Hydrobiologia

September 2022, Volume 849, Pages 4679-4692 https://doi.org/10.1007/s10750-022-05007-3 https://archimer.ifremer.fr/doc/00792/90428/



Thaliacean community responses to distinct thermohaline and circulation patterns in the Western Tropical South Atlantic Ocean

Giachini Tosetto Everton ^{1, 2, 3, *}, Barkokébas Silva Barbara ³, Franchesca García Díaz Xiomara ⁴, Neumann-Leitão Sigrid ³, Bertrand Arnaud ^{1, 2, 3, 5}

- ¹ MARBEC, Univ Montpellier, CNRS, IFREMER, IRD, 34200, Sète, France
- ² Institut de Recherche Pour Le Développement, 34200, Sète, France
- ³ Departamento de Oceanografia, Universidade Federal de Pernambuco, Recife, PE, 50670-901, Brazil
- ⁴ Instituto Socioambiental E Dos Recursos Hídricos, Universidade Federal Rural da Amazônia, Belém, PA, 66077-830, Brazil
- ⁵ Departamento de Pesca E Aquicultura, Universidade Federal Rural de Pernambuco, Recife, PE, 52171-900, Brazil
- * Corresponding author: Everton Giachini Tosetto, email address: evertontosetto@hotmail.com

Abstract:

In western boundary current systems, strong currents transport oligotrophic oceanic waters towards the coast. Thaliaceans may have an advantage in these systems due their ability to filter small particles such as the bacterioplankton, typically responsible for the primary production in oligotrophic waters. Here, we evaluated the structure of the thaliacean community present in the tropical South Atlantic Ocean western boundary current system to test the hypothesis that species distribution and abundance are structured by the circulation and thermohaline features. For that purpose, we used data collected though 40 mm mesopelagic trawls above the slope and around oceanic seamounts and islands. Results reveal distinct patterns in the thaliacean community structure. Over the continental slope, under the influence of the strong North Brazilian Undercurrent, Pyrosoma atlanticum was highly abundant. Soestia zonaria was also present but in a lesser amount. Offshore, around oceanic islands and Seamounts under the influence of the central branch of South Equatorial Current, Doliolida spp. were the dominant thaliacean, co-occurring with P. atlanticum in lower abundance. Mesh selectivity is a potential drawback in these results since the coarse aperture may have lost smaller species and early life stages.

Keywords: Pyrosoma atlanticum, Soestia zonaria, Doliolida, North Brazilian undercurrent, South Equatorial current, Gelatinous zooplankton

Introduction

Holoplanktonic organisms of the class Thaliacea (Chordata: Tunicata) have a global distribution and a rich biodiversity with variations in size, morphology and behaviour, reflecting adaptations to their wide range of habitats (Harbison & Campenot, 1979; Govindarajan et al., 2011; Lucas et al., 2014). Although some species are restricted to polar regions, the greatest diversity and biomass of thaliaceans is found in tropical and sub-tropical waters (Van Soest, 1975; Lucas et al., 2014). Many of these tropical systems are oligotrophic, with primary production typically dominated by bacterioplankton (Hagström et al., 1988; Zubkov et al., 2003). Thaliaceans are efficient filter-feeders, their characteristic barrel-shape body with circular musculature allows them to swim and simultaneously filter pico- and nanoparticles as small as 0.2 µm such as bacteria (Piette & Lemaire, 2015; Dölger et al., 2019). Thanks to this strategy, they flourish and remove a large portion of primary production from the environment even in oligotrophic systems, particularly in tropical western boundary systems (Stone & Steinberg, 2016).

Under specific conditions, such as high temperatures and phytoplankton productivity, upwelling events and/or specific water-mass intrusions, the high rates of asexual reproduction of thaliacieans, producing aggregated zooids, allows rapid population growth and occasional blooms (Deibel, 1982; Gibson & Paffenhöfer, 2000; Henschke et al., 2019). These large aggregations or swarms can compete and overlap other typically more abundant planktonic feeders such as copepods, changing the structure of marine food webs (Harbison & Gilmer, 1976). Historically, due to the high water content in their tissues and low caloric value, thaliaceans and other gelatinous organisms were considered irrelevant and even dead-ends in marine trophic networks. However, this concept has changed in recent decades as it has been observed that these organism are key components in the diet of groups such as crustaceans, fish and turtles (Henschke et al., 2016; Hetherington et al., 2019). Although their energetic content is indeed lower than other zooplankton organisms, due to their slow motion, predators expend less energy in the capture process making them an efficient food source (Henschke et al., 2016; Hetherington et al., 2019). Additionally, due the combination of the efficient filter-feeding lifestyle and eventual large population blooms, thaliaceans are able to transform small particles they into large faecal pellets, which quickly sink, as well as their carcass after death, promoting an important carbon input to the deep ocean (Iseki, 1981; Henschke et al., 2016; Köster & Paffenhöfer, 2016).

In the Western Tropical South Atlantic, the southern South Equatorial Current (sSEC) reaches the continental slope around 10-20°S and bifurcates into two branches. The northern one feeds the strong North Brazil Undercurrent (NBUC), which flows northward along the

Brazilian coast (Dossa et al., 2021). Meanwhile, the central South Equatorial Current (cSEC) transposes the Fernando de Noronha Ridge (FNR), composed by a series of islands and seamounts around 5°S, encountering the NBUC when reaching the Brazilian coast (Dossa et al., 2021). Differences in the intensity and direction of these currents affect the thermohaline structure of the Western Tropical South Atlantic and two areas were previously identified (Assunção et al., 2020): (i) area 1, along the continental slope and under influence of NBUC, presenting thicker thermocline and weaker stratification, hereafter called Western Boundary Current System (WBCS); and (ii) area 2, offshore in the region of FNR under influence of cSEC, with high stratification, hereafter called central South Equatorial Current system (SECS). Although nutrient-rich estuarine plumes and uplift may enhance primary production in western boundary systems, continental drainage is low in the Western Tropical South Atlantic, not reaching beyond 16 km offshore, and uplift events due anti-cyclonic eddies occurring near the shelf break are sporadic (Smith & Demaster, 1996; Castro et al., 2006; Dossa et al., 2021). Therefore, the region presents the lowest primary production along the Brazilian Coast (Ekau & Knoppers, 1999; Castro et al., 2006).

