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Preface: The Oligotrophy to the UTRa-oligotrophy PACific Experiment (OUTPACE cruise, 18 February to 3 April 2015)

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Abstract. The overall goal of OUTPACE (Oligotrophy to UTRa-oligotrophy PACific Experiment) was to obtain a successful representation of the interactions between planktonic organisms and the cycle of biogenic elements in the western tropical South Pacific Ocean across trophic and N₂ fixation gradients. Within the context of climate change, it is necessary to better quantify the ability of the oligotrophic ocean to sequester carbon through biological processes. OUTPACE was organized around three main objectives, which were (1) to perform a zonal characterization of the biogeochemistry and biological diversity of the western tropical South Pacific during austral summer conditions, (2) to study the production and fate of organic matter (including carbon export) in three contrasting trophic regimes (increasing oligotrophy) with a particular emphasis on the role of dinitrogen fixation, and (3) to obtain a representation of the main biogeochemical fluxes and dynamics of the planktonic trophic network. The international OUTPACE cruise took place between 18 February and 3 April 2015 aboard the RV *L'Atalante* and involved 60 scientists (30 onboard). The west–east transect covered ~4000 km from the western part of the Melanesian archipelago (New Caledonia) to the western boundary of the South Pacific gyre (French Polynesia). Following an adaptive strategy, the transect initially designed along the 19° S parallel was adapted along-route to incorporate information coming from satellite measurements of sea surface temperature, chlorophyll *a* concentration, currents, and diazotroph quantification. After providing a general context and describing previous work done in this area, this introductory paper elucidates the objectives of OUTPACE, the implementation

plan of the cruise and water mass and climatological characteristics and concludes with a general overview of the other papers that will be published in this special issue.

1 General context

The additional carbon dioxide (CO₂) in the atmosphere, mainly resulting from fossil fuel emissions linked with human activities (anthropogenic CO₂), is the main cause of global warming (Fifth Assessment Report – Climate Change 2013 – IPCC). The ocean has acted thus far as a major sink of anthropogenic CO₂ (Sabine et al., 2004), preventing greater CO₂ accumulation in the atmosphere and therefore a greater increase in the Earth's temperature. The biological pump (Fig. 1), the process by which carbon (C) is transferred from the upper to the deep ocean by biological processes, provides the main explanation for the vertical gradient of C in the ocean. Its strength and efficiency depends upon the complex balance between organic matter production in the photic zone and its remineralization in both the epipelagic and mesopelagic zones. Before present and for tens of thousands of years, the biological pump was thought to be in an equilibrium state with an associated near-zero net exchange of CO₂ with the atmosphere (Broecker, 1991; Murnane et al., 1999). Climate alterations then started to disrupt this equilibrium and the expected modification of the biological pump will probably considerably influence oceanic C sequestration (and therefore global warming) in future decades (Sarmiento and Gruber, 2006). The long-term decrease in phosphate

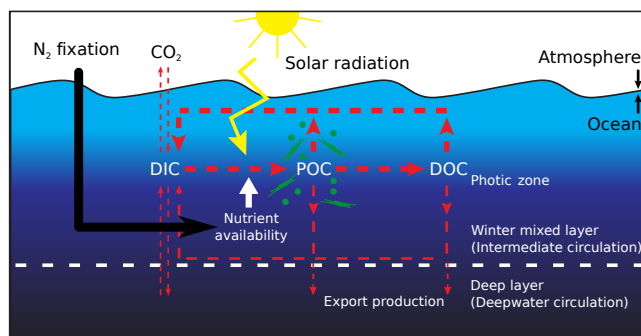


Figure 1. Major C fluxes for a biological pump budget and the main role of N_2 fixation. Biological pump is the C transfer into the ocean interior by biological processes. DIC is dissolved inorganic C, POC is particulate organic C, and DOC is dissolved inorganic C. See Moutin et al. (2012) for a detailed description.

availability, and the shift from primary production previously limited by nitrogen (N) and now by phosphorus (P), associated with increasing inputs of N by dinitrogen (N_2) fixation observed at the Hawaii Ocean Time-series (HOT) station in the north Pacific gyre (NPG) (Karl et al., 1997; Karl, 2014), appear as a first example of biological pump alteration.

The input of new N to the surface ocean through biological N_2 fixation represents a major link between the C and N biogeochemical cycles, relating upper-ocean nutrient availability with the biological pump and ultimately the ocean and climate. This link was recently shown to play a central role in previous natural climate changes over long timescales (Galbraith et al., 2013). It is nevertheless clear that expected climate change due to anthropogenic atmospheric CO_2 input may concern shorter timescales: the increase in atmospheric CO_2 over the past 200 years equals the increase in atmospheric CO_2 between glacial and interglacial periods, which took place over several thousands of years (Sarmiento and Gruber, 2006). Furthermore, enhanced stratification in the tropical and subtropical ocean resulting from global warming (Polovina et al., 2008) might decrease nutrient availability for nutrients such as nitrate (NO_3^-), potentially favoring N_2 fixation in surface waters. It may also decrease phosphate availability (Moutin et al., 2008), and in turn N_2 fixation, net primary production, and export (Karl, 2014). McMahon et al. (2015), covering the past 1000 years, argue that N_2 fixation has increased since the industrial revolution and might provide a negative feedback to rising CO_2 .

The ecosystem changes due to climate change are complex and it is therefore necessary to characterize in detail the interactions between N_2 fixation and the C cycle to obtain a precise representation of the N_2 fixation process in global biogeochemical models, leading eventually to predictions. Even if considerable scientific progress has been made over the last decades (see reviews from Sohm et al., 2011, and Zehr and Kudela, 2011), many questions remain regarding the impact of this process on biogeochemical cycles and C export.

2 The role of N_2 fixation in the oligotrophic ocean and an overview of previous cruises in the western tropical South Pacific Ocean (WTSP)

The efficiency of oceanic C sequestration depends upon many factors, among which is the availability of nutrients to support phytoplankton growth in the photic zone (Fig. 1). Large amounts of N are required for phytoplankton growth, as it is an essential component of proteins, nucleic acids, and other cellular constituents. Fixed N in the form of NO_3^- or ammonium (NH_4^+) is directly usable for growth, but concentrations are low ($< 0.1 \mu\text{mol L}^{-1}$) and often growth limiting in most of the open-ocean euphotic zone (Falkowski et al., 1998; Moore et al., 2013). Dissolved N_2 gas concentrations in seawater are in contrast very high in the euphotic zone (ca. $450 \mu\text{mol L}^{-1}$) and could constitute a nearly inexhaustible N source for the marine biota. However, only some prokaryotic organisms (bacteria, cyanobacteria, archaea), hereafter referred to as N_2 -fixing organisms (or diazotrophs), are able to use this gaseous N source since they possess the nitrogenase enzyme that breaks the triple bond between the two N atoms of the N_2 molecule and converts it into a usable form (i.e., NH_3) (Zehr et al., 1998). On the global scale, they provide $140 \pm 50 \text{ Tg N}$ per year to the surface ocean, contributing more than atmospheric and riverine N inputs (Gruber, 2004). N_2 -fixing organisms thus act as natural fertilizers and contribute to sustaining life and potential C export in coastal and oceanic environments.

