Latitudinal and temporal distributions of diatom populations in the pelagic waters of the Subantarctic and Polar Frontal zones of the Southern Ocean and their role in the biological pump

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Abstract. The Subantarctic and Polar Frontal zones (SAZ and PFZ) represent a large portion of the total area of the Southern Ocean and serve as a strong sink for atmospheric CO₂. These regions are central to hypotheses linking particle fluxes and climate change, yet multi-year records of modern flux and the organisms that control it are, for obvious reasons, rare. In this study, we examine two sediment trap records of the flux of diatoms and bulk components collected by two bottom-tethered sediment traps deployed at mesopelagic depths (∼1 km) in the SAZ (2-year record; July 1999–October 2001) and in the PFZ (6-year record; September 1997–February 1998, July 1999–August 2000, November 2002–October 2004 and December 2005–October 2007) along the 140° E meridian. These traps provide a direct measure of transfer below winter mixed layer depths, i.e. at depths where effective sequestration from the atmosphere occurs, in contrast to study of processes in the surface ocean. Total mass fluxes were about twofold higher in the PFZ (24 ± 13 g m⁻² yr⁻¹) than in the SAZ (14 ± 2 g m⁻² yr⁻¹). Bulk chemical composition of the particle fluxes mirrored the composition of the distinct plankton communities of the surface layer, being dominated by carbonate in the SAZ and by biogenic silica in the PFZ. Particulate organic carbon (POC) export was similar for the annual average at both sites (1.0 ± 0.1 and 0.8 ± 0.4 g m⁻² yr⁻¹ for the PFZ and SAZ, respectively), indicating that the particles in the SAZ were relatively POC rich. Seasonality in the particle export was more pronounced in the PFZ. Peak fluxes occurred during summer in the PFZ and during spring in the SAZ. The strong summer pulses in the PFZ are responsible for a large fraction of the variability in carbon sequestration from the atmosphere in this region. The latitudinal variation of the total diatom flux was found to be in line with the biogenic silica export with an annual flux of 31 ± 5.5 × 10⁸ valves m⁻² yr⁻¹ at the PFZ compared to 0.5 ± 0.4 × 10⁸ m⁻² yr⁻¹ at the SAZ. Fragilaria kerguelensis dominated the annual diatom export at both sites (43 % at the SAZ and 59 % in the PFZ). POC fluxes displayed a strong positive correlation with the relative contribution of a group of weakly silicified and bloom-forming species in the PFZ. Several lines of evidence suggests that the development of these species during the growth season facilitates the formation of aggregates and carbon export. Our results confirm previous work suggesting that F. kerguelensis plays a major role in the decoupling of the carbon and silicic cycles in the high-nutrient low-chlorophyll waters of the Southern Ocean.

1 Introduction

The Southern Ocean is a critical component of the Earth’s ocean–climate system and plays a pivotal role in the global biogeochemical cycles of nutrients and carbon. Due to its unique meridional overturning circulation, deep waters are upwelled south of the polar front supplying the surface waters with nutrients and allowing the ventilation of carbon.

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dioxide accumulated during centuries of deep-sea respiration (Anderson et al., 2002; Pollard et al., 2006). Strong interactions with the atmosphere transform the upwelled deep waters into bottom, intermediate, and mode waters, which spread to lower latitudes renewing the intermediate and abyssal depths of the world ocean (Sarmiento et al., 2004; Sigman et al., 2010). Dissolution of carbon dioxide in these newly formed waters (i.e. the solubility pump) and the transport of photosynthetically fixed carbon to depth in settling particles (i.e. the biological pump) modulate the extent to which the carbon dioxide is transferred to the atmosphere. The balance between these processes determines the role of the Southern Ocean either as a source or sink of atmospheric CO\textsubscript{2} over glacial–interglacial climate cycles (Kohfeld et al., 2005; Anderson et al., 2009; Sigman et al., 2010).