Although the particular characteristic of this system may benefit the thaliaceans filter-feeding behaviour on pico- and nanoparticles, knowledge on the species inhabiting the area is scarce (Carvalho & Bonecker, 2008; Díaz et al., 2008; Neumann-Leitao et al., 2008) and no study evaluated the structure of the thaliacean community. Here, we evaluated the composition, spatial distribution (horizontal and vertical) and abundance of the thaliacean community (Pyrosomatidae, Salpidae and Doliolidae) of the Western Tropical South Atlantic according to the characteristics of the physical environment. In particular, we test the hypothesis according to which differences in the circulation and thermohaline patterns of the WBCS and SECS shape the structure of the thaliacean community in the region.

Material and Methods

Data were collected during the ABRACOS 2 survey (Bertrand, 2017) on board the French R/V Antea from April 9 to May 6, 2017 (rainy season) in the Western Tropical South Atlantic Ocean along the continental slope of Northeastern Brazil (5°S to 9°S) and around the Fernando de Noronha Ridge (between 3°S, 37°W and 5 °S, 31 °W; Fig. 1). Fifty samples were collected in 32 stations at day and night using a micronekton mesopelagic trawl (body mesh: 40 mm, cod-end mesh: 10 mm) at depths ranging from the surface to 1170 m selected according to the presence of sound scattering layers detected with an echosounder (organisms were also captured during net descent and ascent phases). The net was hauled at 2 to 3 knots during aproximatelly 30 minutes once the targeted depth was reached. Additional samples were obtained with a finer

mesh (body mesh: 30 mm, cod-end mesh: 4 mm) around Fernando de Noronha. Specimens collected in these samples were used to improve species composition, only. Thaliaceans were sorted from the entire trawl sample on board and fixed with 4% formaldehyde buffered with sodium tetraborate ($0.5 \, \mathrm{g} \, \mathrm{L}^{-1}$).

At each station, vertical profiles of temperature (°C), salinity, pressure and fluorescence were obtained with a CTD-O2 profiler Seabird SBE911+. Conductivity, temperature, and pressure accuracies were estimated at 0.0003 S m⁻¹, 10-3°C and 0.7 dbar, respectively. Along-track current profiles were recorded with an 'Ocean Surveyor' ship-mounted acoustic Doppler current profiler (SADCP) operating at a frequency of 75 kHz with a depth range of 15-700 m. SADCP data were processed and edited using the Common Ocean Data Access System (CODAS) software package developed at the University of Hawaii (http://currents.soest.hawaii.edu). The relative velocities were rotated from the transducer to the Earth reference frame using the ship gyrocompass. The global positioning system (GPS) was used to retrieve the absolute current velocities. The orientation of the transducer relative to the gyroscopic compass and the amplitude correction factor for the SADCP were determined by standard calibration procedures. Finally, velocity profiles were averaged hourly, providing profiles in the 19–600 m range. SADCP data located over the shelf (bathymetry shallower than 70 m) were often affected by spurious reflections on the bottom, so the data coverage was only partial in these shallow areas. To describe current patterns, data from upper layer (0-70 m depth) were averaged each 0.1 square degree.

To estimate the abundance of organisms, the volume of water filtered by the net was estimated from the distance covered and the area of the net's mouth. In the laboratory, samples with more than 200 thaliacean colonies were fractionated. The colonies of each subsample were identified (mainly following Esnal, 1999; Esnal & Daponte, 1999a, 1999b) and counted. The type of zooid was identified by differentiating oozoids (asexual solitary zooids) and blastozooids (hermaphrodite aggregated zooids) based on the number and arrangement of the body muscle bands, the position and form of the digestive tract and the shape of the tunic (Tavares, 1967). For Doliolidade, only unidentified rigid transparent tunics were recorded. Abundances were standardized in number of individuals or colonies per 10⁵ m⁻³.

Laboratory analyses were performed at the *Laboratório de Ecologia Aquática Tropical* (LECAT-UFRA, Belém-Brazil) and *Laboratorio de Zooplâncton* of *Museu de Oceanografia* of *Universidade Federal de Pernambuco* (LABZOO-UFPE, Recife-Brazil). After laboratory analyses samples and specimens were deposited at the *Museu de Oceanografia* of *Universidade Federal de Pernambuco*.

Data analysis

For data analysis, the 32 stations were aggregated according to Assunção et al. (2020) in two areas with distinct thermohaline structure (Fig. 1): the WBCS, along the continental slope, corresponds to the western boundary current system dominated by the NBUC (Dossa et al., 2021), while the SECS encompasses the Rocas Atoll and part of Fernando de Noronha ridge seamounts and corresponds to the cSEC (Silva et al., 2021). For practical purpose, station 39, which was in the transition zone, was included in the SECS. To dampen effects of dominant species, abundance data was transformed by log (x+1) in all analysis. A bi-factorial Permutational Multivariate Analysis of Variance (PERMANOVA; Anderson et al., 2008) was used to test for spatial (WBCS and SECS) differences among the thaliacean community structure. Additionally, bi-factorial Analysis of Variance (ANOVA) were performed to test for spatial differences among the dominant thaliacean species.

Spatial patterns in thaliacean community abundance were identified by hierarchical cluster analysis using Bray-Curtis similarity matrix. The validity of the groups defined by this analysis was tested though SIMPROF test (5% significance level). A Similarity Percentage (SIMPER) analysis was performed to identify representative species and their contribution to similarity within the groups defined by the cluster analysis.

We performed a constrained ordination analyses to identify associations between the thaliacean species and the environmental variables. The following continuous explanatory variables were used: (i) sea surface temperature (SST), (ii) sea surface salinity (SSS), (iii) maximum value of fluorescence (as an indirect measure of primary productivity) from the surface to maximum sampling depth in each station, (iv) bottom depth, (v) relative presence of Tropical Surface Water ($\sigma_0 < 25.6$), South Atlantic Central Water (σ_0 between 25.6 and 27) and Antarctic Intermediate Water ($\sigma_0 > 27$) from the surface to maximum sampling depth in each station, (vi) the zonal component (westward/eastward) of ADCP data integrated over the first 70 m depth and between 70 and 350 m depth, and (viii) the meridional component (southward/northward) of ADCP data integrated over the first 70 m depth and between 70 and 350 m depth. Detrended Canonical Correspondence Analyses (DCCA) revealed short length of variable gradients (< 3), indicating that a linear method was more appropriate to use on this occasion, and thus Redundancy Analysis (RDA) was selected (Lepš & Šmilauer, 2003).

Environmental and distribution maps were produced in Ocean Data View 5.0 (Schlitzer, 2020) and QGIS 3.4 (QGIS Development Team, 2022). ANOVA was performed in Statistica 10 (StatSoft Inc., 2011). Cluster, SIMPROF, SIMPER and PERMANOVA analysis were performed in Primer v.6 + PERMANOVA (Clarke & Gorley, 2006). DCCA and RDA were performed in CANOCO 4.5 (Lepš & Šmilauer, 2003).

