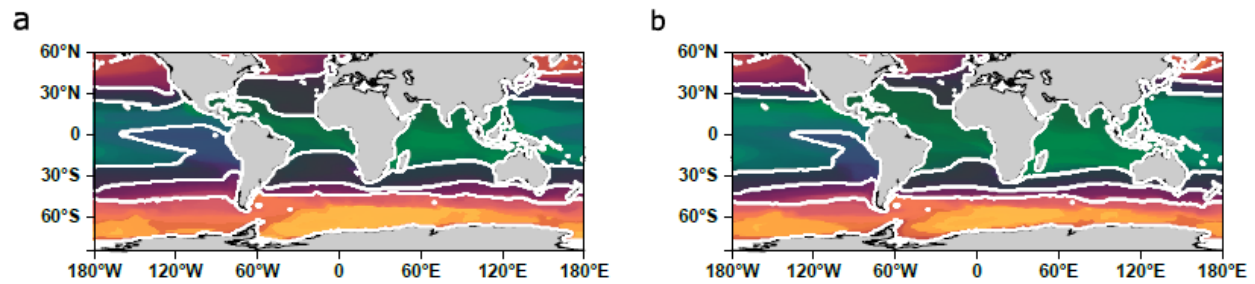
Supplementary Fig. 14 | Global geographical patterns for 180-2000, 5-20, 0.22-3, 0-0.2 μ m plankton size fractions in present day (a, c, e, g) and at the end of the century (b, d, f, h). The dominant province *i.e.* the one predicted to have the highest probability of presence is represented at each grid point of the map. The color transparency is the probability of presence

1048 of the dominant province. Expansion of tropical provinces and shrinkage of temperate
1049 communities is consistently projected in all size fractions.
1050