Most of the surface ocean (60%; Longhurst, 1998) is comprised of low-nutrient, low-biomass oligotrophic areas, which constitute the largest coherent ecosystems on our planet. They support a large part (40%) of the photosynthetic C fixation in the ocean (Antoine et al., 1996). This C fixation is mainly performed by picoplankton (smaller than $2\text{--}3 \mu\text{m}$ in diameter) that are generally thought to represent a negligible fraction of the total particulate organic C (POC) export flux due to their small size, slow individual sinking rates, and tight grazer control that leads to high rates of recycling in the euphotic zone. Consequently, the efficiency of the biological C pump in these oligotrophic systems has long been considered to be low as the greatest proportion of fixed C is thought to be recycled in the surface layer and rapidly reexchanged with the atmosphere.

Recent studies have challenged this view and indicate that all primary producers, including picoplanktonic cells, contribute to export from the surface layer of the ocean at rates proportional to their production rates (Richardson and Jackson, 2007). Export mechanisms differ compared to larger cells as export of picoplankton is mainly due to packaging into larger particles via grazing and/or aggregation processes (Jackson, 2001; Lomas et al., 2010). More recently, Close et al. (2013) pointed out that 40–70% of picoplanktonic cells are small enough to escape detection under the most common definition of suspended particulate organic matter (POM).

In addition to submicron POM export, low $\delta^{15}\text{N}$ signatures in particles from sediment traps at the HOT station suggest that at least part of the production sustained by N_2 fixation is ultimately exported out of the photic zone (Karl et al., 1997, 2012; Scharek et al., 1999a; Scharek et al., 1999b). This may either be direct through sinking of diazotrophs, or indirect, through the transfer of diazotroph-derived N to non-diazotrophic plankton, which are subsequently exported. Karl et al. (2012) reported an efficient summer export flux of C (3 times greater than the mean wintertime particle flux) at the HOT station that was attributed to the direct export of symbiotic N_2 -fixing cyanobacteria associated with diatoms (hereafter referred to as diatom–diazotroph associations or DDAs), which have high sinking and low remineralization rates during downward transit. This result is in accordance with high export fluxes measured in the tropical North Atlantic when the diazotroph community is dominated by DDAs (Subramaniam et al., 2008; White et al., 2012). More recently, Berthelot et al. (2015) and Bonnet et al. (2016a) studied the fate of a bloom of unicellular diazotrophs from Group C (UCYN-C) during a mesocosm experiment in the New Caledonia lagoon after a phosphate enrichment and revealed that $\sim 10\%$ of UCYN-C from the water column was exported to the particle traps daily, representing as much as $22.4 \pm 5.5\%$ of the total POC exported at the height of the UCYN-C bloom. A $\delta^{15}\text{N}$ budget performed in the mesocosms confirmed the high contribution of N_2 fixation (56%; Knapp et al., 2016) to export compared to other tropical and subtropical regions where active N_2 fixation contributes 10 to 25% to export production (e.g., Altabet, 1988; Knapp et al., 2005) and exceptionally up to 92% in the Arabian Sea (Gandhi et al., 2011; Kumar et al., 2017). Mechanistically, the vertical downward flux was enabled by the aggregation of the small ($5.7 \pm 0.8\ \mu\text{m}$) UCYN-C cells into large (100–500 μm) aggregates. In addition to direct export of diazotrophs, the use of nanoSIMS (nanoscale secondary ion mass spectrometry) enabled the tracking of the fate of ^{15}N from both *Trichodesmium* (Bonnet et al., 2016b) and UCYN blooms (Berthelot et al., 2015; Bonnet et al., 2016c) and demonstrated that $\sim 8\%$ of N originating from N_2 fixation is quickly transferred to non-diazotrophic plankton, in particular diatoms, i.e., efficient C exporters to depth (Nelson et al., 1995) during *Trichodesmium* blooms (Bonnet et al., 2016b). This reveals that N_2 fixation can fuel large-size non-diazotrophic plankton growth in the water column, suggesting an indirect export pathway of organic matter sustained by N_2 fixation in the oligotrophic ocean. Most of the above-mentioned studies were performed in microcosms and mesocosms and further open-ocean studies combining the set of complementary approaches described above are needed to better assess the fate of N_2 fixation and its role in C export.

The western tropical South Pacific (WTSP) is an ideal location to study the fate of N fixed by N_2 fixation, as it is considered a hot spot of N_2 fixation in the world ocean (Bonnet et al., 2017a). While average N_2 fixation rates range from

20 to 200 $\mu\text{mol N m}^{-2} \text{d}^{-1}$ in the tropical North Atlantic (Benavides and Voss, 2015) and Pacific (Böttjer et al., 2017; Dore et al., 2002), they reach 30–5400 $\mu\text{mol N m}^{-2} \text{d}^{-1}$ (average $\sim 800 \mu\text{mol N m}^{-2} \text{d}^{-1}$) in the Solomon Sea (western part of the WTSP) (Bonnet et al., 2009, 2015), which is in the upper range of rates reported in the global N_2 fixation MAREDAT database and even surpassed its upper rates (100–1000 $\mu\text{mol N m}^{-2} \text{d}^{-1}$) (Luo et al., 2012). Very high rates have also been recently reported in the Arabian Sea (Gandhi et al., 2011; Kumar et al., 2017). High rates ranging from 151 to 703 $\mu\text{mol N m}^{-2} \text{d}^{-1}$ have also been reported off New Caledonia (Garcia et al., 2007), with seasonal variations closely linked with phosphate availability (Moutin et al., 2005; Van Den Broeck et al., 2004). The seasonal distribution of N_2 fixation is corroborated by in situ and satellite observations (Dupouy et al., 2011) of recurrent large *Trichodesmium* blooms during austral summer conditions (January–March) over the 1998–2010 period in the Melanesian archipelago (MA: New Caledonia, Vanuatu, Fiji Islands). In addition to *Trichodesmium*, which dominates the diazotroph community in the WTSP (Moutin et al., 2005; Bonnet et al., 2015), very high abundances of UCYN-B (up to 106–107 nifH copies L^{-1}) have been reported (Bonnet et al., 2015; Campbell et al., 2005; Moisander et al., 2010). The uncultivated UCYN-A (Zehr et al., 2008) also displays high abundances (105–106 nifH copies L^{-1} ; Bonnet et al., 2015; Moisander et al., 2010) around the MA, but they seem to have different ecological niches compared to *Trichodesmium* and UCYN-B (Bonnet et al., 2015; Moisander et al., 2010).

