Planktic foraminifer and coccolith contribution to carbonate export fluxes over the central Kerguelen Plateau

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A B S T R A C T
We report the contribution of planktic foraminifers and coccoliths to the particulate inorganic carbon (PIC) export fluxes collected over an annual cycle (October 2011/September 2012) on the central Kerguelen Plateau in the Antarctic Zone (AAZ) south of the Polar Front (PF). The seasonality of PIC flux was decoupled from surface chlorophyll a concentration and particulate organic carbon (POC) fluxes and was characterized by a late summer (February) maximum. This peak was concomitant with the highest satellite-derived sea surface PIC and corresponded to a Emiliania huxleyi coccoliths export event that accounted for 85% of the annual PIC export. The foraminifer contribution to the annual PIC flux was much lower (15%) and dominated by Turborotalita quinqueloba and Neogloboquadrina pachyderma. Foraminifer export fluxes were closely related to the surface chlorophyll a concentration, suggesting food availability as an important factor regulating the foraminifer’s biomass. We compared size-normalized test weight (SNW) of the foraminifers with previously published SNW from the Crozet Islands using the same methodology and found no significant difference in SNW between sites for a given species. However, the SNW was significantly species-specific with a threefold increase from T. quinqueloba to Globigerina bulloides. The annual PIC:POC molar ratio of 0.07 was close to the mean ratio for the global ocean and lead to a低碳ate counter pump effect (~5%) compared to a previous study north of the PF (6–32%).

1. Introduction

The Southern Ocean is the largest high nutrient, low chlorophyll (HNLC, Minas et al., 1986) area of the global ocean (Martin et al., 1990; Minas and Minas, 1992). Downstream of Subantarctic island plateaus, iron input from shelf sediments and glacial melt water can alleviate iron limitation and support large scale and long-lasting phytoplankton blooms (Blaın et al., 2001, 2007; Pollard et al., 2007; Tarling et al., 2012). These blooms are dominated by diatoms (Armand et al., 2008; Korb et al., 2008; Quéguiner, 2013) that respond to high macronutrient concentrations, marked turbulence, deep mixed layer depths and usually moderate light levels (Smetacek, 1985; Boyd, 2002; Strzepek et al., 2012). Diatom blooms result in a major contribution of biogenic silica to biomineral production of Southern Ocean waters, although biogenic production of calcium carbonate by calcifying planktonic organisms such as coccolithophores, foraminifers and pteropods can also occur. Although neglected for a long time, the presence of coccolithophores in the Southern Ocean has been diagnosed based on an increasing number of direct observations (Winter et al., 2014) and the development of remote sensing methods (Balch et al., 2005, 2011, 2014). Southern Ocean coccolithophore populations are dominated by the cosmopolitan species Emiliania huxleyi (Saaavedra-Pellitero et al., 2014; Winter et al., 2014) that is thought to be the major component of the "great calcite belt" observed in the vicinity of the Subantarctic Front (SAF) and Polar Front (PF)
(Balch et al., 2014). Several studies have reported modern planktic foraminifer abundances and fluxes in the Southern Ocean from net tows (Asioli and Langone, 1997; Mortyn and Charles, 2003; Bergami et al., 2009; Meilland, 2015) and sediment traps (Donner and Wefer, 1994; King and Howard, 2003; Northcote and Neil, 2005; Salter et al., 2014). Foraminifer assemblages are characterized by a southward dominance of polar species Neogloboquadrina pachyderma. In a review, Hunt et al., (2008) compiled pteropod abundance in the Southern Ocean and reported a switch from a dominance of Limacina retroversa australis north of the PF to Limacina helicina antarctica south of the PF.

The presence of calcareous organisms has important implications not only for food web ecology of the Southern Ocean, but also for the cycling of carbon between the atmospheric, oceanic, and sedimentary reservoirs on various climatically relevant timescales. Two distinct carbon pumps operate to cycle carbon through these different reservoirs (Volk and Hoffert, 1985). The soft tissue scales. Two distinct carbon pumps operate to cycle carbon through these different reservoirs (Volk and Hoffert, 1985). The soft tissue pump transfers particulate organic carbon (POC) originating from photosynthetic production to the ocean interior and plays a key role in the sequestration of atmospheric CO2 (Sarmiento et al., 1988). The carbonate pump exports particulate inorganic carbon (CaCO3, PIC) mainly as detrital calcareous shells (Volk and Hoffert, 1985). Calcification in the mixed layer decreases total alkalinity (TA) and dissolved inorganic carbon (DIC) with a ratio 2:1 and acts as a net source of CO2 to the atmosphere over a seasonal timescale (Frankignoulle et al., 1994). If the PIC production is exported in the deep ocean below the permanent thermocline, the net impact on the atmospheric CO2 occurs at a much longer timescale corresponding to the ocean mixing time (~1000 years, Zeebe, 2012). This phenomenon is known as the “carbonate counter pump” effect. Additionally, it has been suggested that during the last glaciation, lower PIC:POC export ratio due to increased organic carbon export may have contributed to higher dissolution of the deep-ocean carbonate sediments, leading to a decrease in pCO2 compared to the interglacial periods (Archer and Maier-Reimer, 1994; Archer et al., 2000; Sigman and Boyle, 2000). Therefore the PIC:POC ratio of exported particles is likely to have a significant impact on the atmosphere-ocean CO2 fluxes from seasonal to geological timescales (Matsumoto et al., 2002; Sarmiento et al., 2002). More recently, in the Subantarctic Southern Ocean, the strong response of calcifying organisms to natural iron fertilization has been observed to increase the PIC:POC export ratio leading to a strong carbonate counter pump, lowering the efficiency of CO2 sequestration by the biological carbon pump (Salter et al., 2014).