Results

Environmental background

SST ranged from 28.6 to 29.3°C through the area. Shallower stations over the continental slope in the WBCS presented higher surface salinity ranging from 36.5 to 37.5, particularly over the Pernambuco Plateau in the south of the study area, where the highest values were observed (Fig. 2). In the open ocean (SECS), SSS was slightly lower and more stable, ranging from 35.7 to 36 with exception of station 39, in the transition zone. The region of seamounts in the north of the WBCS presented intermediate values around 36 (Fig. 2). Although the SECS presented higher values of maximum fluorescence in the water column, it was quite variable, with both areas presenting high and low values. In the WBCS it ranged from 0.56 to 1 and in the SECS from 0.5 to 1.1 (Fig. 2). For a detailed description of patterns of the thermohaline structure and phytoplankton biomass in the area, see Assunção et al. (2020) and Farias et al. (2022), respectively.

SADCP data integrated from 0 to 70 m depth showed the cSEC flowed westward over the open ocean (with its core around Fernando de Noronha Ridge), feeding the NBUC and surface currents (both flowing north/northwestward) when reaching the slope and continental shelf (Fig. 3). Overall, in the surface waters (0 to 70 m depth) cSEC and NBUC were much more intense north of 7.5°S, where they spread over the continental shelf. South of 7.5°S, surface current speed weakened over the slope and almost dissipated over the continental shelf where some eastward flow was observed (Fig. 3). However, from 70 to 350 m depth, NBUC was intense all over the slope and seamounts of the WBCS (Fig 3). In contrast, in the open ocean cSEC almost dissipated in this depth and eastward flow of the South Equatorial Undercurrent SEUC was observed in the area (Fig. 3). For detailed description of the circulation in the study area see Stramma et al. (2010), Stramma and England (1999), Dossa et al. (2021) and Costa da Silva et al. (2021).

Species composition

Seven species of thaliaceans were observed in pelagic waters of the Weastern Tropical South Atlantic, corresponding to six species from the order Salpida and one from the order Pyrosomatida (Table 1). Only tunics from the order Doliolida were found, which were grouped in a single taxon as Doliolida spp. (Table 1). Overall, the most frequent and abundant species was *Pyrosoma atlanticum* Péron, 1804, which was present in 56.4% of the samples and represented 97.1% of the total specimens collected (Table 1). Although Doliolida spp. was present in 43.6% of the samples, it always occurred in low abundance, representing 2.2% of the

total catch (Table 1). The most frequent and abundant species among Salpida was *Soestia zonaria* (Pallas, 1774) which was present in 30.8% of the samples and represented 0.7% of the total catch (Table 1).

Spatial, vertical, and diel distribution patterns

Overall, the thalicean community presented highly significant differences according to the areas (PERMANOVA Pseudo-F = 27.455, p = 0.001). The dominant species, *P. atlanticum*, presented higher abundance over the continental slope and around oceanic seamounts (WBCS), reaching 700 col. 10^5 m⁻³ (Fig. 4, Table 1). Although also present in oceanic stations of the Fernando de Noronha Ridge (SECS), *P. atlanticum* was significantly (ANOVA F = 18.33, p = 0.0001) less abundant there (max: 3.4 col. 10^5 m⁻³; Fig. 4, Table 1). At daytime, the abundance of *P. atlanticum* was even in the different depths of the sampled water column. Contrastingly, at night, higher abundances were observed in the first 200 m (Figs. 4, 5).

Doliolida spp. occurred in low abundances in both areas, except for station 42A in the SECS, where it reached 27 ind. 10^5 m⁻³ (Fig. 4, Table 1). No significant differences among areas were observed (ANOVA F = 0.069, p = 0.79), however, Doliolida spp. was absent from most stations over the slope in the WBCS (Fig 4). During the day it was more abundant in deeper waters (400 to 850 m) while at night higher abundances occurred in the first 200 m (Figs. 4, 5).

S. zonaria was significantly more abundant in the WBCS (ANOVA F = 8.66, p = 0.005), particularly in the south of the study area where it reached 7.4 ind. 100 km⁻³. Only occasional catches were observed in the SECS. It was more abundant in stations sampled at night where it peaked in the first 200 m of the water column. During the day, *S. zonaria* was more abundant in samples with target depth between 400 and 600 m (Fig 5). Other Salpida species occurred almost exclusively in the first 100 m of water column in stations over the slope in the south of the WBCS (Figs. 4, 5). It is noteworthy that mesh selectivity is a potential drawback in these results since the coarse aperture may have lost smaller species and early life stages.

Community structure

The cluster analysis depicted three groups: A, B and C, with low similarity between each other (Fig. 6). Still, SIMPROF analysis performed on the results of the cluster analysis considered the three groups as valid.

Group A, with 76.3% similarity within group, included most stations located in the WBCS (Fig. 6). High abundances of *P. atlanticum* were the main responsible for the similarity in the group, although *S. zonaria* was also present in the samples of the group but with a much lesser abundance (Table 2). Group B included only offshore stations from the SECS (Fig. 6). The group

had 64.7% similarity and Doliolida spp. contributed for 90% of the similarity within the group (Table 2), although *P. atlanticum* was also present in low abundance. Group C was represented by samples from both areas (although more frequent in the SECS) where thaliaceans were found in very low abundances or completely absent (Figs. 4, 6, Table 2).

Species responses to environmental gradient

The two first canonical axes of the RDA explained 64.3% of species variance (Table 3). Monte Carlo test showed that the first (F-ratio = 40.110, P-value = 0.002) and all canonical axes together (F-ratio = 4.902, P-value = 0.002) were significant. Axis 1 explained 60.7% of the variance and was positively related to northward currents (positive meridional component, representing NBUC) and SSS; and negatively related to westward currents (negative zonal component, representing cSEC), fluorescence and bottom depth (Fig. 7, Table 3). Axis 2 explained additional 3.6% and was positively related to SSS and the relative presence of SACW and AIW in the portion of the water column sampled; and negatively related to the relative presence of TSW, SST and bottom depth (Fig. 7, Table 3).

P. atlanticum and *S. zonaria* were positively related to axis 1, indicating these species were more abundant in shallower samples under the influence of the NBUC with lower fluorescence and higher salinity. Doliolida spp. was related to the negative portion of axis 2, indicating the taxa was more abundant in deeper stations, and related to higher SST and relative presence of TSW (Fig. 7).

