Uncovering the Composition and Diversity of Pelagic Copepods in the Oligotrophic Blue Water of the South Pacific Subtropical Gyre

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The subtropical gyres occupy approximately 40% of the surface of the Earth and are widely recognized as oligotrophic zones. Among them, the South Pacific subtropical gyre (SPSG) shows the lowest chlorophyll-a levels (0.02–0.04 µgL−1), the deepest nutricline (>200 m) and euphotic zone (∼160 m), and the lowest rates of nitrogen fixation. The zooplankton community is poorly known in the SPSG. We report a study focused on the composition and distribution of pelagic copepods within the gyre so as to uncover the diversity and habitat conditions of this special community. Therefore, during the austral spring of 2015, an oceanographic cruise was conducted across the eastern side of the SPSG. Physical and chemical variables were measured in the upper 1000 m, while zooplankton samples were collected by means of vertically stratified hauls using a multiple net sampler for five layers (0–800 m). Satellite data were also used to assess near-surface phytoplankton biomass (Chl-a) and physical-dynamics conditions during the cruise, and 121 species of copepods were identified, which belonged to five taxonomic orders, 24 families, and 50 genera. Calanoida and Cyclopoida were the most frequent orders, containing 57% and 38% of species, respectively, whereas Harpacticoida and Mormonilloida contained 2% of species each, and Siphonostomatoida contained 1% of species. The vertical distribution of copepods revealed an ecological zonation linked to a strongly stratified water column, such that three different vertical habitats were defined: shallow (0–200 m), intermediate (200–400 m), and deep (400–800 m). Both the abundance and diversity of copepods were greater in the shallow habitat and were strongly associated with water temperature, whereas copepods in the subsurface layers subsisted with relatively low oxygen waters (2–3 mL O2 L−1) and presumably originated at the Chilean upwelling zone, being transported offshore by mesoscale eddies. Furthermore, the analysis of species composition revealed a marked dominance of small-sized copepods, which may play a key role in nutrient recycling under an oligotrophic condition, as inferred from their
mostly omnivorous feeding behavior. Our findings also suggested a potentially high endemism within the gyre, although basin-scale circulation and mesoscale eddies, traveling from the coastal upwelling zone and transporting plankton, can also influence the epipelagic fauna.

**INTRODUCTION**

The subtropical gyres are large marine areas moving circularly and are centered in major basins of the world oceans. They form upon the effect of planetary vorticity, the friction with the continents, and the effect of the Coriolis force (Brown et al., 2001; Siedler et al., 2001). These extensive areas may occupy approximately 40% of the surface of the Earth and are characterized by low chlorophyll-a (Chl-a) and extremely reduced biological production; they are widely recognized as ultra-oligotrophic zones (Raimbault and García, 2008; Morel et al., 2010), although their contribution to the global carbon cycle may be significant (Signorini and McClain, 2012). In the North Pacific anticyclonic gyres (ALOHA-Hawaii Station), Sargasso Sea (Bermuda Atlantic Time-Series Study – BATS) and the Atlantic Ocean (Southern Atlantic Transect – AMT), the role of zooplankton in biogeochemical cycles and carbon transport to the deep ocean has been studied with the aim of characterizing and understanding their influence on regional and global climate, as well as to develop the fundamental processes and mechanisms governing their ecology in the ocean (Calbet and Landry, 1999; Schnetzer and Steinberg, 2002; Steinberg et al., 2008; Rees et al., 2015; Goetze, 2017). In contrast, in the South-eastern Pacific gyre (SPSG), the state of knowledge on zooplankton ecology is still incipient, and there are no time series studies as in the North Pacific gyre.

In the SPSG, the circulation system is comprised of the westward South Equatorial Current, which can enhance the levels of chlorophyll-a (Chl-a) due to the equatorial divergence, a narrow poleward western boundary current, the East Australian Current, which has a 29°C surface isotherm and is also known as the warm pool, the eastward South Pacific Current streaming along the South Tropical Front (Subtropical Convergence Zone), and the Humboldt Current System, a broad equatorward eastern boundary current that presents a strong wind-driven upwelling and a very active eddy field (Longhurst, 2007; Schneider et al., 2007). The SPSG is considered as the water mass with the lowest Chl-a concentration (0.02–0.04 µg L⁻¹ in the austral summer) observed in the global ocean. Moreover, this water mass shows the deepest nutricline (>200 m) and euphotic zone (∼160 m) (Bender and Jönsson, 2016). This ultra-oligotrophic condition (Morel et al., 2010) causes the presence of the clearest blue waters of the world ocean. The satellite seasonal Chl-a cycle shows low values in the austral summer and higher values in winter. This seasonal variability is mostly driven by the warming/cooling of surface waters, which involves the mixing of the water column that carries nutrients to the illuminated layer (upper 200 m) in winter (Signorini and McClain, 2012). This stimulates primary production, such as that of phytoplankton, which can occur at low concentrations. In fact, phytoplankton can be found at variable depths with a Chl-a maximum near 180 m depth.

Likewise, only a few studies refer to metazoan zooplankton in the SPSG. Some reports have revealed extremely low mesozooplankton biomass, with a biovolume of ca., and 10% of that were found in the upwelling zones (e.g., González et al., 2019), although in some areas surrounding oceanic islands, such as the Easter Island and Salas and Gómez islands, slight increases in zooplankton biomass have been noted (Palma and Silva, 2006). A few taxonomic descriptions of zooplankton groups have been provided, suggesting that pelagic copepods may be the dominant group (Fisher, 1958). Studies in other groups have received more attention, such as tintinnids, siphonophores, amphipods, and euphausiids (Brinton, 1962; Vinogradov, 1991; Palma, 1999; Robledo and Mujica, 1999; Rojas et al., 2004; Palma and Silva, 2006; Dolan et al., 2007).

In the zooplankton, copepods are the most abundant group (numerically 70–90%). They have a key role in the marine ecosystem as they feed on primary producers and channel C to higher trophic levels. They also contribute to the export of organic carbon toward the deep ocean, thereby providing food for benthic organisms and excreting nutrients that can be recycled by phytoplankton (Harris et al., 2000; Valdés et al., 2018b). These ecological roles of copepods depend considerably on their diversity and distributional patterns (Valdés et al., 2018a; Tutasi and Escribano, 2020). In fact, the diversity patterns are essential to understand the organization and functioning of the organisms present in an ecosystem and their interactions with the environment (Duarte, 2000). However, the understanding of factors explaining the biodiversity patterns of zooplankton remains an open issue (Angel, 2003), and the existing hypotheses are related to timing, spatial heterogeneity, biological interactions, climatic stability, and productivity (Pianka, 1966; Rohde, 1992; Palmer, 1994; Currie et al., 1999). However, more recently, temperature and its spatial gradients have been shown to strongly correlate with zooplankton diversity (González et al., 2020b). In the same context, and with respect to vertical distribution, recent studies suggest that the number of species and abundances of zooplankton tend to decrease with depth, and this decrease appears to be non-linear, with an increase in zooplankton at mid-depths and a decrease at greater depths (Bucklin et al., 2010). Furthermore, several works state that the vertical distribution of zooplankton is determined by prevailing water masses (Longhurst, 1985b; Pages et al., 2001; Hopcroft et al., 2010; Eisner et al., 2013), although other factors affect this as well, such as the diel vertical migration behavior induced by light conditions (Reid et al., 2003; McClellan et al., 2014; Tutasi and Escribano, 2020). This behavior can also occur...
in deep waters where light does not reach, and migration in the dark may be influenced by biochemical factors (Kampa, 1970; Van Haren and Compton, 2013).

