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Optical proxy for particulate organic nitrogen from BGC-Argo floats

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Abstract: Using biogeochemical-Argo float measurements, we propose, for the first time, an optical proxy for particulate organic nitrogen concentration (PON) in the Western Tropical South Pacific, an area influenced by dinitrogen (N2) fixation. Our results show a significant relationship between the backscattering coefficient at 700 nm (b700) and PON, especially when the latter is measured using the wet oxidation method (R2=0.87). b700 may be used to estimate PON concentrations (PONopt) between 0.02 and 0.95 µM, allowing for unprecedented monitoring using autonomous profiling floats. The b700 vs PON relationship can be used to study phytoplanktonic biomass dynamics at relevant seasonal temporal scales, with clear evidence of PONopt as a proxy of phytoplanktonic biomass, at least for this specific area. Temporal analyses of PONopt show significant increases (from 0.16 to 0.80 µM) likely related to new production associated to N2 fixation events measured during stratification periods in the Melanesian Archipelago.

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1. Introduction

The surface waters of the subtropical South Pacific gyre (SPG) are permanently depleted in dissolved inorganic nitrogen (NO3−) [1], making this area the largest oceanic desert in the world’s ocean [2]. However, in some oligotrophic areas, biological N2 fixation offers the marine phytoplankton community a mechanism to relieve nitrogen limitation in the euphotic surface layer [3,4]. In the framework of the Oligotrophy to Ultra-oligotrophy PACific Experiment (OUTPACE) cruise (19 Feb-3 Apr 2015), record N2 fixation rates were recently observed in the upper waters of the Western Tropical South Pacific (WTSP) at the end of austral summer [5]. Despite a NO3−-depleted mixed layer, a significant increase of phytoplanktonic biomass was observed during N2 fixation events. However, the limited observations of seasonal phytoplankton biomass dynamics, hypothesized to be largely driven by N2 fixation [6,7], considerably restricts our understanding of the WTSP’s biogeochemical functioning. To overcome present limitations, in situ observations over a broad range of time scales are required.

Biogeochemical-Argo (BGC-Argo) profiling floats are capable of autonomously observing bio-optical properties such as Chlorophyll-a (Chla) fluorescence and particulate backscattering coefficient (b0.75) at high frequency [8]. The use of bio-optical proxies has been previously shown to have high reliability in the estimation of biogeochemical variables such as Chla and particulate organic carbon (POC), among others [8,9]. While Chla fluorescence is the most commonly used proxy for living phytoplankton cells [10,11], b0.75 variability is instead driven by both the algal
and non-pigmented particle pools including viruses, heterotrophic bacteria, and non-living cells in case 1 oceanic waters [12]. For this reason, \(b_{bp}\) has long been used as a proxy of POC in open ocean water in the absence of mineral particles [9,12–19].

In oligotrophic areas, the carbon hydrogen nitrogen (CHN) method requires a high volume of filtered seawater, up to 10 L, to obtain accurate POC measurements [19]. Filtering such volumes of sea water is extremely time consuming and limits throughput of large numbers of samples. Several factors can lead to large biases in estimated POC in oligotrophic areas, including contamination, adsorption of dissolved organic carbon (DOC) onto filters, particle formation in bottle samples after collection, the contribution of particulate inorganic carbon, as well as particle retention, among others [19–22]. Particulate organic nitrogen and phosphorus measurements obtained by the wet oxidation method (PON\(_{\text{wet}}\)) and POP\(_{\text{wet}}\), respectively) could represent a valuable alternative in oligotrophic areas, since the method’s sensitivity requires smaller volumes of seawater than the CHN method [23], making it less sensitive to the potential contaminations alluded to above. Wet oxidation requires a maximum volume of 1.2 L, even in oligotrophic areas. The smaller volume and the absence of an acidification step to remove inorganic particles on the filter significantly decrease both the time required for each sample and potential particle formation as transparent exopolymer particles (TEP; 22) in bottle samples, thus decreasing potential contamination.

