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**BIOSOPE:  
introduction to the  
special issue**

H. Claustre

# Introduction to the special section Bio-optical and biogeochemical conditions in the South East Pacific in late 2004: the BIOSOPE program

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## Abstract

The objectives of the BIOSOPE (Biogeochemistry and Optics SOuth Pacific Experiment) project was to study, during the austral summer, the biological, biogeochemical and bio-optical properties of different trophic regimes in the South East Pacific: the eutrophic zone associated with the upwelling regime of the Chilean coast, the mesotrophic area associated with the plume of the Marquesas Islands in the HNLC (High Nutrient Low Chlorophyll) waters of this subequatorial area, and the extremely oligotrophic area associated with the central part of the South Pacific Gyre (SPG). At the end of 2004, a 55-day international cruise with 32 scientists on board took place between Tahiti and Chile, crossing the SPG along a North-West South-East transect. This paper describes in detail the objective of the BIOSOPE project, the implementation plan of the cruise, the main hydrological entities encountered along the ~8000 km South East Pacific transect and ends with a general overview of the papers published in this Biogeosciences special issue.

### 1 Prior oceanographic knowledge of the South East Pacific

The South East Pacific (SEP 5° S–40° S, East of 150° W) makes the connection between tropical and high latitudes of the austral ocean. It remains the most sparsely sampled oceanic region of the global ocean from both hydrodynamic (Leth et al., 2004) and biogeochemical (Daneri and Quinones, 2001) points of view.

Large scale investigations have been initially conducted as part of the transpacific SCORPIO sections performed along 43° S and 28° S (Reid, 1973) and the Hawaii-to-Tahiti shuttle experiment (Wyrski and Kilonsky, 1984). More recently, these observations have been supplemented with some WOCE sections (P6 lines, e.g. Wijffels et al., 2001; P19 line Tsuchiya and Talley, 1998) and with the rather intensive deployment of drifters and profiling floats as part of the ARGO program. The general patterns of the surface circulation in the South Equatorial Pacific region can be characterized by

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three main current regimes (Chaigneau and Pizarro, 2005a; Kessler, 2006). On its equator side, the South Pacific Gyre (SPG) is delineated by the South Equatorial Current (SEC) flowing westwards and sometimes embedding the (weaker) South Equatorial Counter Current (SECC) (Wyrтки and Kilonsky, 1984; Eldin, 1983). On its polar side, the (weak) South Pacific Current (SPC), corresponding to the eastern extension of the West Wind Drift, flowing eastward near  $\sim 30^\circ$  S, forms the southern closure of the subtropical gyre circulation (Stramma et al., 1995). Approaching South America, this current turns northwards and contributes to the diffuse surface flows of the broad Peru-Chile (Humboldt) Current (PCC); it can reach punctual velocities of  $15\text{--}20\text{ cm s}^{-1}$  (Chaigneau and Pizarro, 2005b) and sometimes presents complex motions near the coast, with important mesoscale activity (eddies and filaments) associated with the coastal upwelling regime (e.g. Shaffer et al., 1995). Below the PCC, the Peru Chile Undercurrent (PCUC), restricted in a narrow band near and above the shelf break, transports towards the pole the warm and salty subsurface equatorial waters (Silva and Neshyba, 1979).

Even fewer investigations have been dedicated to assess the biological and biogeochemical status of the SEP. Yet, this area, as a result of the hydrodynamical forcing described above presents a remarkable diversity of trophic conditions and even some singularities that do not have any counterpart in others areas of the world ocean. Basically three main biogeochemical regimes can be identified in this large water mass.

The SPG is the largest subtropical anticyclonic gyre and the least described region of the ocean (Longhurst, 1998; Claustre and Maritorena, 2003). Indeed, to our knowledge, the few biological observation reported to date concerns some (very low) chlorophyll concentration along  $105^\circ$  W (Chavez et al., 1995), near surface phytoplankton determination along  $110^\circ$  W (Hardy et al., 1996) and a recent analysis on picophytoplankton distribution in the upper layer along  $32^\circ$  S (Bouman et al., 2006; Grob et al., 2006). Thanks to satellite remote sensing of ocean color, some general surface properties of the SPG have emerged. The SPG is the most oligotrophic of all the sub-tropical gyres. In the vicinity of Easter Island (Rapa Nui), the surface chlorophyll *a* concen-

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tration (Chl-*a*) is the lowest of the global open ocean with annual means as low as 0.019 mg Chl-*a* m<sup>-3</sup> (Fig. 1a). These exceptionally low surface Chl-*a* correspond to very clear waters with deep penetration not only of visible but also of UV radiations (Vasilkov et al., 2001). Interestingly, these hyperoligotrophic characteristics are also closely correlated with exceptional physical features (Fig. 1b). Actually, the pycnocline depth, derived from hydrological database, presents the largest values (>200 m) of the global ocean (Fig. 1b), precisely in the zone where the Chl-*a* concentrations is the lowest. Furthermore the strength of the pycnocline is extremely weak in this zone (Fiedler and Talley, 2006). This apparently tight coupling between the physical and biological fields reveals an extremely deep nutrient source and a weak upward diffusion of nutrient fuelling the phototrophic production in the surface layer. Furthermore, due to the weak source of iron-rich desert dust in the southern hemisphere, atmospheric iron deposition is low, particularly in the SPG. Actually modelling results suggest that the deposition rate could be at least one order of magnitude lower than in the northern hemisphere (Mahowald et al., 2005) (Fig. 1c). Therefore, the upper waters of the SPG are expected to receive, in comparison to other oceanic regions, the lowest nutrients fluxes from deeper layers as well as the lowest iron flux from the atmosphere.

In contrast to the SPG, the equatorial and subequatorial Pacific waters have received considerable attention, particularly during the nineties in the context of the JGOFS program (Murray et al., 1995; Murray et al., 1997; Dandonneau, 1999). The impact of physical (e.g. upwelling strength), chemical (e.g. iron supply) and biological (e.g. grazing pressure) factors on the carbon cycle have been investigated and quantified in details (Murray et al., 1994). Furthermore, the analysis of time series permitted to begin documenting the effect of ENSO on the inter-annual variability of some specific biogeochemical processes (Yoder and Kennelly, 2003). At the regional scale, these subequatorial waters are referred to as High Nutrient Low Chlorophyll waters because they present some moderate oligotrophic characteristics associated with significant amounts of nitrate. However, at a more local scale and in the vicinity of steep islands, these oligotrophic conditions might be temporarily or permanently relieved. The

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enhancement of water productivity has been reported in particular for waters west of the Galapagos Archipelago (Martin et al., 1994), Kiribati Islands (Messié et al., 2006) and Marquesas Archipelago (Signorini et al., 1999; Martinez and Maamaatuaiahutapu, 2004). The reasons behind this enhancement of productivity are still debated and are investigated mostly through modelling or remote sensing. In particular, no in situ investigation has been carried out to date for the remote Marquesas Islands.