Understanding how calcifying communities drive the carbonate counter pump requires a coupled description of the chemical composition and biological properties of different vectors driving CaCO3 export fluxes. Sediment trap studies provide a tractable framework to link detailed analyses of the morphological and physiological properties of exported calcareous particles (e.g. species composition, test size and test weight) with seasonal and annual geochemical budgets. In this context, the study by Salter et al. (2014) quantified a carbonate counter pump effect accounting for 6–32% of measured POC fluxes with a notable contribution from foraminifer species (mainly Globigerina bulloides and N. pachyderma) in iron-fertilized waters downstream of the Crozet Islands. Several studies have reported geochemical transitions in particle stoichiometry across the Polar Front (Trull et al., 2001; Honjo et al., 2008), highlighting the importance of regional variability for a Southern Ocean carbonate counter pump that is partly linked to the biogeography of calcareous organisms (Salter et al., 2014).

The objectives of the present study are to (1) quantify the magnitude of POC export and the carbonate counter pump in an iron fertilized area (the Kerguelen Plateau) south of the Polar Front (Antarctic Zone, AZ), (2) determine the relative contribution of foraminifer and coccolithophores to total PIC export in this regime, and (3) constrain the importance of species composition and test characteristics (size and size-normalized weight) for foraminifer-mediated PIC fluxes in iron fertilized blooms of the Southern Ocean.

2. Materials and methods

2.1. Sediment trap deployment and environmental data

As part of the KEOPS2 project (Kerguelen Ocean and Plateau compared study 2), a sediment trap (Technicap PPS, 2.5 aspect ratio) was moored for 11 months (21 October 2011 to 7 September 2012) at 289 m over the central Kerguelen Plateau (seafloor depth 527 m) at station A3 (50°38.3 ′ S–72°02.6 ′ E, Fig. 1a,b). The carousel comprised 12 sampling cups (250 mL) containing 5% formalin hypersaline solution buffered with sodium tetraborate (pH~8). A detailed description of the sample processing and particulate organic carbon (POC) analyses are provided in Rembauville et al. (2015b). Briefly, swimmers (zooplanktonic organisms actively entering the trap) were manually removed, samples were freeze-dried and the carbonate fraction was dissolved by the addition of acid before the organic carbon content was measured with a CHN analyzer.

Station A3 is characterized by a recurrent and large phytoplankton bloom induced by natural iron fertilization coming from the underlying plateau (Blain et al., 2007). Dissolved iron (dFe) is delivered to the mixed layer through two processes: winter mixing and entrainment of dFe from deeper waters and, to a less extend, vertical diapycnal diffusion of dFe in summer (Bowie et al., 2015). South of the Kerguelen Island, the polar front is permanent and non-motive (Park et al., 2014) and therefore does not impact sediment trap deployment location. At the A3 station, the circulation is weak (< 3 cm s−1) and primarily tidal-driven (Park et al., 2008). Physical data acquired during the sediment trap deployment suggest the record was not subject to major hydrodynamic biases (Rembauville et al., 2015b), allowing a detailed and quantitative discussion of the export fluxes.

Satellite-derived surface chlorophyll a and PIC concentration (MODIS 8 days product, accessed at http://oceancolor.gsfc.nasa.gov/cms/), and sea surface temperature (NOAA OISST, weakly product, Reynolds et al., 2007) were extracted for a 100 km radius around the trap location. Calcite saturation state was calculated in the vicinity of the trap location with the CO2sys toolbox using climatological fields of DIC and Alkalinity (GLODAP, Key et al., 2004) and temperature, salinity, silicate and phosphate (World Ocean Atlas 2013, Garcia et al., 2013). Constants recommended for best practice were used (Dickson et al., 2007) as suggested by Orr et al. (2015).

2.2. Calcium analyses in the bulk and fine fractions

For bulk particulate inorganic carbon analyses, 5 mg of freeze-dried material was weighed (Sartorius MC 210 P balance) into Teflon vials for the mineralization. 1 mL of 65% HNO3 was added and samples were placed in an ultrasonication bath for 20 min. Samples were then dried overnight at 130 °C. 0.5 mL of 65% HNO3 and 0.5 mL of 40% HF were added and samples were ultrasonicated a second time and dried overnight. The resulting residue was dissolved in 10 mL of 0.1 N HNO3 and calcium content analyzed by inductively coupled plasma – optical emission spectrometry (ICP-OES, Perkin-Elmer Optima 2000). The efficiency of the mineralization procedure was estimated using reference material (GBW-07314) and was > 96%.