Understanding the factors and processes controlling the diversity and ecological role of copepods in the SPSG requires basic information, which is clearly not available. To date, only two species of calanoids have been registered in the SPSG: Mesocalanus lighti and M. tenuicornis (Von Dassow and Collado-Fabbi, 2014). The lack of knowledge regarding this important group may be critical in assessing their biological role for the cycling of C and N in this presumably extremely low-productive but large ecosystem of the South Pacific. Copepods, through their feeding and excretion, are important recyclers of N in both highly productive (Valdés et al., 2018b) and oligotrophic systems (Valdés et al., 2018a); therefore, they affect the functioning of the biological pump and the downward flux of C (Tutasi and Escribano, 2020). Moreover, they are the main source of food for larvae, juvenile and adult stages of pelagic fishes, thereby playing a key role in the ocean food web (Steinberg and Landry, 2017). Elucidating such ecological functions in the SPSG requires knowledge on the composition, abundance, and distribution of their species. Therefore, in this study, we present the first description of copepods in the SPSG and their vertical distribution patterns in the epipelagic and upper mesopelagic depths (0–800 m). Based on the existing knowledge regarding the controlling processes of zooplankton distribution in the ocean, we hypothesize that spatial and vertical distribution of copepods and their diversity in the SPSG are associated with the prevailing water masses in this poorly known ecosystem.

**MATERIALS AND METHODS**

**Field Work**

The expedition CIMAR 21-Oceanic Islands was conducted from October to November 2015 onboard the R/V Cabo de Hornos and covered a large area (>4000 km) in the South-east Pacific basin of Chile, between Caldera at the Chilean coast (27°S and 72°W) and Easter Island (27°S and 110°W). For the present study, five stations, located within the SPSG (98°W – 107°W), were selected to assess the copepod community (Figure 1). At each station, measurements of physical and chemical variables were obtained through vertical profiles of a CTD (conductivity-temperature-depth-oxygen) SeaBird SBE 911Plus deployed down to 1000 m. Mesozooplankton samples were collected using vertically stratified hauls with an electronic Multinet Hydrobios, midi type with 0.25 m m mesh-size. Five depth layers were sampled (0–100, 100–200, 200–400, 400–600, and 600–800 m). Single tows of the Multinet were done at each station, resulting in 3 daylight stations and two nighttime stations, depending on the ship itinerary (Supplementary Table 1). The Multinet was equipped with a digital flowmeter to estimate the volume of filtered water. All samples were preserved in 10% buffered formalin. In the laboratory, all copepods were sorted, identified, and counted under stereoscope and compound microscopes. Species identification was based on specialized bibliography and databases (Nishida, 1985; Campos-Hernández and Suárez-Morales, 1994; Bradford-Grieve, 1994; Bradford-Grieve et al., 1999; Boxshall and Halsey, 2004; Razouls et al., 2005/2019). Only adult male and female copepods were identified; therefore, the presence of the early stages of copepods was not considered in this study.

**Analysis of Oceanographic Data**

In order to characterize the near surface bio-physical features, delayed-time sea level anomaly (SLA) from the satellite data was used to represent the geostrophic velocity field for the cruise period (October to November 2015). These data were obtained from the Salto/Duacs AVISO altimetry product1 for 0.25° × 0.25° grids of the central Chile region (20°S–30°S, 98–110°W). Monthly satellite-derived total Chl-a for the study region and for the same period was obtained from version 3.0 of the Ocean Colour Climate Change Initiative (OC-CCI), at processing level 3 and a spatial resolution of 4 km. In addition, the total Chl-a data for each sampling station and for the five depth layers (0–100, 100–200, 200–400, 400–600, and 600–800 m) were obtained from a global biogeochemical reanalysis product from the Copernicus Marine Environment Monitoring Service (CMEMS; GLOBAL_REALANALYSIS_BIO_001_0293).

Field data with regard to temperature, salinity, and dissolved oxygen from the oceanographic cruise were processed with the Gibbs Sea Water (GSW) Oceanographic Toolbox of TEOS-10 (IOC et al., 2010). The vertical profiles of these oceanographic variables were made with the Ocean Data View program. Conservative temperature and absolute salinity were derived from the CTDO, and T-S diagrams were made to identify the water masses. Finally, with the Optimum Multi-parameter (OMP) classical analysis in MATLAB software, we obtained the percentage of contribution of each water mass in the study area (Karstensen and Tomczak, 1999; Karstensen, 2020). Therefore, we used the interactive mode and characteristic water types (Table 1). To improve the water mass selection, the depth range was restricted to the following layers: 0–500, 300–700, and 400–1000 m. The analysis was carried out for each range of depth, and the type values of the water masses were assigned. Subsequently, three ranges were joined and overlapped on purpose to find the scope of each mass. The data were extracted and plotted in the Ocean Data View Program version 5.2.1.

Descriptive statistics (means and range of variations) was obtained for oceanographic data. We grouped oceanographic data according to longitude and depth by using the ordination method of principal coordinates analysis (PCO), based on Euclidean distance with standardized data using PRIMER7 and the PERMANOVA+ add on. The PCO analysis placed the samples onto Euclidean axes using a matrix of inter-points dissimilarities, calculating the distances between centroids for the levels of a selected factor (Anderson et al., 2008). After using the PERMANOVA+ routine, the statistical differences of the

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1 http://www.aviso.altimetry.fr
2 http://www.oceancolour.org/
3 http://marine.copernicus.eu
environmental variables between depth layers (i.e., stratum and water mass) and longitude were tested.