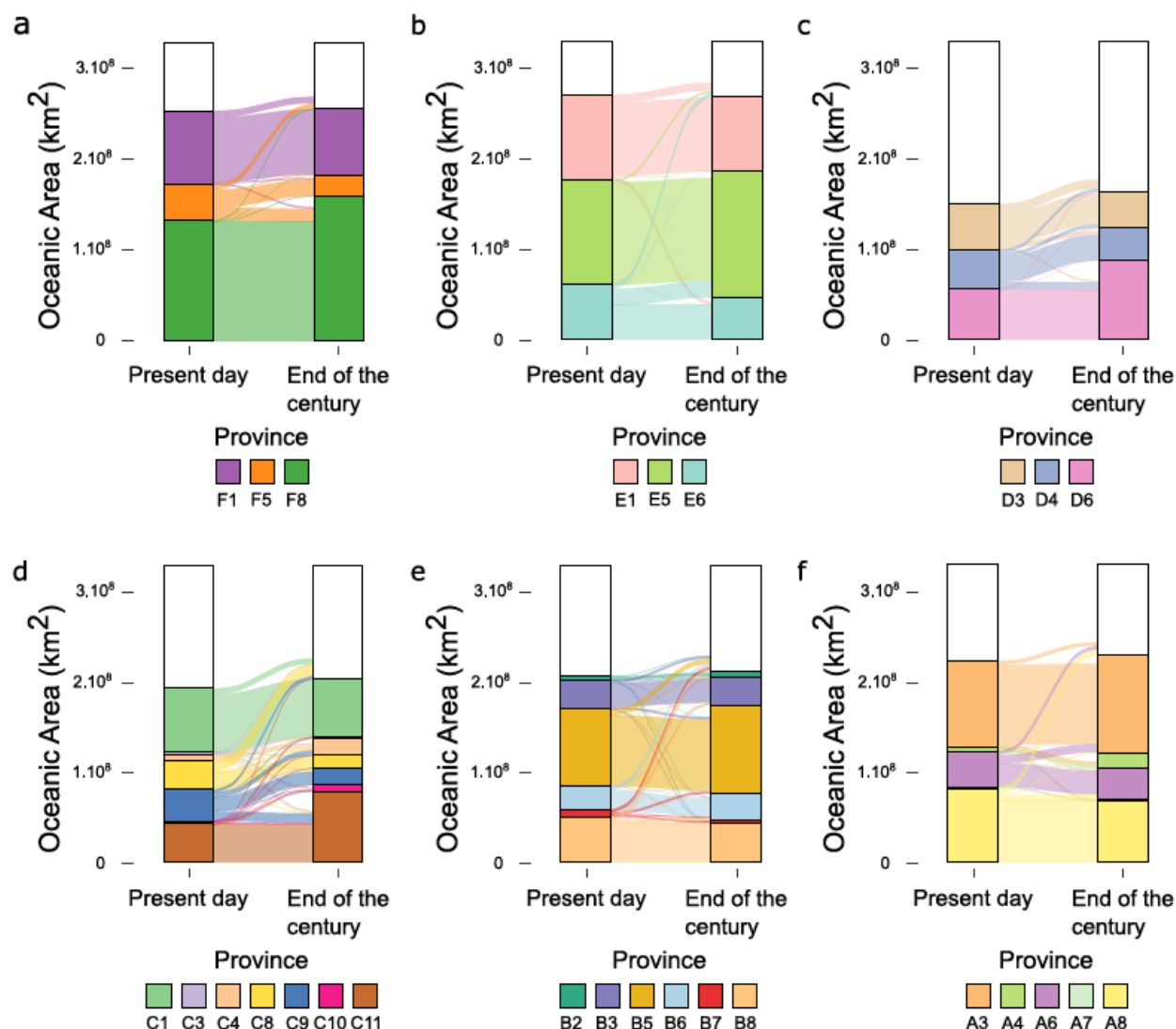




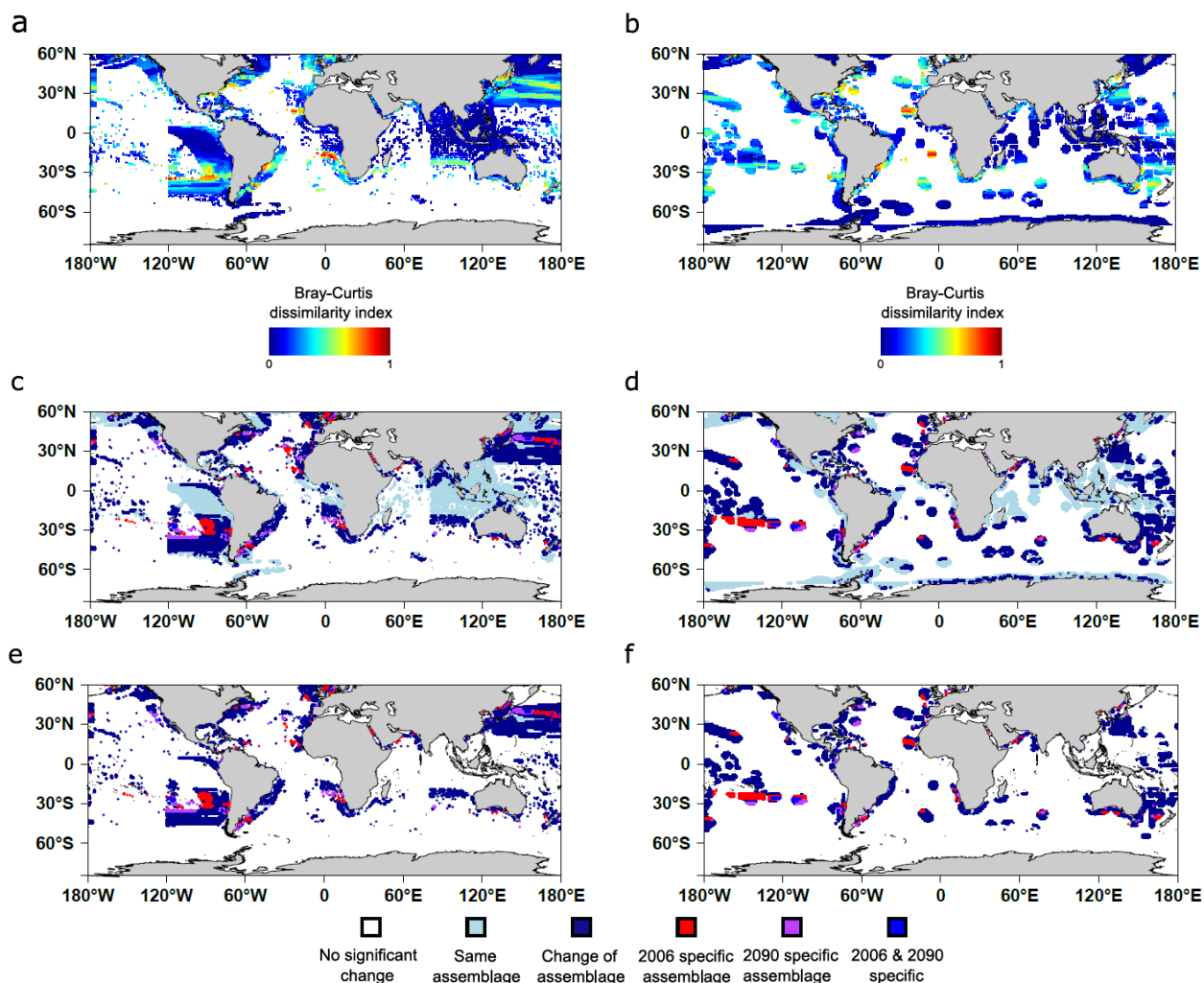
Supplementary Fig. 16 | (a) Projected migration shifts of provinces on the world map. (b) Latitudinal shift distribution (red bars) and longitudinal shift distribution (blue bars). (a) Migration shifts are represented as arrows pointing at the end of century centroid. Arrows are colored according to the size fraction. Some shifts seem to correlate with each other (exemplified with circled arrows). For instance, parallel shifts are projected in the southern pacific equatorial communities of size fractions 0.8-5 μm and 5-20 μm (blue and green circled arrows). All non-poleward arrows belong to small size classes ($<20\mu\text{m}$) showing differential responses to climate change depending on the size class. (b) Some longitudinal shifts are more important than latitudinal shifts with 14 longitudinal shifts superior to 1000 kms. Mean longitudinal shift (around 500 kms) is significantly higher (Student t-test²⁵ p-value <0.01) than mean latitudinal shift (around 290 kms) while medians (longitudinal 190 kms vs latitudinal 230 kms) are not significantly different (Wilcoxon test). The median migration speed is 47 $\text{km}.\text{dec}^{-1}$, (latitudinally 23 $\text{km}.\text{dec}^{-1}$, longitudinally 27 $\text{km}.\text{dec}^{-1}$).