For the fine fractions (20–63 μm and < 20 μm) Ca analyses, the
original 1/8 split samples (Rembauville et al., 2015b) were further split into 1/80 aliquots with a rotary wet-splitter (McLane WSD-10) using purified water (Elix by Millipore purification system) buffered with ammonia as a rinse solution. Coccoliths in sinking particles captured in sediment trap samples may be contained in faecal pellets and/or phytoplankton aggregates. To improve the efficiency of size fraction separation by sieving it is necessary to oxidize the samples to disaggregate particles and retrieve the entire carbonate fraction (Bairbakhish et al., 1999; Broeze et al., 2000; Ziveri et al., 2000; Stoll et al., 2007). The 1/80 aliquots were placed in a 50 mL centrifugation tube for the oxidation steps using a method adapted from Bairbakhish et al. (1999). Samples were centrifuged (5000 rpm, 5 min) and the supernatant withdrawn. Subsequently, 3 mL of Elix water buffered with ammonia, 3 mL of 5% NaClO and 1.5 mL of 30% H2O2 were added and the samples were ultrasonicated for 10 s. Every 10 min, 2 mL of NaClO were added and samples were ultrasonicated for 10 s. This cycle was repeated for one hour. The oxidized aliquot was wet-sieved over a 63 μm and a 20 μm mesh, and the two resulting size fractions (20–63 μm and < 20 μm) were filtered on polycarbonate membranes (0.4 μm pore size, 47 mm diameter). Filters were dried at 40 °C and the residue was leached in 10 mL 1% HNO3, ultrasonicated for 10 min and left 12 h at room temperature before the Ca analysis. Ca concentration was analyzed by inductively coupled plasma-atomic emission spectrometry (ICP-AES, Perkin Elmer, Optima 4300DV). Overall accuracy amounted to better than 2% based on replicate analysis.

For the qualitative analyses of the coccolithophore species composition, samples were prepared in a similar way as for the fine fraction Ca analysis (oxidation and sieving) and then filtered on cellulose acetate membranes (Millipore, 0.45 μm pore size, 47 mm diameter). Filters were dried at 40 °C and observed under a polarized microscope at 1200 magnification.

2.3. Foraminifer carbonate flux estimation

Foraminifer quantification, morphometric measurements and weighing was performed following the methods outlined in Salter et al. (2014). One 1/8 aliquot was sieved on a 63 μm mesh with tap water and the > 63 μm fraction was dried overnight (40 °C). Dried particles were homogeneously placed on a glass tray. Images of the entire 1/8 sample were acquired with a fully automated incident light monocular microscope (Leica Z16 APO), and a motorized stage with a Lstep-PCI controller (Märzhäuser). High-resolution images (1.4 μm^-2 pixel^-1) were taken with a color camera (SIS CC12). Particle size (minimum test diameter, d min) was automatically analyzed using analySIS FIVE software (SIS/Olympus with a MAS software add-in). Foraminifer species were manually counted and classified into morpho-species following the taxonomic concept of Hemleben et al. (1989). Eight species of planktic foraminifer were identified: Neogloboquadrina pachyderma (left coiling), Neogloboquadrina incompta (right coiling), Turborotalita quinqueloba, Globigerinoides uvula, Globigerinella glutinata, Globorotalia inflata, Globigerinoides ruber (sensu stricto) and Trilobatus sacculifer (normal type). Only one empty shell of pteropod (Limacina helicina) was found in the samples and therefore pteropod’s contribution to the passive carbonate flux was considered negligible. However, numerous pteropods were found as swimmers (distinguished by well preserved organic material) actively entering the trap in late summer (Rembauville et al., 2015b). Those shells were withdrawn from the samples as they were considered not to contribute to the passive flux. To determine size-weight relationships, individuals of N. pachyderma (n = 23), N. incompta (n = 10), T. quinqueloba (n = 60) were manually picked from samples representative of different flux conditions (spring, summer and winter). Individuals were placed in aluminium cups and weighed (Mettler Toledo XP2U, 0.1 μg precision). Samples were acclimatized in the weighing room for at least 12 h before the analysis. Once the test weight was determined, the minimal diameter (d min) of each individual was measured with the procedure described above. Size-weight relationships (W = a x d b) were constructed by fitting linear regressions to log-transformed data (Movellan et al., 2012). A species-specific relationship was developed for N. pachyderma, N. incompta and T. quinqueloba. For the other species, an average size-weight relationship was calculated by pooling the entire foraminifer dataset (n = 93). Parameters of the size-weight relationships are given in Table 1. Foraminifer carbonate flux was then calculated using the abundance and size from the whole dataset and species or group-specific size-weight relationships. We refer to the sum of foraminifer and fine fractions (20–63 μm and < 20 μm) PIC as “calculated PIC”.