The wet oxidation method has been shown to be more accurate than the CHN method for PON and POC measurements [23]. Based on the fact that POC and PON generally covary [1,6], one may expect a good correlation between \(b_{bp}\) and PON. As a result, the use of an optical proxy of PON (PON\(_{\text{opt}}\)) could provide a means to estimate seasonal variations of phytoplanktonic biomass and associated particles, especially in oligotrophic areas.

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**Fig. 1.** (a) Trajectory of the BGC-Argo floats deployed in this study (green triangle = \(F_A\), red triangle = \(F_B\), blue triangle = \(F_C\)). The location of the OUTPACE section is represented by black (Western and Eastern Melanesian Archipelago) and grey (South Pacific gyre) circles. Open circles represent the location of available in situ PON measurements in the South Pacific [24]. (b) Zoom on the trajectory of the BGC-Argo floats. Colors represent the time (years) and the black dashed boxes show the “bloom periods” defined in the text.
In areas where nitrogen is the key limiting nutrient, and N₂ fixation is sufficiently favored, both an increase in PON standing stocks and a tight coupling between N₂ fixation and surface water PON accumulation have been observed [24,25]. While PON estimates allow one to follow phytoplanktonic biomass, observations in the South Pacific are sparse [26] (Fig. 1(a)), stressing the need to develop indirect methods for estimating this key variable at regional and pertinent time scales. For this purpose, three BGC-Argo floats were deployed during the OUTPACE cruise. Our study focuses on the mixed layer, where the N₂ fixation process mainly occurs [25]. The main goals of this work are (1) to characterize and discuss, for the first time, bhp vs PON relationships to define an optical proxy of PON (PON[^opt]), and then, (2) to investigate the seasonal dynamics of the particulate organic biomass in the WTSP, an area influenced by N₂ fixation events, using Chlα and PON[^opt] seasonal distributions.

2. Materials and methods

The OUTPACE cruise took place along a West to East transect [Fig. 1(b)]. A total of 18 stations were sampled from the oligotrophic water of the Western and Eastern Melanesian Archipelago (WMA and EMA, respectively) to the clearest ocean waters of the SPG [27].

PON and particulate organic phosphorus (POP) samples were collected at 16 depths between the surface and 500 m from a SBE 911+ CTD-Rosette in polycarbonate bottles. PON and POP concentrations were quantified spectrophotometrically following the wet oxidation method ([PON][wet] and [POP][wet]) based on persulfate digestion at 120 °C [28]. Following this method, a volume of 1.2 L was filtered through a pre-combusted (24 h, 450 °C) 47 mm GF/F filter. The filter was then placed in a Teflon bottle in which 20 mL of milli-Q water and 2.5 mL of the oxidizing reagent were previously dispensed (concentration factor: 1.2/0.0225). Nitrate and phosphate concentrations were then determined in the digested sample using an automated colorimetric procedure on a Technicon auto-analyzer [29]. The repeatability, calculated as the coefficient of variation (CV) for [PON][wet] and [POP][wet] field-collected replicates (n=10), was 2% and 3%, respectively. The accuracy, linked to the uncertainty of the calibration curve’s slope (calculated as the CV of the slope) of [PON][wet] and [POP][wet] was 1.46% and 1.36% (n=14), respectively. For each station, pre-combusted GF/F filters were used on board as sample blanks. The blank consisted of adding the same volume of oxidizing reagent to 20 mL of milli-Q water, in which a pre-combusted GF/F filter was previously added. The means of filter blanks of [PON][wet] and [POP][wet] were 0.036 ± 0.002 μM and 0.0021 ± 0.0001 μM. The quantification limits of [PON][wet] and [POP][wet], calculated as ten times the standard deviation of 10 blank measurements [30], were 0.02 μM and 0.001 μM, respectively. The maximum quantification limits for [PON][wet] and [POP][wet] were [PON]max = [standard NO₃⁻][max]/(concentration factor) = 0.95 μM and [POP]max = [standard PO₄³⁻][max]/(concentration factor) = 0.060 μM, respectively. Mineralization efficiencies measured daily with P-choline and urea standards were 100 ± 2% for N and 99 ± 1% for P (n=27).