When going eastward towards the South Pacific gyre (GY), Halm et al. (2012) reported rates of 12–190 $\mu\text{mol N m}^{-2} \text{d}^{-1}$ on the western border of the gyre, and Raimbault and Garcia (2008) and Moutin et al. (2008) reported rates of $60 \pm 30 \mu\text{mol N m}^{-2} \text{d}^{-1}$ in the central gyre, indicating a decreasing gradient of N_2 fixation from west to east and low N_2 fixation rates relative to other ocean gyre ecosystems. The organisms responsible for these fluxes are different from common autotrophic diazotrophs such as *Trichodesmium* or UCYN-B, and are mainly affiliated with heterotrophic proteobacteria and low abundances of UCYN-A (Bonnet et al., 2008; Halm et al., 2012).

The west to east zonal gradient of N_2 fixation and the distinct diversity of N_2 -fixing organisms along this gradient together provide a unique opportunity to study how production, mineralization, and export of organic matter depends upon N_2 fixation in contrasting oligotrophic regimes (from oligotrophy to ultra-oligotrophy). Comparisons between different systems along a zonal gradient of trophic status and N_2 fixation will provide new insights for identifying and understanding fundamental interactions between marine biogeochemical C, N, P, silica (Si), and iron (Fe) cycles in oligotrophic ecosystems.

3 Objectives of OUTPACE

The overall goal of OUTPACE was to obtain a precise representation of the complex interactions between planktonic organisms and the cycle of biogenic elements (C, N, P, Si, Fe), considering a variety of scales, from single-cell processes to the whole WTSP Ocean. To meet this goal, the three specific objectives of OUTPACE were

1. to perform a zonal characterization of the biogeochemistry and biological diversity of the WTSP during the strongest stratified period (austral summer).
2. to study the production and fate of organic matter (including C export) of three contrasting environments (from oligotrophy to ultra-oligotrophy) with a particular emphasis on N_2 fixation.
3. to obtain a representation of the main biogeochemical fluxes (C, N, P, Si, Fe) and the dynamics of the planktonic trophic network, both in situ and by using microcosm experiments.

The detailed study of biological production and its subsequent fate at a given site implied a combination of adaptive and Lagrangian strategies. Indeed, as pointed out by d'Ovidio et al. (2015), the spatiotemporal domain of an oceanographic cruise is also one in which horizontal stirring generated by ocean circulation at the mesoscale induces strong variability in biogeochemical tracer distributions. Consequently, ephemeral and local gradients due to mesoscale activity can easily mask the large-scale gradients. Following d'Ovidio et al. (2015), this problem can be overcome by adopting an adaptive sampling strategy (described in Sect. 4.1.) based on information on sea surface temperature (SST), chlorophyll *a* (chl *a*) concentrations, and currents provided by satellite products analyzed in real time during the entire cruise.

4 Implementation of the OUTPACE cruise

The OUTPACE cruise was conducted during austral summer conditions from 18 February to 3 April 2015 aboard the RV *L'Atalante*. We performed a ~ 4000 km zonal transect from the north of New Caledonia to the western part of the GY, finally reaching French Polynesia (Fig. 2). Along this transect, two types of stations were sampled: 15 short-duration (SD, 8 h) stations dedicated to a large-scale description of biogeochemical and biodiversity gradients, and three long-duration (LD, 7 days) stations for Lagrangian process studies.

4.1 Adaptive strategy

Following the planned adaptive strategy, the initial transect designed to approximatively follow 19° S was modified along-route thanks to the information coming from satellite

images. The regions along the vessel route were first characterized on a large scale through the analysis of satellite (altimetry, SST, ocean color) data. These data were automatically retrieved and processed to derive Eulerian and Lagrangian diagnostics of ocean circulation and biogeochemistry: Okubo–Weiss parameter maps, Lagrangian coherent structures (LCSs), and chl *a* maps (d'Ovidio et al., 2015).

The satellite data were treated in near real time on land and the obtained data were transmitted onboard together with a daily bulletin containing the analysis of remote sensing information, along with suggestions for LD station positions (the complete series of the 42 bulletins is available on the OUTPACE website at <https://outpace.mio.univ-amu.fr>, section “Adaptive Strategy”).

Two main criteria were adopted in suggesting LD station positions:

1. The areas for the LD A and LD B (LD C) stations were characterized by local maxima (minima) of sea surface chl *a* concentration to sample MA (GY) conditions.
2. Local minima of surface current intensity for all LD stations were considered to increase the chance of sampling a homogeneous water mass.

Once the suggested positions for LD stations were relayed via the daily bulletin, at one of these locations real-time analysis of water samples using quantitative polymerase chain reaction (qPCR) was conducted to measure the abundances of six groups of diazotrophs (Stenengren et al., 2017). In this way, we located regions with diazotrophs, while also resolving the contrasting role *Trichodesmium* spp. or UCYN-dominated diazotroph communities have on biogeochemical cycles.

Finally, the exact locations of the three LD stations were then determined onboard in real time from a rapid survey using a moving vessel profiler (MVP) equipped with conductivity–temperature–depth (CTD) and fluorimeter sensors, accompanied by the hull-mounted thermosalinograph and acoustic Doppler current profiler.

During this rapid survey, it was planned to follow two different sampling routes: a cross of about 40 km each side, followed by a zigzag route covering an area of 25 km each side at the center of the cross. This strategy was applied as planned for the LD A and LD C stations.

The LD A station was performed east of the northern extremity of New Caledonia in an anticyclonic recirculation characterized by a relatively high surface chl *a* concentration. The LD C station was performed in a cyclonic eddy in the most oligotrophic part of the OUTPACE transect (GY) close to the Cook Islands.