2.4. Test size and size normalized weight comparison with assemblages from Crozet

Discrete measurements of the test size and weight of
foraminiferal individuals calculate the calculation of size-normalized weight (SNW), a commonly used descriptor of test density/thickness (Bijma et al., 1999; Beer et al., 2010a; Marshall et al., 2013). The SNW was calculated for each individual by dividing the weight by the minimum test diameter (SNW (µg µm⁻¹) = W/dmin). Given the good relationship between area and minimum diameter, this method is considered as an appropriate mean to characterize the test density (Beer et al., 2010b). We compared the Kerguelen data for three species independently (SNW, a commonly used descriptor of test density/thickness (Bijma et al., 1999; Beer et al., 2010a; Marshall et al., 2013). Given the good relationship between area and minimum diameter, this method is considered as an appropriate mean to characterize the test density (Beer et al., 2010b). We compared the Kerguelen dataset (station A3, AAZ) with previously published size and weight data using the same methodology from the Crozet Islands (Salter et al., 2014). Stations M10 and M5 are located in the PFZ (Pollard et al., 2002; Planquette et al., 2007; Salter et al., 2014). Statistical differences in minimum diameter (dmin) and size-normalized weight (SNW) between the four study sites were tested for three species independently (N. pachyderma, T. quinqueloba, G. bulloides) using a non-parametric Kruskall-Wallis test. If the four sites constituted a significantly different groups, a post-hoc Tuckey test was performed to identify which sites were significantly different from the others. If the four sites constituted a significantly homogeneous group, the data from the four sites were pooled for each species and differences between the three species were tested using a Kruskall-Wallis test followed by a Tuckey post-hoc test. All tests were performed at a significance level of 5%.

3. Results

3.1. Seasonality of POC and bulk PIC fluxes

Surface chlorophyll a concentration displayed two peaks (Fig. 2a). The major peak (2.5 µg L⁻¹) occurred during spring at the onset of thermal stratification (November 2011) and a second moderate peak (1 µg L⁻¹) in summer (January 2012). PIC fluxes were characterized by two short (<15 days) and intense (∼1.5 mmol m⁻² d⁻¹) export events lagging the chlorophyll a peaks by one month. These two PIC export events comprised primarily Thalassiosira antarctica and Chaetoceros Hyalocheae resting spores (Rembauville et al., 2015a).

The satellite-derived mixed layer PIC concentration displayed a clear seasonal pattern (Fig. 2a) with moderate values in spring (0.4 µmol L⁻¹ in October/November 2011) and a strong increase in summer to reach nearly 1 µmol L⁻¹ in end January 2012. The PIC concentration decreased gradually after this summer peak to reach low values of 0.2 µmol L⁻¹ in winter 2012. Total bulk PIC fluxes displayed a similar seasonality as the surface satellite-derived PIC concentration (Fig. 2b). A moderate peak of 33 µmol m⁻² d⁻¹ in the first cup (21 October to 4 November 2011) was followed by very low fluxes for the remainder of spring (<10 µmol m⁻² d⁻¹). PIC fluxes gradually increased in the summer to 30 µmol m⁻² d⁻¹ before a clear maximum in late summer (110–120 µmol m⁻² d⁻¹) that persisted for one month (25 January to 22 February 2015). Autumn and winter fluxes were very low (<12 µmol m⁻² d⁻¹). Assuming negligible PIC flux out of the collecting period (corresponding to the months of September and October characterized by low chlorophyll a concentration), the annual PIC export was low (6.6 mmol m⁻² yr⁻¹). The annually-integrated PIC:POC molar ratio was equal to 0.07.

3.2. Seasonal dynamics of foraminifer and coccolith export fluxes

The seasonality of total foraminifer test flux closely followed chlorophyll a dynamics (Fig. 3a). A major peak of 800 indiv. m⁻² d⁻¹ was observed in spring. In December, when surface chlorophyll a concentrations were low, the total foraminifer flux was very low (15 indiv. m⁻² d⁻¹). During the second surface chlorophyll a increase (January to mid-February), the total foraminifer flux increased again to reach values of 450–550 indiv. m⁻² d⁻¹. Foraminifer flux was very low in autumn (30 indiv. m⁻² d⁻¹) and negligible in winter. There was no major seasonal change in the foraminifer assemblage throughout the year. At an annual scale, 4 species dominated (>95%) the foraminifer flux. The community assemblage was dominated by T. quinqueloba (31.8%), closely followed by N. pachyderma (30.8%) with lower contributions of N. incompta (18%) and G. uvula (15.3%) (Table 2).

Total and fine fractions (20–63 µm and < 20 µm) PIC fluxes are presented in Fig. 3c. The 20–63 µm fine fraction displayed very low fluxes (<15 µmol m⁻² d⁻¹) throughout the year with maximum in February 2012. The fine fraction <20 µm fluxes followed a similar seasonal pattern as total PIC fluxes. Spring and summer (October to mid-January) were characterized by low fluxes with values <25 µmol m⁻² d⁻¹ and peaked to the highest values ∼100 µmol m⁻² d⁻¹ in late summer (February). In autumn and winter, the PIC fine fraction <20 µm fluxes were <15 µmol m⁻² d⁻¹.

<table>
<thead>
<tr>
<th>Species</th>
<th>dmin range (µm)</th>
<th>W range (µg)</th>
<th>a</th>
<th>b</th>
<th>R²</th>
</tr>
</thead>
<tbody>
<tr>
<td>N. pachyderma (n=23)</td>
<td>102–300</td>
<td>0.3–5.5</td>
<td>5.26 × 10⁻⁷</td>
<td>2.90</td>
<td>0.71</td>
</tr>
<tr>
<td>N. incompta (n=10)</td>
<td>128–230</td>
<td>0.9–3.0</td>
<td>3.98 × 10⁻⁴</td>
<td>1.61</td>
<td>0.77</td>
</tr>
<tr>
<td>T. quinqueloba (n=60)</td>
<td>132–340</td>
<td>0.3–4.9</td>
<td>3.54 × 10⁻⁹</td>
<td>3.85</td>
<td>0.71</td>
</tr>
<tr>
<td>Global (n=93)</td>
<td>102–340</td>
<td>0.3–5.5</td>
<td>1.25 × 10⁻⁷</td>
<td>3.16</td>
<td>0.67</td>
</tr>
</tbody>
</table>