The permanent upwelling associated with the PCC represents one of the most productive area of the global ocean (Carr, 2002) fuelling some of the largest fisheries. Despite this important biological and trophic impact, the PCC remains the least well known eastern boundary current system, both from the dynamical and the biogeochemical points of view (Leth et al., 2004; Daneri and Quinones, 2001). Upwelling conditions along the Chilean coast present clear singularities. High biomass is recorded out of phase with a priori upwelling favourable winds (towards the equator) (Thomas, 1999). This biomass sometimes presents extreme westwards filamentous extension (200–300 km). These extensions are more important than what would be expected from the sole local dynamics of “wind driven” upwelling. The importance of mesoscale dynamics (meanders and gyres) and their interaction with the complex large scale circulation has therefore been proposed (Leth and Shaffer, 2001; Thomas et al., 1994) as regional specificities of SEP that might account for large offshore biomasses associated with these filaments.

## 2 Objectives of the BIOSOPE program

In 2001, after a whole decade of rather intensive field observations in various oceanic provinces as part of the JGOFS program, Daneri and Quinones wrote a contribution in the US JGOFS newsletter with a title clearly summarizing their concerns “Under sampled ocean systems : a plea for an international study of biogeochemical cycles in the Southern Pacific Gyre and its boundaries”. The BIOSOPE (Biogeochemistry and Optics South Pacific Experiment) program, jointly endorsed by the IMBER and SOLAS

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programs, was completely in line with this plea. Its overall goal was to explore and describe the biological, biogeochemical and optical characteristics of the South East Pacific. Using core measurements (in line with the former JGOFS core parameters) as well as new ones (e.g. based on the use of molecular biology and isotopic techniques, novel optical devices), two major goals were set.

1. Perform detailed studies in a certain number of oceanic provinces of the SEP in order to quantify the variables and processes that are essential to the understanding of trophic relationships, biogeochemical cycles of carbon and related elements as well as water optical signatures. The primary motivation of BIOSOPE was to study the South Pacific Gyre, expected to be the end member of oligotrophic conditions in the global ocean. But other SEP oceanic provinces were also of interest. The water masses west of Marquesas (local biomass enhancement visible from satellite) deserved to be investigated in the more general context of the HLNC conditions associated with the subequatorial area. Similarly, the upwelling zone extending offshore the Chilean coast was of great interest because of its particular filamentous patterns.
2. Understand in details the relationships linking optical properties of SEP waters to their biological and to biogeochemical characteristics. Besides surface Chl-*a*, an increasing number of biogeochemical or biological properties begin to be accessible from remote sensing (Ciotti and Bricaud, 2006; Uitz et al., 2006; Loisel et al., 2006; Siegel et al., 2002). The bio-optical models allowing the extraction of such “new products” still require validation and eventually refinement. A strong component of BIOSOPE was thus dedicated to optical and bio-optical studies. It was planned to elaborate a self consistent data base covering the complete range of trophic, biogeochemical and optical conditions that can be observable in open-ocean waters. Such a database would be invaluable to test and refine bio-optical models and eventually will allow identifying any peculiarity of the SEP with regard to other open ocean environments.

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### 3 Implementation of the cruise

The BIOSOPE cruise took place during austral summer of 2004 (26 October–11 December), during a moderate phase of the El Niño Southern Oscillation ENSO (see: <http://www.ncdc.noaa.gov/oa/climate/research/2004/ann/enso-monitoring.html>). The ~8000 km transect, investigated with the French Research Vessel l'Atalante, started west of the Marquesas archipelago and ended off coastal waters of Chile (Fig. 2). Along this transect, two main types of stations (Table 1) were occupied, the so-called “short” and “long” stations.

The occupation of short stations, on a daily basis, had to cope with two main constraints. Sampling for biogeochemical flux measurements performed under simulated in situ conditions (e.g. production of the various biological stocks, nutrient assimilation) had to be performed two hours before sunrise while optical measurements had to be performed around noon in phase with satellite overpasses so that they could be used for ocean colour satellite validation. Consequently, the short stations generally consisted of two sub-stations, station StA (before sunrise) and StB (around noon), generally spaced by ~40 miles (~4 sailing hours apart). Certain measurements (CTD, other sensors, and some chemical measurements, e.g. nutrient and pigment concentrations) were systematically performed at both stations StA and StB. Their analysis confirm that, over the whole transect, the variability of StA vs StB remains extremely small (except for stations StA20 and StB20 in the vicinity of the Chilean coast) in comparison to the inter-station variability.

Six long stations were investigated for period longer than two days (Table 1), allowing the deployment of drifting mooring for sediment traps and production lines, and high frequency (3 h) repetitive sampling through the CTD-rosette. The position of four long stations was determined using real-time ocean colour data (SeaWiFS, MODIS, MERIS) looking for the highest (MAR, west of Marquesas Island; UPW, UPX for the upwelling conditions off the Chile), and the lowest (GYR in the center of the South Pacific Gyre) surface Chl-*a* concentrations. It should be noted that the GYR station (114° W, 26° S)

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is extremely close to the location (115° W, 26° S) of the most oligotrophic area of the global ocean identified from an historical analysis of SeaWiFS ocean colour data . The reference station for High Nutrient Low Chl-*a* waters (HLN) in the subequatorial zone was set at the latitude of the Marquesas Archipelago, but east of it. Finally, the EGY station, located at the eastern border of the Gyre, was assumed to be representative of the transition zone between the South Pacific Gyre and the meridian branch of the anticyclonic circulation.

## 4 Water masses in the South East Pacific

### 4.1 The Sub Equatorial area: Marquesas to St 2 (142° W–132° W)

North of 14° S, waters are under the influence of the equatorial regime. The eastward flowing South Equatorial Counter Current (SECC, ~141° W; 8° S) is embedded in the westwards flowing South Equatorial Current (SEC, ~138° W; 9° S) (Fig. 3). At 143° W, 11° S a vein of the SEC is also recorded in the subsurface water (data not shown). The post-cruise trajectories of 3 profiling floats deployed west of 132° W confirm that the area is clearly under the influence of the SEC (Table 2). Surface waters are warm (up to 27° C) and relatively fresh (~35.6) (Fig. 4). In the Marquesas area, a rather homothermous layer of 70–100 m develops delineated by a steep thermocline, and associated with weak water column stability (Brunt-Vaisala frequency, Fig. 3). Nutrient concentrations are significant (nitrates=1.88  $\mu\text{M}$ ) in surface around the Marquesas Islands (Fig. 5) and, in spite of a strong decreasing gradient, remain detectable until 132° W (nitrates >0.3 $\mu\text{M}$ ). The distribution of in vivo fluorescence, a proxy of Chl-*a*, and of particle attenuation coefficient ( $c_p$ ), a proxy of particulate organic carbon (POC) are also uniform in this layer (Fig. 6). Moving eastwards, a sub-surface Chl-*a* maximum develops at ~70m that lies on the pycnocline (station HLN-St1), while  $c_p$  remains homogenous in this layer. Oligotrophic conditions start at St2 with the presence of a deep chlorophyll maximum (DCM) located at ~120 m. Below (>300 m), a noticeable signal of

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Chl-*a* fluorescence higher than the other surface values observed during the transect is clearly associated with suboxic ( $<50 \mu\text{mole kg}^{-1}$ ) conditions (Fig. 6b). This relative oxygen minimum likely reflects the signature of a north-westwards propagation of the oxygen minimum zone developing along South America (Fiedler and Talley, 2006). At this location, these suboxic conditions might also affect the degradation of particulate matter sinking from the upper layers.