Discussion

Our results reveal that in the Western Tropical South Atlantic off Northeast Brazil, the thaliacean community structure presents different patterns according to the current system and associated thermohaline structure. This was not necessarily expected since in other gelatinous organisms, such as planktonic cnidarians, the spread of oligotrophic oceanic waters over the coast led to typically homogeneous oceanic communities in the study area and other regions (Thibault-Botha et al., 2004; Tosetto et al., 2021).

In the WBCS, *P. atlanticum* and in a lesser extent *S. zonaria* were the most representative thaliacean species. These species were highly related to the NBUC, the strong western boundary subsurface current that reaches up to 1.2 ms⁻¹ and flows parallel to the coast (Dossa et al., 2021). In this area, *P. atlanticum* reached the highest abundances observed in the study (up to 700 col. 10⁵ m⁻³). Massive occurrences of *P. atlanticum* were reported previously in areas with sloping topography under influence of strong currents, such as the Northern California current and Guinea current. These studies associated the massive occurrences of the species with local

changes in water temperature and salinity (Lebrato & Jones, 2009; Miller et al., 2019; Schram et al., 2020), but the relation of *P. atlanticum* with water circulation were not addressed properly by the authors. Tunic tissues of *P. atlanticum* are typically thicker than other thaliaceans (Hirose et al., 1999). This trait, associated to the larger sizes of the colonies may grant them resistance to survive in strong boundary currents and the possibility to take advantage from local conditions of water and additional food supply. These organisms may also have been pushed by the coastward currents and accumulated over the slope. Although this process was never observed in thaliaceans, many studies reported the aggregation of other planktonic organisms in sloping topography induced by oceanic currents and species vertical behavior (Cotté & Simard, 2005; Sourisseau et al., 2006; Hazen et al., 2009).

Additionally, *P. atlanticum* and *S. zonaria* where negatively related to fluorescence. Fluorescence is an indirect proxy of primary production and food availability. These species are efficient filter-feeders consuming pico- and nanoparticles including the bacterioplankton typically responsible for most primary production in oligotrophic systems such as the Western Tropical South Atlantic. Thus, the large abundance of *P. atlanticum* observed in this study may have contributed to the reduction of the chlorophyll stocks as previously observed in other ocean basins (Drits et al., 1992; Perissinotto et al., 2007).

Interestingly, the Salpida species *lasis cylindrical* (Cuvier, 1804), *Salpa aspera* Chamisso, 1819 and *Cyclosalpa pinnata* (Forskål, 1775) occurred almost exclusively in stations in the south of the WBCS, where the Pernambuco Plateau is located (Buarque et al., 2016). The particular topography and circulation in the region reduces influence of western boundary currents over the shelf and oceanward transport of coastal water is observed (Dossa et al., 2021; Tosetto et al., 2021). This hydrographic configuration has significant implications in the structure of pelagic communities and possibly the entire ecosystem (Eduardo et al., 2018; Tosetto et al., 2021).

Doliolida spp. dominated stations from group B, located exclusively in the SECS. Unfortunately, only tunics were found in our samples, thus, identification to species level was not possible. However, common doliolid species in waters of the Western Tropical South Atlantic are *Dolioletta gegenbauri* (Uljanin, 1884), *Dolioloides rarum* (Grobben, 1882), *Doliolum denticulatum* Quoy & Gaimard, 1834 and *Doliolum nationalis* Borgert, 1893 (Esnal & Daponte, 1999a; Díaz et al., 2009). These are typical epipelagic warm water species (Esnal & Daponte, 1999a), thus, the close positive relation of doliolid species observed in this study with tropical surface water mass (Fig. 7) is likely expected.

Although we did not perform depth stratified samples, our results suggest the three more frequent and abundant taxa performed diel vertical migrations. Higher abundances of *P. atlanticum* occurred in samplings carried out at night between 100 and 200 m depth. At daytime

this species was steadly abundant in the sampled water column. Since no tows were made below 1170 m depth, we cannot infer whether during daytime these organisms were aggregated in deeper waters or were distributed evenly throughout the water column. Both hypotheses are possible since P. atlanticum is known to undertake large vertical migrations below 1000 m depth (Andersen & Sardou, 1994; Henschke et al., 2019). Both Doliolida spp. and S. zonaria were more abundant in samplings carried out down to 200 m depth during night, and below 400 during daytime. S. zonaria vertical migration was not studied previously, however, this is a common behaviour in other Salpida species (e.g. Wiebe et al., 1979; Madin et al., 1996; Nishikawa & Tsuda, 2001) and thus expected for S. zonaria. Differently, large vertical migration, as suggested by the larger abundances occurring in distinct depths during day and night, were never observed for doliolids, which often are reported in epipelagic waters (Esnal & Daponte, 1999a). Vertical migration is an important mechanism transporting carbon assimilated in the surface by epipelagic phytoplankton to deeper water masses across the thermocline, particularly when considering large swarms as observed herein for P. atlanticum. Thus, future studies should use closing net devices to better describe the vertical migration of thaliacean species and quantify carbon fluxes.

Mesh selectivity is always a potential drawback when sampling pelagic organisms. While a coarse mesh may lose smaller species and early life stages, a small mesh may filter high rates of unwished particles, obstructing water passage and rapidly clogging the mesh (Riccardi, 2010; Tosetto et al., 2019). Studies focussing on thaliancean communities use wide range of mesh sizes, from fine plankton nets to large micronekton meshes (e.g. Li et al., 2011; Henschke et al., 2019; Miller et al., 2019). In this study, we used a relatively coarse mesh to sample the thaliacean community (40 mm in the main body and 10 mm in the cod-end). Still, with this gear we could filter a large volume of water reaching deeper strata. The coarse meshes also reduce clogging and bow wave effect, enabling the catch of large specimens and colonies (Vannucci, 1968). Thaliacea encompasses species with a wide range of sizes and no single method would allow a sampling all of them. The additional trawls we performed with the finer mesh (30 mm in the main body and 3 mm in the cod-end) collected two additional salp species (Table 1). Unfortunately, few tows were performed with this mesh, and we could not properly compare the effects of mesh selectivity on the structure and abundance of the thaliacean community in the Western Tropical South Atlantic, an important subject to be addressed in future studies. Additionally, studies carried out with samples collected with a 300 μm mesh size from 200 m depth to surface at the Archipelago of Saint Peter and Saint Paul, close to FNR, recorded eight thaliaceans species: P. atlanticum, D. rarum, D. denticulatum, I. cylindrica (as Weelia cylindrical), Thalia democratica (Forskål, 1775), Thalia cicar van Soest, 1973, Pegea socia (Bosc, 1802) and *Thalia* sp. (Díaz et al., 2009). Thus, at least three of the species collected with the finer mesh in Saint Peter and Saint Paul were not present in our samples.