A second set of PON measurements used a PerkinElmer 2400 CHN analyzer ([PON][CHN]), with standards prepared with a 20 g L⁻¹ glycine (VWR C14037000) solution (N range: 0.15-10 μM). Seawater samples (2 L) were filtered through pre-combusted (4 h, 450°C) 25 mm GF/F filters, dried at 60°C and stored in 1.5 mL Eppendorf PE tubes. The repeatability for [PON][CHN] field-collected replicates (n=6) was 4.60% and the calibration curve slope CV was 2.83% (n=13). For each station, pre-combusted GF/F filters were used on board as sample blanks. These filter blanks were processed in the same way as sample filters without the filtration step. The mean of filter blanks of [PON][CHN] was 0.021 ± 0.014 μM. The quantification limit of [PON][CHN] was 0.13 μM. [PON][wet] and [PON][CHN] concentrations showed an excellent agreement (R²=0.92; slope = 1.03 ± 0.03). The recovery between [PON][CHN] and [PON][wet] is close to 100% (95% C.I of the slope = [0.96 1.11]; 95% C.I of the intercept = [-0.03 0.028]) with a very low standard deviation over the range of concentrations measured as part of this study, highlighting that the
Three BGC-Argo floats (F_A, F_B, and F_C) were deployed in March/April 2015 near the stations LDA, LDB, and LDC respectively [Fig. 1(b)], and their collected data were downloaded from the Coriolis database website (ftp://ftp.ifremer.fr/ifremer/argo/dac/coriolis/). These floats were equipped with a Sea-Bird Electronics (SBE41CP) conductivity-temperature-depth (CTD) sensor (Seabird Inc., USA) and an additional sensor package: the WETLabs Environmental Characterization Optics triplet puck (ECO3, Seabird Inc., USA) measuring the fluorescence of Chla at excitation/emission wavelengths of 470/695 nm and the angular backward scattering coefficient of particles at 700 nm (Table 1).

Table 1. Equipment details for each float used in this study, (the first fifteen, nineteen, and eleven profiles were recorded every day for floats F_A, F_B, and F_C, respectively, before starting to sample every five days).

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<td>9.7</td>
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</tbody>
</table>

Parameters used in this study: Latitude, Longitude, Time (days), Pressure (dbar), Salinity, Temperature (°C), Chla (mg m⁻³), bbg (m⁻³)

Measurements were collected every 1 or 5 days between 1,000 dbar and the surface, with a sampling resolution of 1 dbar between the surface (0.1-1 dbar) and 250 dbar and 10 dbar between 250 and 500 dbar, respectively. Growth rate of diazotrophs like *Trichodesmium* (the main diazotroph in the WTSP) are generally low and blooms last for months, indicating that 5 days is a correct sampling time to capture a diazotroph bloom. The data were quality controlled following the standard Argo protocol [32–34].

Mixed layer depth (MLD) was calculated using a threshold density of 0.03 kg m⁻³ deviation from the reference value at 10 m depth [35].

The fluorescence measurements of Chla were converted to Chla concentrations according to the procedure detailed by [33]. Chla concentrations were cleaned from out-of-range values and, following the recommendations of [36], adjusted Chla concentrations were divided by a factor of two. A non-photochemical quenching correction was applied following the standard Argo protocol [33]. Vertical profiles of Chla concentrations showed that most of the Chla values observed in deep waters (>200 dbar) were negative (figure not shown). To correct the negative deep Chla concentrations, we removed a constant value (the deepest Chla fluorescence value from the profile, i.e., so-called “deep-offset correction”). During the OUTPACE cruise, a “Fluorimeter, Chelsea Aquatracka MKIII” attached to a SeaBird CTD rosette was used to measure the Chla fluorescence. Calibration of the fluorimeter was carried out using HPLC Chla measurements from 13 OUTPACE stations (stations SD1, LDA, SD6, SD8, SD10, SD12, LDB, LDC, SD14, SD15), hence just before the BGC-Argo deployments. Very good agreement (p>0.01, Fig. 2) was observed between the first vertical profile of adjusted Chla concentrations and the vertical profile of Chla concentrations measured *in situ* at the closest OUTPACE CTDs in time and space (Table 1).
The backscattering sensors measure the angular scattering coefficient at 124° relative to the
direction of light propagation at a wavelength of 700 nm. This measurement is then transformed
into the $b_{bp}$ following [34], using the conversion factor of [37] ($\chi = 1.076$). Negative values of $b_{bp}$
were removed and vertical profiles were quality-controlled following the standard Argo protocol
[34]. Both datasets (Chla and $b_{bp}$) have indeed been checked qualitatively and no sensor drift or
bio-fouling have been observed.