4.1.1 The transition zone between the sub Equatorial area and the SPG (St 2 to St 6:  $132^\circ \text{W}$ – $123^\circ \text{W}$ ).

In the surface/sub-surface layer of this area, the so-called South Tropical Surface Waters (STSW) (Fiedler and Talley, 2006) are clearly signed by a very high salinity (Figs. 4b and 7) actually the highest of the Pacific (Tomczak and Godfrey, 2001), with the 36.5 isohaline nearly outcropping the surface at  $128^\circ \text{W}$ ,  $15^\circ \text{S}$ . These salty waters result from evaporation strongly exceeding precipitation (Tomczak and Godfrey, 2001). The area does not present any noticeable currents (Fig. 3), which is confirmed by the moderate displacement of the two profiling floats deployed in this region (Table 2, floats #2341, #2342). This zone is also characterized by increasing oligotrophic conditions with the deepening of the nutricline. Nitrates are absolutely depleted ( $<3 \text{ nM}$ ) in the 0–100 m water column, while phosphate and silicate are still detectable (0.1 and  $1 \mu\text{M}$ , respectively). The DCM deepens very markedly eastward ( $\sim 170 \text{ m}$  at station 6). This is associated with a strengthening of the permanent thermocline, a consequence of the deepening of the sub-surface isotherms (e.g. the  $15^\circ \text{C}$  isotherms deepens from 240 to 320 m). In the deeper layer ( $>300 \text{ m}$ ) the fluorescence signal as well as the suboxic conditions observed north-westwards are clearly vanishing.

4.1.2 The central part of the SPG: (St 6 to St 13:  $123^\circ \text{W}$ – $101^\circ \text{W}$ )

The central part of the gyre is characterized by the strongly stratified Eastern South Pacific Central Waters (ESPCW) (Emery and Meincke, 1986) that cover a wide range

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of temperature and salinity values (Fig. 7) and correspond to the water masses associated with the permanent thermocline (Tomczak and Godfrey, 2001). This area is delineated by extremely low levels of Chl-*a* fluorescence in the surface layer ( $\sim 0.02 \text{ mg Chl-}a \text{ m}^{-3}$ ) (Fig. 6) as well as by an extremely deep DCM which lies in the 160–200 m range over a distance of  $\sim 2500 \text{ km}$ . The DCM position appears to be mainly driven by the density field (the base of the DCM follows the  $26 \text{ kg m}^{-3}$  isopycnal). The lowest levels of  $c_p$  in the 0–200 m layer are recorded in this region, especially at station 6, 7 and 8. No inorganic nitrogen is a priori available for the biological production throughout the 0–150 m water column (nitrates  $< 3 \text{ nM}$ ). In contrast, both phosphates and silicates are always present at significant concentrations ( $0.1$  and  $1 \mu\text{M}$ , respectively). In the eastern part of the zone, on each side of  $100^\circ \text{ W}$ , two eastward flowing current veins probably represent the signature of the South Pacific Current (west wind drift) which splits in two bands (Stramma et al., 1995) and delineates the northern limit of the Subtropical Front, also identified by the strong salinity gradient at  $\sim 100^\circ \text{ W}$ . Mesoscale features related to Rossby waves (e.g. Wang et al., 1998) may also explain these current patterns. The six profiling floats deployed in this zone remained in a very restricted region (displacement of only a few degrees in latitude or longitude) over a two-year period (Table 2).

#### 4.1.3 The transition zone between the SPG and the coastal upwelling area (St 13 to St 19: $100\text{W}^\circ$ – $81^\circ \text{ W}$ )

East of  $100^\circ \text{ W}$ , the transition zone between the salty ESPCW and the waters influenced by fresher Subantarctic Surface Waters (SASW) (Emery and Meincke, 1986; Tomczak and Godfrey, 2001) (Fig. 7) clearly delineates the core of the subtropical front (Chaigneau and Pizarro, 2005b). This zone also corresponds to a shoaling of the DCM which is located at 80 m at EGY station ( $91^\circ \text{ W}$ ;  $32^\circ \text{ S}$ ) and of the nutriclines (e.g. the  $0.1 \mu\text{M NO}_3$  isoline rises from 160 m at St 13 to 30 m at EGY). East of EGY (stations 17 to 19) HLNC conditions are observed with surface nitrates  $> 2.5 \mu\text{M}$  and surface Chl-*a*  $< 0.2 \text{ mg m}^{-3}$  ( $< 0.1$  for St 19). At this location, a tongue of low salinity waters observed at a depth of 300 m progressively uplifts in an easterly direction and outcrops

at the surface (salinity 34) at 78° W 33° S. Here, the waters present the highest oxygen saturation levels of the whole transect. This location, at the south-eastern edge of the SPG, corresponds to the source of the South Pacific Intermediate Water, ESPIW, (Emery and Meincke, 1986), which then spreads north-westwards into the intermediate layer of the SPG (Schneider et al., 2003) and transfers the surface water properties of the waters off central Chile to tropical latitudes.

#### 4.1.4 The coastal upwelling area (St 19 to UPX: East of 81° W)

East of 78° W, the ESPIW lies above the relatively saltier (>34.5) Equatorial Subsurface Water (EESW) (Blanco et al., 2001) which extends in the 100–400 m range and is part, especially when approaching the Chilean coast, of the poleward Peru-Chile undercurrent (PCUC). The PCUC presents two veins at this location (Fig. 3) that have their velocity maximum at ~250 m. Two (0–500 m) veins of the equatorward PCC are also embedded with the PCUC illustrating the complexity of the circulation in the water off Chile (Shaffer et al., 1995). The only float deployed in this area (~76° W; #2358, Table 2) presented a clear northern drift during the December 2004–December 2006 period. In the surface layer, the shoaling and the narrowing of the isotherms allow the delineation of a reduced mixed layer (~15–20 m) where the highest nutrient concentrations (nitrates up to 15  $\mu\text{M}$  in surface) as well as the highest Chl-*a* fluorescence (corresponding to 3 mg Chl-*a*  $\text{m}^{-3}$ ) and  $c_p$  signal are recorded. The intermediate layer where PCUC predominates is clearly associated with suboxic conditions and with a very significant signal of “deep” Chl-*a* fluorescence.