In conclusion, we observed clearly distinct patterns in the thaliacean community in the Western Tropical South Atlantic likely related to the circulation and thermohaline structure in the area. Over the continental slope, where the strong NBUC flows northward, high abundances of *P. atlanticum*, and, to a lesser degree, *S. zonaria* were observed. Over the Pernambuco plateau, in the South of this area and where current intensity was lower, other less abundant salp species, such as *I. cyindrica*, *C. pinnata* and *S. aspera*, were present as well. Meanwhile, offshore around the Fernando de Noronha Ridge, Doliolida spp. were the dominant thaliacean, co-occurring with *P. atlanticum* in lower abundance.

Acknowledgements

We are grateful to the French oceanographic fleet for funding the survey ABRAÇOS 1 and the officers, crew and scientific team of the R/V Antea for their contribution to the success of the operations. The present study was not possible without the support of all members from LABZOO and other laboratories from UFPE and UFRPE. We thank to CAPES (Coordenação de Aperfeiçoamento de Pessoal de Nível Superior) and CNPq (Brazilian National Council for Scientific and Technological Development), which provided Research Scholarships to E.G.T., B.B.S. and S.N.L. This work is a contribution to the LMI TAPIOCA (www.tapioca.ird.fr), CAPES/COFECUB program (88881.142689/2017-01), the European Union's Horizon 2020 projects PADDLE (grant agreement No. 73427) and TRIATLAS (grant agreement No. 817578).

References

Andersen, V., & J. Sardou, 1994. *Pyrosoma atlanticum* (Tunicata, Thaliacea): diel migration and vertical distribution as a function of colony size. Journal of Plankton Research 16: 337–349.

Anderson, M. J., R. N. Gorley, & K. R. Clarke, 2008. Permanova For Primer: Guide to Software and Statiscal Methods. PRIMER-E, Plymouth.

Assunção, R. V., A. C. Silva, A. Roy, B. Bourlès, C. H. S. Silva, J.-F. Ternon, M. Araujo, & A. Bertrand, 2020. 3D characterisation of the thermohaline structure in the southwestern tropical Atlantic derived from functional data analysis of in situ profiles. Progress in Oceanography 187: 102399.

Buarque, B. V., J. A. Barbosa, J. R. G. Magalhães, J. T. Cruz Oliveira, & O. J. C. Filho, 2016. Postrift volcanic structures of the Pernambuco Plateau, northeastern Brazil. Journal of South American Earth Sciences 70: 251–267.

Carvalho, P. F. de, & S. L. C. Bonecker, 2008. Tunicata, Thaliacea, Pyrosomatidae, Pyrosomella verticillata (Neumann, 1909): first record from the southwest Atlantic Ocean. Check List 4: 272.

Castro, B. M., F. P. Brandini, A. M. S. Pires-Vanin, & L. B. Miranda, 2006. Multidisciplinary oceanographic processes on the Western Atlantic continental shelf between 4°N and 34°S. The Sea 14: 1–39.

Clarke, K. R., & R. N. Gorley, 2006. PRIMER 6 + PERMANOVA. .

Cotté, C., & Y. Simard, 2005. Formation of dense krill patches under tidal forcing at whale feeding hot spots in the St. Lawrence Estuary. Marine Ecology Progress Series 288: 199–210.

Deibel, D., 1982. Laboratory determined mortality, fecundity and growth rates of *Thalia democratica* Forskal and *Dolioletta gegenbauri* Uljanin (Tunicata, Thaliacea). Journal of Plankton Research 4: 143–153.

Díaz, X. F. G., L. M. de O. Gusmão, & S. Neumann-Leitão, 2008. New record of Thalia cicar van Soest 1973 (Urochordata: Thaliacea) in the Equatorial Atlantic. Biota Neotropica 8: 99–104.

Díaz, X. F. G., L. M. de O. Gusmão, & S. Neumann-Leitao, 2009. Biodiversidade e dinâmica espaço-temporal do zooplâncton In Hazin, F. H. V. (ed), O arquipélago de São Pedro e São Paulo: 10 anos de estação cientíica. SECIRM, Brasília: 138–147.

Dölger, J., T. Kiørboe, & A. Andersen, 2019. Dense Dwarfs versus Gelatinous Giants: The Trade-Offs and Physiological Limits Determining the Body Plan of Planktonic Filter Feeders. The American Naturalist 194: E30–E40.

Dossa, A. N., A. C. Silva, A. Chaigneau, G. Eldin, M. Araujo, & A. Bertrand, 2021. Near-surface western boundary circulation off Northeast Brazil. Progress in Oceanography 190: 102475.

Drits, A. V., E. G. Arashkevich, & T. N. Semenova, 1992. *Pyrosoma atlanticum* (Tunicata, Thaliacea): grazing impact on phytoplankton standing stock and role in organic carbon flux. Journal of Plankton Research 14: 799–809.

Eduardo, L. N., T. Frédou, A. S. Lira, B. P. Ferreira, A. Bertrand, F. Ménard, & F. L. Frédou, 2018. Identifying key habitat and spatial patterns of fish biodiversity in the tropical Brazilian continental shelf. Continental Shelf Research 166: 108–118.

Ekau, W., & B. Knoppers, 1999. An introduction to the pelagic system of the Northeast and East Brazilian shelf. Archive of Fishery and Marine Research 21.

Esnal, G. B., 1999. Pyrosomatida In Boltovskoy, D. (ed), South Atlantic Zooplankton. Backhuys Publishers, Leiden: 1401–1408.

Esnal, G. B., & M. C. Daponte, 1999a. Doliolida In Boltovskoy, D. (ed), South Atlantic Zooplankton. Backhuys Publishers, Leiden: 1409–1421.

Esnal, G. B., & M. C. Daponte, 1999b. Salpida In Boltovskoy, D. (ed), South Atlantic Zooplankton. Backhuys Publishers, Leiden: 1423–1444.

Farias, G. B., J.-C. Molinero, C. Carré, A. Bertrand, B. Bec, & P. A. M. de C. Melo, 2022. Uncoupled changes in phytoplankton biomass and size structure in the western tropical Atlantic. Journal of Marine Systems 227: 103696.

Gibson, D. M., & G.-A. Paffenhöfer, 2000. Feeding and growth rates of the doliolid, Dolioletta gegenbauri Uljanin (Tunicata, Thaliacea). Journal of Plankton Research 22: 1485–1500.

Govindarajan, A. F., A. Bucklin, & L. P. Madin, 2011. A molecular phylogeny of the Thaliacea. Journal of Plankton Research 33: 843–853.