Table 1 Parameters of the size-weight relationship (W (µg) = a × dmin (µm)³) for the different foraminifer groups OR species considered. All the regressions are highly significant (p < 0.01).
seasonality comparable to the surface chlorophyll a with a strong peak in early spring (18 \( \mu \text{mol m}^{-2} \text{d}^{-1} \) in October 2011) and a secondary increase in late summer (11 \( \mu \text{mol m}^{-2} \text{d}^{-1} \) in January 2011). Fluxes were much lower the remainder of the year (<5 \( \mu \text{mol m}^{-2} \text{d}^{-1} \)). The relative contribution of each foraminifer species/group to the total foraminifer PIC and the calculated PIC annual flux is reported in Table 2. The relative contribution of the major foraminifer species to the total foraminifer PIC fluxes was comparable to their contribution to numerical fluxes, and a notable fraction (19%) of foraminifer PIC was exported as unclassified test fragments. T. quinqueloba displayed the highest contribution to the calculated PIC (4.9%), followed by N. pachyderma (3.3%) and N. incompta (1.8%). The contribution of G. uvula was very low (0.5%). Microscopic observations of the fine size fractions after the organic oxidation step revealed the absence of juvenile foraminifers and calcareous dinophytes in the 20–63 \( \mu \text{m} \) size fraction and the presence of coccoliths aggregated to diatom frustules and unidentified CaCO\(_3\) fragments. Therefore, the <20 \( \mu \text{m} \) fine fraction represents a slight underestimation of coccolith calcite fluxes (Ziveri et al., 2007). The total contribution of foraminifer tests to the annual calculated PIC export was 14.8%. Conversely, the contribution of the coccolith fine fractions (<20 \( \mu \text{m} \) and 20–63 \( \mu \text{m} \)) to the annual calculated PIC flux was high (85.2%), primarily due to their major contribution in the late summer export peak.

The relationship between the bulk and calculated PIC flux is presented in Fig. 4. Data points are close to the 1:1 relationship. A highly significant linear correlation (Pearson, \( n=12, p<0.01 \)) existed between the bulk and calculated PIC. Regression suggested a slope close to 1 (0.94, \( R^2=0.99 \)) and the annual calculated PIC export (6.5 mmol m\(^{-2} \text{d}^{-1} \)) was very close to the annual bulk PIC flux measured (6.6 mmol m\(^{-2} \text{d}^{-1} \)). These statistics ensure the analytical method was robust and the partitioning of PIC fluxes among the quantified biological vectors accounted for the majority of total PIC measured in the samples.

### Table 2
Relative contribution of foraminifer species to the annual numerical export and annual foraminifer PIC. Relative contribution of foraminifers and fine fractions (<63 \( \mu \text{m} \)) to the calculated annual PIC export.

<table>
<thead>
<tr>
<th>Species/group</th>
<th>Numerical foraminifer flux (%)</th>
<th>Foraminifer PIC (%)</th>
<th>Calculated PIC (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>N. pachyderma</td>
<td>30.8</td>
<td>22.6</td>
<td>3.3</td>
</tr>
<tr>
<td>N. incompta</td>
<td>18.0</td>
<td>11.9</td>
<td>1.8</td>
</tr>
<tr>
<td>T. quinqueloba</td>
<td>31.8</td>
<td>32.8</td>
<td>4.9</td>
</tr>
<tr>
<td>G. uvula</td>
<td>15.3</td>
<td>3.4</td>
<td>0.5</td>
</tr>
<tr>
<td>Other foraminifer species</td>
<td>4.1</td>
<td>10.3</td>
<td>1.4</td>
</tr>
<tr>
<td>Foraminifer fragments</td>
<td>19.0</td>
<td>2.8</td>
<td></td>
</tr>
<tr>
<td>Total foraminifers</td>
<td>14.8</td>
<td>85.2</td>
<td></td>
</tr>
<tr>
<td>&lt;63 ( \mu \text{m} )</td>
<td>10</td>
<td></td>
<td></td>
</tr>
<tr>
<td>20–63 ( \mu \text{m} )</td>
<td>75.2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>&lt;20 ( \mu \text{m} )</td>
<td></td>
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</tr>
</tbody>
</table>

### 3.3. Relative contribution of foraminifers and coccoliths to carbonate export

The individual size-weight relationships were considered sufficiently reliable to calculate the contribution of each foraminifer species to the PIC export (all fits were highly significant, \( R^2 > 0.66 \), Table 1). The total foraminifer-mediated PIC export showed a
assemblages

Probability histograms of size distribution at each site for \textit{N. pachyderma}, \textit{T. quinqueloba} and \textit{G. bulloides} are presented in Figs. 5a, b and c, respectively. All the density functions displayed quasi-unimodal distributions. For \textit{N. pachyderma}, \( d_{\text{min}} \) was significantly higher in the AAZ (M6 and A3 sites, \( 195 \pm 39 \mu m \), mean \pm standard deviation) than the PFZ (M5 and M10 sites, \( 151 \pm 30 \mu m \)). For \textit{T. quinqueloba}, \( d_{\text{min}} \) was significantly higher at A3 (\( 206 \pm 51 \mu m \)) than at the three other sites (M5, M10 and M6, \( 167 \pm 29 \mu m \)) that constituted a significantly homogeneous group. Only 5 g. \textit{bulloides} were observed at A3 and therefore were not taken into account in the analysis. For \textit{G. bulloides}, \( d_{\text{min}} \) was significantly homogeneous at the three Crozet sites (M5, M10 and M6, \( 244 \pm 65 \mu m \)).