The severe meteorological conditions due to the development of tropical cyclone Pam (a category 5 storm) that hit Vanuatu, forced us to perform the LD B station at a more easterly location than initially planned. Satellite imagery allowed for the targeting of a large filament of high surface

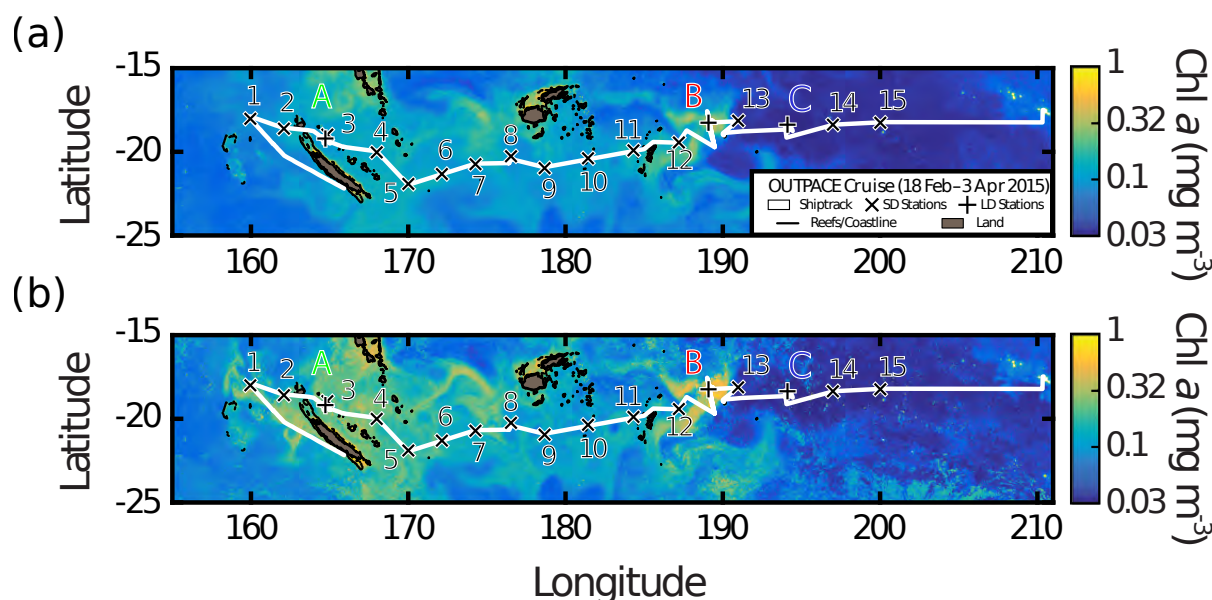


Figure 2. Transect of the OUTPACE cruise superimposed on (a) arithmetic mean surface chl *a* and (b) quasi-Lagrangian weighted mean chl *a* of the WTSP during OUTPACE. The two types of station, short duration (X) and long (+) duration, investigated for a period longer than 7 days, are indicated. The satellite data are weighted in time by each pixel's distance from the ship's average daily position for the entire cruise. The white line shows the vessel route (data from the hull-mounted acoustic Doppler current profiler (ADCP) positioning system). Coral reefs and coastlines are shown in black, land is grey, and areas of no data are left white. The ocean color satellite products are produced by CLS with support from CNES.

chl *a* concentration close to Niue. Due to the circulation patterns associated with the bloom, the rapid survey strategy was adapted in order to perform four sections across the structure (see details in de Verneil et al., 2017a).

SD station positions were chosen in relation to the LD stations, so that they were roughly equidistant from each other, respected territorial waters, and incorporated the changing conditions during the cruise.

4.2 General program at each station

Every station (Table 1) was investigated from the surface to 2000 m.

4.2.1 SD stations

Each of the 15 SD stations was investigated for 8 h. Specific operations during the SD station occupation consisted of

1. two 0–200 m CTD casts and Niskin bottle sampling and one 200–2000 m CTD cast and Niskin bottle sampling using the classical SBE 9plus CTD rosette (C-R) for measurements of core parameters (dissolved oxygen; dissolved inorganic carbon; total alkalinity; nutrients; chl *a*; particulate and dissolved organic C, N, P, and Si) and some more specific ones (for example primary production rates, N_2 fixation rates, and diazotroph abundance). An underwater vision profiler (UVP) was

attached to the CTD rosette to quantify and visualize suspended particulate material;

2. one 0–500 m CTD cast and bottle sampling using the trace metal clean SBE 9plus CTD rosette (TM-R) equipped with 24 teflon-lined GO-FLO bottles to sample for trace metal analyses;
3. optical sensors casts in which integrated measurements of bio-optical properties and pigments were made with instruments measuring hyperspectral radiometry in the UV–visible domain with UV–VIS TriOS spectroradiometers, and a MicroPro free-fall profiler (Satlantic) was used for downward irradiance measurements;
4. hauls for phytoplankton and zooplankton sampling with specific nets; and
5. a profile of turbulence measurements using a VMP1000 equipped with microsensors for temperature and shear that enable accurate estimates of the eddy diffusion coefficient K_z .

The specific configurations of the two CTD rosettes are available here: <https://outpace.mio.univ-amu.fr/spip.php?article137>.

Table 1. Date, location, and general characteristics of the stations investigated along the OUTPACE transect. Distance in kilometers from the first SD station (SD1).

Station	CTD cast (0–2000 m)	Arrival date (UTC)	Departure date (UTC)	Latitude (deg min)		Longitude (deg min)		Latitude (deg)	Longitude (deg)	Cumulative distance (km)	Bottom depth (m)
Nouméa		x	19/02/2015 21:00	22	14	166	28	–22.23	166.47	x	0
SD 1	007	21/02/2015 20:00	22/02/2015 09:30	18	0	S 159	54	E –18.00	159.90	0	4068
SD 2	016	22/02/2015 21:45	23/02/2015 10:00	18	38	S 162	8	E –18.63	162.12	210	2567
SD 3	020	24/02/2015 03:45	24/02/2015 10:00	19	0	S 164	54	E –19.00	164.90	488	3252
LD A	067	25/02/2015 13:00	02/03/2015 23:15	19	13	S 164	41	E –19.21	164.69	528	3491
SD 4	071	04/03/2015 08:30	04/03/2015 15:15	20	0	S 168	0	E –20.00	168.00	839	4995
SD 5	075	05/03/2015 05:45	05/03/2015 13:30	22	0	S 170	0	E –22.00	170.00	1103	4405
SD 6	079	06/03/2015 03:15	06/03/2015 12:30	21	22	S 172	8	E –21.37	172.13	1303	2509
SD 7	083	07/03/2015 00:15	07/03/2015 09:30	20	44	S 174	16	E –20.73	174.27	1506	2451
SD 8	087	07/03/2015 21:00	08/03/2015 07:15	20	42	S 176	24	E –20.70	176.40	1694	2028
SD 9	091	08/03/2015 22:15	09/03/2015 07:45	20	57	S 178	39	E –20.95	178.65	1927	3864
SD 10	095	10/03/2015 00:00	10/03/2015 07:15	20	28	S 178	31	W –20.47	181.48	2187	819
SD 11	099	10/03/2015 21:45	11/03/2015 05:15	19	59	S 175	40	W –19.98	184.33	2449	2234
SD 12	103	11/03/2015 21:00	12/03/2015 05:00	19	29	S 172	50	W –19.48	187.17	2711	7717
LD B	151	15/03/2015 12:00	20/03/2015 22:30	18	14	S 170	52	W –18.24	189.14	2985	4912
SD 13	152 (0–500 m)	21/03/2015 10:30	21/03/2015 11:00	18	12	S 169	4	W –18.20	190.93	3096	4598
LD C	199	23/03/2015 12:00	28/03/2015 22:00	18	25	S 165	56	W –18.42	194.06	3371	5277
SD 14	210	30/03/2015 01:30	30/03/2015 09:15	18	25	S 163	0	W –18.42	197.00	3640	4998
LD 15	213	31/03/2015 00:30	31/03/2015 08:30	18	16	S 160	0	W –18.27	200.00	3916	4965
Papeete		02/04/2015 21:00	x	17	34	S 149	36	W –17.57	210.40	x	0