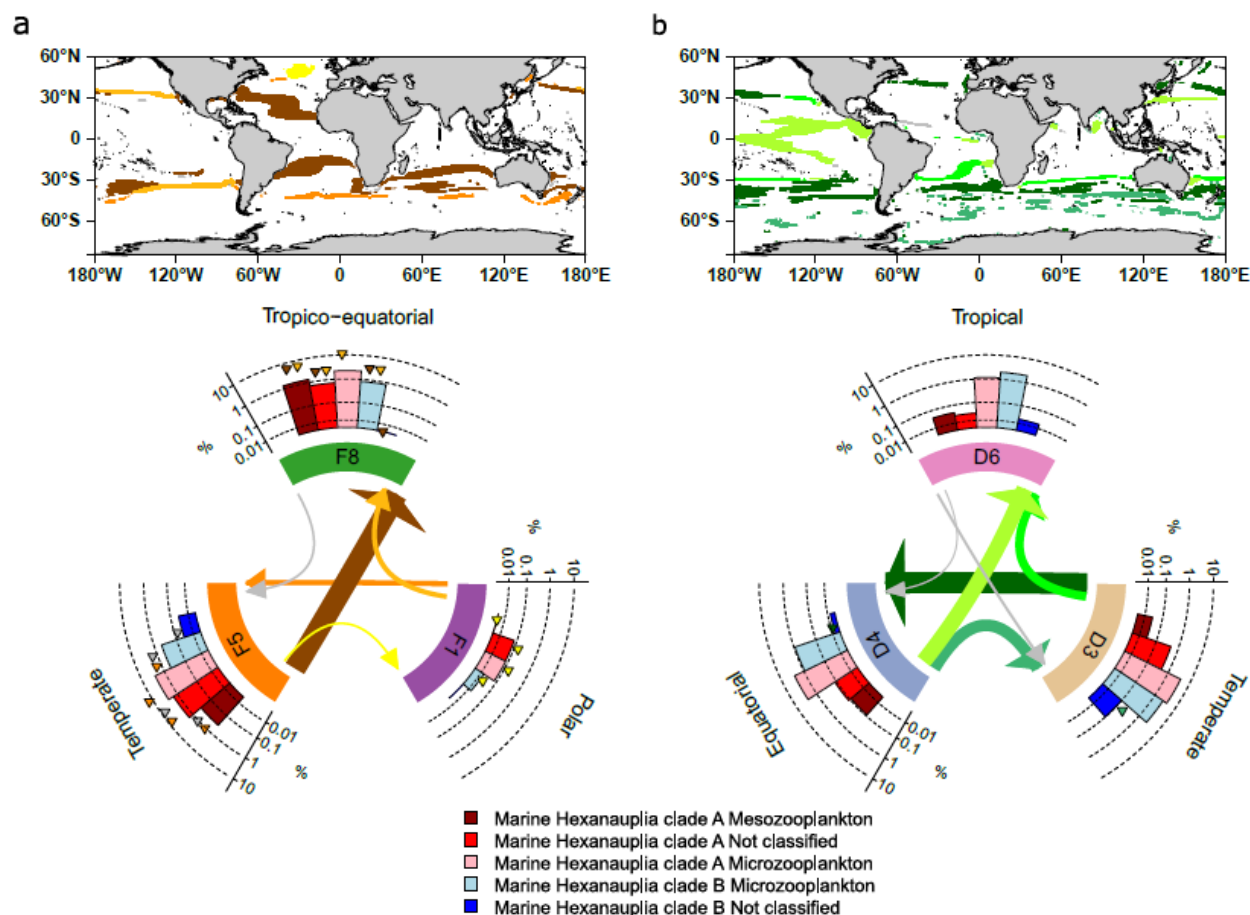


Supplementary Fig. 17 | (a) Present day and (b) end of century combined size class biogeographies using the PHATE algorithm⁷. Each grid point is associated with a color depending on its PHATE coordinates (using three axes). Each coordinate is respectively assigned to a given degree of color between 0 and 255 according to equation 2 (either red green or blue depending on the axis). Boundaries of clusters using k-medoids (k=4) are represented in white.