Hagström, Å., F. Azam, A. Andersson, J. Wikner, & F. Rassoulzadegan, 1988. Microbial loop in an oligotrophic pelagic marine ecosystem: possible roles of cyanobacteria and nanoflagellates in the organic fluxes. Marine Ecology Progress Series 49: 171–178.

Harbison, G. R., & R. B. Campenot, 1979. Effects of temperature on the swimming of salps (Tunicata, Thaliacea): Implications for vertical migration1: Temperature effects on salps. Limnology and Oceanography 24: 1081–1091.

Harbison, G. R., & R. W. Gilmer, 1976. The feeding rates of the pelagic tunicate Pegea confederata and two other salps1: Salp feeding rates. Limnology and Oceanography 21: 517–528.

Hazen, E., A. Friedlaender, M. Thompson, C. Ware, M. Weinrich, P. Halpin, & D. Wiley, 2009. Fine-scale prey aggregations and foraging ecology of humpback whales *Megaptera novaeangliae*. Marine Ecology Progress Series 395: 75–89.

Henschke, N., J. D. Everett, A. J. Richardson, & I. M. Suthers, 2016. Rethinking the Role of Salps in the Ocean. Trends in Ecology & Evolution 31: 720–733.

Henschke, N., E. A. Pakhomov, L. E. Kwong, J. D. Everett, L. Laiolo, A. R. Coghlan, & I. M. Suthers, 2019. Large Vertical Migrations of *Pyrosoma atlanticum* Play an Important Role in Active Carbon Transport. Journal of Geophysical Research: Biogeosciences 124: 1056–1070.

Hetherington, E., C. Kurle, S. Benson, T. Jones, & J. Seminoff, 2019. Re-examining trophic dead ends: stable isotope values link gelatinous zooplankton to leatherback turtles in the California Current. Marine Ecology Progress Series 632: 205–219.

Hirose, E., S. Kimura, T. Itoh, & J. Nishikawa, 1999. Tunic Morphology and Cellulosic Components of Pyrosomas, Doliolids, and Salps (Thaliacea, Urochordata). The Biological Bulletin 196: 113–120.

Iseki, K., 1981. Particulate Organic Matter Transport to the Deep Sea by Salp Fecal Pellets. Marine Ecology Progress Series 5: 55–60.

Köster, M., & G.-A. Paffenhöfer, 2016. How efficiently can doliolids (Tunicata, Thaliacea) utilize phytoplankton and their own fecal pellets?. Journal of Plankton Research plankt;fbw089v1.

Lebrato, M., & D. O. B. Jones, 2009. Mass deposition event of *Pyrosoma atlanticum* carcasses off Ivory Coast (West Africa). Limnology and Oceanography 54: 1197–1209.

Lepš, J., & P. Šmilauer, 2003. Multivariate Analysis of Ecological Data using CANOCO. Cambridge University Press, Cambridge.

Li, K., J. Yin, L. Huang, J. Zhang, S. Lian, & C. Liu, 2011. Distribution and abundance of thaliaceans in the northwest continental shelf of South China Sea, with response to environmental factors driven by monsoon. Continental Shelf Research 31: 979–989.

Lucas, C. H., D. O. B. Jones, C. J. Hollyhead, R. H. Condon, C. M. Duarte, W. M. Graham, K. L. Robinson, K. A. Pitt, M. Schildhauer, & J. Regetz, 2014. Gelatinous zooplankton biomass in the

global oceans: geographic variation and environmental drivers: Global gelatinous biomass. Global Ecology and Biogeography 23: 701–714.

Madin, L. P., P. Kremer, & S. Hacker, 1996. Distribution and vertical migration of salps (Tunicata, Thaliacea) near Bermuda. Journal of Plankton Research 18: 747–755.

Miller, R. R., K. M. Sakuma, B. K. Wells, J. C. Field, M. Way, & S. Cruz, 2019. Distribution of pelagic thaliaceans, <i>Thetys vagina</i and *Pyrosoma atlanticum*, during a period of mass occurrence within the California current. California Cooperative Oceanic Fisheries Investigations Reports 60: 94–108.

Neumann-Leitao, S., E. M. E. Sant'anna, L. M. D. O. Gusmao, D. A. Do Nascimento-Vieira, M. N. Paranagua, & R. Schwamborn, 2008. Diversity and distribution of the mesozooplankton in the tropical Southwestern Atlantic. Journal of Plankton Research 30: 795–805.

Nishikawa, J., & A. Tsuda, 2001. Diel vertical migration of the tunicate Salpa thompsoni in the Southern Ocean during summer. Polar Biology 24: 299–302.

Perissinotto, R., P. Mayzaud, P. Nichols, & J. Labat, 2007. Grazing by Pyrosoma atlanticum (Tunicata, Thaliacea) in the south Indian Ocean. Marine Ecology Progress Series 330: 1–11.

Piette, J., & P. Lemaire, 2015. Thaliaceans, The Neglected Pelagic Relatives of Ascidians: A Developmental and Evolutionary Enigma. The Quarterly Review of Biology 90: 117–145.

QGIS Development Team, 2022. QGIS Geographic Information System. .

Riccardi, N., 2010. Selectivity of plankton nets over mesozooplankton taxa: implications for abundance, biomass and diversity estimation. Journal of Limnology 69: 287.

Schlitzer, R., 2020. Ocean Data View. .

Schram, J., H. Sorensen, R. Brodeur, A. Galloway, & K. Sutherland, 2020. Abundance, distribution, and feeding ecology of Pyrosoma atlanticum in the Northern California Current. Marine Ecology Progress Series 651: 97–110.

Silva, A. C., A. Chaigneau, A. N. Dossa, G. Eldin, M. Araujo, & A. Bertrand, 2021. Surface Circulation and Vertical Structure of Upper Ocean Variability Around Fernando de Noronha Archipelago and Rocas Atoll During Spring 2015 and Fall 2017. Frontiers in Marine Science 8: 598101.

Smith, W. O., & D. J. Demaster, 1996. Phytoplankton biomass and productivity in the Amazon River plume: correlation with seasonal river discharge. Continental Shelf Research 16: 291–319.

Sourisseau, M., Y. Simard, & F. Saucier, 2006. Krill aggregation in the St. Lawrence system, and supply of krill to the whale feeding grounds in the estuary from the gulf. Marine Ecology Progress Series 314: 257–270.

StatSoft Inc., 2011. Statistica, version 10. Tulsa.

Stone, J. P., & D. K. Steinberg, 2016. Salp contributions to vertical carbon flux in the Sargasso Sea. Deep Sea Research Part I: Oceanographic Research Papers 113: 90–100.