4.2.2 LD stations

Each of the three LD stations was investigated with a drifting array (see below) that was deployed for 7 days. A series of CTD (C-R) casts (0–500 m) were performed every 3 h near the actual position of the drifting array while numerous specific operations (see below) were carried out in between CTD casts.

A total of 13 surface velocity program (SVP) drifters anchored at 15 m depth were deployed in order to study relative surface dispersion. The drifters were launched with three at LD A, six at LD B, and four at LD C. A drifting array (equipped with three PPS5 sediment traps, current meters, specific oxygen sensors, and specific high-frequency temperature sensors; please consult <https://outpace.mio.univ-amu.fr/spip.php?article75> for details) was then deployed at the chosen station position to start the process study. The drifting array was recovered at the end of each LD station occupation, immediately following the last CTD cast. An additional CTD cast from the surface to the bottom (5000 m) was undertaken at the LD B station.

Specific operations during LD stations were identical to those performed at SD stations, along with other operations like in situ production measurements, in situ particulate material sedimentation measurements, trace metal clean pumping for process experiments, and profiles of current measurements. Finally, at each LD station, a drifting PROVOR-type Argo float (ProBio equipped with sensors to measure chl *a* (fluorescence), chromophoric dissolved organic matter (CDOM) (fluorescence), PAR, irradiance at three wavelengths, backscattering, and dissolved oxygen (optode)) was deployed (Table 2).

The following color code was proposed to present data from the different LD stations: LD A is green, LD B is red, and LD C is blue.

All details regarding the OUTPACE cruise are available on the OUTPACE website: <https://outpace.mio.univ-amu.fr/>.

The general scheduled and realized programs are available here: http://www.obs-vlfr.fr/proof/php/outpace/outpace_log_and_basic_files.php.

5 General characteristics of the upper water masses in the WTSP

Water characteristics (temperature, salinity, density, chl *a* fluorescence) for the upper 700 m as measured by the C-R are presented in Fig. 3. The deep CTD casts from all SD and LD stations are presented subsequent to post-cruise processing using Sea-Bird Seasoft software, adopting the TEOS-10 standard. The upper surface layer (0–30 m) observed during the OUTPACE transect was characterized by warm waters, with temperatures between 26.18 and 29.93 °C (Fig. 3a), and relatively low salinity, i.e., absolute salinity between 35.03 and 35.81 g kg^{–1} (Fig. 3b). Density anomalies of 21.72–22.91 kg m^{–3} between 0 and 30 m increased gradually between 30 and 200 m to reach 24.89–25.38 kg m^{–3} (Fig. 3c). The salinity increase in subsurface waters (100–200 m) from 187° W longitude (Fig. 3b) indicated the geographical border between waters under the influence of the MA (SD1 to SD12) and waters from the GY (SD13 to SD15, LD C). LD B was not classified here and required a further analysis (de Verneil et al., 2017a). The classification between MA and GY waters will be helpful to describe general biogeochemical and bio-

Table 2. PROVOR Argo floats deployed along the OUTPACE transect.

Station	Float number	Deployment location		Date of	Time (UTC)	Closest CTD cast
	WMO id.	Latitude	Longitude	deployment		
LD A	ProBio075b	19° 13.00 S	164° 29.40 W	03/03/2015	22:45:00	OUT_C_067
SD 11	6901663	19° 59.56 S	175° 38.59 W	11/03/2015	05:03:00	OUT_C_099
SD 12	6901664	19° 32.61 S	172° 46.47 W	12/03/2015	05:00:00	OUT_C_103
LD B	6901666	19° 44.50 S	170° 31.31 W	13/03/2015	01:15:00	
LD B	6901667	17° 38.39 S	170° 59.47 W	13/03/2015	18:49:00	
LD B	6901668	18° 13.90 S	170° 44.20 W	20/03/2015	21:36:00	OUT_C_151
LD B	ProBio077b	18° 16.09 S	170° 43.80 W	20/03/2015	22:00:00	OUT_C_151
Before LD C	6901669	18° 46.92 S	168° 09.06 W	22/03/2015	07:55:00	
LD C	6901670	18° 41.20 S	165° 45.18 W	22/03/2015	19:48:00	
LD C	6901671	18° 28.16 S	165° 46.21 W	28/03/2015	21:25:00	OUT_C_199
LD C	ProBio079b	18° 28.16 S	165° 46.21 W	28/03/2015	21:30:00	OUT_C_151
SD 14	6901679	18° 24.26 S	162° 59.34 W	30/03/2015	09:45:00	OUT_C_210
SD 15	6901680	18° 15.29 S	159° 59.23 W	31/03/2015	08:15:00	OUT_C_213