At present, the Southern Ocean biological pump is not operating at its full capacity (De La Rocha, 2010). Low sun angles, deep wind-mixed surface waters and lack of the micronutrient iron restrict phytoplankton growth (Boyd et al., 2007; Venables and Moore, 2010) making the Southern Ocean the largest high-nutrient low-chlorophyll (HNLC) region in the world ocean (Martin, 1990). However, there are exceptions to this situation with areas of higher phytoplankton abundance occurring along oceanographic fronts (Laubscher et al., 1993; Moore and Abbott, 2000), downstream of some islands (Blain et al., 2001; Park et al., 2010), in the wake of the retreating seasonal sea ice (Smith Jr. et al., 1988; Brzezinski et al., 2001), in coastal polynyas (Arrigo and van Dijken, 2003) and in coastal systems of Antarctica (Sedwick et al., 2000; Brzezinski et al., 2001). In these areas iron fertilises the surface layer triggering phytoplankton blooms in spring and summer.

Diatoms are one of the most abundant primary producers in the Southern Ocean and make a major contribution to the biogenic silica (BSi) content of deep-sea sediments. Extensive diatom blooms occasionally develop in the Antarctic Circumpolar Current (ACC; e.g. Kopeczynska et al., 2001; Green and Sambrotto, 2006; Alvin et al., 2008; Grigorov et al., 2014) which results in the formation of a band of siliceous ooze that encircles Antarctica (DeMaster, 1981; Buesseler et al., 2001; DeMaster, 2002). This “diatom ooze belt” constitutes the world’s largest sedimentary sink for BSi accounting for about one-third of the global BSi accumulation (Tréguer and De La Rocha, 2013; Tréguer, 2014). Therefore, the production and export of diatoms in the Southern Ocean are critical components of the global cycling of silica, and potentially the biological carbon pump. Indeed there is evidence from silicon isotopes and other palaeo-proxies that silica export in the Southern Ocean has varied, in concert with other biogeochemical changes and atmospheric CO\textsubscript{2} variations, over a range of timescales (Brzezinski et al., 2002; Matsumoto et al., 2002; Sarmiento et al., 2004).

Most of our current knowledge about the temporal and spatial dynamics of phytoplankton in the Southern Ocean waters derives from satellite observations and biogeochemical models (e.g. Moore et al., 1999; Moore and Abbott, 2000; Arrigo et al., 2008; Gregg and Rousseaux, 2014). Recently, advances in the interpretation of optical signals have allowed the determination of the specific contribution of major phytoplankton groups (e.g. coccolithophores, phaeocystis-like, diatoms) to phytoplankton abundance on a global and regional scale (e.g. Alvain et al., 2005; Raitos et al., 2008; Rousseaux and Gregg, 2012; Alvain et al., 2013). However, as a more complete picture of the structure of the planktonic communities emerges, it becomes evident that in order to determine the role of phytoplankton in the biological pump and biogeochemical cycles, it is of critical importance to distinguish not just among major taxonomic groups but also within them. In particular, recent studies have shown how variations in the population of diatom communities, which exhibit a wide range of competitive strategies, contribute to the regulation of the stoichiometric relationship between elements such as carbon and silicon in the global ocean (Boyd et al., 2010; Assmy et al., 2013; Boyd, 2013; Quéguiner, 2013).