Boxplots of SNW are presented for the three species in Fig. 6. For each species, there was no significant difference in SNW among sites. Therefore, the data from all the sites were pooled by species. Each species SNW constituted a significantly homogeneous group different from the two others. \textit{G. bulloides} SNW (\( 31 \pm 14 \times 10^{-3} \mu g \mu m^{-1} \), mean \pm standard deviation) was significantly higher than \textit{N. pachyderma} SNW (\( 18 \pm 11 \times 10^{-3} \mu g \mu m^{-1} \)) that was also significantly higher than \textit{T. quinqueloba} SNW (\( 10 \pm 4 \times 10^{-3} \mu g \mu m^{-1} \)) (Fig. 6).

4. Discussion

4.1. Foraminifer test flux amplitude and seasonality

We observed moderate planktic foraminifer test fluxes of 500–1000 indiv. m\(^{-2}\) d\(^{-1}\) despite high primary production levels in this naturally iron-fertilized area. The low test fluxes we report over the central Kerguelen Plateau, and the dominance of \textit{N. pachyderma} and \textit{T. quinqueloba} are consistent with the general decrease in flux from the SAZ to the AAZ that goes with a switch from a mixture of subpolar and polar water species to a dominance of the two aforementioned species. Donner and Wefer (1994) reported very low fluxes (\(~50\) indiv. m\(^{-2}\) d\(^{-1}\)) in the Northern Weddell Sea and Bransfield Strait (AAZ) whereas fluxes where much higher at the Maud Rise (\(~1 \times 10^3\) indiv. m\(^{-2}\) d\(^{-1}\)) where \textit{N. pachyderma} dominated the community assemblage, followed by \textit{T. quinqueloba}. King and Howard (2003) reported foraminifer export fluxes south of Tasmania with highest numerical fluxes of \(~1 \times 10^4\) indiv. m\(^{-2}\) d\(^{-1}\) in the SAZ very close to the SAF and lower values (\(4 \times 10^3\) indiv. m\(^{-2}\) d\(^{-1}\)) in the PFZ. The transition from SAZ to PFZ was associated with a switch from temperate species to a dominance of \textit{N. pachyderma} and \textit{T. quinqueloba}. South of New Zealand, Northcote and Neil (2005) described fluxes of \(5 \times 10^3\) indiv. m\(^{-2}\) d\(^{-1}\) with a major contribution of \textit{G. inflata} in the SAZ. In the PFZ North of the Crozet Islands, foraminifer numerical export fluxes were
~1 × 10^4 indiv. m^-2 d^-1 and mostly represented by N. pachyderma with a notable contribution of the larger temperate species G. bulloides and G. inflata (Salter et al., 2014).

The seasonal dynamics of foraminifer test export flux at station A3 was characterized by two peaks in spring and summer closely related with surface chlorophyll a concentration, but were not particularly associated with SST dynamics. Jonkers and Kučera (2015) have analyzed the phenomenology of foraminifer export fluxes at global scale and demonstrated that a group composed of temperature and cold water species (comprising N. pachyderma, N. incompta and T. quinqueloba) displayed two export peaks in spring and summer. Our results are highly consistent with this general scheme and support the close link between primary production (assessed from surface chlorophyll a) and foraminifer production (Hemleben et al., 1989; Kaas, 2001; Schieber et al., 2001; Kuruyanagi and Kawahata, 2004; Lombard et al., 2011) and subsequent export (Schiebel, 2002). At Crozet (M5 and M10 sites in PFZ) foraminifer test export occurred in one continuous event in summer from January to March (Salter et al., 2014) when SST was generally highest (> 8 °C) and chlorophyll a concentration was low (0.5 µg L^-1, Salter et al., 2012). This strongly contrasts with the close link we observe between the chlorophyll a concentration and the foraminifer test export at A3. However, the comparison of flux seasonality must be treated with caution because of the different sediment trap deployment depths (289 m at A3 versus > 2500 m for the M5 and M10 sites), increasing water depth might dampen seasonal particle flux signal. Our results from a shallow sediment trap at A3 suggest that food availability might be the major controlling factor for low temperature communities of the AAZ.

4.2. Foraminifer test size and SNW distribution

The calculation of the calcite saturation state is strongly dependent on the input variables of DIC and Alkalinity (a 1% change in one of these variables can drive a 10% change in saturations state, Orr et al., 2015). Given this uncertainty, the climatological field suggests that all of the sediment trap deployments around Crozet and Kerguelen were located in waters oversaturated with respect to CO_3^{2-} with a calcite saturation state > 1 (Fig. 1b,c). Therefore it is unlikely that seawater carbonate chemistry has strongly affected test weight and size through dissolution during particle sinking. However, test dissolution would lead to an underestimation of the weight in the sediment trap material and therefore the SWN should be considered as a lower estimate compared to living individuals.

