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Three major mesoplanktonic communities resolved by in situ imaging in the upper 500 m of the global ocean

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Abstract

Aim: The distribution of mesoplankton communities has been poorly studied at global scale, especially from in situ instruments. This study aims to (1) describe the global distribution of mesoplankton communities in relation to their environment and (2) assess the ability of various environmental-based ocean regionalizations to explain the distribution of these communities.

Location: Global ocean, 0–500 m depth.

Time Period: 2008-2019.

Major Taxa Studied: Twenty-eight groups of large mesoplanktonic and macroplanktonic organisms, covering Metazoa, Rhizaria and Cyanobacteria.

Methods: From a global data set of 2500 vertical profiles making use of the Underwater Vision Profiler 5 (UVP5), an in situ imaging instrument, we studied the global distribution of large (>600 μ m) mesoplanktonic organisms. Among the 6.8 million imaged objects, 330,000 were large zooplanktonic organisms and phytoplankton colonies, the rest consisting of marine snow particles. Multivariate ordination (PCA) and clustering were used to describe patterns in community composition, while comparison with existing regionalizations was performed with regression methods (RDA).

[Correction added on 10 August 2023, after first online publication: new affiliation has been added to the author, Helena Hauss, in this version.]

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Results: Within the observed size range, epipelagic plankton communities were *Trichodesmium*-enriched in the intertropical Atlantic, Copepoda-enriched at high latitudes and in upwelling areas, and Rhizaria-enriched in oligotrophic areas. In the mesopelagic layer, Copepoda-enriched communities were also found at high latitudes and in the Atlantic Ocean, while Rhizaria-enriched communities prevailed in the Peruvian upwelling system and a few mixed communities were found elsewhere. The comparison between the distribution of these communities and a set of existing regionalizations of the ocean suggested that the structure of plankton communities described above is mostly driven by basin-level environmental conditions.

Main Conclusions: In both layers, three types of plankton communities emerged and seemed to be mostly driven by regional environmental conditions. This work sheds light on the role not only of metazoans, but also of unexpected large protists and cy-anobacteria in structuring large mesoplankton communities.

KEYWORDS

biogeography, Copepoda, global ocean, in situ imagery, plankton communities, Rhizaria, spatial distribution, *Trichodesmium*

1 | INTRODUCTION

Plankton is defined as organisms unable to swim against currents. This definition based on habitat rather than taxonomy thus results in a large taxonomic diversity (Caron et al., 2012; Ibarbalz et al., 2019), encompassing bacteria, protists, algae, as well as drifting animals. Therefore, planktonic organisms cover a large size range, spanning 6 orders of magnitude, from micrometre to metre (Lombard et al., 2019). Plankton supports oceanic food webs (Falkowski, 2012; Ware & Thomson, 2005) and plays a major role in biogeochemical cycles through the biological pump (Longhurst & Glen Harrison, 1989). As drifters, planktonic organisms are distributed worldwide but their distribution is shaped by the conditions of the water mass they are embedded in (Hays et al., 2005). Because these conditions strongly vary with latitude, both plankton diversity (Ibarbalz et al., 2019; Rombouts et al., 2009; Rutherford et al., 1999; Tittensor et al., 2010) and biomass (Ikeda, 1985) vary with latitude: high diversity and low biomass at low latitudes; low diversity and high biomass at high latitudes. Indeed, at low latitudes, higher temperatures increase the rate of metabolism, resulting in both shorter generation times and higher mutation rates, causing a higher diversity (Brown, 2014). Conversely, at high latitudes, more nutrients are available, enabling higher production and higher biomass. Overall, plankton distribution is governed by both water mass movements (through dispersal) and environmental conditions, and the contribution of each effect might depend on the size of organisms (Sommeria-Klein et al., 2021; Villarino et al., 2018). Because they are sensitive to environmental conditions, planktonic organisms are also global change sentinels (Beaugrand et al., 2002, 2009; Hays et al., 2005). Studying plankton biography thus is relevant to understand anthropocene pelagic ecosystems.

In terrestrial biogeography, biomes rest on vegetation types, but also coincide with climatic zones and soil type distribution constraining

plant growth (Brown & Lomolino, 1998). Compared to the terrestrial realm, assessing oceanic biogeography presents inherent difficulties: costly global scale offshore sampling, observing distribution varying in time and space in a three-dimensional and opaque environment. Early ocean biogeographies considered various taxa's distribution, including copepods, euphausiids, Rhizaria or phytoplankton (Spalding et al., 2012). Simultaneously, non-biological regionalizations were based on the physical environment: ocean currents, temperature, salinity, ice conditions (Spalding et al., 2012). Novel technologies, such as satellites, fostered ecological ocean geography: using surface chlorophyll a concentration computed from ocean colour (a proxy for phytoplankton concentration) new regionalizations emerged. However, most new approaches ignore organisms distribution: the 56 Longhurst Provinces (Longhurst, 1995, 2010) considered physical forcing (sea surface temperature, mixed layer depth...) as phytoplankton distribution regulator. Another widely used global synthetic regionalization is based on latitudinal bands (Spalding et al., 2012). As explained above, it correlates with major environmental variables (temperature, light intensity...). Other regionalizations, the Word Marine Ecoregions (Spalding et al., 2007) or the Large Marine Ecosystems (Sherman, 2005) include biotic data, but focus on coastal areas only. In contrast, Costello et al. (2017) delineated marine biogeographic realms using the distribution of marine animals and plants, while Hofmann Elizondo et al. (2021) defined biomes using phytoplankton community composition as well as species cooccurrence patterns. Furthermore, these regionalizations focus on the epipelagic layer and are not suitable for less described deeper ocean, harder to sample and not necessarily linked to surface characteristics (Costello et al., 2018; Spalding et al., 2012). Few regionalizations targeted the mesopelagic: Reygondeau et al. (2018) suggested dividing it into 13 provinces, based on environmental variables, while defining the top and bottom boundaries of the mesopelagic layer dynamically, based on environmental conditions (light, density, carbon flux).