**Biological Data**

**Composition, Abundance, and Diversity**

The composition of species was compared with the global database of the diversity and geographic distribution of marine planktonic copepods in the South Pacific Ocean. In particular, it was compared with the areas associated with the SPSG, such as the Central Tropical Pacific (Zone 19), the waters in front of Chile (Zone 26), and the waters around Australia (Zone 18), following the areas defined in the copepods database of Razouls et al. (2005/2019). To assign a few ecological attributes of the species present in the systematic list, horizontal distribution in the South Pacific Ocean (East and West coast and open ocean), vertical position in the column water (epipelagic, mesopelagic and bathypelagic), and trophic behavior (omnivorous, carnivorous, herbivorous) were extracted from this database, which complemented the data from the literature.

The abundance (N), species richness (S), and diversity indices of Shannon-Wiener (H’) and Simpson (1- λ’) were analyzed using descriptive statistics (mean, ranges and variation).

The indices used the following equations:

\[
H' = \sum_{i=1}^{s} p_i \log_2 p_i
\]

\[
1 - \lambda' = 1 - \sum_{i=1}^{s} \frac{n_i (n_i - 1)}{N(N - 1)}
\]

Where \(H'\) = Shannon diversity index, \(p_i\) = the proportion of each \(i\) species, and \(s\) = total number of species. \(1 - \lambda'\) = Simpson diversity index, \(n_i\) = the number of individuals of species \(i\), and \(N\) = total number of species.

The Shannon-Wiener index ranges between 1 and 5, and the Simpson index ranges between 0 and 1. In this study, we decided to use both indices. Although the first index is more common in ecological studies, it is much more dependent on

**TABLE 1** | Source water types used for the basic OMP analysis.

| Water mass | Temperature (°C) | Salinity | Oxygen (µmol kg⁻¹) | References |
|------------|-----------------|----------|-------------------|------------|
| STW        | 20.80           | 35.52    | 240.65            | Schneider et al., 2003; Silva et al., 2009; Llanillo et al., 2012; Llanillo et al., 2013; Vergara et al., 2016 |
| ESPW       | 12.50           | 34.21    | 199.86            | Schneider et al., 2003 |
| ESSW       | 10.00           | 34.80    | 13.60             | Schneider et al., 2003; Silva et al., 2009; Llanillo et al., 2012; Llanillo et al., 2013; Vergara et al., 2016 |
| AAIW       | 3.00            | 34.20    | 156.00            | Schneider et al., 2003; Silva et al., 2009; Llanillo et al., 2012; Llanillo et al., 2013; Vergara et al., 2016 |

Subtropical Water (STW), Eastern South Pacific Intermediate Water (ESPW), Equatorial Subsurface Water (ESSW), Antarctic Intermediate Water (AAIW).
The taxonomic path length of branch between i and j species (Clarke and Warwick, 1998). The average taxonomic distinctness ($\Delta^+$) and variation in taxonomic distinctness ($\Lambda^+$). These indices show an average and an expected deviation for the study zone, when compared with the inventory of 546 species registered for the Central Tropical Pacific (Zone 19) by Razouls et al. (2005/2019). With the list of found species, a taxonomic aggregation matrix was made at the class, order, family, genus, and species levels. Thereafter, in the PRIMER7 program, with the TAXDTEST routine, the aggregation matrix was organized, and a classification tree was built from a random selection ($n = 1000$) of species from the general species inventory. Thus, comparing the indices and establishing confidence were limited to 95%.

The average ($\bar{3}$) and variation ($\bar{4}$) in taxonomic diversity are defined according the following equations:

$$\Delta^+ = \frac{2}{N(N-1)} \sum_{i} \sum_{j < i} w_{ij}$$

(3)

$$\Lambda^+ = \frac{2}{N(N-1)} \sum_{i} \sum_{j < i} (\omega_{ij} - \Delta^+)^2$$

(4)

Where the double summation is applied over all species $i, j$, $N$ is the species number in the sample, and $\omega_{ij}$ is the assigned taxonomic path length of branch between $i$ and $j$ species (Clarke and Warwick, 1998). $\Delta^+$ is the variance of the taxonomic distances ($\omega_{ij}$) between each pair of species $i$ and $j$ from their mean value ($\Delta^+$).

**Copepods Community Analysis**

Multivariate analyses of the copepod community were conducted with PRIMER7 and PERMANOVA+ add on software (Anderson et al., 2008). The data were transformed with log $x + 1$ according to the Power Law of Taylor for aggregated data (Taylor, 1961). Thereafter, a similarity matrix for multivariate analysis of the composition and abundance was built. We first applied a cluster analysis using the Bray-Curtis index and average linking method and an ordination analysis of nonmetric multidimensional scaling (nMDS). Two factors were examined: longitude effects (98°–110°W) and depth effects (five levels). The ANOSIM and permutational multivariate analysis of variance (PERMANOVA+) routines were applied to test these effects upon the hypothesis that community composition and the abundance of copepods did not differ significantly between a short range of longitude, but that they certainly differed between depth strata, considering the distribution of water masses. In the PERMANOVA+, the factor levels of depth and longitude (stations) were pooled as duplicates to assess the individual effect of each single factor. Since no differences in abundance, richness, or diversity of copepods were found (Kruskal-Wallis test $P > 0.05$, Supplementary Table 7), the day/night factor was not considered for these multivariate analyses. Thereafter, the differences or similarities between groups were analyzed with the SIMPER module (similarity percentages), with the aim of finding the contribution of individual species to similarity or dissimilarity between the analyzed groups. We also calculated the weighted mean depth (WMD) or gravity center for species with the highest contribution from SIMPER analysis:

$$WMD = \frac{\sum ni * zi * di}{\sum ni * zi}$$

Where $di$ is the depth of a sample $i$ (center of the depth interval), $zi$ is the thickness of the stratum, and $ni$ is the number of individuals per 100 m$^3$ at that depth (Worthington, 1931; Roe et al., 1984; Andersen et al., 2004). When vertical distribution of a species was bimodal, WMD was calculated for two portions of the water column (surface = 0–200; deep = 200–800 m).

Finally, the biological and environmental (BIOENV) analysis based on Spearman’s correlation matrix was used to determine which oceanographic variable (temperature, salinity, oxygen, and Chl-a) affected the abundance and distribution of copepods. The biological data matrix was log ($x + 1$) transformed, and the oceanographic data was normalized (subtracting the mean and dividing by the standard deviation for each variable). To determine whether the composition and abundance of zooplankton were related to oceanographic variables, a distance-based redundancy analysis (dbRDA) and a distance-based linear model (DistLM) (Clarke and Warwick, 2001; Clarke and Gorley, 2006; Anderson et al., 2008) were performed. DistLM uses permutation methods to test the plausibility of the null hypothesis (i.e., no relationship between the resemblance matrix of the principal component scores of zooplankton data and one or more environmental predictor variables). To find the best explanatory model from all possible combinations of the predictor variables, the analysis followed a stepwise procedure selection (Anderson et al., 2008) and the Akaike’s information criterion (AIC) (Akaike, 1973). The smallest achieved value of AIC indicated the best model, i.e., the best combination of the environmental predictor variables that explains the largest amount of variation in the response variable (copepods species). Additionally, further marginal tests can account for the percentage of variance of copepods when each environmental variable is considered individually through a Pseudo-F value, which is a direct multivariate analog of the Fisher’s F ratio used for standard analysis of variance. Moreover, we used a multiple linear regression model to evaluate the relationship between the total abundance, richness, and diversity indices (Shannon-Wiener and Simpson) of copepods and environmental variables (Temperature, Salinity, Oxygen, and Chl-a). Multicollinearity between variables was tested with a
multiple Spearman correlation, temperature and salinity were treated separately due to their high correlation ($R > 0.97$) (Supplementary Table 6).