The compilation of the large dataset generated with the automated microscope from Crozet and Kerguelen samples revealed that location relative to the Polar Front had a significant impact on the size of N. pachyderma with smaller individuals in the PFZ (Fig. 5). This pattern was not evident for T. quinqueloba and G. bulloides. When food is not limiting, temperature is presumed a fundamental factor influencing foraminifer growth rate at the species level (Lombard et al., 2009). An explanation of the Bergmann’s rule (larger individuals in colder environments) in plankton is that lower growth rate due to lower temperature leads to larger individuals at sexual maturity (von Bertalanffy, 1960; Atkinson, 1994). Under this hypothesis, colder SST south of the Polar Front might explain larger individuals of N. pachyderma at M6 and A3 sites. However the fact that Crozet communities of T quinqueloba and G. bulloides have a significantly homogeneous size in the PFZ and AAZ suggests that temperature is not the only factor at play and that population dynamics (Schiebel et al., 1997) and the availability of prey (Schmidt et al., 2004) as well as genetic diversity within a given morphospecies (Weiner et al., 2015) might also constrain planktonic foraminifer size.

SNW was originally considered as a proxy for CO_3^{2-} (Lohmann, 1995; Bijma et al., 1999; Broecker and Clark, 2001; Barker and Elderfield, 2002; Bijma et al., 2002). Additionally, the comparison of foraminifer tests from modern sediment traps samples and Holocene sediments demonstrated the impact of ocean acidification and the lowering of CO_3^{2-} on the reduction of the test weight at high southern latitudes (Moy et al., 2009). However, there is a growing number of observations suggesting that the relationship between the SNW and the CO_3^{2-} is not homogeneous among foraminifer species (Beer et al., 2010a; Meiland, 2015), and the relationship is more robust for certain species than for others (Marshall et al., 2013). Our results show that for a given species, SNW is not statistically different regarding the hydrography but that SNW varies significantly between the dominant species N. pachyderma, T. quinqueloba and G. bulloides. This suggests that ecological conditions other than the carbon chemistry of ambient seawater at long (Weinkauf et al., 2013) and short time scale (de Villiers, 2004; Marshall et al., 2013), and species physiological characteristics and metabolism might be responsible for the three-fold SNW increase between T. quinqueloba and G. bulloides. This has potentially important implications for the carbon pumps because it implies that planktic foraminifer community composition together with the magnitude of the numerical flux (number of individuals) plays a role in the foraminifer-mediated PIC flux.

4.3. Seasonality and magnitude of the coccolith fine fraction export

The sediment trap record represents the first annual record of coccolith calcite export south of the Polar Front. Over the central Kerguelen Plateau, we observe a clear decoupling between the two chlorophyll a peaks (November and January) and the coccolith fine fraction (~20 µm) export peak (February). The algorithm used to calculate PIC concentration based on satellite remote sensing reflectance is associated with a root mean square error (RMSE) of 1.2 µmol L^-1 (Balch et al., 2005). The maximum satellite-derived PIC concentration we report is ~1 µmol L^-1 which is lower than the RMSE. Additionally, the sunlight penetration depth constraining satellite data is <20 m in such a productive area (Gordon and McCluney, 1975), preventing the detection of subsurface features. For this reason, we only consider the satellite-derived PIC as qualitative data product. The uncertainty on satellite-derived PIC concentration, the shallow sediment trap depth (289 m) and the sampling temporal resolution (15 days) prevent a robust calculation of coccolith sinking speed or turnover time. However, the satellite-derived PIC concentration displays a clear seasonal signal tightly coupled to the coccolith fine fraction export. This result suggests that the algorithm used to derive coccolithophore presence from satellite data (Gordon et al., 2001; Balch et al., 2005) is sensitive, if not quantitative, over the central Kerguelen Plateau.

Historical observations suggest a diatom to coccolithophore succession from spring to summer in various locations of the global ocean (Margalef, 1978; Holligan et al., 1983; Llochte et al., 1993; Ziveri et al., 1995; Thunell et al., 1996; Ziveri and Thunell, 2000; Schiebel et al., 2011). Using satellite hyperspectral measurements and the PhytoDOAS method, Sadeghi et al. (2012) built a climatology of coccolithophore biomass in the Southern Atlantic. They reported a recurrent coccolithophore bloom in February/March, in good agreement with our measurement of maximum fine fraction (~20 µm) export flux in February. Sadeghi et al. (2012) highlighted the importance of SST maxima for the origination of a coccolithophore bloom in the high latitude ocean. Similarly, we report the highest coccolith calcite export flux during the period of highest SST (~5 °C), in agreement with the hypothesis of a temperature control on the coccolithophore bloom. More recently, Hopkins et al. (2015) used satellite-derived PIC as a proxy of coccolithophore biomass and concluded to a co-
occurrence of chlorophyll $a$ and coccolithophore peaks in the Southern Ocean. The results at large spatial and temporal scales differ somewhat from the uncoupling we observe at our specific location. Such differences may be attributed to inter-annual variability in the seasonality of chlorophyll $a$ concentrations and/or the timing of coccolithophore production.