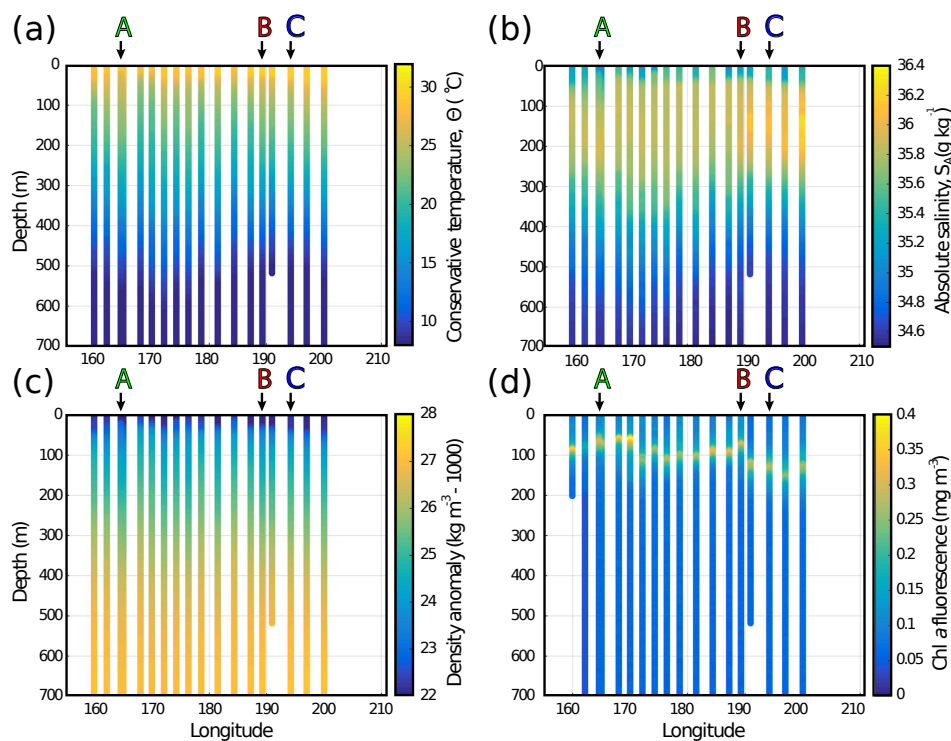


Figure 3. Zonal sections of (a) conservative temperature Θ , (b) absolute salinity SA, (c) density anomaly, and (d) fluorescence in the upper 700 m from the classic CTD rosette at SD and LD stations along the OUTPACE transect. The three LD stations are highlighted by their color-coded letter and corresponding arrow.

logical features. Between 200 and 700 m, a decreasing gradient of temperature and salinity indicated the presence of permanent thermocline waters. Temperature fell from 19.27 to 22.08 °C at 200 m to 5.48–6.91 °C at 700 m (Fig. 3a). Absolute salinities of 35.76–36.21 g kg⁻¹ at 200 m decreased to 34.49–34.61 g kg⁻¹ (Fig. 3b). The density anomalies of 24.89–25.38 kg m⁻³ at 200 m increased largely to 26.97–27.13 kg m⁻³ at 700 m (Fig. 3c). The maximum fluorescence

depth, considered here as the main indicator of the trophic state, increased from ~ 100 m depth in the MA waters to ~ 115–150 m depth in the GY waters, which allowed us to sample the oligotrophic to ultra-oligotrophic transition in the WTSP for the purpose of the OUTPACE project.

6 The climatological context of the campaign

The OUTPACE cruise took place in the WTSP, a region impacted by the El Niño–Southern Oscillation (ENSO), known to be the most important mode of SST variability on inter-annual to decadal timescales (Sarmiento and Gruber, 2006). ENSO-related SST anomalies are caused by a combination of changes in ocean circulation (mainly changes in the strength and source of equatorial upwelling) and anomalous local air–sea heat exchanges. The most dramatic effects of ENSO in the surface ocean are well documented in the eastern tropical Pacific, where seasonal upwelling conditions can be suppressed with severe economic consequences for fisheries (Chavez et al., 2003). During El Niño phases (negative Southern Oscillation Index (SOI), defined below), the warm pool normally positioned in the western Pacific is found farther east, resulting in the aforementioned suppression of upwelling conditions off Peru. La Niña, the opposite phase with a positive SOI, reverses this situation. After decades of intense study, ENSO is still an active field of research (Takahashi et al., 2011).

Given the known importance of ENSO for the tropical South Pacific, it is worthwhile to determine in which climatological conditions the cruise was performed. To achieve this goal, we identified years of opposing ENSO phases and analyzed the corresponding WTSP conditions with available satellite data. ENSO phases were identified using the monthly time series of SOI provided by NCEP (<http://www.cpc.ncep.noaa.gov/>, downloaded 5 December 2016). The SOI metric uses differences in standardized sea level air pressure between Papeete (Tahiti) and Darwin (Australia) to represent the ENSO phase, as previously mentioned. The OUTPACE region lies between these two locations (Fig. 2), highlighting again ENSO's possible influence on the cruise.

WTSP conditions were estimated with SST and surface chl *a* concentration measured by the MODIS Aqua satellite mission and available at the NASA Ocean Color Data website (<https://oceandata.sci.gsfc.nasa.gov/>, downloaded 14 December 2016). Global annual and monthly (March) averages of both SST and chl *a* were provided by NASA at level 3 (i.e., mapped) with 4 km satellite pixel resolution, resulting in four separate datasets from 2003 to present. The data within the OUTPACE region, defined between 25° S, 155° E, and 15° S, 149° W, to envelope the cruise track as in Fig. 2, were extracted. March was chosen as the month of study because it was the central month of the cruise. Plots of these four datasets can be found on the OUTPACE dataset website (http://www.obs-vlfr.fr/proof/php/outpace/outpace_figures.php). Probability density distributions were generated for each dataset in its entirety. Additional probability distributions were also calculated on data subsets, for years 2003 and 2011 as chosen by SOI to represent opposite ENSO phases, along with 2015, the year of the cruise. In order to gauge significance between probability distributions of different years, the temporal standard deviation of each pixel

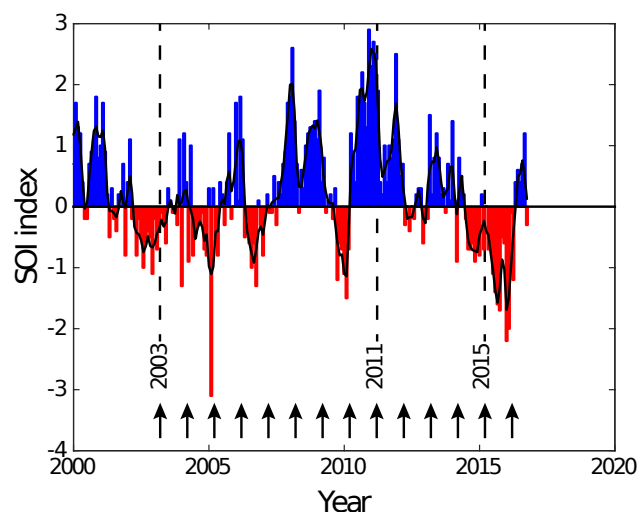


Figure 4. Time series of the monthly Southern Oscillation Index (SOI) from January 2000 to October 2016. Negative and positive values are shaded red and blue to signify El Niño and La Niña, respectively. SOI smoothed by a lowess filter with a five-point window is shown as a solid black line. Black arrows indicate March for the years when satellite data are available. Dashed vertical lines indicate March 2003, 2011, and 2015.

was calculated for each of the four datasets. This resulted in a distribution of standard deviations, the median of which was taken to represent inherent interannual variability.