**RESULTS**

**Bio-Physical Near-Surface Features and Oceanographic Characterization**

The total Chl-a concentration in the study area, located between 2800 and 3800 km from the coastal zone of Chile, during October and November of 2015 was extremely low, with mean values lower than 0.1 mg m$^{-3}$ (Figure 1B). The sea level anomaly and estimated geostrophic currents derived from satellite information showed a moderate to low level of eddy kinetic energy (EKE), prevailing positive or zero SLA values around the sampling stations (Figure 1C), contrasting with the observed patterns described in the coastal zone with higher EKE activity (Hormazabal et al., 2013).

The vertical profiles of temperature and salinity between 0 and 800 m showed a highly stratified water column. High temperature values (ca. 20°C) in the upper layer (0–150 m) delimited the mixed layer, below which the temperature decreased to 7°C at the depth of 450 m, defining the permanent thermocline. Between 450 and 800 m depth, the temperature decreased uniformly down to 5°C (Figure 2A). The salinity showed high values (35.5 – 36.0) in the upper layer, down to approximately 150 m depth, after which it decreased rapidly to 34 at 350 m, maintaining this value down to 800 m depth (Figure 2A).

The water column was well oxygenated at most depths (~5 ml L$^{-1}$), and the highest concentrations of oxygen were found between the surface and 350 m depth. Thereafter, oxygen declined in the layer between 350 and 500 m (Figure 2A). In four stations, the oxygen decreased down to 3.7 ml L$^{-1}$, and one station (98°W) showed the lowest value (2.98 ml L$^{-1}$). From 500 to 750 m depth, the oxygen increased again up to 5 ml L$^{-1}$, although the station located at 98°W showed a different pattern, where oxygen increased up to 4.5 ml L$^{-1}$. Below 750 m depth, the oxygen decreased to ~3 ml L$^{-1}$ (Figure 2A). Predominantly, this vertical profile is a characteristic for signaling the presence of a potential intra-thermocline anticyclonic eddy (ITE; Hormazabal et al., 2013; Cornejo D’Ottone et al., 2016). Complete data on oxygenation of the water column can be found in the oxygen section reported as Supplementary Figure 1.

In the SPSG at 27°S, in oceanic waters, between 98 and 107°W, the T-S diagram and OMP analysis showed four water masses: Subtropical Water (STW) with 100% of contribution between 0 and 200 m depth, the Eastern South Pacific Intermediate Water (ESPIW) with 60–80%, the Equatorial Subsurface Water (ESSW) with 20–40% at 200–400 m depth, and the Antarctic Intermediate Water (AAIW) with 20–60% between 400 and 600 m and 80–100% of contribution at 800–1000 m depth (Figures 2B,C).

Regarding the environmental variables, the PCO analysis allowed us to distinguish some clusters associated with depth (Supplementary Figure 2). The PCO1 and PCO2 components explained 98.82% of variability in the oceanographic data (temperature, salinity, and oxygen). Three groups were differentiated. The first group represented all the stations from the upper stratum between 0 and 200 m, the second group had data of the intermediate stratum between 200 and 400 m depth, and a third group had the data of the deeper strata (400–600 m and 600–800 m depth). The scatter plot of the second group...
had a minor distance with the third group (deep) as compared to the first group (upper), indicating that the environmental characteristics of the second group (200–400 m) are more similar to those of deep waters, as compared to those of the surface layer.

**Composition of Species**

A total of 121 species of copepods were found in the SPSG, and some collected species are shown in Figure 3. These species belong to five orders, 24 families, and 50 genera (Figure 4 and Supplementary Table 2). The orders Calanoida and Cyclopoida were the most represented, with 57% and 38% of the species, respectively, whereas Harpacticoida and Mormonilloida were represented with 2% of the species each, and Siphonostomatoida was represented with 1% of the species. Cyclopoida showed a higher proportion than Calanoida in the upper layer (0–200 m) but this proportion changed below 200 m depth, with an

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**FIGURE 3** | Examples of common species of copepods collected during October to November 2015 in the SPSG. (A) Aegisthus mucronatus, (B) Agetus limbatus, (C) Agetus typicus, (D) Centropages brachiatus, (E) Farranula curta, (F) Farranula gibbula, (G) Lucicutia gaussae, (H) Monocorycaeus robustus, (I) Oncaea media, (J) Oncaea mediterranea, (K) Sapphirina intestinata, (L) Sapphirina metalina, (M) Triconia conifera, (N) Vettoria parva, (O) Order Siphonostomatoidea (Family Pandaridae). The black scale is 500 μm.
In the order Calanoida, the families with most species were Clausocalanidae (12), Aetideidae (9), Paracalanidae (8), and Metridinidae (7). The most abundant calanoid genera were Clausocalaus, Pleuromamma, Paracalanus, and Lucicutia. In Cyclopoida, the families were Corycaeidae (17), Oncaeidae (10), and Sapphirinidae (10). The most abundant copepods were the genera Corycaeus, Farranula, Oithona, Oncaea, and Sapphirina.

According to Razouls et al. (2005/2019), 62% of these species exhibit a wide geographic range in the South Pacific Ocean, with 15% occurring in the oceanic South Pacific zone and West coast, while only 4% can be found in oceanic and coastal zones (East and West). Regarding the vertical distribution, 34% of the species were found in the epi- and mesopelagic depths, 25% were discovered only in the epipelagic zone, 19% were found in epi-, meso-, and bathypelagic depths, and 6% were observed in meso- and bathypelagic depths. However, 13% of these species did not show a clearly defined vertical habitat. On the other hand, the ecological role of copepods species of SPSG was assessed by assigning the feeding habits of various species in accordance with Benedetti et al. (2016). From the 121 species found in the SPSG, it may be stated that 21% of the species exhibit omnivorous and herbivorous behaviors, while 17% are omnivorous, 14% are carnivorous, and 10% are omnivores and detrivores. However, 36% of the species remain unknown.