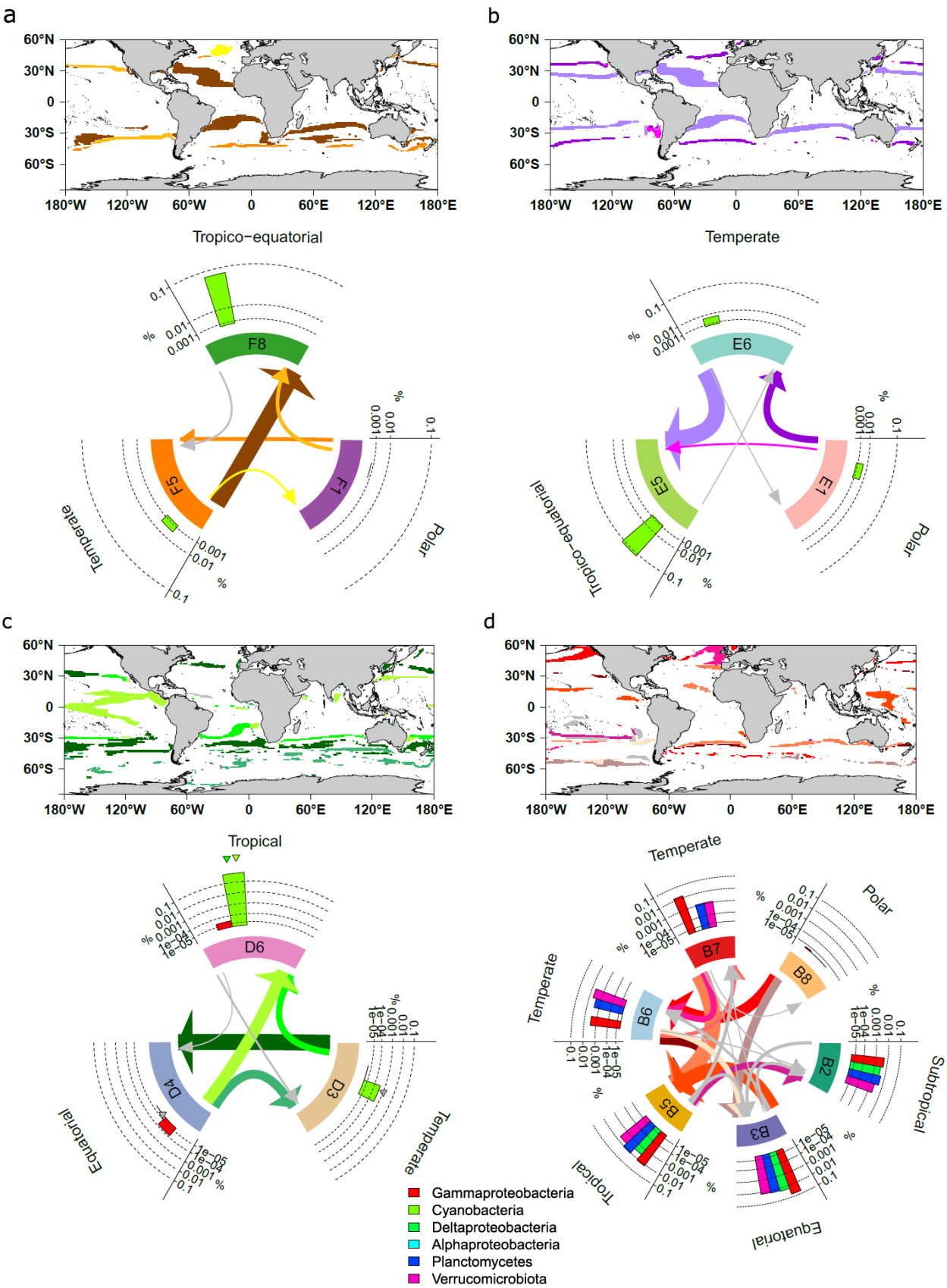


Supplementary Fig. 18 | Probabilistic covered areas of the provinces projected in present day and at the end of the century. (a) 180-2000 μm (b) 20-180 μm (c) 5-20 μm (d) 0.8-5 μm (e) 0.22-3 μm (f) 0-0.2 μm . The area covered by a province is defined as the area in which this province is dominant and weighted by its probability of presence at each point and grid cell area. Areas not covered by the provinces are represented in white.