## 5 Special issue presentation

The goal of this special issue is to present the knowledge gained concerning the South East Pacific based on the large dataset acquired during the BIOSOPE cruise. Although a disciplinary approach around three main topics (biology, biogeochemistry and optics)

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was adopted for data acquisition, many questions relevant to this project have benefited from multidisciplinary. The cruise strategy as well as on-board experimental design has been organized to promote and favour such a synergic approach. Therefore, even if papers from this special issue (and for some others published elsewhere) are introduced below according to disciplines, most of the contributions have taken advantage of this collective effort so that paper scopes are often multidisciplinary too. Finally, it should be noted that some papers compares data acquired within the SEP with data acquired with similar techniques in other oceanic provinces. These papers represent a first step in revealing the existence (or the lack of) specificities of SEP waters with respect to other environments.

## 5.1 Biology and biodiversity

The very large trophic gradient that was extensively sampled during the BIOSOPE cruise offered a unique opportunity to understand better how the structure of biological communities in the open ocean adapts to varying nutrient conditions, with a specific focus on the extremely oligotrophic conditions of the central SPG. It is expected that under such conditions, the community structure shifts towards very small cell sizes and that the microbial loop becomes dominant. However, whether the extreme conditions (e.g. depth of the DCM and relative isolation of the SPG waters) have favoured the establishment of yet unknown communities and micro-organisms is an intriguing possibility.

The autotrophic community was analysed in detail using different approaches, some of which are quite novel. The overall distribution of phytoplankton communities was assessed from HPLC pigment signatures and compared to distribution modelled from statistical relationships for the global ocean in order to highlight SEP peculiarities (Ras et al., 2007). Grob et al. (2007) relied on flow cytometry to determine the distribution in picoplankton and its impact on POC and on the particle attenuation coefficient. Masquelier and Vaulot (2007) used epifluorescence microscopy to look at picoplanktonic groups such as cyanobacteria (in particular those forming colonies), autotrophic

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picoeukaryotes, but also at larger protests such as ciliates and dinoflagellates (in particular some displaying intriguing green autofluorescence that are particularly abundant in the area). The diversity of picophytoplankton was assessed by novel culturing approaches (Le Gall et al., 2007). Beaufort et al. (2007) investigated another important autotrophic group, the coccolithophorids, and in particular their impact on calcite production in the SEP.

With respect to larger autotrophs, the relation between the physico-chemical conditions and the composition of the diatom assemblage in two different HNLC conditions, the tropical central Pacific and the offshore Peru-Chile Current, was examined by Gómez et al. (2007). The contributions to the knowledge of dinoflagellates, a group of especial relevance in warm oligotrophic ocean, have been numerous and has been presented in other journals than this special issue (Gómez, 2006, 2007b, d, e; Gómez and Furuya, 2007).

During BIOSOPE, heterotrophic bacteria have been studied in great details from the point of view of their production relative to primary production along the trophic gradient (Van Wambeke et al., 2007b) and of the factor(s) regulating this production both spatially (Van Wambeke et al., 2007a) and temporally, with a special focus on the diel cycle an UV impact at long stations (Van Wambeke et al., 2007c). The relationship between bacteria and available sugars was investigated by Sempéré et al. (2008), while a new approach, based on the incorporation of labelled phosphorus into phospholipids, allowed showing that heterotrophic bacteria play a critical role in the phosphorus cycle, especially in the SPG (Van Mooy et al., 2008). The very specific bacterial community of the surface microlayer has been assessed both from the phylogenetic and activity points of view by (Obernosterer et al., 2007).

Heterotrophic eukaryotes play a key role in recycling the organic matter from the microbial food web. The diversity of tintinnids was investigated with respect to the phytoplankton resource by (Dolan et al., 2007) and follows on other recent descriptions of this microzooplankton group in the SEP (Dolan, 2006). The symbiotic association between the diatom *Chaetoceros* and the tintinnid *Eutintinnus* was investigated as an

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example of the microplankton adaptation to the severe oligotrophic conditions (Gómez, 2007a). The microscope observations have also allowed elucidating the mysterious nature of the consortia of the protozoan *Solenicola* and the frustule of the diatom *Lep-  
5 tocyllindrus mediterraneus* (Gómez, 2007c). Larger mesozooplankton is the focus of the work by Carlotti (2007).

### 5.1.1 Biogeochemistry

Two “greenhouse” gases have been the subject of dedicated studies during BIOSOPE. Azouzi et al. (2007) provides an analysis of anthropogenic CO<sub>2</sub> penetration in the SEP and compares it with earlier estimates of anthropogenic tracers. Charpentier  
10 et al. (2007) focuses on the processes of N<sub>2</sub>O production and shows differences in the saturation level as well as in the source of this gas according to the hydrodynamic and associated trophic regimes considered. Nitrogen cycle was further addressed in Raimbault and Garcia (2007) who shows, in particular, that nitrogen fixation, while extremely low in the SPG, nevertheless sustains most of new production in this area.  
15 This extremely weak nitrogen fixation is supported by extremely low abundance of the cyanobacterial nifH gene (Bonnet et al., 2007). Iron which is an essential element for life, in particular for nitrogen fixation, is vanishing low in the top 350 m (~0.1 nM) (Blain et al., 2007) which confirms the extremely low atmospheric deposition recorded during the transect by Wagener et al. (2007). As a result of low iron concentration and low  
20 nitrogen fixation phosphate concentration in surface layers was always above a threshold of 0.1 μM (Raimbault et al., 2007) and thus never constituted a limiting element for phytoplankton growth (Moutin et al., 2008), even in the hyperoligotrophic conditions of the SPG. A new method based on simultaneous spiking of water samples with <sup>33</sup>P and <sup>14</sup>C allowed to measure autotrophic and bacterial fraction in various size fraction and  
25 suggest that the microbial community turns over very slowly (Duhamel et al., 2007). Primary production measurement rates using <sup>13</sup>C labelling confirms extremely low rates in the core of the SPG (100 mg C<sup>-2</sup> d<sup>-1</sup>). These results are however partly contradictory with the bacterial carbon demand estimated by Van Wambeke et al. (2007b).

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An optical technique based on the diel cycle in the attenuation coefficient (Claustre et al., 2007) suggests that rates of community production are much higher than expected and could be due to exceptional DOC release during photosynthetic processes. This DOC release, unfortunately not measured during the cruise, might explain part of the discrepancies between bacterial carbon demand and production rates using various methods.