Bottom-tethered sediment traps have contributed significantly to the characterisation of the spatial and temporal variability of biochemical and microorganism fluxes in the Southern Ocean (Romero and Armand, 2010). For example, sediment trap experiments have revealed that the particle export in this region is highly seasonal and that maximum fluxes of particulate matter occur in areas under the influence of seasonal sea ice where diatoms are most abundant (e.g. Fischer et al., 2002; Pilskaln et al., 2004; Grigorov et al., 2014). Other studies have revealed the crucial role of particular diatom species in driving the biological pump in naturally iron-fertilised waters, such as Chaetoceros and Thalassiosira resting spores around the Kerguelen Plateau (Rembauville et al., 2015) or the resting stages of Eucampia antarctica in the Crozet Islands system (Salter et al., 2012). Most of these studies have been carried out in areas of relatively high primary production such as coastal systems and areas under the influence of seasonal sea ice. However, very few sediment trap experiments have been conducted in the pelagic province that, despite its relatively low phytoplankton abundance, is responsible for approximately 90 % of the annual primary production in the Southern Ocean due to its large size (Arrigo et al., 2008).

In this work we document the seasonal and inter-annual variability of the chemical (total mass, BSi, carbonate and particulate organic carbon – POC) and biological (diatom assemblages) composition of the material captured at two mooring sites along the 140°E meridian, representative of a large proportion of the Subantarctic Zone (SAZ) and Polar Frontal Zone (PFZ). The main objectives of this study are

1. to document the latitudinal and seasonal variations in the composition of the particle fluxes and diatom communities across sites;
2. to assess the role of the seasonal variability of diatom communities on the biological pump and cycling of silica;
3. to provide annual estimates of biogenic silica, carbonate, POC and diatom-valve fluxes to the deep ocean for the SAZ and PFZ.

2 Oceanographic and biological setting

The Southern Ocean is divided into concentric zones surrounding Antarctica by a series of frontal systems (Fig. 1), which are characterized by large geostrophic surface velocities (Orsi et al., 1995) and are linked to contours of sea surface height (SSH; Sokolov and Rintoul, 2002, 2009b, a). Between these fronts lie zones of weak flow that tend to have relatively uniform water mass properties (Zentara and Kamykowski, 1981; Rintoul and Bullister, 1999) and biological characteristics (Boyd, 2002; Thomalla et al., 2011). The SAZ extends from the subtropical front (STF) to the Subantarctic Front (SAF) and represents a transition zone between the subtropical gyres to the north and the ACC to the south (Rintoul and Bullister, 1999). SAZ surface waters along ∼140°E have summer sea surface temperatures (SSTs) ranging between 11 and 12°C (Fig. 2), whilst the mixed layer depth during winter can exceed 600 m (Rintoul and Trull, 2001). The SAF is characterised by a marked latitudinal surface gradient in temperature and salinity and, in the Australian sector, is split into two branches or filaments at mean latitudes of 50.5 and 52°S (Sokolov and Rintoul, 2002). The PFZ lies just south of the SAZ and represents the northernmost extent of the Antarctic waters. PFZ surface waters have summer SSTs typically between 5 and 6°C in summer (Fig. 3), but the winter mixed layer is shallower (less than 200 m) than at the SAZ (Rintoul and Trull, 2001).

Together the SAZ and PFZ make up the sub-Antarctic region (Fig. 1), which is the site of the formation of the subantarctic mode and Antarctic Intermediate waters (SAMW and AAIW, respectively; McCartney, 1977). Both SAMW and AAIW are subducted northward beneath the subtropical gyres ventilating their lower thermocline (Sallée et al., 2006; Downes et al., 2009) and eventually supplying with nutrients the surface waters across the oceans of the Southern Hemisphere and North Atlantic (Sarmiento et al., 2004).

In terms of biogeochemical distributions, the SAZ and PFZ can be defined as HNLC regimes but with a remarkable difference between them. While PFZ surface waters are replete with phosphate, nitrate and silicate until at least midsummer, in the SAZ silicate remains at low levels throughout the year (Rintoul and Trull, 2001; Wang et al., 2001). Dissolved iron concentrations in the mixed layer along the 140°E longitude transect are low and exhibit a decreasing trend with increasing latitude, with 0.27 in the SAZ and 0.22 ±0.02 nmol L⁻¹ in the PFZ (Lannuzel et al., 2011). Primary production is thought to be co-limited by iron supply and light in the PFZ, and by iron supply and low silicic acid concentration in the case of the SAZ (Boyd et al., 1999; Boyd et al., 2001; Lannuzel et al., 2011).