Stramma, L., & M. England, 1999. On the water masses and mean circulation of the South Atlantic Ocean. Journal of Geophysical Research: Oceans 104: 20863–20883.

Stramma, L., S. Schmidtko, L. A. Levin, & G. C. Johnson, 2010. Ocean oxygen minima expansions and their biological impacts. Deep Sea Research Part I: Oceanographic Research Papers 57: 587–595.

Tavares, D. Q., 1967. Occurrence of doliolios and salps during 1958, 1959, and 1960 off the São Paulo coast. Boletim do Instituto Oceanográfico de São Paulo 16: 87–97.

Thibault-Botha, D., J. R. E. Lutjeharms, & M. J. Gibbons, 2004. Siphonophore assemblages along the east coast of South Africa; mesoscale distribution and temporal variations. Journal of Plankton Research 26: 1115–1128.

Tosetto, E. G., A. Bertrand, S. Neumann-Leitão, A. Costa da Silva, & M. Nogueira Júnior, 2021. Spatial patterns in planktonic cnidarian distribution in the western boundary current system of the tropical South Atlantic Ocean. Journal of Plankton Research 43: 270–287.

Tosetto, E. G., S. Neumann-Leitão, & M. Nogueira Júnior, 2019. Sampling planktonic cnidarians with paired nets: Implications of mesh size on community structure and abundance. Estuarine, Coastal and Shelf Science 220: 48–53.

Van Soest, R. W. M., 1975. Zoogeography and speciation in the Salpidae. Beaufortia 23: 181–2015.

Vannucci, M., 1968. Loss of organisms through the meshes Zooplankton sampling. Unesco, Paris: 77–86.

Wiebe, P. H., L. P. Madin, L. R. Haury, G. R. Harbison, & L. M. Philbin, 1979. Diel vertical migration bySalpa aspera and its potential for large-scale particulate organic matter transport to the deep-sea. Marine Biology 53: 249–255.

Zubkov, M. V., B. M. Fuchs, G. A. Tarran, P. H. Burkill, & R. Amann, 2003. High Rate of Uptake of Organic Nitrogen Compounds by Prochlorococcus Cyanobacteria as a Key to Their Dominance in Oligotrophic Oceanic Waters. Applied and Environmental Microbiology 69: 1299–1304.

Figures

Figure 1. Geographic location of the study area in the Western Tropical South Atlantic, showing the sampled stations.

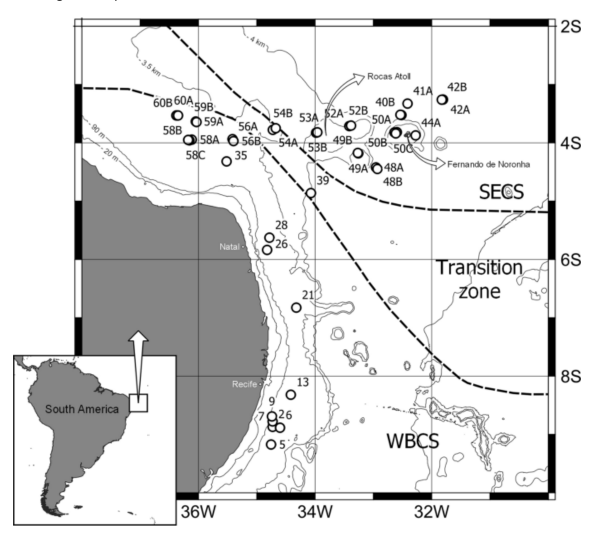


Figure 2. Sea-surface temperature (SST), sea-surface salinity (SSS) and maximum fluorescence during autumn 2017 in the Western Tropical South Atlantic.

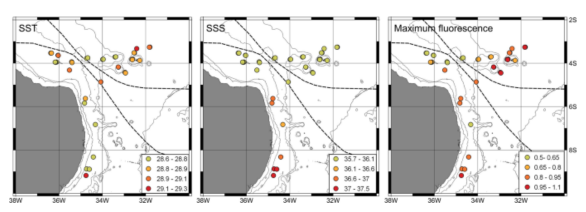


Figure 3. Current vectors of ADCP data integrating 0 - 70 (a) and 70 - 350 (b) m depth. Large arrows indicate the predominant currents in the area (NBUC = North Brazil Undercurrent; cSEC = Central branch of the South Equatorial Current; SEUC = South Equatorial Undercurrent).

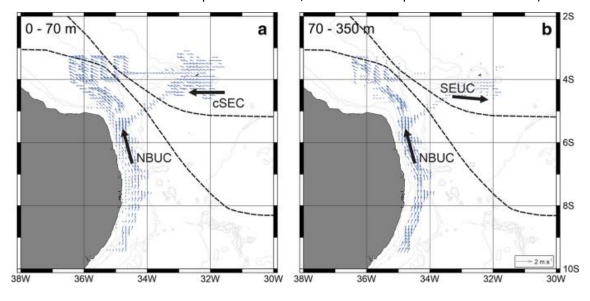


Figure 4. Geographic distribution of the abundance of *Pyrosoma atlaticum*, Doliolida spp. and Salpida species. (s) = solitary individuals, (a) = aggregates. Values standardized in number of individuals or aggregates per $10^5 \,\mathrm{m}^{-3}$.

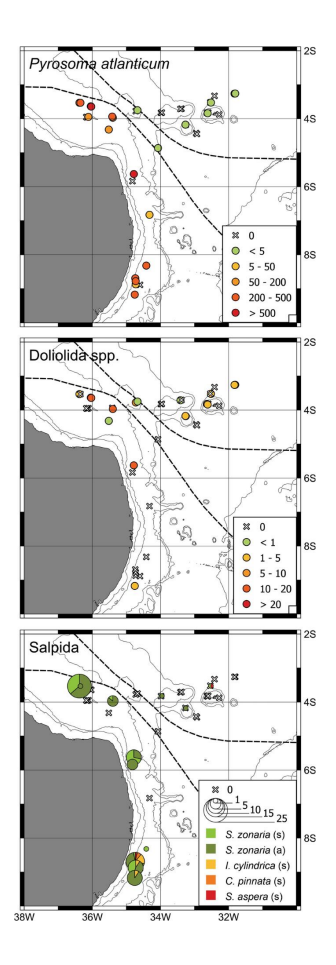


Figure 5. Average relative abundance (individuals and aggregates per $10^5 \, \text{m}^{-3}$) per target depth strata (including net descent and ascent phases) and day period of *Pyrosoma atlaticum*, Doliolida spp. and Salpida species in the Western Tropical South Atlantic. x = Depth strata not sampled.