### Abundance and Diversity of Copepods

In the oceanic zone, total copepods abundance ranged between 6 and 3000 ind. 100 m$^{-3}$, with a mean of 570 ind. 100 m$^{-3}$ (Figures 6A, B and Supplementary Figure 3). The mean richness by station was 15 species, with a maximum of 36 (E:64A, 105 W, stratum 0–100 m) and a minimum of two species (E:26, 101 W, stratum 400–600 m) (Figures 6A, B). The diversity index of Shannon showed a mean of 1.90, and the highest value was 3.80 (E:28, 104 W, stratum 100–200 m), while the lowest was 0.81 (E:26, 101 W, stratum 400–600 m). The range of variation of the Shannon index indicates that changes in species composition can strongly vary over space, revealing a heterogeneous copepod community, although most variance was accounted for by depth stratum. The Simpson index, on the other hand, indicates the degree of dominance by a single species. The average Simpson index was 0.78, with a range of 0.91–0.42 (E:26, 101 W, stratum 0–100 m and E:26, 101 W, stratum 400–600 m, respectively) (Figures 6A, B). This range shows a low level of dominance and a high level of diversity, especially in surface waters (0–100 m). However, diversity and abundance significantly decreased with depth, and the two epipelagic strata (0–100, 100–200 m) showed the highest abundances and species richness and diversity compared to the deepest layer (200–400, 400–600, and 600–800 m) (Figure 6B). Over the longitude axis, low variability was found without significant differences in abundance, richness, and diversity. In contrast, depth strata showed significant differences (Table 2).

With the inventory of 541 species of copepods recorded for the oligotrophic zones in the Pacific Ocean (Razouls et al., 2005/2019) and the list of 121 species identified in this study, the average and variation taxonomic distinctness (delta and lambda+) shown in the funnel plot (Figure 6C) were calculated. Generally, the taxonomic distinctness (delta+) values fell within the 95% confidence limits, indicating that the copepod community contained the expected level of species diversity for

**FIGURE 4** Pie charts to illustrate the distribution of orders of the subclass Copepoda in five stations and different depth strata in the oligotrophic water of the south Pacific subtropical gyre during the CIMAR-21 cruise (October–November 2015).
FIGURE 5 | Pie charts showing the horizontal and vertical distribution of pelagic copepods in the South Pacific Ocean (EC = East Coast, WC = West Coast, O = Oceanic; E = Epipelagic, M = Mesopelagic, B = Bathypelagic, A = Abisopelagic) and trophic regime (Om = Omnivore, D = Detritivore, H = Herbivore, C = Carnivore [Razouls et al., 2005/2019; Benedetti et al., 2016].

FIGURE 6 | (A) Boxplot of abundance (N, log scale), richness (S) and diversity index (Shannon-Wiener and Simpson) of the copepod community in the Pacific Southeast, Chile, (~27°S) during October and November of 2015 by longitude. (B) Boxplot of the same biological descriptors by depth strata and the boundary of the box indicates the 25th and 75th percentile, the line within the box indicates the median, and the whiskers above and below the box indicate the 90th and 10th percentile. (C) Average taxonomic distinction index (delta+) and average taxonomic variation index (lambda+) for the copepod community in the SPSG. The central line shows the mean, while the continuous lines are the distribution of probability at 95%.

the region, albeit with high variability. The taxonomic distinction index (delta+) had an expected mean of 65, and the expected taxonomic distinction variation (lambda+) had a value of 100 (Figure 6C). In the delta+, the highest value was found at 107°W at 400–600 m depth, while most samples were greater than average, with only two falling below average (Figure 6C).
The lowest values were observed at 101°W at the 0–100 and 400–600 m depth layers. The most oceanic stations (107°W and 105°W), especially in the 200–400 and 400–600 m depth layers, fell outside and above of the confidence limit, showing that average taxonomic distinctness was higher than what was randomly expected. Moreover, in the lambda+, all the stations of the upper layer (0–100 m) and three stations (105, 104, and 98°W) of the second surface layer (100–200 m) were found outside and above the confidence limit, thus displaying a much higher variability (Figure 6C).

**Multivariate Analysis and Environmental Correlates**

Five groups with 20% similarity (80% dissimilarity) were found. One group was formed with the two first levels of depth (0–200 m), along with some samples of the third strata at 200–400 m of depth. Four groups with stations were formed with the last two levels of depth (400–600 m and 600–800 m) (Figure 7A). The similarity analysis (ANOSIM) showed significant differences in the composition and abundance of copepods by depth stratum ($R = 0.398$, $p$-Value = 0.005). No differences were found by longitude ($R = 0.104$, $p$-Value = 0.198). The main differences by depth were found between the deepest and shallowest strata (i.e., 400–600 m and 600–800 m vs. 0–100 m and 100–200 m). The intermediate stratum (200–400 m) showed differences with the shallowest stratum (0–100 m) ($R = 0.556$, $p$-Value = 0.008), but not with the second (100–200 m) or the deepest (400–600 m and 600–800 m) ($R = 0.174$, $p$-Value = 0.095 and $R = 0.136$, $p$-Value = 0.159), even though the second fell on the rejection limit ($R = 0.238$, $p$-Value = 0.056).

In the percentage similarity analysis (SIMPER), the composition and abundance of copepod species showed a higher average value of Bray-Curtis dissimilarity (70–95) than of similarity (8–32) (Supplementary Tables 3, 4). The average of similarity decreased with depth, being higher in the first two strata (32.47 and 32.88). The species with the highest percentage of contribution were *Lucicutia flavicornis* and *Farranula gibbula* in the first stratum (0–100 m) and *Oithona setigera* and *Oncaea media* in the second (100–200 m). The intermediate stratum (200–400 m) showed an average similarity of 24.50, and the most important species were *Haloptilus longicornis*, *Oncaea media*, and *Oncaea mediterranea*. In the deepest strata, with an average Bray-Curtis similarity of 11.25 (400–600 m) and 8.02 (600–800 m), the species with the highest contribution were *Pleuronema gracilis* and *Aegisthus mucronatus* at the depth of 400–600 m and *Mormonilla phasma* and *Oncaea media* at the depth of 600–800 m (Supplementary Table 3).

A high level of dissimilarity in the community structure among depth strata was found (Supplementary Table 4). However, seven dominant species reached a higher percentage of contribution (38–32%), especially between the upper (0–200 m) and the deeper strata (600–800 m). The contribution of single taxa to spatial dissimilarity was relatively low (<8.5%), suggesting that changes in abundance can occur rather than changes in presence/absence of species, thereby explaining the high level of dissimilarity between the depth layers (Supplementary Table 4).