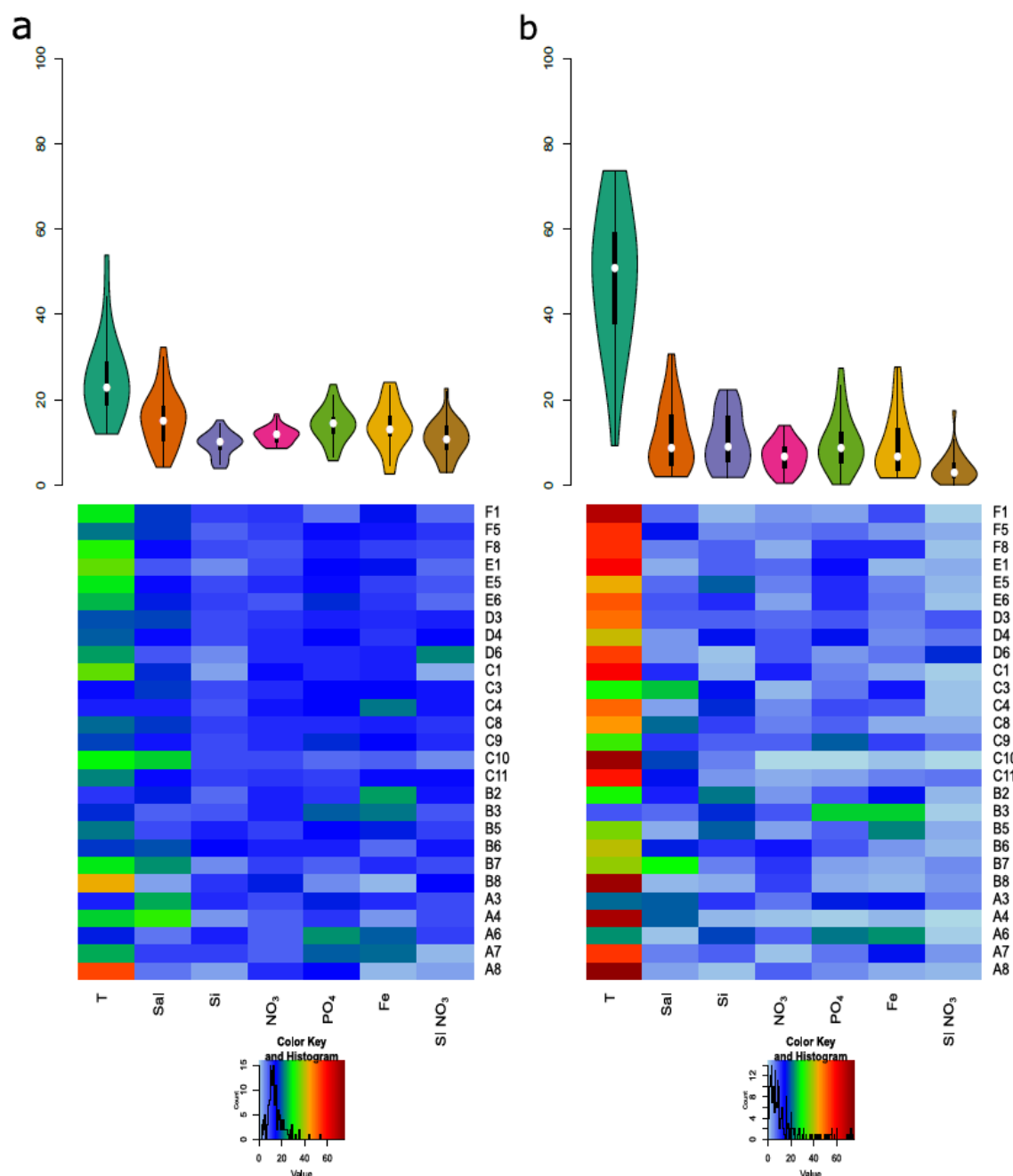


Supplementary Fig. 20 | Projected compositional change in marine hexanauplia in areas of dominant community change in the five major basins in size fraction (a) 180-2000 μm and (b) 5-20 μm . Top: Areas of dominant community change are highlighted using different colors depending on the dominant community transition. Bottom: Circular plots summarizing significant compositional shifts in marine hexanauplia: each type of transition is colored differently and according to the one on the map or in grey if they represent less than 2% of the transitions. Barplots represent mean relative abundances in 5 types of Marine Hexanauplia (copepods) based on genome abundance²⁸ in the given province. Arrows represent dominant community shifts pointing towards the end of the century projected province and their widths are proportional to the area of change. Significant compositional changes in a type of genome (e.g. mesozooplankton from clade A) are represented by triangles of the associated transition color above the corresponding bar (e.g. dark red for mesozooplankton from clade A) of the barplot of the province projected at the end of the century (which is at the tip of the arrow).