In summary, ocean biogeography was described through regionalizations, but mostly in the epipelagic layer or coastal areas. In comparison, widespread offshore areas, although crucial for biogeochemical cycles (Emerson et al., 1997) and the target of conservation through developing marine protected areas, were sampled much less, due to technical constraints. In the context of global warming, mesoplankton communities are facing changes in their abundance, size and diversity (Richardson, 2008), susceptible to affect higher trophic levels, with consequences for exploited marine resources such as fisheries. Thus, it is not only essential to describe mesoplankton communities at global scale, but also to assess the consistency between these and biogeochemistry-based regionalizations, typically used for the conservation and management of marine ecosystems (Rice et al., 2011).

Although a few organisms groups' spatial distribution – copepods (Beaugrand et al., 2013; Rombouts et al., 2009; Woodd-Walker et al., 2002), microorganisms (Fuhrman et al., 2008; Hörstmann et al., 2022; Li et al., 2004) or larger species assemblages (Brandão et al., 2021; Rutherford et al., 1999; Soviadan et al., 2022; Tittensor et al., 2010) – were described previously, consistency between these and biogeochemistry-based regionalizations remains unexplored.

Limited quantitative and basin-scale data about offshore planktonic organism distribution is available, especially for in situ imaging data. Traditional sampling tools (nets, pumps...) require lengthy taxonomic identification (Benfield et al., 2007), partly subjective (Culverhouse et al., 2003), therefore not scaling well to large spatiotemporal scales. Besides, they may damage fragile organisms (Remsen et al., 2004). New in situ cameras now image planktonic organisms in their natural environment and resolve their fine-scale vertical distribution (Stemmann et al., 2008), while generating large datasets (Kiko et al., 2022), homogenized by reviewing images (Irisson et al., 2022). These tools also allow studying fragile taxonomic groups such as Rhizaria, whose contribution to global planktonic biomass was underestimated (Biard et al., 2016; Dennett et al., 2002; Drago et al., 2022). These approaches lack in taxonomic identification fineness, but compensate with identifications and data quantity consistency. Among these imaging systems, the Underwater Vision Profiler 5 (UVP5) images planktonic organisms and marine snow particles larger than 600 µm Equivalent Spherical Diameter (ESD) along vertical profiles (Picheral et al., 2010), therefore sampling large meso- and small macroplankton, mostly comprising animals and some large phytoplankton colonies (this assemblage is referred as "plankton" in the following, for simplicity). Concentration and biovolume estimates from the UVP5 proved coherent with those from net samples for large (>1 mm ESD) Arctic copepods, while smaller organisms were underestimated (Forest et al., 2012). Data from UVP5 was already used to estimate organic carbon vertical particle flux (Guidi et al., 2015) or study zooplankton distribution (Biard et al., 2016; Drago et al., 2022; Forest et al., 2012; Stemmann et al., 2008). Leveraging net data from the Tara Oceans expedition, Soviadan et al. (2022) detected a lower particle flux attenuation in Oxygen

Minimum Zones (OMZ). UVP5 can also inform on planktonic organisms' individual traits, highlighting different size and activity patterns around Arctic ice melt zones (Vilgrain et al., 2021).

We study global-scale plankton biogeography, leveraging several UVP5 datasets. We address the following questions: which types of large mesoplankton and macroplankton communities exist in the open ocean; how these communities differ between epipelagic and upper-mesopelagic layers; are these communities driven by rather large or small-scale processes; do they correlate with environmental conditions? We first describe plankton community structures; their relation to their immediate physical and biogeochemical environment. We then assess the ability of various physics and biogeochemistry-based regionalizations to describe these planktonic communities distribution.

2 | MATERIALS AND METHODS

2.1 | Data collection

Data from multiple oceanographic campaigns (Figure S1, Table S1) (2008–2019) – when UVP5 vertical profiles were performed – was aggregated, creating a large dataset covering world's oceans. This in situ imaging system captures objects within an approximately 1L volume, up to 20 Hz frequency during a CTD cast descending part (Conductivity, Temperature, Depth sensor) (Picheral et al., 2010). All objects larger than 100 μ m ESD were measured for area and grey level. Images were saved for objects larger than 600 μ m ESD; this paper focuses on the latter part. In the following, we thus use the word "plankton" to refer to all the organisms detected by the UVP5, consisting mostly in mesozooplankton and macrozooplankton, as well as large phytoplankton colonies. The CTD provided temperature, salinity and also chlorophyll *a* fluorescence and oxygen profiles.

Satellite GlobColour¹ products completed the environmental dataset: averaged over 1 month on a 100×100 km area centred on UVP5's sampling site. Although this data resolution is low in terms of time and space, using higher resolution data provided a too great missing data proportion. Satellite data provided: surface chlorophyll *a* concentration; particulate backscattering coefficient (bbp); photosynthetically active radiation (PAR); diffuse attenuation coefficient (K_{dPAR}); particulate organic carbon (POC); particulate inorganic carbon (PIC). With these data, organisms were associated with specific environmental conditions.

2.2 | Data processing

All UVP5 images entered the EcoTaxa web application (Picheral et al., 2017). They were classified as marine snow, artefact, badfocus, unidentified, or into several taxonomic groups. All objects

¹http://globcolour.info

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were manually validated or corrected. Striving for consistency, a taxonomic sorting guide was published and circulated; difficult groups were reviewed by a single operator across all cruises. Human operators fully checked the resulting dataset, often several for each image. Data from 2500 fully validated profiles was retained (6.8 M objects, 330,000 classified as plankton). Differences in classification taxonomic depth among cruises caused some groups to be merged, obtaining a lower common denominator. Then, other groups were merged because they exhibited similar patterns in preliminary analysis (e.g., Copepoda and Copepodalike), for a final list of 28 taxa (Figure 1). Finally, the same operator reviewed a random subsample of images from each final taxonomic group, with error rates at <10% for all groups and <2.5% for most (Drago et al., 2022). Although a normalized biomass size spectrum revealed an underestimation in the 600μ m-1 mm size

range, 80% of organisms in our dataset were over 1mm in ESD, therefore accurately detected by UVP5. Furthermore, provided underestimation was consistent across taxonomic groups (we checked by inspecting dominant taxa's per-taxon size spectra), the community composition is little affected by the absolute concentrations underestimation.