The vertical distribution of 13 species selected by SIMPER analysis revealed that five species occurred at all depth strata (*Aegisthus mucronatus*, *Farranula gibbula*, *Oncaea media*, *O. mediterranea*, and *Pleuronema abdominalis*; Figure 7B). Three species occurred between 0 and 600 m depth (*Oithona setigera*, *Lucicutia flavicornis*, and *Pleuronema gracilis*), four species were restricted to the upper 400 m of the water column.

**TABLE 2** | Kruskal-Wallis to test the influence of longitude and depth stratum on abundance, species richness, and diversity of copepods in the SPSG.

| Longitude        | df | chi-Square | $P$-value |
|------------------|----|------------|-----------|
| N (log x+1)      | 4  | 4.328      | 0.363     |
| S                | 4  | 4.828      | 0.305     |
| H′               | 4  | 3.065      | 0.547     |
| Lambda’          | 4  | 1.7525     | 0.781     |

**Deep stratum** | $df$ | chi-Square | $P$-value |
|-----------------|------|------------|-----------|
| N (log x+1)     | 4    | 14.784     | 0.00517** |
| S               | 4    | 15.123     | 0.00445** |
| H′              | 4    | 16.529     | 0.00351** |
| Lambda’         | 4    | 15.66      | 0.00351** |

N: Log abundance (ind. 100 m$^{-3}$). S: Richness, H′: Shannon-Wiener diversity index and Lambda: Simpson diversity index. ** = Highly significant ($P < 0.010$).

**FIGURE 7** | Multivariate analysis of the composition and abundance of copepods species in the SPSG (~27°S) during October–November 2015. (A) Non-metric multidimensional scaling (nMDS) plot on the first two axes based on the Bray-Curtis similarities of the log (x+1) transformed copepod species abundance data for depth strata. (B) The weighed vertical distribution (WMD) for species selected by SIMPER analysis (black dot), depth stratum with the lowest abundance (red stripe), depth for the highest abundance (blue triangle), and WMD for species with bimodal distribution (gray dot).

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showed significant linear effects on the diversity indices and variables, it was found that water temperature and salinity (diversity indices and total abundance) with environmental conditions (0–400 m) as compared to the deeper ones (400–600 and 600–800 m). Higher temperature values in the shallow and intermediate strata by the procedure. The bubbles show the vertical gradient with results by means of a dbRDA plot, with the temperature retained as the best predictor variable (Pseudo-$F = 4.605$, $p = 0.001$, AICc = 195.85), accounting for 17.3% of the total variation observed in the copepod community composition across five depth strata (Table 3). Figure 8B shows the DistLM results by means of a dbRDA plot, with the temperature retained by the procedure. The bubbles show the vertical gradient with higher temperature values in the shallow and intermediate strata (0–400 m) as compared to the deeper ones (400–600 and 600–800 m).

When assessing the association of community descriptors (diversity indices and total abundance) with environmental variables, it was found that water temperature and salinity showed significant linear effects on the diversity indices and species richness in most cases, but not oxygen concentration or Chl-a (Figure 9). Temperature and salinity explained most of the variance of the diversity indices after a multiple linear regression model (Table 4). Temperature was the best predictor ($R^2 > 0.69$), especially for the Shannon-Wiener diversity index ($R^2 = 0.75$). The contribution of each environmental variable to the total variance of the community descriptors is shown in Table 4.

### DISCUSSION

#### Hydrographic Conditions in the SPSG

Three water masses have been previously registered in oceanic waters of the Southeast Pacific in the region located at 27°S and between 98 and 110°W: The Subtropical Water (STW), Antarctic

![Figure 8](image-url)
FIGURE 9 | The relationship between species richness (S), abundance (log N), and the Shannon-Wiener (H’) and Simpson (λ’) indices of diversity of the copepod community in the SPSG during the austral spring of 2015 and environmental (bio-physical) drivers (temperature, salinity, oxygen, and Chl-a).

TABLE 4 | Multiple linear regression (stepwise) to assess the relation between community descriptors (N = abundance, S = Richness, H = Shannon-Wiener diversity index and Lamda-1 = Simpson diversity index) and environmental conditions.

| Response variable | Predictor variables | AIC | Select Variable | R² | Intercept | Slope | Standard error of bk | t | P-value |
|-------------------|---------------------|-----|----------------|----|-----------|-------|----------------------|---|---------|
| N log(x+1)        | Temp, Oxy, Chi-a    | 5.27| Temp.          | 0.7002 | 1.706     | 0.261 | 0.036                | 7.168 | 3.48e-07 *** |
|                   | Sal, Oxy, Chi-a     | 7.86| Sal            | 0.6661 | (-77.38) | 2.362 | 0.357                | 6.625 | 1.16e-06 *** |
| S                 | Temp, Oxy, Chi-a    | 85.94| Temp.         | 0.6931 | (-2.65)  | 1.379 | 0.196                | 7.049 | 4.51e-07 *** |
|                   | Sal, Oxy, Chi-a     | 89.65| Sal, Chi-a    | 0.6704 | (-320.92) | 9.565 | 2.710                | 3.529 | 0.00199 **   |
|                   | Sal, Chi-a          |       | Sal           | 0.6704 | 59.918   | 4.216 | 1.421                | 0.170 n.s. |
| H                 | Temp, Oxy, Chi-a    | 35.47| Temp.         | 0.7506 | 1.112     | 0.127 | 0.016                | 8.136 | 4.45e-08 *** |
|                   | Sal, Oxy, Chi-a     | -31.71| Sal        | 0.7083 | (-37.148) | 1.143 | 0.156                | 7.309 | 2.56e-07 *** |
| Lamda-1           | Temp, Oxy, Chi-a    | -119.02| Temp    | 0.5228 | 0.609     | 0.013 | 0.003                | 4.909 | 6.57e-05 *** |
|                   | Sal, Oxy, Chi-a     | -116.69| Sal     | 0.474   | (-3.356) | 0.119 | 0.027                | 4.453 | 0.00020 *** |

AIC = Akaike’s information criterion, R² = Determination coefficient. Significance (** Highly significant P < 0.001, ** Significant P < 0.01, n.s. non-significant P > 0.05).

Intermediate Water (AAIW), and the Pacific Deep Water (PDW) (Fuenzalida et al., 2007; Silva et al., 2009). As proposed by Silva et al. (2009), there are two water masses in this subtropical area, which may not extend beyond 95°W. The Sub-Antarctic Water (SAAW) extends offshore and occupies the upper 200 m, and the Equatorial Sub-surface Water (ESSW) advects westward in a similar fashion, but with its core centered at ca. 400 m depth at 95°W. However, according to our data, at a latitude of 27°S, the presence of SAAW was unclear, and it did not show a strong contribution in its characteristic layer (200–400 m) at the offshore area (95–108°W). SAAW did not coincide with the area either and suggested limits of its distribution for the Southeast Pacific (Llanillo et al., 2012). An additional water mass was later described for the region by Schneider et al. (2003); Hernández-Vaca et al. (2017). This is the Eastern South Pacific Intermediate Water (ESPIW), with its core in the subtropical convergence approximately at 33–38°S, between the Chilean coast, and at 90°W. This water mass is formed by subduction southeastward of the wind-driven gyre circulation, or subduction of SAAW forming the ESPIW, which continues advecting toward the equator following the direction of the Humboldt current system. ESPIW is located at 50–400 m of depth, and it spreads within the subtropical gyre as far as 150°W. In the present study, we confirmed the presence of the ESPIW water mass located at 200–300 m depth within 80–95°W and the deepest at 300–400 m in the SPSG (95–110°W).