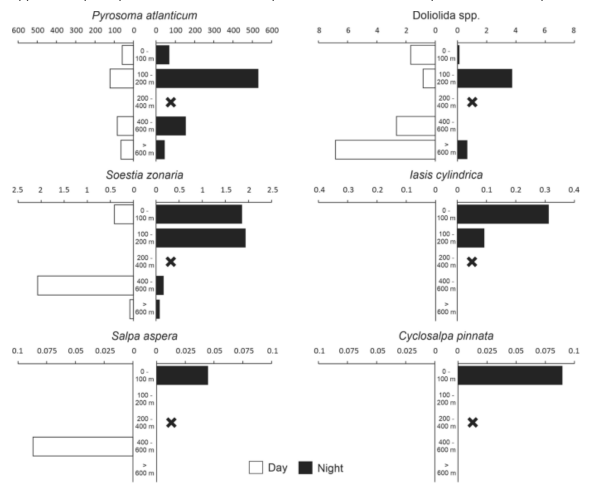
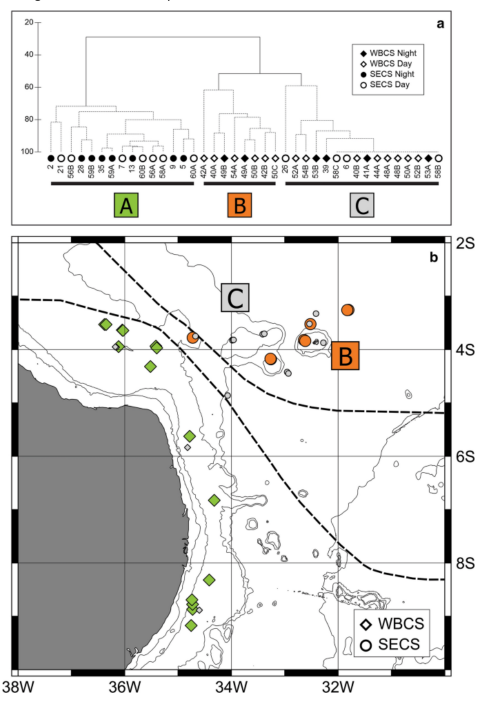
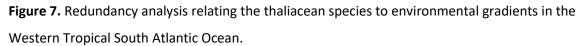


Figure 6. (a) Cluster analysis dendrogram of data indicating three groups samples with similar thaliacean communities in the Western Tropical South Atlantic Ocean, dashed lines are significant groups in the SIMPROF analysis. (b) Map indicating distribution of the groups arranged in the cluster analysis.





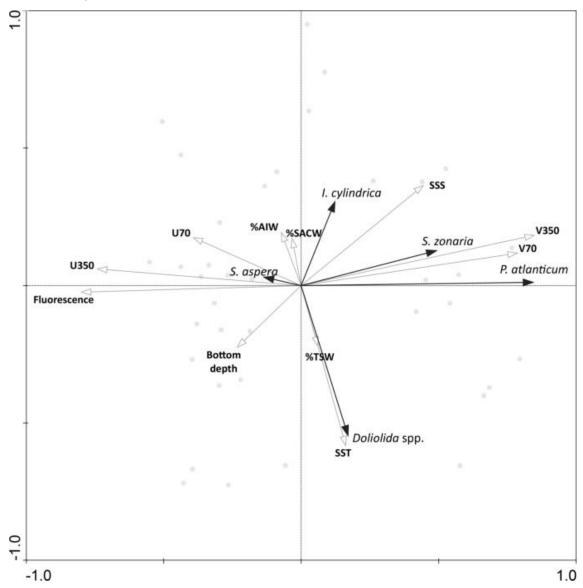


Table 1. Species composition and basic statistics of thaliaceans from two areas (Assunção *et al.*, 2020) in the Western Tropical South Atlantic Ocean off Northeast Brazil. Mean abundance (colonies or individuals $10^5 \, \text{m}^{-3}$) per station and standard deviation, range of abundance, frequency of occurrence (f; considering both areas). X = Species present only in the additional samples and not included in the analysis.

	WBCS		SECS		
Species	Mean ± SD	Range of non- zero abundances	Mean ± SD	Range of non- zero abundances	f (%)
Pyrosoma atlanticum (colony)	206.42 ± 209.67	27.65 - 699.96	0.36 ± 0.79	0.2 - 3.38	56.41
Doliolida spp.	2.06 ± 4.17	0.33 - 12.42	2.5 ± 5.89	0.37 - 27.04	43.59
Soestia zonaria (aggregate)	1.01 ± 1.44	0.54 - 4.7	0.04 ± 0.11	0.33 - 0.39	25.64
Soestia zonaria (solitary)	0.39 ± 0.81	0.18 - 2.69	0.02 ± 0.09	0.43	17.95
lasis cylindrica (solitary)	0.08 ± 0.22	0.28 - 0.9	-	-	7.69
Salpa aspera (solitary)	0.01 ± 0.04	0.18 - 0.18	0.02 ± 0.09	0.43	5.12
Salpa aspera (aggregate)			Χ		
Cyclosalpa pinnata (solitary)	0.02 ± 0.08	0.36 - 0.36	-	-	2.56
Salpa fusiformis (solitary)			Χ		
Cyclosalpa polae (solitary)			X		

Table 2. Results of SIMPER analysis, showing the relative contribution of thaliaceans taxa in the formation of the groups defined in the Cluster analysis

Species	Α	В	С
Pyrosoma atlanticum	93.1	9.4	30.5
Doliolida spp.		89.8	45.7
Soestia zonaria	4		23.8

Table 3. Summary of the Redundancy Analysis (RDA) performed between the thaliacean taxa and environmental explanatory variables from the Western Tropical South Atlantic

	Axis 1	Axis 2
Eigenvalues	0.607	0.036
Species-environment correlations	0.843	0.578
Cumulative variance (%):		
Of species data	60.7	64.3
Of species-environment relationships	92.9	98.3
Correlations of explanatory variables:		
0-70 m currents zonal component	-0.3304	0.1006
0-70 m currents meridional component	0.6625	0.0678
70-350 m currents zonal component	-0.6243	0.0347
70-350 m currents meridional component	0.7143	0.1054
Bottom depth	-0.1959	-0.1299
Sea surface temperature	0.1365	-0.3367
Sea surface salinity	0.3739	0.2105
Fluorescence	-0.6734	-0.0143
% of Tropical Surface Water	0.0506	-0.1276
% of South Atlantic Central Water	-0.028	0.099
% of Antarctic intermediate water	-0.0608	0.1115