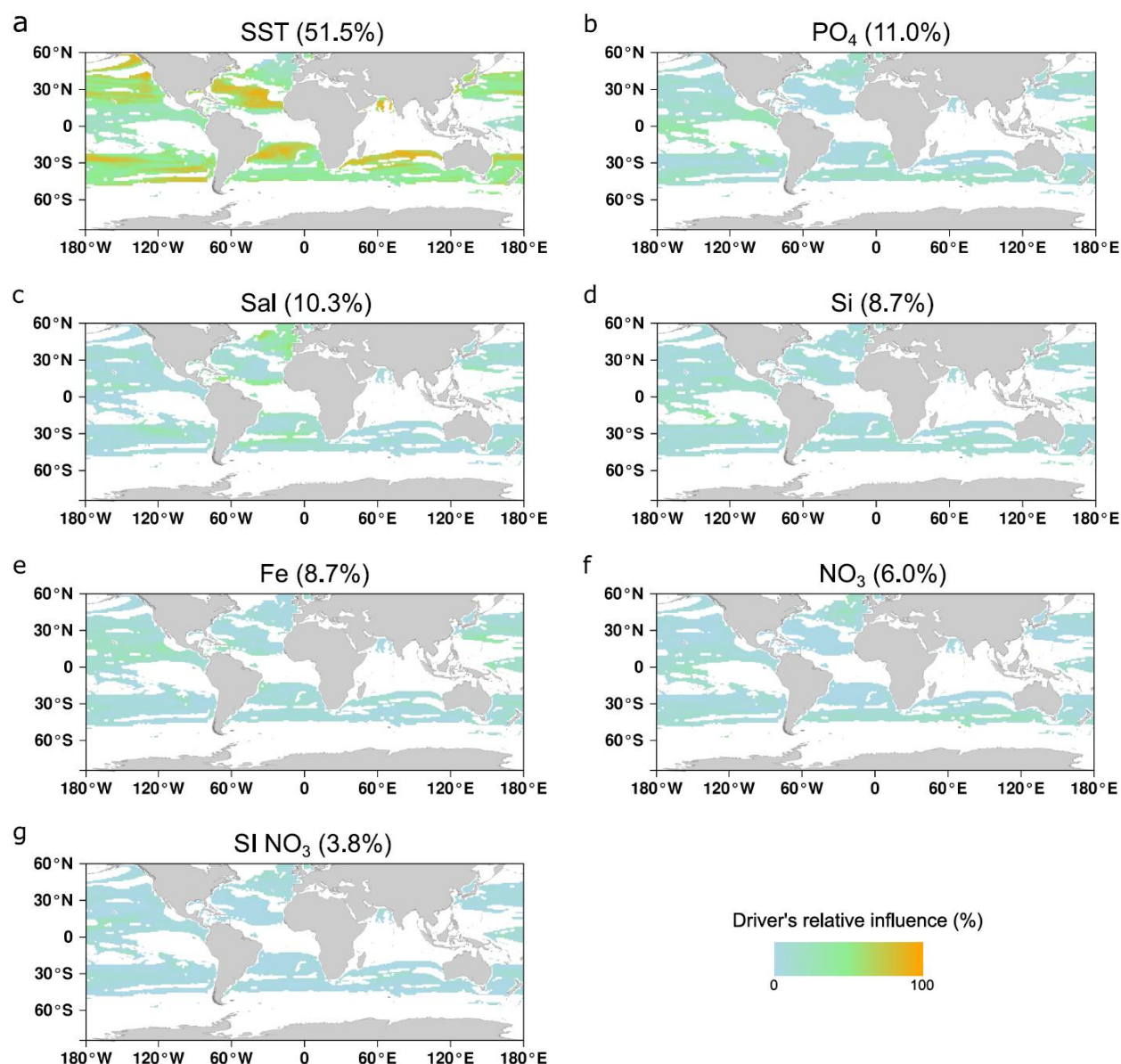
Supplementary Fig. 21 | Projected compositional change in bacterial diazotrophs

1108 **in areas of dominant community change in the five major basins in size fraction**
1109 **(a) 180-2000 μm (b) 20-180 μm (c) 5-20 μm and (d) 0.22-3 μm .** Top: Areas of
1110 dominant community change are highlighted using different colors depending on the
1111 dominant community transition. Bottom: Circular plots summarizing significant
1112 compositional shifts in marine diazotrophs: each type of transition is colored differently
1113 and according to the one on the map or in grey if they represent less than 2% of the
1114 transitions. Barplots represent mean relative abundances in main clades of marine
1115 diazotrophs²² based on genome abundance in the given province. Arrows represent
1116 dominant community shifts pointing towards the end of the century projected province
1117 and their widths are proportional to the area of change. Significant compositional
1118 changes in a type of genome (e.g. cyanobacteria) are represented by triangles of the
1119 associated transition color above the corresponding bar (e.g. yellowgreen for
1120 cyanobacteria) of the barplot of the province projected at the end of the century (which
1121 is at the tip of the arrow).