Concentrations (L⁻¹) were computed per 5m bins along each profile from object counts per class and imaged water volume. Concentration and biovolume were also computed for marine snow (objects >600 μ m identified as aggregates) and bulk particulate matter (all objects >100 μ m ESD imaged by the UVP, including both plankton and marine snow). Marine snow and bulk concentrations and biovolumes were considered environmental variables, as proxies for, respectively, organic matter amounts sinking from the upper layers to ocean depths and the overall trophic water mass state.

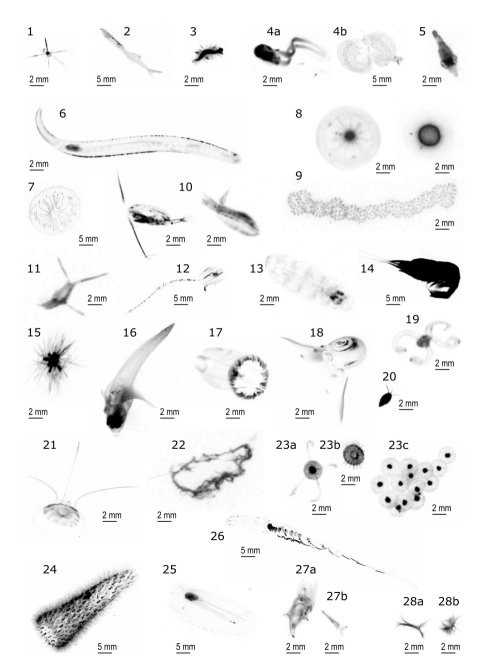


FIGURE 1 Examples of UVP5 images for selected taxonomic groups. 1: Acantharia, 2: Actinopterygii, 3: Annelida, 4: Appendicularia (4a: Appendicularia body, 4b: Appendicularia house), 5: Cephalopoda, 6: Chaetognatha, 7: Cnidaria others, 8: Collodaria, 9: colonial Collodaria, 10: Copepoda, 11: Crustacea others, 12: Ctenophora, 13: Doliolida, 14: Eumalacostraca, 15: Foraminifera, 16: Gymnosomata, 17: Hydrozoa others, 18: Limacinidae, 19: Mollusca others, 20: Ostracoda, 21: Narcomedusae, 22: Nostocales, 23: Phaeodaria (23a: Coelodendridae, 23b: Aulacantha, 23c: colonial Aulosphaeridae), 24: Pyrosoma, 25: Salpida, 26: Siphonophorae, 27: Thecosomata (27a: Cavoliniidae, 27b: Creseidae), 28: Trichodesmium (28a: tuft, 28b: puff).

After removing abnormal values (codes for missing data, negative salinity...), profiles with more than 20% of missing data for any variable were ignored. All variables were linearly interpolated at a 1m vertical resolution. Outliers were detected by computing the absolute deviation around a moving median along the profile (Leys et al., 2013) and removed. Smoothing was performed using a moving average. Potential density and apparent oxygen utilization (AOU) were computed from temperature, salinity and oxygen concentration. Thermocline, halocline and pycnocline depths were calculated as depth of the largest variation in the relevant variable computed in a 5 m sliding window. The mixed layer depth (MLD) was computed at depth where density differed by more than 0.03 kg/m³ from the reference density in the 0-5m surface layer (de Boyer Montégut et al., 2004). The deep chlorophyll maximum (DCM) and euphotic zone (Z_{au}) depths were computed from the chlorophyll profile (Morel & Maritorena, 2001). The stratification index was computed as the difference in potential density between the surface and 250m (below the pycnocline in most profiles). All 1m precision profiles were binned over 5m to match plankton data bins. Rare instances (1.7%) of missing satellite data were replaced by the corresponding variable average value.

Plankton and environmental data were averaged over two layers: epipelagic and upper mesopelagic. Instead of the commonly used fixed boundary (200 m) between these two layers (Costello & Breyer, 2017), we applied a dynamic definition. It was modified from Reygondeau et al. (2018) and meant to better represent the functional difference between the two layers. It was set as the deepest value among the mixed layer depth and the euphotic depth, hence delimiting the zone above which photosynthetic plankton activity is highest, resulting in an 88-m median value for the epi-mesopelagic boundary (Q1=52m, Q3=121m, Figure S2). As numerous UVP5 profiles stopped at 500m, we set the upper mesopelagic zone bottom at that depth. Any profile covering less than 80% of both layers' thickness was removed (3% and 29% of profiles for epipelagic and mesopelagic layers, respectively).

2.3 | Global plankton community distribution analysis

First, to synthetically describe plankton communities, a principal component analysis (PCA) was conducted on plankton concentration values. In each layer, the Hellinger transformation was applied to averaged plankton concentrations, as a compromise between absolute and relative concentrations, to focus on community composition differences among profiles (Legendre & Legendre, 2012). This helped go beyond the well-known pattern of high latitudes higher concentrations and lower ones around the equator, hence optimizing the taxonomic identification effort, while reducing very high abundances importance. Environmental variables were projected into the PCA space according to their correlation with plankton concentrations, after a log n+1 transformation for marine snow and bulk concentrations, to avoid over-representing some very high values. This allowed

to visualize correlations directly on the PCA biplot and help with axes interpretation. Each profile's scores on the first five principal components (PC) helped perform a synoptic hierarchical agglomerative clustering (HAC), using the Euclidean distance and Ward's criterion (Legendre & Legendre, 2012), in order to get a synoptic view of plankton communities. Using the PC scores, not the original data, preserved most of the variance, while removing noise. The resulting dendrogram (Figure S3) was separated into some main branches, based on inertia jumps, identifying broad plankton community types. Taxa proportions were computed for each plankton cluster.