The variability of $b_{bp}$ with PON is investigated between the surface and 500 dbar using the
OUTPACE CTD-rosette water samples, which were the closest in time and space to the float
profiles where $b_{bp}$ was measured (Table 1). To avoid the effect of internal waves, each bottle data
value was paired with a $b_{bp}$ value at the same density coordinate. Differences in density between
the bottle data and associated $b_{bp}$ values were found to be less than 0.005 kg m$^{-3}$.

3. Results

3.1. From backscattering to particulate organic matter

The $b_{bp}$ vs PON relationship is investigated to understand the extent to which $b_{bp}$ can be used as
a proxy of PON to better assess the impact of N$_2$ fixation events at the relevant seasonal temporal
scales. Here we investigate the $b_{bp}$ vs PON relationship (1) in the mixed layer (ML), (2) in the
ML and the deep Chla maximum (DCM) [0-150 dbar, Fig. 3(b)], and (3) in the whole sampled
water column (0-500 dbar), before deriving PON$_{opt}$ and POP$_{opt}$.

All PON$_{wet}$ concentrations and most of the PON$_{CHN}$ concentrations are above their respective
quantification limits. By considering these data, a significant relationship is obtained between
$b_{bp}$ and PON, regardless of the experimental method used to measure PON (wet vs CHN) (Fig. 3;
Table 2). The $b_{bp}$ vs PON$_{wet}$ relationship is better than that of $b_{bp}$ vs PON$_{CHN}$ between the
surface and 150 (500) dbar [Figs. 3(c), 3(d); Table 2]. Methodologically based variability could
explain this discrepancy. Indeed, the wet oxidation method achieves a sensitive measurement
from a smaller volume (1.2 L) of seawater than the CHN method (2-10 L), making this method
highly sensitive and suitable for PON analyses in oligotrophic waters [6,28].
Fig. 3. (a) Ratio of $b_{bp}/PON_{(wet)}$ vs Pressure, (b) Chl concentrations vs Pressure, and scatter plots between $b_{bp}$ (m$^{-1}$) measured during the first profiles of the $F_A$, $F_B$, and $F_C$ floats and (c) $PON_{(wet)}$, (d) $PON_{(CHN)}$, and (e) $POP_{(wet)}$. (The yellow, grey, and black lines represent the best linear regression fits for the data points in the ML, 0-150 dbar, and 0-500 dbar, respectively). Mixed layer = yellow markers/yellow lines, 0-150 dbar = yellow + grey markers/ grey lines, 0-500 dbar = yellow + grey + black markers/ black lines. (f) $b_{bp}$ vs $PON_{(wet)}$ slopes in the mixed layer, between the surface and 150 dbar, and between the surface and 500 dbar (note: circle = $F_A$/LDA, square = $F_B$/LDB, triangle = $F_C$/LDC). Red lines correspond to quantification limits. Statistical parameters are in Table 2.
Table 2. Statistical parameters for b<sub>bp</sub>/PON<sub>(wet)</sub>, PON<sub>(CHN)</sub>, and POP<sub>(wet)</sub> for the three depth categories: 0-500 dbar, 0-150 dbar, and Mixed Layer. n is the number of data points, R<sup>2</sup> is the determination coefficient, RMSE the Roots Mean Square Error (µM), Median of residuals (residuals = observed minus fitted values), and 75<sup>th</sup> and 25<sup>th</sup> percentile represents the interquartile range of residuals.