As a consequence of these different physical and biochemical properties, the SAZ and PFZ exhibit two distinct phytoplankton communities. SAZ surface waters are dominated by coccolithophores, other flagellates and cyanobacteria with lower abundances of diatoms. PFZ waters are also rich in coccolithophores and flagellates, but contain few cyanobacteria, whereas diatoms are more abundant and of larger size (Popp et al., 1999; Kopczynska et al., 2001; de Salas et al., 2011). Finally, it is worthy to note a feature present in the PFZ but not in the SAZ; a subsurface chlorophyll maximum (SCM) dominated by large diatom species (Kopczynska et al., 2001) has been consistently reported during summer in the PFZ within or beneath the seasonal pycnocline. The formation and maintenance of this SCM is most likely due to the settling of phytoplankton cells as a response to iron and silicate co-limitation in the mixed layer during summer (Popp et al., 1999; Parslow et al., 2001).
Figure 2. (a) Available mean sea surface temperature (SST), photosynthetically available radiation (PAR) and chlorophyll \(a\) concentration for the 47\(^\circ\) S site. The dashed intervals represent the studied periods. (b) Temporal variability of the total and major component fluxes, and their relative contributions to the total mass flux for the < 1 mm fraction at 1000 m water depth at the 47\(^\circ\) S site for the period July 1999–October 2001. Biogenic silica, carbonate and particulate organic carbon (POC) were directly measured. “Other” indicates un-characterized mass contributions (organic components other than carbon and small amounts of lithogenic material; Trull et al., 2001). Grey horizontal bars highlight the summer period (December to February) of each year.

### 3 Material and methods

#### 3.1 Field experiment

A series of deep-moored sediment trap deployments was initiated in 1997 by the Australian SAZ program (Trull et al., 2001b) and now continues as a component of the Australian Integrated Marine Observing System Southern Ocean Time Series (Trull et al., 2010; Shadwick et al., 2015). Two sites representative of a large proportion of the SAZ and PFZ were occupied quasi-continuously for the decade 1997–2007. Both sites were located along the 140\(^\circ\) E longitude: station 47\(^\circ\) S was set on the abyssal plain of the central SAZ, whereas station 54\(^\circ\) S was placed on a bathymetric high of the Southeast Indian Ridge in the PFZ (Fig. 1, Table 1). Additionally, two other sites were instrumented over a 1-year period, beneath the SAF (site 51\(^\circ\) S, 1997–1998) and within the southern Antarctic Zone (AZ) (site 61\(^\circ\) S, 2001–2002). Here, we present data from the 47\(^\circ\) S 1000 m trap between 1999 and 2001 (2-year record) and from the 54\(^\circ\) S 800 m trap between the following years: 1997–1998, 1999–2000, 2002–2004 and 2005–2007 (6-year record). Biogenic particle flux data of sites 47, 51 and 54\(^\circ\) S for the first year deployment (1997–1998) and of site 61\(^\circ\) S for the year 2001–2002 have already been published in Trull et al. (2001a) and Rigual-Hernández et al. (2015), respectively.

All traps were MacLane Parflux sediment traps: conical in shape with a 0.5 m\(^2\) opening area and equipped with a carrousel of 13 or 21 sampling cups. Cup rotation intervals...
Figure 3. (a) Available mean sea surface temperature (SST), photosynthetically available radiation (PAR) and chlorophyll $a$ concentration for the 54° S site. The dashed intervals represent the studied periods. (b) Temporal variability of the total and major component fluxes, and their relative contributions to the total mass flux for the < 1 mm fraction at 800 m water depth at the 54° S site for the periods July 1999–August 2000, November 2002–October 2004 and December 2005–October 2007. Biogenic silica, carbonate and particulate organic carbon (POC) were directly measured. “Other” indicates un-characterised mass contributions (organic components other than carbon and small amounts of lithogenic material; Trull et al., 2001). Grey horizontal bars highlight the summer period (December to February) of each year.