Testing for potential diurnal and seasonal biases, we computed the variance portion in plankton community composition, explained by acquisition time or season. We used a redundancy analysis (RDA), with Hellinger-transformed concentrations as response variables, and a binary variable as explanatory, either day/night or productive/non-productive season, defined based on latitudes and sampling months (Table S2) (Lalli & Parsons, 1997). Diurnal effects were tested in both layers; while seasonality was not tested in the mesopelagic layer since seasonal changes are weaker at depth (Costello et al., 2018). In both cases, explained variance was computed as R^2 .

2.4 | Correspondence with ocean regionalizations

In the second step, we aimed to capture the link between plankton communities and their environment. To assess various regionalizations ability to capture processes driving plankton ecology, the part of variance in plankton community composition explained by each was computed. This was performed analogously to the circadian or seasonal effects test. In each layer, a RDA was performed. with Hellinger-transformed concentrations as response variables and a qualitative variable with regions from a given regionalization as explanatory variable. To ensure adequate representativeness, each region in each regionalization had to contain at least 25 profiles for inclusion. This limited the number of regions used for each regionalization, but resulted in similar region numbers throughout all tested regionalizations: all contained 13-18 regions for the epipelagic and 10-12 regions regarding the mesopelagic zone (Table 1). This is important for comparison as the variance portion explained by a categorical variable often increases with the number of modalities.

To quantify how much variance is explainable by a categorical variable with a similar cardinality, a maximal model was built by computing a regionalization on plankton concentrations themselves, with a similar groups number. We used the PCA on Hellinger-transformed concentration data, followed by a *k*-means clustering on the first five PCs. Here, the group's number was set as a middle ground between cardinalities of other regionalizations (epipelagic: 15; mesopelagic: 12). Note that in the epipelagic layer maximal model, the difference between 13 and 18 modalities is inconsequential: 13 modalities explain 55.7% of variance, while 18 modalities explain 59.2% (+3.5% of variance). A RDA was performed with this explanatory variable, and, since response and /ILEY

explanatory variables are built with the same data, this RDA captures the maximum part of explainable variance.

Other tested regionalizations included: Longhurst provinces (Longhurst, 2010), 10° latitudinal bands and mesopelagic provinces (Reygondeau et al., 2018) (tested for the mesopelagic layer only). Besides these regionalizations, often based on climatological averages, a regionalization based on each profile's immediate environment was generated with a PCA performed on environmental data in both layers. This PCA included layer wise data: CTD, UVP5 bulk and marine snow data; as well as data shared between layers: depth of clines and satellite data. The latter were also included in the mesopelagic PCA as phytoplankton present in the epipelagic layer can influence processes in the mesopelagic layer (Guidi et al., 2009; Hernández-León et al., 2020). Projections on the first 5 PCs were used to generate a kmeans clustering. For comparison purposes, the number of modalities was set to be similar to the other regionalizations (13 for the epipelagic, 10 for the mesopelagic). Corresponding PCAs and maps can be found in supplementary (Figures S4 and S5). Finally, we also compared these regionalizations to a null model: profiles were randomly grouped into a similar number of clusters. If the variance portion explained by a given regionalization is similar to the null model portion, this regionalization does not capture plankton community composition variations.

All analyses were conducted with R version 4.0.3 and the 'vegan' package version 2.5.7 (Oksanen et al., 2018).

3 | RESULTS

3.1 | Spatial distribution of plankton communities

Within the UVP5 size range ($600 \mu m$ to a few cm), more than 330,000 organisms were detected. Three groups dominated the number of individual images: Copepoda (metazoan), *Trichodesmium* (cyanobacteria) and Phaeodaria (Rhizaria subgroup, unicellular eukaryote) (Figure 2). Apart from Rhizaria, *Trichodesmium* and Nostocales (the latter both Cyanobacteria), all other imaged organisms belonged to the Metazoa kingdom.

3.1.1 | Epipelagic layer

To describe epipelagic plankton communities, 2517 profiles were included. The first two PCs captured 41.8% of variance. The first PC distinguished between Trichodesmium-rich communities and copepod-rich ones. Trichosdesmium-rich communities were associated with warm and stratified waters. The proportion of copepods was higher in cold, high chlorophyll and particle-rich waters (Figure 3a). On the second PC, a third pole emerged, represented by Collodaria (Rhizaria), extant in oligotrophic waters (with deep DCM and Z_{eu}). The HAC dendrogram was separated into 3 clusters (Figure S3). Cluster 1 was characterized by a co-dominance of multiple Rhizaria groups (Acantharia, Collodaria and Phaeodaria), copepods, nostocales and Trichodesmium (Figure 3c), although Collodaria emerged as the third structuring ones in PCA space. This community type was widely distributed in oceans but detected at lower frequencies in high latitudes (Figure 3b). Cluster 2, enriched in copepods, also had a widespread distribution but dominated the subpolar North Atlantic and Arctic shelf seas, and upwelling areas (California Current, Peruvian and Benguela upwellings). Cluster 3, enriched in Trichodesmium, existed in the Atlantic Ocean's intertropical band.