A new approach of particle flux determination, based on the optical quantification of particles forming the stock of large particulate matter (LPM $>90\ \mu\text{m}$ ) was validated against the sediment trap fluxes at the six long stations and then applied along the whole BIOSOPE transect (Guidi et al., 2007). Another approach developed by Carlotti et al. (2007) make use of mesozooplankton biomass size structure determined from optical plankton counter to derive size-based fluxes associated to this community. Significant diel variations in concentration and spectral slope of the particle size distribution were observed for particles smaller than  $100\ \mu\text{m}$ , but not for LPM (Stemmann et al., 2007). Finally, detailed characterization of the nature of the particle material was achieved using a combination of lipid biomarkers associated with their carbon isotopic ( $\delta^{13}$ ) composition (Tolosa et al., 2007).

## 6 Optics and bio-optics

The hyperoligotrophic conditions encountered in the centre of the gyre during BIOSOPE offered the opportunity to improve our knowledge of pure water optical properties by (1) setting upper limits to pure water absorption, especially in the UV range (Morel et al., 2007b) and (2) investigating in great detail backscattering properties by pure water (Twardowski et al., 2007) and permitting the evaluation of different values proposed for this coefficient. Furthermore (Morel et al., 2007a) highlight optical singularities of Pacific waters when compared to Mediterranean waters, especially in the short wavelength domain including UV and for low Chl-*a*. The high penetration of UV radiation in the SPG was also confirmed by (Tedetti et al., 2007). Additionally,

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previously published model of the so-called bi-directional effects in water upwelling radiances was validated for the first time in these extremely clear waters (Voss et al., 2007). Supplementing the BIOSOPE data-base with data from the eastern Atlantic (Stramski et al., 2007) evaluate several approaches for quantifying particulate organic carbon concentration in surface waters from the determination of certain optical properties. Huot et al. (2007b) built upon the unique optical measurements of Stramski et al. (2008) and Twardowski et al. (2007) to establish a relationship between the spectral backscattering coefficient and the Chl-*a* concentration and to examine the variations of the backscattering ratio with trophic status. Huot et al. (2007a) evaluates the validity of various optical proxies of phytoplankton biomass, either derived from space or from in situ measurement, for the estimation of photo-physiological parameters used in bio-optical modelling of primary production. Finally optical measurements made during BIOSOPE were also used by Uitz et al. (2007) to describe the photosynthetic characteristics of different phytoplankton groups and parameterize them for use in primary production models, and by Brown et al. (2007) to evaluate a new approach to estimate, from remotely sensed ocean color, the concentration of substances other than Chl-*a*.<sup>1</sup>

## 7 Final note: the evaluation process and the role of guest editors

All papers published in this special issue have been evaluated according to the normal procedure used for regular issues of Biogeosciences. In particular, none of the three BIOSOPE guest editors did intervene at any stage of manuscript evaluation. Their role was restricted to the revision of manuscripts prior to submission and to the coordination of the various submissions (ensuring consistency between the various manuscripts). For each manuscript, the BIOSOPE guest editors have provided a list of four to five potential referees which may or may not have been used by the regular Biogeosciences

<sup>1</sup>Brown, C., Huot, Y., Werdell, P. J., Gentili, B., and Claustre, H.: The origin and global distribution of second order variability in satellite ocean color, Remote Sensing of Environment, submitted, 2007.

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editors. Therefore, papers published in this BIOSOPE special issue comply with the general quality standards of Biogeosciences.

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**Table 1.** Location and date of the various stations investigated along the BIOSOPE transect.

Station	Date	Longitude	Latitude
MAR1	26 Oct 2004	-141.23	-8.42
MAR2	27 Oct 2004	-141.27	-8.39
MAR3	28 Oct 2004	-141.26	-8.34
MAR4	29 Oct 2004	-141.27	-8.32
HNL1	31 Oct 2004	-136.86	-9.00
HNL2	1 Nov 2004	-136.89	-9.01
HNL3	2 Nov 2004	-136.98	-9.06
STA1	3 Nov 2004	-134.35	-11.51
STB1	3 Nov 2004	-134.10	-11.74
STA2	4 Nov 2004	-132.39	-13.31
STB2	4 Nov 2004	-132.11	-13.55
STA3	5 Nov 2004	-130.38	-15.13
STB3	5 Nov 2004	-129.93	-15.53
STA4	6 Nov 2004	-128.38	-16.87
STB4	6 Nov 2004	-127.97	-17.23
STA5	7 Nov 2004	-125.95	-18.51
STB5	7 Nov 2004	-125.55	-18.75
STA6	8 Nov 2004	-123.41	-20.13
STB6	8 Nov 2004	-122.89	-20.45
STA7	9 Nov 2004	-120.86	-21.75
STB7	9 Nov 2004	-120.38	-22.05
STA8	10 Nov 2004	-118.33	-23.29
STB8	10 Nov 2004	-117.89	-23.55
STA9	11 Nov 2004	-116.02	-24.71
GYR1	12 Nov 2004	-114.00	-26.00
GYR2	12 Nov 2004	-113.99	-26.00
GYR3	13 Nov 2004	-114.02	-26.02

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**Table 1.** Continued.

Station	Date	Longitude	Latitude
GYR4	14 Nov 2004	-114.02	-26.03
GYR5	15 Nov 2004	-114.01	-26.06
GYR6	16 Nov 2004	-113.99	-26.07
STA10	17 Nov 2004	-110.67	-26.85
STA11	20 Nov 2004	-107.59	-27.70
STB11	20 Nov 2004	-107.29	-27.77
STA12	21 Nov 2004	-104.75	-28.44
STB12	21 Nov 2004	-104.31	-28.54
STA13	22 Nov 2004	-101.83	-29.15
STB13	22 Nov 2004	-101.48	-29.23
STA14	23 Nov 2004	-98.87	-29.92
STB14	23 Nov 2004	-98.39	-30.04
STA15	24 Nov 2004	-95.83	-30.70
STB15	24 Nov 2004	-95.43	-30.79
STA16	25 Nov 2004	-93.00	-31.42
EGY1	25 Nov 2004	-91.47	-31.82
EGY2	26 Nov 2004	-91.47	-31.82
EGY3	27 Nov 2004	-91.44	-31.85
EGY4	28 Nov 2004	-91.41	-31.86
EGY5	29 Nov 2004	-91.41	-31.90
EGY6	30 Nov 2004	-91.41	-31.90
STA17	1 Dec 2004	-87.43	-32.30
STB17	1 Dec 2004	-86.78	-32.40
STA18	2 Dec 2004	-84.21	-32.67
STB18	2 Dec 2004	-84.07	-32.68

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**Table 1.** Continued.