The qualitative microscopic observation of the $< 20 \mu$m and 20–63 $\mu$m fractions indicate that *Emiliania huxleyi* represents $> 95\%$ of the coccolithophores assemblage with a minor contribution of *Helicocphora cartieri*. This finding is consistent with previous observations of a strict dominance of *E. huxleyi* with low abundances south of the PF (Saaedra-Pellitero et al., 2014; Winter et al., 2014). *E. huxleyi* is reported to bloom in waters with generally low silicic acid concentration resulting by its consumption by diatoms (Holligan et al., 1983; Townsend et al., 1994; Tyrrell and Merico, 2004). Additionally, this species has been shown to be tolerant to low iron concentration (Brand et al., 1983; Sund and Huntsman, 1995; Muggli and Harrison, 1997; Findlay and Giraudeu, 2000; Holligan et al., 2010). In January, the silicic acid concentration at the station A3 reaches $< 2 \mu$mol L$^{-1}$ (Mosseri et al., 2008) and iron concentration is $\sim 0.1 \mu$mol L$^{-1}$ (Blain et al., 2008). Moreover, the high nitrate, phosphate and ammonium concentrations (Mosseri et al., 2008) and the highest SST in late summer might be favorable conditions for a *E. huxleyi* bloom. Nevertheless, despite the summer stratification, the SST of 5 °C is still in the lower end of the thermal niche of *E. huxleyi* (1–31 °C, McIntyre et al., 1970). This temperature is likely to result in relatively low growth rate (Fisher and Honjo, 1989; Fielding, 2013). This may explain why the magnitude of the bloom is weak and corresponds to low surface chlorophyll $a$ concentration at this period of the season. This weak coccolithophore bloom drives most (85.2%) of the annual PIC export that appears very low (6.6 mmol m$^{-2}$ y$^{-1}$) compared to coccolith fine fraction export from the temperate ocean (0.2–0.8 mol m$^{-2}$ y$^{-1}$, Ziveri et al., 2007).

4.4 Southern Ocean carbonate counter pump affected by different planktonic calcifying organisms

The annually-integrated PIC:POC export ratio of 0.07 (mol:mol) is close to the mean ratio for the global ocean (0.06 ± 0.03, Sarmentio et al., 2002) and appears much lower than the ratio found in sediment traps of the PFZ and the SAZ (1, from a data compilation by Salter et al., 2014). The annual POC export (98.2 mmol m$^{-2}$ yr$^{-1}$, Rembauville et al., 2015b) and the annual PIC export (6.6 mmol m$^{-2}$ yr$^{-1}$) at station A3 allow us to estimate the strength of the carbonate counter pump: the reduction of the CO$_2$ drawdown by the biological pump due to the CO$_2$ production during the calcification process in the mixed layer (Frankignouille et al., 1994; Zeebe, 2012; Salter et al., 2014). As the trap depth (289 m) was close to the winter mixed layer depth (220 m in this region of the Southern Ocean (Park et al., 1998; de Boyer Montégut et al., 2004)), POC fluxes were not corrected for attenuation with depth. The carbonate counter pump effect (CC$_{pump}$,%) was calculated from the annual fluxes as CC$_{pump}$ = (PIC$_{flux}$ × $\Psi$)/POC$_{flux}$ × 100. $\Psi$ is the mole of CO$_2$ emitted by mole of CO$_3^{2-}$ precipitated during the calcification process and ranges 0.7–0.8 for seawater at 5 °C and a pCO$_2$ of 300–400 μatm (Frankignouille et al., 1994). The calculation leads to a CC$_{pump}$ of 4.7–5.4% at station A3. This value is consistent with the previous reported value at the M6 site also located in the AAZ (1–4%) and is significantly lower that the values in the PFZ at the M5 and M10 sites (6–32%) reported in Salter et al. (2014).

In the PFZ downstream Crozet, foraminifiers were significant contributors to the production and export of PIC (30–50%), with a lower contribution of coccoliths (20%) and pteropods (5%, Salter et al., 2014). Conversely, foraminifiers are minor contributors over the central Kerguelen plateau in the AAZ (<15%, Table 2). The similarity of the CC$_{pump}$ between the M6 and A3 sites in the AAZ supports the idea that the position of productivity relative to the Polar Front (Salter et al., 2014) exerts a major control on the magnitude of the CC$_{pump}$ through two processes: (1) changes in the relative abundance of heterotrophic calcifiers foraminifers/pteropods to autotrophic coccolithophores, and (2) a change in the contribution of foraminifer species with different SNWs.

During the last two million years the glaciations have been characterized by lower CO$_2$ concentration in the atmosphere that has been explained by a combination of both biology (strengthening of the biological pump) and physics of the Southern Ocean (Sigman and Boyle, 2000; Kohfeld et al., 2005; Robinson et al., 2005; Martínez-Botí et al., 2015). The higher efficiency of the biological pump was likely linked to higher deposition of eolian iron and more complete utilization of nutrients at high latitudes (Mahowald et al., 2006; Martínez-Garcia et al., 2014). Our results from naturally fertilized Southern Ocean blooms suggest that the magnitude of the associated carbonate counter pump (Salter et al., 2014) depends not only on the dominant calcifying planktonic organisms (foraminifers versus coccolithophores), but also on the species assemblage that responds to the increase in primary production.

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