3.1.2 | Mesopelagic layer

In the mesopelagic layer, 1747 profiles described plankton communities. The first two PCs captured a 39.6% variance. The first PC separated copepod-rich waters from waters with Phaeodaria (Figure 4a). Copepod-enriched waters were oxygen-rich, while Phaeodariarich waters showed high AOU and significant stratification of the water column top. On the second PC, a third Eumalacostraca pole emerged, associated with warmer, saltier, more oligotrophic waters. The HAC dendrogram was again split into 3 clusters (Figure 53). The first cluster, with a higher proportion of phaeodarians and fewer copepods (Figure 4c), was emblematic of the Peruvian upwelling, but also present in Mediterranean Sea and Pacific Ocean profiles. The Copepoda-enriched cluster 2 existed at high northern hemisphere

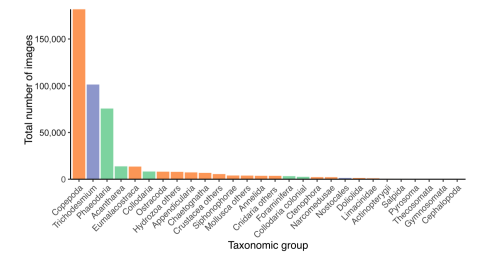
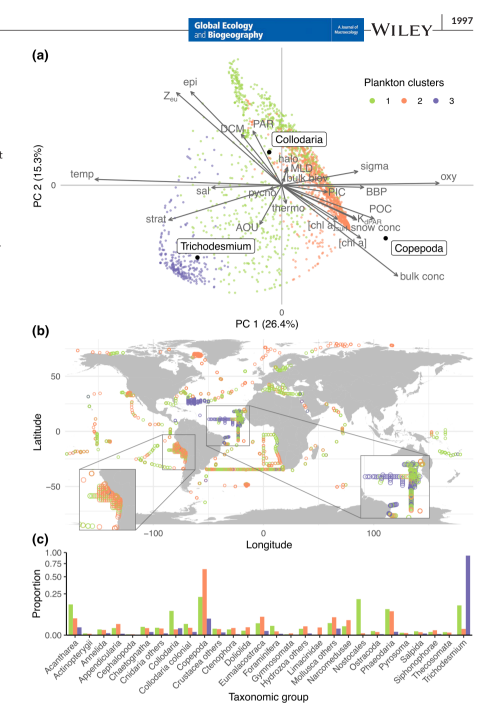


FIGURE 2 Dataset composition: total number of images per taxonomic group. Rhizaria (unicellular eukaryotes) are highlighted in green, Cyanobacteria are in purple and Metazoa in orange. FIGURE 3 Plankton clusters within the epipelagic layer. (a) PCA performed on Hellinger-transformed plankton concentrations, illustrated by a biplot in scaling 2. Only taxa with a contribution higher than average are represented. Environment variables are projected as supplementary variables. Points represent profiles and are coloured according to the cluster defined by the HAC. (b) Map of epipelagic profiles, coloured as in A. (c) Relative composition of epipelagic plankton clusters, coloured as in A. Note that the Y axis is square root transformed.



latitudes; also present in the intertropical band and south of the Atlantic (Figure 4b). Finally, cluster 3 was not dominated by any taxon but had a diversified composition: Copepoda, Eumalacostraca, Foraminifera, Phaeodaria and *Trichodesmium*. Fewer profiles were in this cluster but broadly distributed.

3.1.3 | Comparison between epipelagic and mesopelagic layers

We compared each station's plankton community type in epipelagic and mesopelagic layers (Figure 5). Some stations, included in the epipelagic analysis, had no corresponding mesopelagic samples as the sea bottom or UVP maximum depth was too shallow. These stations are "NA" in Figure 5 mesopelagic row. Among the 1122 profiles where the epipelagic part contained a mixed-type community (cluster 1 epipelagic), the mesopelagic layer contained a phaeodarian-enriched community (cluster 1 mesopelagic) in 49% of cases and a copepod-enriched community (cluster 2 mesopelagic) in 33% of cases. In the copepod-rich epipelagic community (cluster 2 epipelagic), most profiles displayed a mesopelagic copepod-enriched community (52%, cluster 2 mesopelagic). Finally, most profiles with a *Trichodesmium*-enriched epipelagic layer (cluster 3 epipelagic) had a phaeodarian-enriched mesopelagic community (75%, cluster 2 mesopelagic). Overall, this analysis highlights only an incomplete similarity between epipelagic and mesopelagic plankton communities' compositions, suggesting they are driven by different processes disjoining the surface and deep communities.

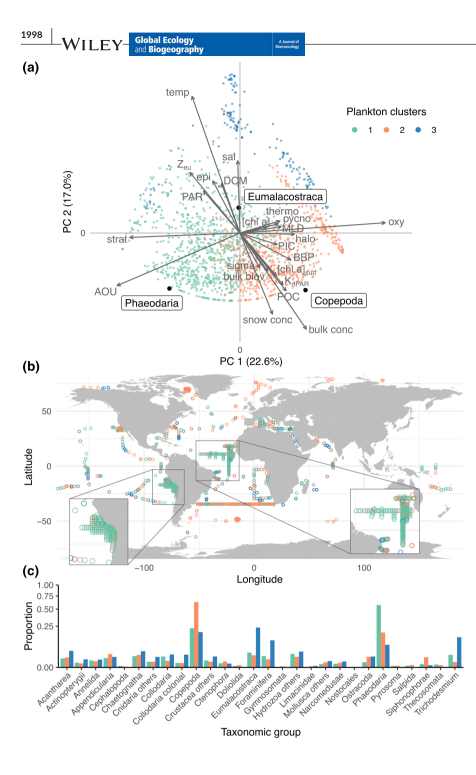


FIGURE 4 Plankton clusters within the mesopelagic layer. (a) PCA performed on Hellinger-transformed plankton concentrations, illustrated by a biplot in scaling 2. Only taxa with a contribution higher than average are represented. Environment variables are projected as supplementary variables. Points represent profiles and are coloured according to the cluster defined by the HAC. (b) Map of mesopelagic profiles, coloured as in A. (c) Relative composition of mesopelagic plankton clusters, coloured as in A. Note that the Y axis is square root transformed.

3.2 | Representativity of existing ocean regionalizations

In both layers, the clustering built on the plankton data itself (maximal model) explained about half of the variance in community composition (epipelagic: 56.7%; mesopelagic: 46.7%; Table 1). The limited amount of explained variance unsurprisingly confirms plankton communities' diversity cannot be summarized in only 12–15 groups, but also highlights that other regionalizations should be gauged relatively to these figures, not to 100% of variance explained. All regionalizations explained more variance than the null model (<1% of variance explained in both layers).