The time series of SOI is presented in Fig. 4, with El Niño (La Niña) values shaded red (blue). During OUTPACE cruise sampling in austral summer 2015, a strong El Niño was observed. The other El Niño year considered here was 2003, chosen because of its relative strength and duration similar to 2015; for similar reasons of intensity and duration, 2011 was designated a representative La Niña year.

The probability density distributions for the annual and March means of both SST and chl *a* are presented in Fig. 5. In SST, the March mean showed a larger proportion of warmer temperatures in relation to the annual mean (white lines in Fig. 5a, c), reflecting the austral summer season. The 2011 La Niña distributions showed a greater proportion of warm temperatures in both annual and March distributions (blue lines in Fig. 5a, c), which is consistent with the idea of La Niña accentuating the warm pool present in the WTSP. The probability peak of March 2011 SST was much more localized, between 26 and 30 °C, than 2003, 2015, and the mean (Fig. 5c). The 2003 March SST (red line in Fig. 5c) was the widest distribution, with a slight rightward skew. By contrast, both the mean and 2015 March probabilities (white and green lines, respectively) had a left skew. Considering the median 95 % confidence interval for the March SST time series, the central peaks for these distributions could not be considered distinct.

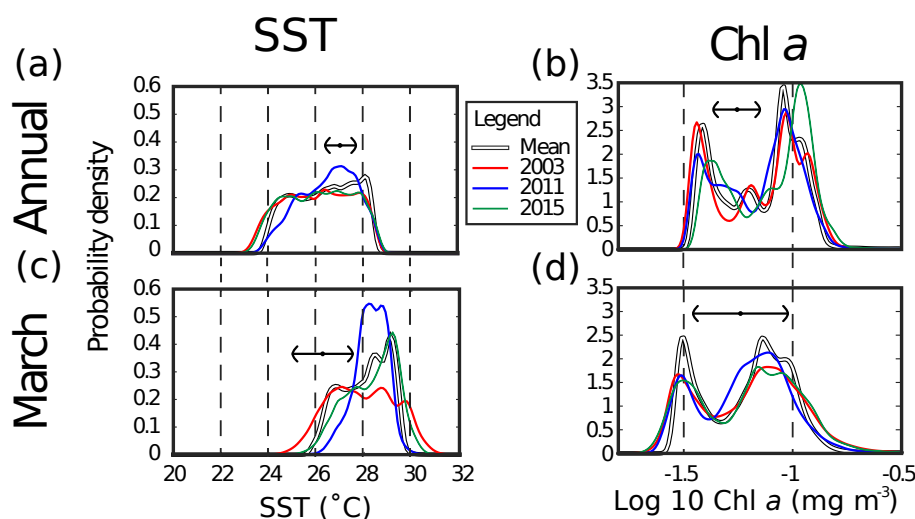


Figure 5. Probability density estimates for MODIS Aqua data in the OUTPACE region, using mean (a) annual SST, (b) annual chl *a*, (c) March SST, and (d) March chl *a*. Probability densities for the ensemble of all years is shown in white, while densities for 2003, 2011, and 2015 are in red, blue, and green, respectively. Also plotted are 95 % confidence intervals (2 standard deviations) for each subset, estimated using the median variance of pixel interannual variability. Note the logarithmic scale for chl *a*.

Satellite chl *a* showed distributions completely different from SST. For all quantities considered, the annual and March means (Fig. 5b, d), as well as the years 2003, 2011, and 2015, showed a bimodal distribution, where the two peaks were distinct enough from each other to be significant from the median pixel variability. Interestingly, 2003 and 2011 (red and blue lines, Fig. 5b) annual chl *a* distributions overlapped considerably more than 2015 (green line), which had its entire probability distribution shifted to the right, though this shift may not have been significant. The March mean for 2003 and 2015 chl *a* (red and green lines, respectively, Fig. 5d), however, almost entirely overlapped, possibly signifying that El Niño chl *a* distributions were more alike than La Niña years. The annual 2015 mean of chl *a* being slightly different from other years, and yet the March 2015 mean resembling 2003, might indicate that 2015 was indeed a slightly different year from a surface chlorophyll perspective, but this slight difference was concentrated in other parts of the year than late austral summer when OUTPACE took place.

In summary, the SOI metric of ENSO showed that 2015 was an El Niño event, and both the SST and chl *a* satellite data in the WTSP partially reflected this. The March 2003 and 2015 SST distributions resembled each other, but they also resembled the entire dataset more than the 2011 La Niña distribution. Looking at SST, perhaps La Niña events impact the region more than El Niño. Chl *a* distributions for the El Niño years also overlapped more than with La Niña, but the variability inherent in the time series precluded the declaration of significant differences. Overall, in the WTSP both SST and chl *a* were not atypical from what one would expect;

thus, we determined that climatological effects upon the results of OUTPACE were minimized.

7 Special issue presentation

The goal of this special issue is to present the knowledge obtained concerning the functioning of WTSP ecosystems and associated biogeochemical cycles based on the datasets acquired during the OUTPACE experiment. The cruise strategy was organized to promote collaboration between physicists, biologists, and biogeochemists with expertise including marine physics, chemistry, optics, biogeochemistry, microbiology, molecular ecology, genetics, and modeling. Most of the contributions to this volume have benefited from this collective effort and are presented according to the main objectives of the OUTPACE experiment.

The hydrological and dynamical context of biogeochemical sampling is described for the entire cruise route (Fumonia et al., 2017) and specifically at the three long-duration stations, where low physical variability validated the quasi-Lagrangian sampling strategy employed (de Verneil et al., 2017b). Turbulence measurements revealed an interesting longitudinal gradient with higher turbulence levels in the west, i.e., the Coral Sea, compared to the eastern part within the gyre, consistent with the increasing oligotrophy (Bouruet-Aubertot et al., 2017). The large-scale circulation was dominant even though the mesoscale and sub-mesoscale circulation can have a strong influence (Rousselet et al., 2017), in particular on the bloom observed at station LD B (de Verneil et al., 2017a).

An important focus of OUTPACE was on dinitrogen fixation and its fate in the ecosystem (Caffin et al., 2017a, b).