<table>
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<th>b&lt;sub&gt;bp&lt;/sub&gt;/PON&lt;sub&gt;(wet)&lt;/sub&gt;</th>
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<td>25th percentile</td>
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The b<sub>bp</sub>/PON<sub>(wet)</sub> ratios exhibit a low variability with depth, with an average value of 1.9 × 10<sup>−3</sup> ± 5.7 × 10<sup>−4</sup> m<sup>−1</sup> µM<sup>−1</sup> [Fig. 3(a)] in the layer including the ML and the DCM, [Fig. 3(b)]. In the ML, the slope derived from the b<sub>bp</sub>/PON<sub>(wet)</sub> relationship [1285 ± 114; Figs. 3(c), 3(f)] is not significantly different from that observed in the 0-150 dbar layer [1280 ± 98; Figs. 3(c), 3(f)]. In contrast, below the DCM (>150 dbar), an increase in b<sub>bp</sub>/PON<sub>(wet)</sub> ratios is observed, with an average value reaching 4.1 × 10<sup>−3</sup> ± 9.5 × 10<sup>−4</sup> m<sup>−1</sup> µM<sup>−1</sup> [Fig. 3(a)]. The slope from the b<sub>bp</sub>/PON<sub>(wet)</sub> relationship obtained between 0 and 500 dbar is significantly lower than those observed in both the ML and between the surface and 150 dbar [Fig. 3(f)].

Hereafter, to investigate the temporal variability of phytoplanktonic biomass and associated particles in the WTSP, the b<sub>bp</sub> measurements will be discussed in terms of PON using the relationship established from the b<sub>bp</sub> and PON<sub>(wet)</sub> data set measured between the surface and 150 dbar [Fig. 3(c)] [Eq. (1)].

\[
PON^{opt} = 1280 \times b_{bp}(700) - 0.38 \, (\mu M)
\]  

(1)

Studying each float separately in the 0-150 m layer, the b<sub>bp</sub>/PON<sub>(wet)</sub> relationships are only significant for F<sub>A</sub> and F<sub>B</sub>, with similar slopes (p<0.001) (Table 3). The F<sub>C</sub>/LDC data are very
few (n=9) and F_C exhibits low and quasi-constant PON_{wet} values. We therefore decide not to calculate the b_{pp} vs PON relationship for the LDC station individually. Nevertheless, including F_C/LDC data in the global b_{pp} vs PON_{wet} relationship does not significantly affect the slope (Table 3), so the global relationship is therefore calculated with all the data in the 0-150 m layer.

### Table 3. Slopes and intercepts for the b_{pp} vs PON_{wet} and b_{pp} vs POP_{wet} relationships for F_A/F_B/LDA-LDB, and F_A/F_B/F_C in the 0-150 m layer. (R^2 = the determination coefficient, p is the p-value).

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<td>1280 ± 98</td>
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<tr>
<td></td>
<td>F_A/F_B/F_C</td>
<td>59 ± 3</td>
<td>-0.01 ± 0.002</td>
<td>0.92</td>
<td>p &lt; 0.001</td>
</tr>
</tbody>
</table>

We consider the intercept of the PON_{wet} vs b_{pp} relationship as the lower detection limit of b_{pp} to derive PON_{opt}. The highest value of b_{pp} for which this equation is valid corresponds to the maximum value measured. Values of b_{pp} for which the b_{pp} vs PON_{wet} relationship is valid range between 3.4×10^{-4} and 9.5×10^{-4} m^{-1}. Additionally, values of b_{pp} for which the b_{pp} vs POP_{wet} relationship is valid range between 2.7×10^{-4} and 9.5×10^{-4} m^{-1}.

### 3.2. Mixed layer depth and chlorophyll-a seasonal distribution

In general, the seasonal Chla patterns mimic the MLD seasonal variability, with low mixed layer values observed during the summer (January/February/March) and high values during the winter (July/August) (Fig. 4). However, in the WMA (F_A), increases of Chla at shorter time scales were superimposed onto the seasonal pattern, with values reaching 0.09 mg m^{-3} in March/April 2015 and 0.10 mg m^{-3} in October/November 2015. These episodes were not related to abrupt changes in the MLD but occurred during conditions of shallow MLD (< 60 dbar) [dashed frame on Fig. 4(a)]. In the EMA (F_B), Chla followed the same seasonal variations as observed in 2016 and 2017 in the WMA (F_A), except for the summer period: despite a shallow MLD (< 45 dbar), Chla showed relatively high values, reaching 0.14 mg m^{-3} in March/April 2015, 0.08 mg m^{-3} in January/February 2016, and 0.10 mg m^{-3} in February 2017 [dashed frame on Fig. 4(c)]. The absence of intense vertical mixing and relatively high light levels (figure not shown) during these stratified periods should result in a diminution of the Chla concentrations. Observing the opposite, we hypothesize that the high Chla concentrations point to an additional controlling factor of the variability of Chla dynamics in the area, probably linked to diazotroph blooms.