Thus, all regionalizations were relevant for explaining plankton communities' distribution. Among the tested regionalizations, Longhurst provinces explained more variance than others (epipelagic: 26.0%; mesopelagic: 13.2%), corresponding to about half of the explainable variance in the epipelagic layer and 1/3 in the mesopelagic zone. In the epipelagic, latitudinal bands explained similar variance to the local environment; in the mesopelagic, Reygondeau's mesopelagic provinces explained some more variance than the local environment and latitudinal bands. In both layers, plankton community composition was better explained by biogeochemical-based provinces than by the immediate and local environment the plankton was sampled in.

288

n = 775

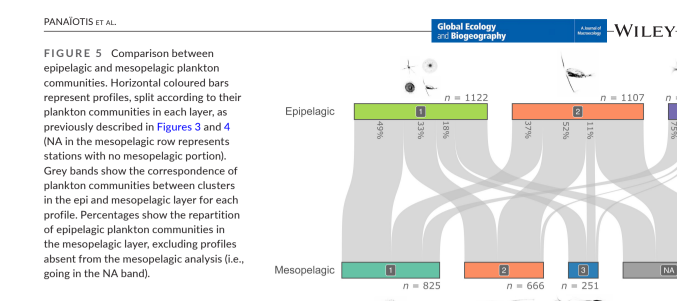


TABLE 1 Variance in community composition explained by different regionalizations for the epipelagic and mesopelagic layers. 2203 and 1193 profiles were included in the epipelagic and mesopelagic layers respectively.

	Epipelagic		Mesopelagic	
Regionalization	n	R ²	n	R ²
Maximal model	15	0.567	12	0.467
Null model	15	0.007	11	0.007
Longhurst provinces	18	0.260	12	0.132
Latitude bands	13	0.173	11	0.104
Local environment	13	0.168	10	0.092
Mesopelagic provinces	-	-	11	0.116

Abbreviation: *n*, number of groups for each regionalization.

4 | DISCUSSION

In summary, three meso/macro plankton communities' types were detected in both epipelagic and mesopelagic layers. Their composition was better explained by basin-scale environmental conditions than by local ones. Below, we first briefly discuss methodological aspects to assess our results' robustness (these are discussed in more detail in Appendix S1) and then discuss our findings' consequences in the existing knowledge context.

4.1 | Potential biases

Plankton concentrations vary temporally, with seasons, but also between day and night because of the diel vertical migrations (DVM) (Lampert, 1989), conducted by many – mostly pluricellular – plankton taxa. Here, both diel and seasonal effects explained very little variance (Table S3), showing that they barely impact community composition and seemed negligible compared to spatial effects, although DVM was detected on raw concentrations (Figure S6). Although UVP5 profiles were distributed unevenly in space (Figure S1) and in time (Figure S7), our analyses were robust to downsampling, showing that our results were not solely representative of oversampled areas (see procedure description in Appendix S1). Moreover, UVP5 samples were shown to cover diverse enough environmental conditions, representative of worldwide oceans (Figures S8 and S9 and text in Appendix S1).

Finally, the UVP5's detection capabilities are limited, both in terms of size (>600 μ m for detection and >1 mm for quantitative measurements) and taxonomy (only broad taxa can be identified). Thus, if poorly taxonomically resolved taxa (e.g. Copepoda, Eumalacostraca) could have been better resolved, the resulting lower level taxa might have emerged as structuring mesoplankton communities. Similarly, it cannot be excluded that the detection of smaller organisms could have revealed further structuring taxa (e.g. diatoms at high latitudes, Dutkiewicz et al., 2020). Indeed, smaller plankton are typically more abundant than larger organisms. These elements could also impact our findings regarding how the distribution of plankton communities can be explained by various regionalizations: assuming a strong geographic signature, this could have increased the ability of one given regionalization in explaining the distribution of plankton communities.

4.2 | Plankton communities general structure

4.2.1 | Epipelagic layer

Three plankton community types emerged in the epipelagic layer, mostly driven by water masses' temperature and trophic statuses: copepod-enriched communities in cold and productive waters; *Trichodesmium*-enriched communities in warm waters; and mixedtype communities in oligotrophic ones.

Copepods' dominance in almost all pelagic ecosystems is already documented (Brandão et al., 2021; Ibarbalz et al., 2019; Rombouts

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et al., 2009; Soviadan et al., 2022), more so in the rich and productive Arctic waters (Brandão et al., 2021; Drago et al., 2022; Forest et al., 2008; Soviadan et al., 2022; Trudnowska et al., 2020). Similarly to other zooplankton, copepods' diversity decreases with latitude (Brandão et al., 2021; Ibarbalz et al., 2019) and conversely for size (Brun et al., 2016; Horne et al., 2016). Brandão et al., 2021 found a 400-500 µm median ESD for copepods between 60°N and 60°S. As UVP5 only detects copepods over 1 mm in ESD (Forest et al., 2012), tropical and temperate copepods' concentrations were likely underestimated but in various situations, conclusions on community composition are little affected, since all taxa concentrations are similarly underestimated near the UVP target size range limits. Large changes in the biogeography, community composition and diversity of calanoid copepods in the North Atlantic Ocean are detected, as a result of global warming (Beaugrand et al., 2009; Beaugrand et al., 2010). As copepods act as a trophic link between primary producers and higher trophic levels (Rombouts et al., 2009), these changes might prove detrimental to marine resources, like exploited fish stocks (Beaugrand et al., 2010). We show that, within the 600μ m range to a few cm, copepods are proportionally more abundant in polar waters and, to a lesser extent, in temperate ones, but clearly not in tropical ones.