Station	Date	Longitude	Latitude
STA19	3 Dec 2004	-81.64	-32.95
STB19	3 Dec 2004	-81.20	-33.02
STA20	4 Dec 2004	-78.37	-33.32
STB20	4 Dec 2004	-78.12	-33.35
STA21	5 Dec 2004	-75.84	-33.58
UPW1	6 Dec 2004	-73.37	-34.00
UPW2	7 Dec 2004	-73.39	-33.98
UPW3	8 Dec 2004	-73.34	-33.86
UPX1	9 Dec 2004	-72.42	-34.51
UPX2	10 Dec 2004	-72.43	-34.58
UPX3	11 Dec 2004	-72.49	-34.69

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**Table 2.** ARGO floats deployed along the BIOSOPE transect. The left column refers to the 5 main zones defines in the paper.

	Float number/ http link	Date of deployment	Deployment location	
			Latitude	Longitude
Zone 1	<a href="http://sio-argo.ucsd.edu/0049910c.html">2339/http://sio-argo.ucsd.edu/0049910c.html</a>	5 Nov 2004	13.58	132.10
Zone 1	<a href="http://sio-argo.ucsd.edu/0049909c.html">2338/http://sio-argo.ucsd.edu/0049909c.html</a>	4 Nov 2004	12.00	134.09
Zone 1	<a href="http://sio-argo.ucsd.edu/0045842c.html">2336/http://sio-argo.ucsd.edu/0045842c.html</a>	26 Okt 2004	10.05	142.28
Zone 2	<a href="http://sio-argo.ucsd.edu/0049913c.html">2342/http://sio-argo.ucsd.edu/0049913c.html</a>	8 Nov 2004	18.82	125.45
Zone 2	<a href="http://sio-argo.ucsd.edu/0049913c.html">2341/http://sio-argo.ucsd.edu/0049913c.html</a>	7 Nov 2004	17.23	128.09
Zone 3	<a href="http://sio-argo.ucsd.edu/0049919c.html">2348/http://sio-argo.ucsd.edu/0049919c.html</a>	23 Nov 2004	29.30	101.50
Zone 3	<a href="http://sio-argo.ucsd.edu/0049918c.html">2347/http://sio-argo.ucsd.edu/0049918c.html</a>	21 Nov 2004	27.73	107.40
Zone 3	<a href="http://sio-argo.ucsd.edu/0049917c.html">2346/http://sio-argo.ucsd.edu/0049917c.html</a>	17 Nov 2004	26.14	113.93
Zone 3	<a href="http://sio-argo.ucsd.edu/0049916c.html">2345/http://sio-argo.ucsd.edu/0049916c.html</a>	11 Nov 2004	23.62	117.95
Zone 3	<a href="http://sio-argo.ucsd.edu/0049915c.html">2344/http://sio-argo.ucsd.edu/0049915c.html</a>	10 Nov 2004	22.11	120.36
Zone 3	<a href="http://sio-argo.ucsd.edu/0049914c.html">2343/http://sio-argo.ucsd.edu/0049914c.html</a>	9 Nov 2004	20.38	122.83
Zone 4	<a href="http://sio-argo.ucsd.edu/0049927c.html">2356/http://sio-argo.ucsd.edu/0049927c.html</a>	4 Dec 2004	33.06	81.18
Zone 4	<a href="http://sio-argo.ucsd.edu/0049926c.html">2355/http://sio-argo.ucsd.edu/0049926c.html</a>	3 Dec 2004	32.79	84.04
Zone 4	<a href="http://sio-argo.ucsd.edu/0049925c.html">2354/http://sio-argo.ucsd.edu/0049925c.html</a>	2 Dec 2004	32.30	86.79
Zone 4	<a href="http://sio-argo.ucsd.edu/0049924c.html">2353/http://sio-argo.ucsd.edu/0049924c.html</a>	1 Dec 2004	31.86	91.41
Zone 4	<a href="http://sio-argo.ucsd.edu/0049923c.html">2352/http://sio-argo.ucsd.edu/0049923c.html</a>	26 Nov 2004	31.43	92.88
Zone 4	<a href="http://sio-argo.ucsd.edu/0049921c.html">2350/http://sio-argo.ucsd.edu/0049921c.html</a>	25 Nov 2004	30.77	95.45
Zone 4	<a href="http://sio-argo.ucsd.edu/0049920c.html">2349/http://sio-argo.ucsd.edu/0049920c.html</a>	24 Nov 2004	30.09	98.32
Zone 5	<a href="http://sio-argo.ucsd.edu/0049929c.html">2358/http://sio-argo.ucsd.edu/0049929c.html</a>	6 Dec 2004	33.62	75.95

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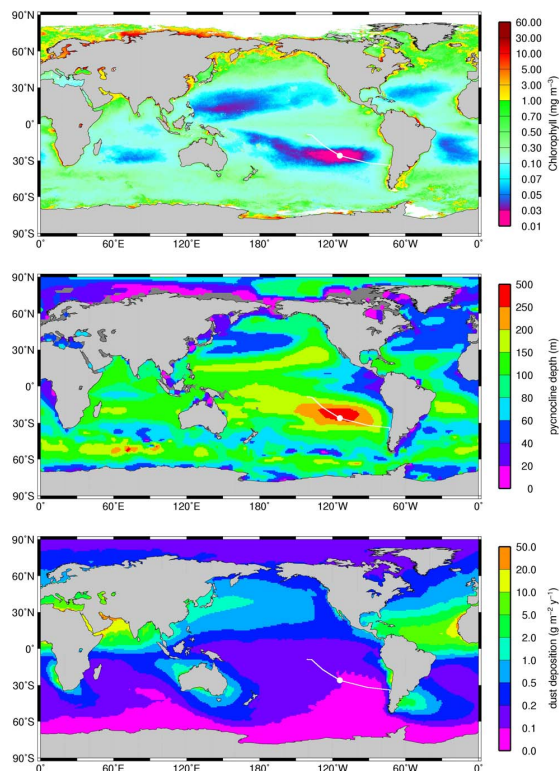
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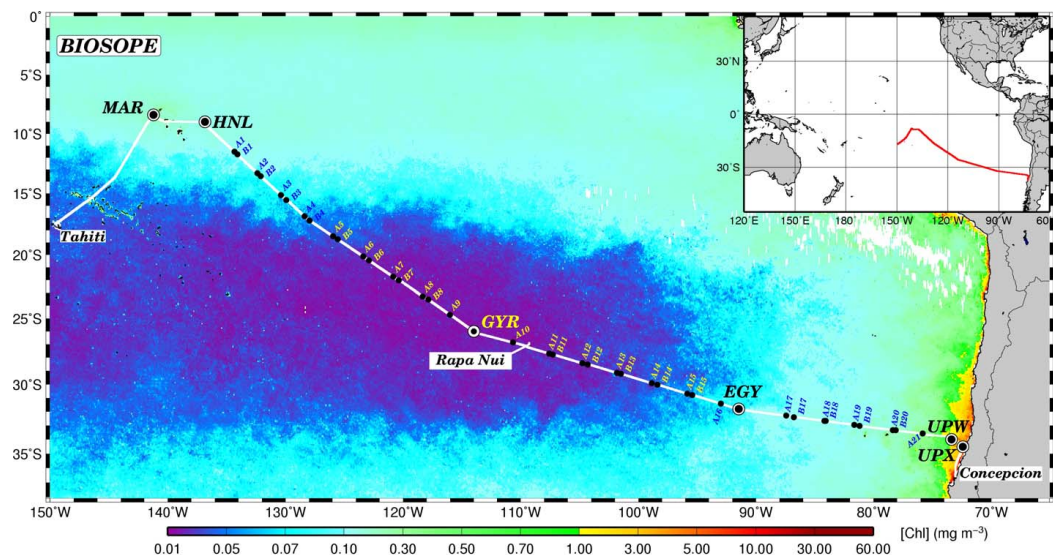
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**Fig. 1.** Global context of the BIOSOPE cruise in the South East Pacific. **(a)** Annual composite image of SeaWiFS derived Chl-*a* for year 2000. **(b)** Distribution of the pycnocline depth (adapted from Fiedler and Talley, 2006). **(c)** Dust deposition flux at the ocean surface (adapted from Mahowald et al., 2005). The white line identifies the cruise track of the BIOSOPE cruise. The white circle corresponds to the GYR station located in the core of the SPG and investigated over a 5-day period.