N_2 fixation was detected at all 18 sampled stations and the transect could be divided into two main characteristic subareas (Bonnet et al., 2017b): (i) the MA waters (160° W to 170° E) exhibiting very high N_2 fixation rates ($631 \pm 286 \mu\text{mol N m}^{-2} \text{d}^{-1}$, i.e., among the highest reported for the global ocean; Luo et al., 2012) and dominated by *Trichodesmium* (Stenengren et al., 2017), and (ii) the GY waters (170 – 160° E) exhibiting low N_2 fixation rates (average $85 \pm 79 \mu\text{mol N m}^{-2} \text{d}^{-1}$) dominated by UCYN-B (Stenengren et al., 2017). The differing $\delta^{15}\text{N}$ signature of suspended particles measured over the photic layer of MA (-0.41‰) and GY waters (8.06‰) confirms the presence of two contrasting regions in terms of N_2 fixation. Thanks to the Lagrangian strategy followed at the LD stations and the low dispersion measured, showing that we sampled the same water masses (de Verneil et al., 2017b), N budgets were established (Caffin et al., 2017b). N_2 fixation was the major external source of N, representing more than 90 % of new N input into the photic layer, and the e-ratio quantifying the efficiency of a system to export particulate organic matter was higher in MA waters than in GY waters (Caffin et al., 2017b). Caffin et al. (2017a) revealed that the diazotroph-derived nitrogen (DDN) was efficiently transferred from diazotrophs (*Trichodesmium* and UCYN) to non-diazotrophic phytoplankton, both autotrophs and heterotrophs. Hunt et al. (2017) report an efficient transfer of DDN in zooplankton. The fate of C and N was under the influence of programmed cell death in diazotrophs (Berman-Frank et al., 2017) but diazotrophs were poorly exported directly, and we suspect that this transfer of DDN fueled indirect export associated with N_2 fixation. By using nitrogen isotope budgets, Knapp et al. (2017) confirmed that $> 50\%$ of export production was supported by N_2 fixation in MA waters. Stenengren et al. (2017) identified a clear niche separation between a subsurface (UCYN-A1 and A2 with their hosts) and a surface group (*Trichodesmium*, UCYN-B, and the heterotrophic group, het-group) based on a temperature–depth gradient. They also found discrepancies between the UCYN-A and their hosts in both abundance and distribution, which suggests that the UCYN-A could be living freely or with a wider diversity of hosts than previously believed. Finally, the assemblage of epibiotic microorganisms associated with *Trichodesmium* were characterized in relation with environmental parameters (Frischkorn et al., 2017). N_2 fixation in the ocean does not only occur in tropical sunlit surface waters but also in less obvious environments such as temperate latitudes and aphotic waters. Here, N_2 fixation was also measured in the mesopelagic zone along the OUTPACE transect and the diazotroph community present were identified. Deep N_2 fixation rates were low but measurable and recurrently found along the transect, with the exception of the easternmost stations located in the ultra-oligotrophic subtropical Pacific gyre (Benavides et al., 2017). N_2 fixation activity was presumably driven by the dominating gamma-proteobacterial community and fueled by the presence of labile organic matter compounds. Benavides et

al. (2017) provided further evidence that N_2 fixation in the deep ocean is not negligible and likely impacts global nitrogen inputs in a significant manner.

The dynamics of phytoplankton (Bock et al., 2017; Leblanc et al., 2017; Lefevre et al., 2017; Guidi, 2017), heterotrophic bacterioplankton (Van Wambeke et al., 2017), and zooplankton (Carlotti et al., 2017; Hunt et al., 2017) along the zonal gradient of diazotroph diversity and activity are described together with the composition and distribution of dissolved organic carbon (Panagiotopoulos et al., 2017) and the changes in inorganic carbon content along the longitudinal transect (Wagener et al., 2017). Stations sampled during the OUTPACE cruise were characterized by a highly stratified community structure, with significant contributions of Prochlorococcus and picophytoeukaryote populations to biomass (Bock et al., 2017). Size-fractionated results show a non-negligible contribution of the pico-sized fraction (< 2 – $3 \mu\text{m}$) to both Si biomass and uptake, which could confirm the previous hypothesis of Si assimilation by Synechococcus populations or reflect the presence of an overlooked Si group such as Parmales (Leblanc et al., 2017). Surface DOC concentrations varied little (50 – $75 \mu\text{M}$) across the transect, with slightly higher values observed at LD B ($78 \mu\text{M}$), and labile organic matter (sugars were used as a good proxy) closely followed DOC patterns ranging from 1.5 to $3 \mu\text{M}$ with higher values also recorded at LD B ($3.5 \mu\text{M}$). Labile organic matter accounted for about 3 – 5% of DOC, with glucose being the dominant sugar ($> 60\%$ of total sugars) (Panagiotopoulos et al., 2017). Valdes et al. (2017) suggest that copepods can retain N and P compounds obtained from feeding in the upper layer, preventing the rapid loss of these nutrients. Copepods were able to sustain and modify the composition of microbial communities and could provide P for further development of cyanobacterial blooms.

Optical properties of the WTSP waters are presented, with a focus on the cyanobacterial (diazotroph) impact upon bio-optical properties, UV–VIS light attenuation (Dupouy et al., 2017), and chl *a* algorithms (Frouin et al., 2017). Operational NASA bio-optical algorithms (OC4v6; color index, CI) substantially underestimated surface chl *a* concentration, but a normalized reflectance difference index, robust to atmospheric correction errors, performed well over the range of chl *a* values encountered across the transect (Frouin et al., 2017). *Trichodesmium* is considered the main nitrogen-fixing species, especially in the South Pacific region. Due to the paucity of in situ observations, alternative methods for estimating the presence of *Trichodesmium* must be sought to evaluate the global impact of these species on primary production. Rousset et al. (2017) elaborate a new satellite-based algorithm and use that algorithm to estimate the extent of *Trichodesmium* surface blooms and their dynamics during the OUTPACE experiment. Finally, the main processes controlling the biological carbon pump in the WTSP were investigated using a unidimensional vertical (Gimenez et al., 2017) and regional (Dutheil et al., 2017) biogeochemical–physical

coupled models. The new knowledge gained on the interactions between planktonic organisms and the cycle of biogenic elements is then used to propose a new scheme for the biological carbon pump function and its role, at the present time and in the near future, in the oligotrophic Pacific Ocean (Moutin et al., 2017).

Data availability. All data and metadata are available at the French INSU/CNRS LEFE CYBER database (Scientific Coordinator: Hervé Claustre, Data Manager, Webmaster : Catherine Schmechtig) at the following web address: <http://www.obs-vlfr.fr/proof/php/outpace/outpace.php> (INSU/CNRS LEFE CYBER, 2017).

Competing interests. The authors declare that they have no conflict of interest.

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