Epipelagic plankton communities were also shaped by Trichodesmium, a filamentous cyanobacteria found in subtropical and tropical regions (Capone et al., 1997; Westberry & Siegel, 2006) and previously observed with UVP5 in tropical waters (Guidi et al., 2012; Sandel et al., 2015). Given that phytoplanktonic organisms are rather uncommon in the size range observed by the UVP5 (>600 µm) compared to zooplanktonic organisms, the role of Trichodesmium as structuring mesoplankton communities was quite unexpected. Trichodesmium contribute to primary production and fixation of atmospheric nitrogen (Capone et al., 1997) and can grow in nitrogen-limited environments, unlike other phytoplankton types (Westberry & Siegel, 2006). Their toxicity to several zooplankton species (Hawser et al., 1992) might contribute to the guasi-exclusion of other types of planktonic organisms in Trichodesmium-enriched communities, provided that both Trichodesmium and other organisms concentrations are sufficiently high to cause encounters.

Finally, the third cluster revealed a mixed plankton community, characterized by copepods' or Trichodesmium's non-dominance. This cluster contained profiles from diverse environments with varying conditions, thus very diversified composition-wise. Its average composition (Figure 3) is skewed by high concentrations from a few profiles, therefore not representative of every profile's composition aggregated in this cluster. Although Collodaria emerged as a structuring group in the PCA, it did not dominate the cluster's relative composition. Actually, six groups were found, accounting for 85% of the composition: Acantharia (Rhizaria), Collodaria (Rhizaria), Copepoda, Nostocales, Phaeodaria (Rhizaria) and Trichodesmium, highlighting the importance of various rhizarian groups, recently highlighted by other in situ imaging-based studies (Biard et al., 2016; Dennett et al., 2002). Acantharia and Collodaria are symbiontsbearing Rhizaria, widely distributed in oceans, but more abundant in tropical oligotrophic surface waters, where their symbionts are

photosynthetically active (Suzuki & Not, 2015), hence coherent with our observations. Collodaria contributes significantly to the total organic matter in these environments (Biard et al., 2016; Suzuki & Not, 2015). Conversely, Phaeodaria are heterotrophic Rhizaria, lacking symbionts (Kling & Boltovskoy, 1999), flux-feeders and thus usually found below the epipelagic layer, where they feed on sinking particles (Biard & Ohman, 2020). However, some species are common in surface layers (Nakamura & Suzuki, 2015). Because Phaeodaria's mineral skeleton is made of silica, they can act as major biogenic silica exporters (Biard et al., 2018) and their distribution could be restricted by silica availability (Biard & Ohman, 2020; Nakamura & Suzuki, 2015). Finally, nostocales (Aphanizomenon, Dolichospermum, and Nodularia) were identified contextually (i.e., not just based on aspect on images), only in the Baltic Sea, and at very high concentrations. As a consequence, they account for 19% of the mixed-type community, though they were found in only a few profiles.

4.2.2 | Mesopelagic layer

In the mesopelagic layer, three types of plankton communities also emerged: copepod-enriched in cold and oxygenated waters, Phaeodaria-enriched in cold and oxygen-depleted waters, and, again, a mixed community in warmer waters.

In the latter, four groups accounted for 65% of organisms: Copepoda, Eumalacostraca, Foraminifera, and *Trichodesmium*. Foraminifera, heterotrophic rhizarians, feed on mesopelagic plankton and are typical of deep and rather poorly oxygenated waters (Biard & Ohman, 2020). Various organisms were identified within the broad Eumalacostraca group in our dataset, depending on the ecosystem sampled, resulting in a very heterogeneous group. Thus, Eumalacostraca are not representative of any typical environment, restricting ecological interpretations, and most likely under-sampled.

In cold waters, the plankton community type was linked to oxygen availability: a higher proportion of copepods was found when oxygen was available and Phaeodaria in water masses, with high AOU and low oxygen concentration, typical of Oxygen Minimum Zones (OMZ). Major OMZs are found below eastern-boundary upwelling systems, with particularly high primary production. Many planktonic taxa' concentrations, such as calanoid copepods', is reduced in OMZs (Auel & Verheye, 2007; Hoving et al., 2020; Kiko et al., 2020; Soviadan et al., 2022). Associated with the low oxygen and low plankton concentrations, and possibly caused by it, the downward flux of particulate carbon is intense and less attenuated than elsewhere, resulting in high vertical export of the organic matter photosynthesis produces at the surface (Cavan et al., 2017; Engel et al., 2022). Conversely, Phaeodaria are typical of deep, low oxygenated waters (Biard & Ohman, 2020; Ikenoue et al., 2019) and already detected in OMZs (Hoving et al., 2020). Indeed, protists might prove more tolerant to hypoxia, as their passive feeding mode requires less oxygen than active feeding. In OMZs, they may therefore play a disproportionate role in the regulation of the vertical flux compared to elsewhere.

Although unexpected at first glance, the detection of *Trichodesmium* in the mesopelagic layer is consistent with previous observations (Benavides et al., 2022; Sellner, 1992; Walsby, 1978). Here, their presence in the mesopelagic layer could be partly explained by our dynamic definition of the epi-mesopelagic boundary: it started at shallower than 50m for 25% of profiles, a depth where the presence of *Trichodesmium* would not be surprising. However, those were found mostly at higher latitudes (Figure S2), where *Trichodesmium* is absent. Its presence at a great depth could also result from downwelling and subduction events, bringing colonies to deeper waters (Guidi et al., 2012), or even simply represent dead colonies sinking down.

4.2.3 | Comparison between epipelagic and mesopelagic plankton communities

Conditions in the epipelagic layer constrain those in the mesopelagic: i.e., the epipelagic phytoplankton type influences particle sizes in the mesopelagic (Guidi et al., 2009). Similarly, mesopelagic zooplankton biomass is conditioned by the net primary production in the euphotic layer, since it feeds on its remnants (Hernández-León et al., 2020). However, the results above show a low similarity of plankton communities between epipelagic and mesopelagic layers. Such a result is to be linked to the functional definition of the epimesopelagic boundary that was used, based on hydrological characteristics, so that different water masses and their potential inherent plankton community are not mixed together.