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**Fig. 2.** Transect of the BIOSOPE cruise superimposed on a SeaWiFS composite image of Chl-*a* concentration in the upper layer for November–December 2004. The two main types of station, long and short, are indicated. The six long stations investigated for a period longer than 2 days are identified by a 3 letter code (see text). The 21 short stations are generally split into substations **(a)** (early morning) and **(b)** (around noon). Four stations (referred as A9, A10, A16 and A21) did not follow this scheme. Note that the actual denomination for the short stations begins with St (Table 1), but the code has been simplified on this map to facilitate reading.

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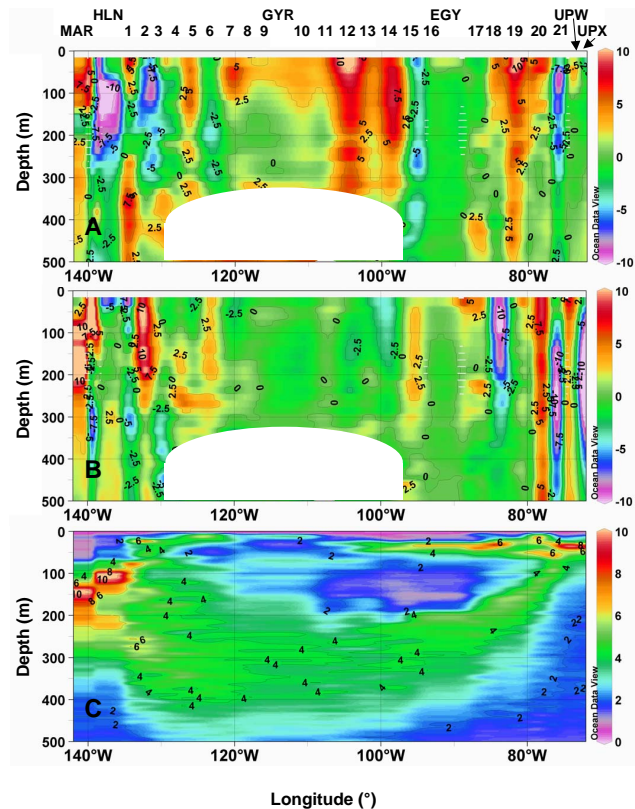
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**Fig. 3.** Currents and Brunt-Väisälä frequency along the BIOSOPE transect. **(a)** Zonal component ( $\text{cm s}^{-1}$ ): purple is westward and red is eastward. **(b)** Meridional component ( $\text{cm s}^{-1}$ ): red is northward and purple is southward. **(c)** Brunt-Väisälä frequency ( $\text{cyc h}^{-1}$ ). No current was monitored in the masked area (no echo for the 75 KHz RDI ADCP because of the very low level of particles in this layer).

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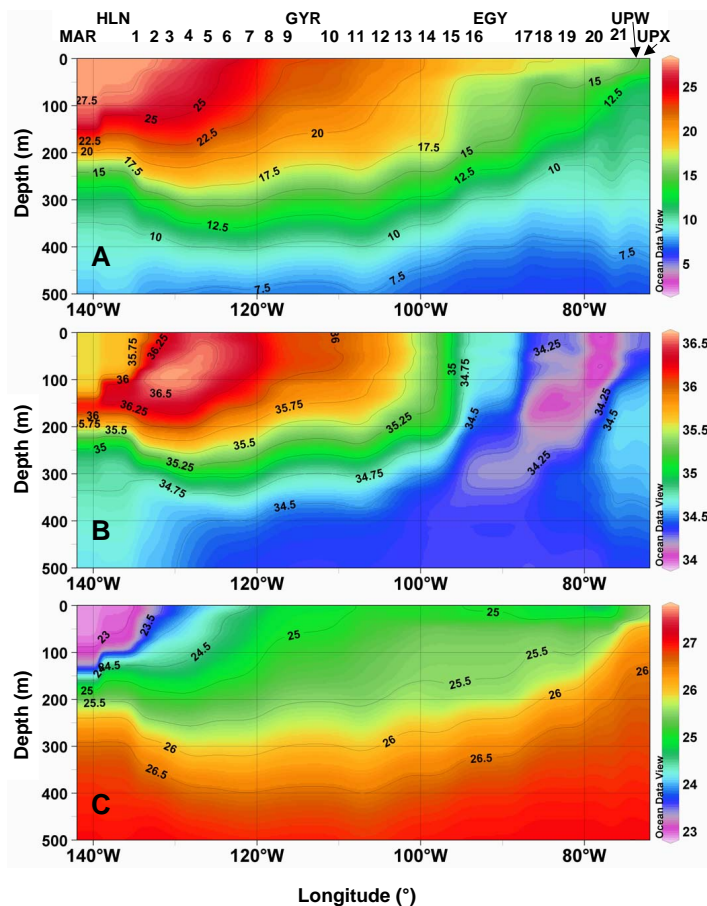
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**Fig. 4.** Hydrological section along the BIOSOPE transect. **(a)** Potential temperature ( $^{\circ}\text{C}$ ). **(b)** Salinity. **(c)** Potential density ( $\text{kg m}^{-3}$ ).