The presence and contribution of the different water masses may also control prevailing conditions for near-surface and subsurface oxygenation at the SPSG. For instance, our vertical profiles of oxygen showed one of the stations (27°S, 98°W)
with low concentrations of oxygen at depth (2–3 ml L\(^{-1}\)). This low oxygen was also evident in the T-S-O\(_2\) diagram, indicating the contribution of the ESSW at the offshore region (20–40\%). In fact, ESSW was present in the entire coast-ocean section, with a nucleus at 300–500 m depth, and became more evident in the coastal zone with a contribution of 40–50\%. Recently, Cornejo D’Ottone et al. (2016) also registered low oxygen concentrations at the subsurface level (200–400 m) in oceanic waters (~900 km from shore) in the southeastern Pacific (30\(^{\circ}\)S, 81\(^{\circ}\)W), associated with an anticyclonic intra-thermocline eddy (ITE), which exhibited properties linked to ESSW (Chaigneau et al., 2011). This ESSW in the coastal zone is the main source for the upwelling process, but it also forms the oxygen minimum oxygen zone (OMZ), whereas ITEs are mesoscale structures (10–100 km, months) that can transport heat, salt, and the physicochemical and biological properties of such ESSW toward the open ocean (Pizarro et al., 2006; Hormazabal et al., 2013; Cornejo D’Ottone et al., 2016). Therefore, the offshore advection of ESSW from the coastal zone driven by mesoscale eddies may also transport nutrients to the subsurface water of the SPSG, and this process may sustain some of the low primary production that has taken place at the deeper depths of the euphotic zone. Although the primary production remains low, this can provide food resources for the highly diverse copepod community at the SPSG.

### Composition of Copepods in the SPSG

This study provides the first glimpse at the copepod community composition in the SPSG, identifying 121 species. This number should be considered high, provided that only a few locations were sampled compared to the records from the coastal zone of north and central-south of Chile, where 76–118 species have been identified (Hidalgo et al., 2010; Fierro, 2014; Pino-Pruer et al., 2014). However, it is relatively low when compared to the global database of the diversity and geographic distribution of planktonic marine copepods, in which 546 species of copepods have been registered for the oceanic zone of the Pacific, an extensive area named as the Central Tropical Pacific (Zone 19), which includes the North Subtropical Gyre, the tropical area with the north and south equatorial countercurrent, and the South Subtropical Gyre. However, when focusing solely on the SPSG, as one of the most unexplored areas of the ocean, 122 records of epipelagic species of copepods have been reported for the western zone (155\(^{\circ}\)W). These data come from cruises in the 1960s, and were published by Williamson and McGowan (2010). The number of species is similar to those found in our study in the eastern zone (121 species), although the data cannot be easily compared due to different sampling methods, such as larger mesh size (505 \(\mu\)m) and a different sampling gear that may sample another type of community (macrozooplankton).

Another comparable region is the North Pacific Central Gyre. In this region, a considerably high sampling effort has been invested in the ALOHA’s time series, in which the richness of species shows a value of 158 species within the North Pacific Gyre (Williamson and McGowan, 2010). In contrast, a higher species richness has been found in the Atlantic central gyres, such as 243 species at North Atlantic Gyre (Vereshchaka et al., 2017) and 300 species at South Atlantic Gyre (Center of gyre) (Piontkovski et al., 2003) (time series BATS and MAT). Although these marked differences may be caused by variable sampling efforts, it may also be linked to differences in the levels of biological production between the Pacific and Atlantic oceanic regions.

Despite the fact that the SPSG appears to be inhabited by fewer species than all the other subtropical gyres, previous studies based on morphological taxonomy (Williamson and McGowan, 2010) and metagenetic analysis (Hirai and Tsuda, 2015) indicate that, at the very least, the species composition and the dominant species are similar between the North and the South Pacific gyres, suggesting that both systems may share common ecological features. In this context, the circulation patterns of the large basins in each hemisphere and the high stability described for the central gyre ecosystems (Longhurst, 1998; Miller, 2004) may be thought to promote a high level of endemism. In fact, Hirai et al. (2015); González et al. (2020a) revealed a unique lineage of the copepod Pleuromamma abdominalis in the SPSG, suggesting limited connectivity with surrounding regions. However, in this study, some species typically seen in the coastal regions of Chile were found within the SPSG, such as Atrophia minuta, Centropages brachiatius, and Drepanopus forcipatus, implying the possibility that mesoscale eddies may reach the SPSG, carrying in zooplankton species from the upwelling zone (Morales et al., 2010). Other species similarly found within the SPSG include those with distribution in the western zone in Australian waters, such as Copilia hendorffi, Farranula orbisa, and Oncaea scottodicarloi, as well as species present in both coastal zones (East and West), such as Calanoides patagoniensis, Clausocalanus jobei, Ctenocalanus cier, Farranula curta, and Paracalanus indicus. Hence, it seems that the pelagic fauna of the SPSG is subject to the dynamics of currents of the South Pacific basin; this may explain the presence of species from both coasts in SPSG, which may result from larval dispersal processes by large-scale currents or mesoscale structures (Mackas et al., 2005; Morales et al., 2010; Chunellet et al., 2015).

According to the multivariate analysis, the most important species present in the SPSG of the epipelagic zone (0–200 m) were principally cycloponds, such as Oithona setigera, Oncaea media, O. mediterranea, and Farranula gibbula and the Calanoid L. lucicutia flavicornis. However, previous studies in similar systems report a greater abundance of calanoids, followed by poecilostomatoids and cycloponds (Paffenhofer and Mazzocchi, 2003; Dong et al., 2019). However, in recent phylogenetic reviews, the order Cyclopoida now includes the order Poecilostomatoida (Martinez-Arbizu, 2000; Khodami et al., 2017), and the union of these two considerably exceeds the Calanoida order, which presents as dominant groups of small-sized species in oligotrophic waters, such as those in the families of Clausocalanidae, Calocalanidae, and Paracalanidae (Paffenhofer and Mazzocchi, 2003). Nevertheless, the higher abundance and dominance of small-sized copepods species in oligotrophic areas is remarkable (Supplementary Figure 4). Small copepods may associate with a complex and efficient nutrient recycling microbial food web in the oligotrophic ocean (Armengol et al., 2019). In the same context, small copepods
include species with trophic regime omnivores-herbivores that feed on smaller-sized organisms, such as heterotrophic and autotrophic protists and copepod nauplii. For this reason, they are considered to have a relevant position in the food web, acting as a key component of the microbial loop and, hence, controlling the carbon recycling (Hwang et al., 2010). In this regard, the abundance of the smallest-sized copepods, such as *Oithona* spp., may have even been underestimated due to the use of the 200 μm mesh-size nets. Some studies made with 100 μm mesh-size have previously demonstrated the numerical importance of this genus (Hwang et al., 2007).