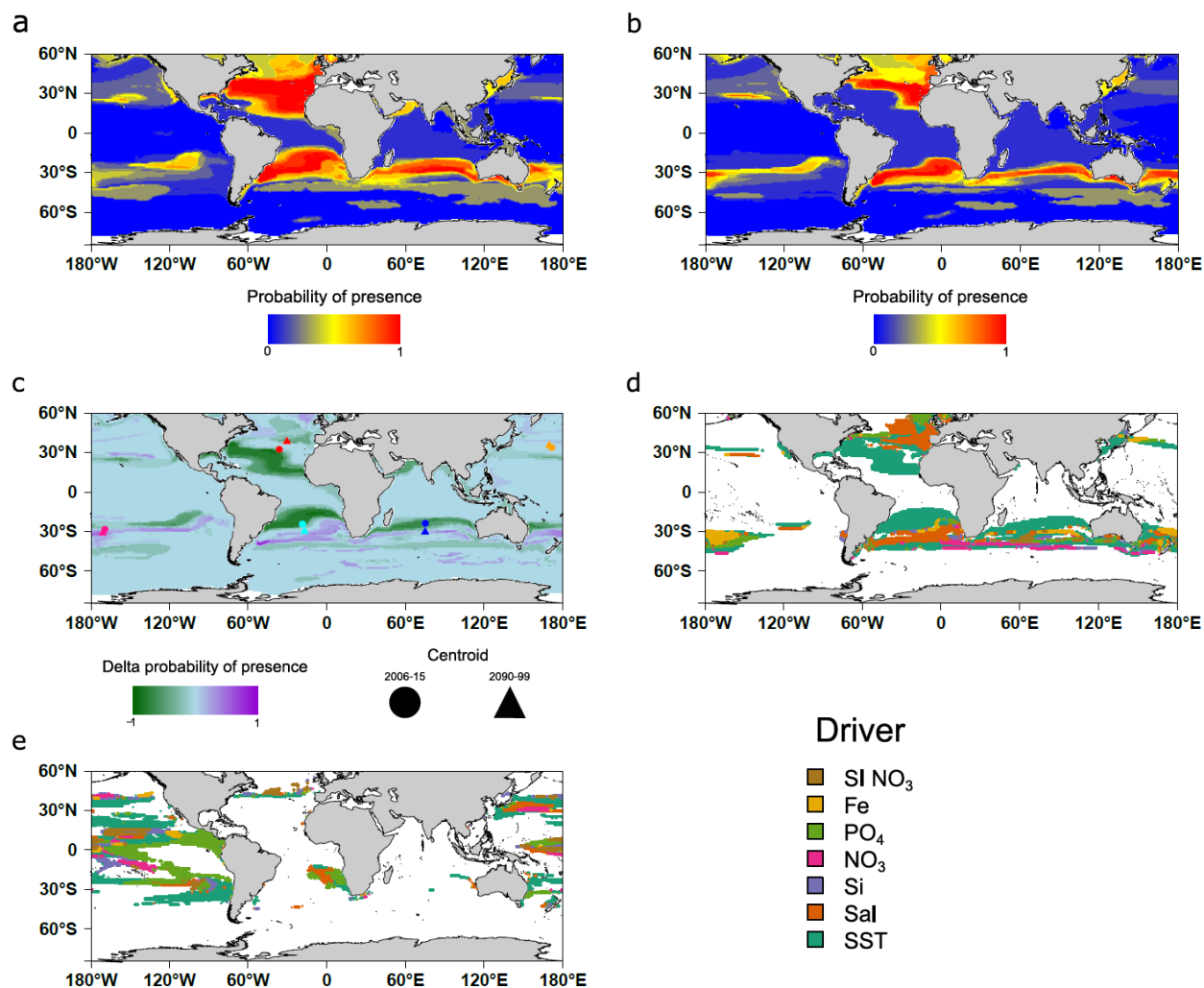


Supplementary Fig. 22 | Distributions and heat maps of the relative influences of the different drivers in (a) defining single niches associated with the provinces from DALEX R package²⁹ (b) driving climate change associated reorganization of single provinces. Median relative influence of Temperature is significantly higher than for all other environmental parameters (Pairwise Wilcoxon test $p < 0.01$ for all parameters) in (a) defining the niches and (b) driving province reorganization. This is also the case within individual size fractions for (b) but not for (a). Respectively, Salinity (Sal) and Phosphate (PO₄), have second and third highest median relative influences far behind Temperature whereas dissolved Silica and seasonality

1130 index of Nitrate (SI NO₃) have the lowest median relative importance in (a). Numbers above
 1131 violins are median (and mean) relative influences.
 1132



Supplementary Fig. 23 | Maps of environmental drivers' relative influences in driving province reorganization. Each environmental parameter's relative influence is quantified by considering only the variation of each parameter individually between present day and end of the century as defined in Barton et al.³⁰ (*Materials and Methods*) and where a significant change is projected. Importantly the mean impact of SST (51.5%) is to a great extent the highest. PO₄ is the second most impacting driver (11.0%). Contrary to supplementary Fig. 24, relative influence is calculated here by combining all provinces together. Therefore, mean relative influences slightly differ especially for dissolved silica (Si).



Supplementary Fig. 24 | Projection maps of province F5 of size fraction 180-2000 μ m in present day (a), at the end of the century (b) and their difference (c); Drivers of changes for province F5 (d) and C9 (e). At each grid point, the probability of presence of the province is computed as the average of the predicted probability of each of the four machine learning techniques (gbm, nn, rf and gam). Red color indicates a high probability of presence. (a) The projected province corresponds to the sampled province (North Atlantic and South Atlantic) but several other places have high probabilities of presence such as South Australia where no sampling is available. (b) At the end of the century, the province is projected to reduce significantly in size. (c) Delta probability of presence map (2006/15 – 2090/99) and core range shift in the 5 major oceanic basins of province F5 of size fraction 180-2000 μ m. In all the basins, the centroid of the province is projected to migrate poleward. (d) Main drivers associated with the projected changes. Changes are mainly driven by sea surface temperature (56%) followed by salinity (16%). (e) Main drivers associated with the shrinkage of the equatorial cluster C9 of