### 3.3. PON seasonal distribution

In contrast to Chla, the temporal evolution of PON_{opt} was relatively even, with low values generally below 0.20 µM (Fig. 4). However, as for Chla, increases of PON_{opt} were observed during the first year in the WMA (F_A) and each year at particular times, between January and April, in the EMA (F_B). Indeed, in the WMA (F_A), between January 2016 and July 2017, despite the seasonal cycle of Chla, PON_{opt} was constant over time [Fig. 4(b)], with an average value of 0.18 ± 0.06 µM over this period [Fig. 4(b)]. Conversely, PON_{opt} values were significantly higher (p<0.01, t-test) in 2015 than those observed in 2016 and 2017 [Fig. 4(b)]. Maximum values were recorded during March/April 2015 and October/November 2015, averaging 0.41 ± 0.09 µM during both periods [dashed frames on Fig. 4(b)], simultaneous with increases of Chla [dashed frames on Fig. 4(a)].
Fig. 4. Temporal variations of the mixed layer depth (dbar) (black line) and the mixed layer average Chla (mg m\(^{-3}\)) concentration for floats (a) F\(_A\) (green line), (c) F\(_B\) (red line), and (e) F\(_C\) (blue line), along with temporal variations of (b) the mixed layer average PON\(^{opt}\) concentration (µM) (± sd) for float F\(_A\) (green line), (d) float F\(_B\) (red line), and (f) float F\(_C\) (blue line). Lighter colors show standard deviation around the mean (the uncertainty in the regression has been incorporated into the error propagation). Black dashed frames show the “bloom periods” as defined in the text.
In the EMA (F_B) during the summer periods, the average PON_{opt} values reached 0.70 ± 0.10 μM during March/April 2015, 0.50 ± 0.20 μM during January/February 2016 and 0.60 ± 0.03 μM during February 2017 [dashed frames on Fig. 4(d)], coinciding with relatively strong increases of Chl_a [dashed frames on Fig. 4(e)]. Outside of these summer periods, PON_{opt} values were mostly constant with an average value of 0.17 ± 0.06 μM [Fig. 4(d)], similar to F_A in 2016/2017. In the SPG (F_C), no increases of PON_{opt} were observed during the summer period. Instead, PON_{opt} was constant throughout the study period, with an average value of 0.16 ± 0.04 μM [Fig. 4(f)].

Hereafter, periods with both significant increases of PON_{opt} by a factor of 5 (from 0.16 to 0.80 μM) and Chl_a by a factor of 4 to 5 (from 0.03 to 0.14 mg m^{-3}) are called ‘bloom periods’ [dashed frames on Fig. 1(b) and Fig. 4]. These periods are: March/April 2015 and October/November 2015 for F_A; and March/April 2015 and January/February 2016 and February 2017 for F_B.