Another interesting aspect of copepods from the SPSG is the presence of blue coloration. For instance, in this study, *Farranula gibbula* was an abundant species in surface waters. This species shows blue coloration, which is associated with a complex blend of the pigment astaxanthin and a carotenoprotein coming from dietary sources. This has been interpreted as a strategy for protection from strong solar and/or UV radiation photosensitivity, antioxidant activity, and camouflage in the intense blue water (Mojib et al., 2014). The ultra-oligotrophic waters of SPSG are characterized by the highest UVR penetration ever reported for the marine environment (Tedetti et al., 2007). However, there is very little information in the study area on ever reported for the marine environment (Tedetti et al., 2007). High diversity in oligotrophic waters is also a characteristic of microzooplankton (Pierrot-Bults, 2003). The abundance and diversity found in this study were lower than in other studies conducted in oligotrophic areas (McGowan and Walker, 1979; Piontkovski et al., 2003; Vereshchaka et al., 2017), although comparisons with other studies is difficult due to variable sampling efforts and the dependence of the Shannon-Wiener diversity index on sample size and the type of logarithm used for its calculation (Gray, 2000). In any case, the SPSG community of copepods is described for the first time in the present study, and it should be considered a unique copepod assemblage, as those in other central gyres. For example, Karl and Church (2017) analyzed a long time series (since 1988) in the North Pacific Gyre (NPSG) in the ALOHA station, applying new techniques (genomics), and discovered new microorganisms and processes, showing that the NPSG system does not have a relatively stable plankton community, as initially believed. The findings revealed a considerable high temporal variability in various biological processes and on variable time scales, although these authors stated that despite the importance of this unique ecosystem, it is still understudied and poorly characterized with respect to its ecosystem structure. In this context, the key point to be emphasized is that such central gyres’ ecosystem may function based on their own dynamics and, consequently, be unique in terms of their biological attributes, including biodiversity patterns and species composition.

Our community analysis revealed that copepod assemblages are structured by depth, and therefore, they exhibit an ecological vertical zonation. In this respect, it has been proposed that the vertical distribution of zooplankton is mainly determined by variable water masses (Longhurst, 1985a; Pagès et al., 2001; Hopcroft et al., 2010; Eisner et al., 2013). However, water masses may not determine the distribution of zooplankton in all cases. For example, diel vertical migration (DVM) can modify the vertical distribution of zooplankton (Lalli and Parsons, 1997; Miller and Wheeler, 2012). In this study, no significant DVM could be found after the comparison of abundance and community descriptors between daytime and nighttime samples. However, our sampled layers may not have sufficient resolution to detect day-night exchanges for dominant small-size copepods whose migration ranges may be smaller than 100 m. Nevertheless, DVM behavior seems more pronounced in larger zooplankton, such as euphausiids or large-sized copepods, which were rarely found in our samples.

The use of two diversity indices, Shannon and Simpson, helped us interpret the variable copepod community. The Shannon index revealed a strong variable diversity over the space while being majorly associated with different depth strata, whereas the Simpson index indicated a low level of dominance
by a single species, confirming that copepod diversity in this ecosystem is much higher than in most coastal systems, where single or few species often numerically control the community (González et al., 2020b). In diversity studies, the Shannon index is mostly used, but we found the Simpson index to be useful for comparisons with other ecosystems or regions. In the same context, and regarding the influence of environmental factors on copepod diversity, it has been suggested that vertical environmental changes are considered an important factor in maintaining the diversity of planktonic taxa and ecological vertical zonation (Rutherford et al., 1999; Sommer et al., 2017). In this respect, we found a significant and positive correlation between composition, richness and diversity indices, and temperature and salinity. Our interpretation of these findings is that temperature and salinity are indeed reflecting the changes in distribution from distinct water masses, which, in turn, shape the copepod composition and diversity. Therefore, the observed vertical patterns of copepod distributions and diversity in the SPSG obey the variable contribution of water masses. This view, however, does not discard the influence of temperature on copepod diversity due to its regulating effects on eco-physiological traits and population dynamics (Huntley and Boyd, 1984; Escribano et al., 2014). Other studies provide further evidence that temperature has a crucial role in shaping the diversity patterns of zooplankton (Tittensor et al., 2010) and copepods in the ocean (Woodd-Walker et al., 2002; González et al., 2020b).

CONCLUSION

The oligotrophic environment of the SPSG as well as the oceanographic conditions are reflected in a strongly stratified and highly oxygenated water column. Satellite data show an extremely low concentration of Chl-a (∼0.01 mg m⁻³). At depth, four water masses STW, ESPIW, ESSW (originated from an intra-thermocline gyre), and AAIW were identified. Under these environmental conditions, our results provide some of the first insights into the hidden diversity in copepod assemblages across the epipelagic and mesopelagic zones of SPSG. This copepod community shows a high diversity with dominance of small-sized species, mainly omnivores in surface layers, being mostly cyclopoids (i.e., Fur anomala, Oithona, and Oncaea) and larger at depths (i.e., Pleuromamma, Halopilus). The shallow (epipelagic) stratum showed greater abundance and diversity compared to the deeper strata (mesopelagic). In the horizontal scale, less variation was found with regard to abundance and richness, while water temperature and salinity were the variables that best explained the copepod assemblages over both the vertical and horizontal axes, revealing the importance of the contributing water masses in the SPSG region.

DATA AVAILABILITY STATEMENT

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found below: https://www.pangaea.de/, PDI-23795 (Johanna et al., 2020).

AUTHOR CONTRIBUTIONS

JM-M contributed to taxonomic identification of copepod species, and data analysis and writing. AC-A contributed with the satellite data, review of physical data, and writing of results and discussion. RE integrated all results and developed the direction of the manuscript. JM-M and RE participated in discussion and data analyses and commented on the manuscript. PH contributed with confirm some copepod species and review of the manuscript. WS contributed with the review of the physical data of water masses. All authors contributed to the article and approved the submitted version.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmars.2021.625842/full#supplementary-material
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