Where the plankton community is enriched in copepods and the environment productive in the epipelagic layer, the mesopelagic plankton community is usually copepod-rich as well. This is consistent with high secondary production in the mesopelagic, below subpolar surface waters hosting high primary production, which is not true at subtropical latitudes (Robinson et al., 2010). Below oligotrophic Rhizaria-rich epipelagic communities, the mesopelagic community was mostly Phaeodaria-enriched in the eastern tropical South Pacific OMZs but was copepod-enriched in the South Atlantic. The split between these two mesopelagic communities therefore seems to be driven by copepods' oxygen limitation in OMZs (Engel et al., 2022). In the South Atlantic gyre, both epipelagic (Signorini et al., 2015) and mesopelagic (Sutton et al., 2017) layers are considered as oligotrophic, consistent with a Rhizaria-enriched community in the epipelagic (Suzuki & Not, 2015). In the Peruvian upwelling system, the epipelagic community was mostly Rhizaria-enriched; surprising, since this environment is very productive (Ayón et al., 2008). Still, the presence of Rhizaria was already reported there by Santander Bueno, 1981, within a diverse zooplankton community over the region. This productive upwelling area drives an OMZ in deeper waters (Engel et al., 2022; Kiko & Hauss, 2019), which imposes a Phaeodaria-enriched community in the mesopelagic. Within Trichodesmium-rich stations, only equatorial Atlantic stations were sampled deep enough to be included in the epi/mesopelagic

The forcing environmental conditions associated with communities in both layers are not the same: oxygen plays a role at depth but less near the surface, while light structures life near the surface but not in the mesopelagic layer. Besides, the conditions that remain structuring ones are less variable with increasing depth, leading to a more homogeneous habitat with depth (Costello et al., 2018; Costello & Breyer, 2017). This also shapes the plankton community, which becomes less spatially contrasted in the mesopelagic (Soviadan et al., 2022).

4.3 | Plankton communities distribution was driven by regional conditions

Among the regionalizations tested, the plankton communities distribution was better explained by Longhurst provinces (Longhurst, 2010), a regionalization based on physical forcings as drivers of phytoplankton distribution, which might drive zooplankton communities too. Although plankton diversity and biomass follows a latitudinal gradient (Ikeda, 1985; Rombouts et al., 2009), mediated by joint effects of light availability and temperature, the regionalization based on 10° latitudinal bands explained less variance than Longhurst provinces. In the mesopelagic, the regionalization computed by Reygondeau et al., 2018 - specifically for this layer and mainly based on annual climatologies of biogeochemical variables - did not explain plankton community distribution better than Longhurst provinces, even though it was supposed to be more appropriate. Yet, more importantly, all these basin-scale regionalizations explained plankton community distribution as well, or better, than a regionalization based on local conditions, at sampling time. This suggests that plankton communities' spatial structure for our 28 taxonomic groups is driven by regional environmental conditions more than by very local and immediate conditions and processes. These findings agree with those of Stemmann et al., 2008, who showed that mesopelagic macro-zooplankton communities were structured by large, basin-scale processes. Genomic analyses also underlined regional scales processes' importance in structuring plankton communities (Richter et al., 2020), especially for mesozooplankton (Sommeria-Klein et al., 2021).

Among the many other existing regionalizations of the oceans, the one proposed by Hofmann Elizondo et al., 2021 and based on phytoplankton biogeography would have constituted an interesting point of comparison to assess how it correlates with mesoplankton communities. However, such comparison was not possible because of the low cardinality of this regionalization. Still, Hofmann Elizondo et al., 2021's tropical biome matches well our *Trichodesmium*enriched community in the Atlantic ocean, while the high latitude biome coincides well with communities rich in copepods.

5 | CONCLUSION AND PERSPECTIVES

In both layers, three plankton community types emerged in the observed size range (> 600μ m) and seemed mostly driven by basinlevel environmental conditions. Following on studies investigating plankton distribution and diversity across life kingdoms – from viruses to metazoans (de Vargas et al., 2015; Ibarbalz et al., 2019; Sunagawa et al., 2015) – this work highlights the role not only of metazoans, but also of unexpected large protists and cyanobacteria in structuring meso and megaplankton communities. This confirms underwater imaging relevance to reveal the importance of otherwise overlooked plankton groups, such as Rhizaria (Biard et al., 2016; Dennett et al., 2002).

The development of regionalizations – either based on biotic or abiotic data – for wide-ranging offshore areas is highly desired for conservation purposes, like the creation of protected marine areas. However, they should not be restricted to oceans' upper layers. Indeed, biological activity in the mesopelagic layer is key to mediate the flux of organic carbon from the surface and the deep seafloor is also heavily impacted by human activities (bottom fishing, seafloor mining...) (Watling et al., 2013). Therefore, oceans' deeper layer biogeographies, currently rare, are also required to balance between human exploitation and ecological conservation.

AUTHOR CONTRIBUTIONS

Thelma Panaïotis, Jean-Olivier Irisson and Lars Stemmann conceived the study; Marcel Babin, Tristan Biard, François Carlotti, Laurent Coppola, Lionel Guidi, Helena Hauss, Lee Karp-Boss, Rainer Kiko, Fabien Lombard, Andrew M. P. McDonnell, Marc Picheral, Andreas Rogge, Anya M Waite and Lars Stemmann contributed to data acquisition and image validation; Thelma Panaïotis conducted the analyses under the supervision of Jean-Olivier Irisson and Lars Stemmann; Thelma Panaïotis wrote the manuscript; all authors contributed to the discussion of the results, supported manuscript preparation and approved the final submitted manuscript.

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CONFLICT OF INTEREST STATEMENT

The authors declare no conflicts of interest.

DATA AVAILABILITY STATEMENT

The code supporting this study is available at: https://github.com/ ThelmaPana/planktyp. The dataset is made available on SEANOE: 10.17882/95809.

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BIOSKETCH

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SUPPORTING INFORMATION

Additional supporting information can be found online in the Supporting Information section at the end of this article.

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