4. Discussion

At first order, b_{bp} variability is driven by the concentration of bulk particulate matter, with changes in the particulate size distribution, refractive index, and particle morphology, among other factors, generally acting at second order [12]. The inherent optical properties (IOP) values relative to the concentration (or mass) of biogeochemical parameters (here b_{bp}/PON_{wet}) are sensitive to these second-order effects. While a relatively constant b_{bp}/PON_{wet} ratio indicates homogeneous bulk particulate matter in terms of backscattering efficiency in the surface layer, an increase at depth indicates modification and/or heterogeneity of bulk particulate matter. Our result suggests that the influence of different particle composition and/or a potential presence of distinct microbial communities observed during the OUTPACE cruise (diazotrophic organisms in the ML and DCM microbial communities below the ML, [25,38]) was not significant compared to changes in organic particulate concentration. Thus, it seems that biomass changes are a dominant source of b_{bp} variability, as previously reported by [39]. These observations are also in agreement with those of [19], who reported that the POC vs b_{bp} relationship was relatively insensitive to community composition. In contrast, below the DCM (>150 dbar), an increase in b_{bp}/PON_{wet} ratios is observed. This finding could reflect the presence of an assemblage of particles dominated by detritus and heterotrophs (but requires further analysis based on a better characterization of the bulk particulate matter below the DCM). The main conclusion of these observations is that the highly sensitive wet oxidation method provides a good b_{bp} vs PON_{wet} relationship for this area between the surface and 150 dbar, and valid between 0.02 and 0.95 µM (~50-fold biomass increase), which thanks to float-based measurements of b_{bp} can be leveraged to expand both the number of observations and the spatiotemporal scales resolved.

During the periods of no bloom in the Melanesian Archipelago, and throughout the study period in the SPG, PON_{opt} concentrations were remarkably stable and low (<0.20 μM), displaying a net balance between production (or supply) and remineralization (or removal) processes [24]. The relative increase of Chl_a and the constant values of PON_{opt} during the winter periods suggest a seasonal change in Chl_a cell quota due to a lower light intensity in winter coupled with deeper mixed layer depths [40,41]. As a result, the constant PON_{opt} suggests that the marked Chl_a seasonal cycle, which can be falsely interpreted as a biomass variation, is in fact not related to new production but is instead a steady phytoplankton biomass. In our study region, the absence of phytoplanktonic blooms could be linked to nitrogen limitation [1], while N_2 fixation could be constrained by lower iron availability [42,43].

During the bloom periods of the austral summer, coinciding with the increase of Chl_a concentrations, PON_{opt} concentrations increased by a factor of two to five only in the Melanesian Archipelago. A co-variation between Chl_a concentrations and b_{bp} values (or PON_{opt} in our case) is linked to phytoplankton biomass production [44–47]. The results obtained in this study extend this observation to subtropical areas, pointing to outstanding increases of living phytoplanktonic
biomass and associated particles, even in oligotrophic regions. To explain the recurrent increases of phytoplanktonic biomass during austral summer periods, it is necessary to identify the nitrogen sources in the mixed layer. The two first blooms (March/April 2015 for both FA and FB) reported in this study occurred during the OUTPACE cruise. During those blooms, high N\textsubscript{2} fixation rates were reported while NO\textsubscript{3} was extremely low. Regarding the diazotroph bloom observed at station LDB, mesoscale vertical fluxes were too weak to displace the nitracline [48]. As a result, delivered nitrogen to the surface from mesoscale activity could not sustain the observed primary production. Regarding the blooms observed in our study after the cruise, except during February 2017 in the EMA no significant decreases of salinity were measured in parallel to PON\textsuperscript{opt} increases over the study period [Fig. 5], suggesting no direct influence of precipitation or riverine inputs.

Furthermore, atmospheric particle deposition fluxes are very low in the region [49]. The top of the nitracline was 90 dbar in the area during the summer [6], significantly deeper than the measured MLD (<60 dbar, Fig. 4). However, physical processes could vertically displace isopycnals tens of meters along with the nitracline [50]. Nevertheless, extreme events such as cyclones reported in this area perturbed the phosphacline but not the deeper nitracline, resulting in surface increases of phosphate but not of nitrate [51]. Therefore, in the absence of vertical N supply, the variations of PON\textsuperscript{opt} appear to be a valuable proxy of new production related to intense N\textsubscript{2} fixation events during austral summer conditions. In this specific environment characterized by active biological N\textsubscript{2} fixation, most of the new nitrogen is rapidly integrated into the phytoplanktonic biomass. Even if part of the nitrogen is released to the labile dissolved organic and inorganic pools, it is rapidly re-assimilated by the nitrogen-starved organisms [52,53]. We therefore conclude that the increases of PON\textsuperscript{opt} are most likely related to living POM, i.e. phytoplankton cells, and, to a lesser extent detritus.