1156 size fraction 0.8-5. Considering only latitudes between the two tropics, changes are mainly
1157 driven by decreases in PO_4 (24%) in addition to SST (27%) (overall 34% STT and 20% PO_4).
1158

1159

Model	Reference
CESM1-BGC	Gent et al., 2011 ¹³
GFDL-ESM2G	Dunne et al., 2013 ¹⁴
GFDL-ESM2M	Dunne et al., 2013 ¹⁴
HadGEM2-ES	Collins et al., 2011 ¹⁵
IPSL-CM5A-LR	Dufresne et al., 2013 ¹⁶
IPSL-CM5A-MR	Dufresne et al., 2013 ¹⁶
MPI-ESM-LR	Giorgetta et al., 2013 ¹⁷
MPI-ESM-MR	Giorgetta et al., 2013 ¹⁷
NorESM1-ME	Bentsen et al., 2013 ¹⁸

1160 **Supplementary Table 1 | Earth System models used to compute the mean model.**

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1162

Fraction	Province	Climatic annotation	Area 2006-15 (MKm ²)	Area 2090-99 (MKm ²)	Delta area (MKm ² (%))
180-2000	F1	polar	82	73	-9 (-11%)
180-2000	F5	temperate	44	26	-18 (-41%)
180-2000	F8	tropico-equatorial	140	169	+29 (+21%)
20-180	E1	polar	97	84	-13 (-13%)
20-180	E5	tropico-equatorial	119	145	+26 (+22%)
20-180	E6	temperate	65	49	-15 (-23%)
5-20	D3	temperate	51	41	-10 (-20%)
5-20	D4	equatorial	46	37	-8 (-18%)
5-20	D6	tropical	65	97	+32 (+48%)
0.8-5	C1	polar	71	64	-6 (-8%)
0.8-5	C3	subtropical	3,6	1,4	-2,2 (-61%)
0.8-5	C4	subtropical	6	19	+13 (+217%)
0.8-5	C8	temperate	33	15	-18 (-54%)
0.8-5	C9	equatorial	35	19	-16 (-45%)
0.8-5	C10	subtropical	3.8	7.4	+3,6 (+95%)
0.8-5	C11	tropical	49	86	+37 (+75%)
0.22-3	B2	subtropical	5,1	6,7	+1,6 (+31%)
0.22-3	B3	equatorial	33	32	-1 (-0.3%)
0.22-3	B5	tropical	88	102	+13 (+15%)
0.22-3	B6	temperate	30	35	+5 (+17%)
0.22-3	B7	temperate	8	2	-6 (-75%)
0.22-3	B8	polar	51	44	-7 (-14%)
0-0.2	A3	tropical	101	114	+13 (+13%)
0-0.2	A4	subtropical	6	19	+13 (+216%)
0-0.2	A6	equatorial	40	36	-4 (-10%)
0-0.2	A7	temperate	1,7	1	-0,7 (-41%)
0-0.2	A8	polar	81	67	-14 (-17%)

1163 **Supplementary Table 2 | Genomic provinces climatic annotations and oceanic surfaces**
1164 **covered in present day (2006-15) and at the end of the century (2090-990).**

1165

1166

	Fraction (μm)	SST	Sal	Si	NO3	PO4	Fe	SI NO3
Niche definition	180-2000	27,7	16,8	10,1	10,3	11,4	14	9,7
	20-180	29,7	13,3	8,9	10,7	16,3	13,2	7,9
	5-20	21,5	14,2	8,4	12,7	13,7	12,6	17
	0,8-5	22,6	17,3	9,2	12,3	13,4	14,5	10,8
	0,22-3	23,9	13,7	10,5	13	12,5	14,2	12,2
	0-0,2	27,1	16,4	9,3	9,5	17,9	12,1	7,6
	all	25	15,5	9,5	11,6	14,2	13,6	10,8
Climate change	180-2000	75,2	8,4	2,5	2,6	4,1	5,8	1,4
	20-180	71,7	4,2	7,3	3,9	8,7	3,3	1,0
	5-20	57,9	6,1	4,4	6,6	9,0	6,5	1,0
	0,8-5	54,9	13,4	7,2	5,5	9,7	5,9	3,6
	0,22-3	42,6	9,9	13,5	6,4	9,1	15,2	3,3
	0-0,2	42,3	18,9	6,6	6,4	12,1	10,8	2,9
	all	51,5	10,3	8,7	6	11	8,7	3,8

1167 **Supplementary Table 3 | Summary table of the relative importance of each environmental**
1168 **driver in niche definition and in driving geographical reorganization in response to**
1169 **climate change.** Note that in both cases (niche definition and climate change), the row 'all' is
1170 not the mean over the size fractions. In the case of niche definition, this is due to a different
1171 number of niches in each size class. In the case of climate change, relative influence is either
1172 calculated for single provinces at a given grid point then recalculated for individual size class or
1173 calculated for all provinces together (row 'all').

1174

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