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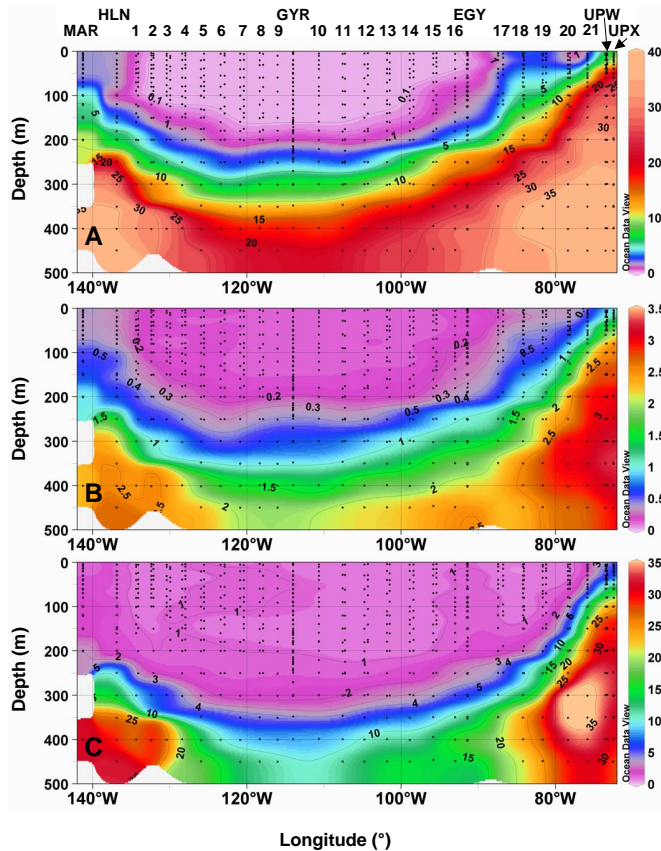
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**Fig. 5.** Nutrient concentrations ( $\mu\text{M}$ ) along the BIOSOPE transect. **(a)** Nitrate ( $\text{NO}_3$ ). **(b)** Phosphates ( $\text{PO}_4$ ). **(c)** Silicates ( $\text{Si}(\text{OH})_4$ ).

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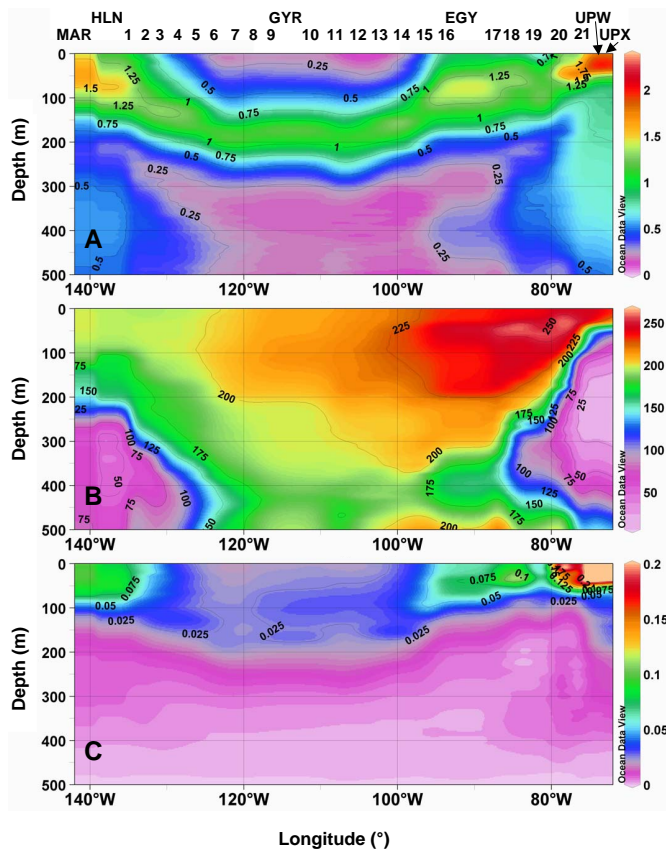
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**Fig. 6.** Distribution of specific biogeochemical and optical properties along the BIOSOPE transect. **(a)** In vivo fluorescence of Chl-*a* (V). **(b)** Oxygen concentration ( $\mu\text{M kg}^{-1}$ ). **(c)** attenuation coefficient,  $c_p$  ( $\text{m}^{-1}$ ). The attenuation coefficient data have been processed as described in Claustre et al. (2007), by correcting for deep (450–500 m) values.

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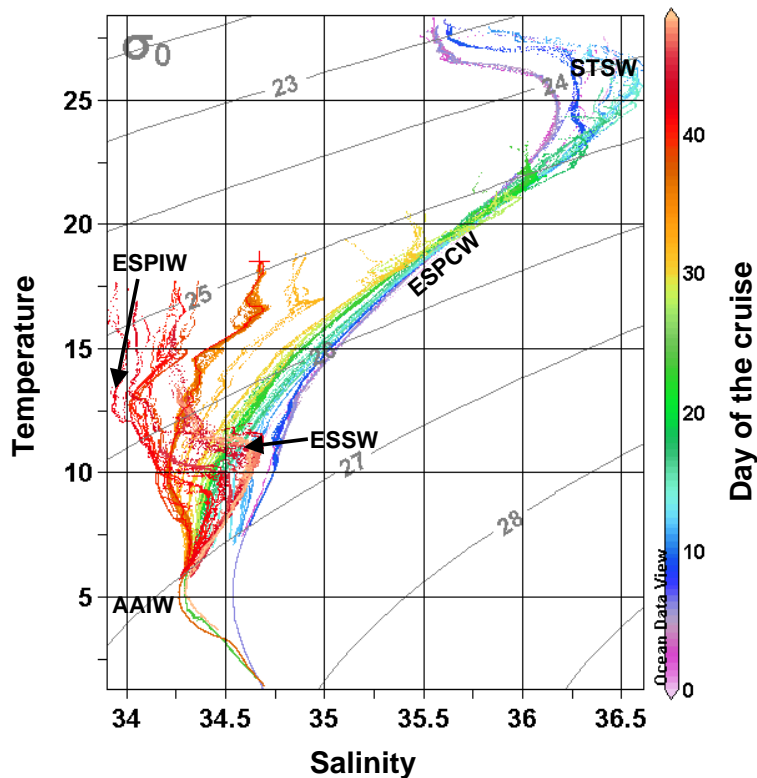
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**Fig. 7.** Temperature-salinity diagram and identification of the main waters masses encountered along the BIOSOPE transect. STSW: South Tropical Surface Waters; ESPCW: Eastern South Pacific Central Waters; ESPIW: South Pacific Intermediate Water; ESSW: Equatorial Subsurface Water; AAIW: Antarctic Intermediate Waters. The colour code refers to the day of the mission starting near Marquesas Archipelago (purple) and ending along the Chilean Coast (red). 224 CTD casts are used for this plot. Except for four deep casts (down to the sea floor), the other ones were performed down to 500 m.

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