In our study, we propose to use an optical proxy of PON rather than the traditional optical proxy of POC for several reasons. First, POC values show concentrations below 4 µM in oligotrophic gyres [54]. This limit value is close to the uncertainty reported in previous studies predicting POC from b\textsubscript{bp} [9,19]. The use of PON would be a superior alternative as the wet oxidation procedure, coupled with spectrophotometric measurements, allows accurate quantification of
PON (quantification limit = 0.02 µM which correspond to 0.13 µM in terms of POC following the Redfield proportion), with reasonable volumes of filtered seawater (1.2 L). Second, oceanic primary production is primarily controlled by nitrogen availability [55,56]. Thus, it would be helpful, in order to better understand controls of primary production, to gain insight on nitrogen cycling, particularly on the seasonal cycle of nitrogen pools directly, rather than carbon pools. This argument is even more important to consider in environments characterized by active biological N\textsubscript{2} fixation [5] where N budgets are needed [6]. Theoretically, all essential elements may be used to track primary production rates and biomass accumulations. Relative to carbon, the ~30 essential nutrient elements display a range of plasticity in their cellular requirements [57], with N being the least plastic by varying in cellular N:C molar ratios from ~1/5 to 1/10, a factor of two [58]. This variability is low compared to changes in biomass at the scale of the world Ocean, and even in the oligotrophic WTSP [6]. Therefore, if biomass is dominated by phytoplankton, a close link between PON and POC is to be expected. Similarities notwithstanding, the relative ease of PON measurements in oligotrophic areas favors the use of a PON optical proxy.

In addition to PON\textsubscript{(wet)}, the wet oxidation method can simultaneously measure POP\textsubscript{(wet)}. Interestingly, the b\textsubscript{bp} vs POP\textsubscript{(wet)} relationship was statistically better than b\textsubscript{bp} vs PON\textsubscript{(wet)} [Fig. 3; Table 2], highlighting the possible use of POP as a biomass proxy, which would drive an accurate estimation of standing POP stocks and associated biogeochemical fluxes. The better relationship between POP and b\textsubscript{bp} than PON and b\textsubscript{bp} suggests that the bulk particulate matter of POP could be less optically variable than PON-bulk particulate matter. This result could be linked to the variability of particle composition between both PON and POP pools. One main signature of a given particulate organic matter pool is the relative contributions of living particles and detrital material. Turnover rates of POP have been previously shown to be significantly higher than POC in the subtropical south Pacific [59], with the carbon and nitrogen pools containing more refractory material than the phosphorus pool [60–62]. Consequently, POP was considered to contain a higher contribution of living particles than PON or POC [59]. Therefore, the significant b\textsubscript{bp} vs POP\textsubscript{(wet)} relationship obtained [Fig. 3(c); Table 2] suggests that the variability of b\textsubscript{bp} could be linked, at least for this study area, more to the abundance of phytoplanktonic biomass than to associated material, as previously reported by [63] and [64]. Consequently, the optical proxy of POP estimated from b\textsubscript{bp} could also be a living biomass indicator and offer its own unique perspectives.

5. Conclusion

The combined use of in situ PON (and POP) measurements using the wet oxidation method, and optical properties measured by BGC-Argo floats close to in situ measurements, allowed to highlight, for the very first time, the excellent b\textsubscript{bp} vs PON (and POP) relationships between 0 and 150 dbar in under-sampled oligotrophic waters. In such areas, the quantification of POP\textsuperscript{opt}, in the range between 0.02 and 0.95 µM, documents phytoplanktonic biomass dynamics (and associated properties) at the relevant seasonal temporal scale. Increases of POP\textsuperscript{opt} by a factor of 5 (from 0.16 to 0.80 µM) were observed in the mixed layer of the WTSP during stratified conditions in the absence of significant nitrogen sources other than N\textsubscript{2} fixation. The pertinence of the b\textsubscript{bp} vs PON (POP) relationships should be investigated in other oligotrophic areas, but also in other trophic regimes. This new relationship also opens a promising avenue to assess PON (POP) from ocean color remote sensing using various existing inverse methods [65–67].

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Disclosures

The authors declare no conflicts of interest.

References