Table 1. Deployment summary of sediment traps at stations 47, 54 and 61° S.

| Site and trap designations | Hydrographic zone | Latitude ° S | Longitude ° E | Water column depth (m) | Trap depth (m) |
|---------------------------|-------------------|--------------|---------------|------------------------|---------------|
| 47° S_1000                | SAZ               | 46°46′S      | 142°4′E       | 4540                   | 1060          |
| 54° S_800                 | PFZ               | 53°45′S      | 141°45′E      | 2280                   | 830           |
| 61° S_2000                | AZ                | 60°44′S      | 139°54′E      | 4393                   | 2000          |

were established based on anticipated mass fluxes. The shortest intervals corresponded with the austral summer and autumn ranging typically between 4.25 and 10 days, whereas the longest intervals were 60 days, corresponding with winter (Table 2). Each trap was paired with an Aanderaa current metre and temperature sensor. The 250 mL collection cups were filled with a buffered solution of sodium tetraborate (1 g L$^{-1}$), sodium chloride (5 g L$^{-1}$) and mercury chloride (3 g L$^{-1}$) in unfiltered deep seawater from the region (collected at 1200 m depth, 49°17′S, 153°58′E). Full details of the mooring designs can be found in Bray et al. (2000) and Trull et al. (2001a).
3.2 Quality check of downward particle fluxes

Current speeds largely influence the efficiency with which sediment traps collect the particles sinking in the water column (Baker et al., 1988; Yu et al., 2001). The threshold of current velocity above which sinking particles are no longer directed for possible under trapping in the present study. Additionally, radioisotope analyses of material from the first year deployment by Trull et al. (2001) provide some extra insights to assess the collection efficiency of the traps. The $^{230}$Th flux/production ratios for the 1997–1998 deployment were $0.6 \pm 0.1$ and $0.7 \pm 0.1$ for the 47 and 54° S traps, respectively. These values suggest that some degree of under trapping is likely to have occurred at both sites. However, as these values are almost identical for both traps, it can be assumed that the trapping efficiency did not account for the observed latitudinal variations in the magnitude of the particle export between sites. Taking into consideration all the above and the fact that the assessment of trapping efficiency from $^{230}$Th alone is fraught with uncertainties (Trull et al., 2001a; Buesseler et al., 2007), trap fluxes were not corrected for possible under trapping in the present study.

3.3 Determination of major constituents of the flux

A detailed description of the methodology used for the determination of the flux intensity and composition of settling particles for the first mooring deployments in 1997–1998 can be found in Bray et al. (2000) and Trull et al. (2001a). After
recovery, sediment trap cups were allowed to settle before supernatant was drawn off with a syringe for salinity, nutrients and pH measurements. The remaining sample slurries were sieved through a 1 mm sieve and then split into 10 fractions using a rotary splitter (McLane, Inc.). Three of these splits were filtered onto Nucleopore filters (0.45 pore size), removed from the filter as a wet cake of material, oven-dried at 60 °C and ground in a mortar. This material was used to determine the dry mass flux and the major components of the flux (particulate inorganic carbon (PIC), POC and biogenic silica). PIC was determined by closed system acidification with phosphoric acid and coulometry. Particulate total carbon (PC) was determined by unacidified combustion using a carbon-hydrogen-nitrogen (CHN) elemental analyzer. POC was calculated from PC by subtraction of PIC. Total silicon and aluminium contents were estimated by HF–HNO₃ microwave digestion and inductively coupled plasma emissions spectrometry following the methodology described by Bray et al. (2000). Biogenic silica was determined from total silica by subtracting lithogenic silica estimated by assuming a lithogenic Al:Si mass ratio of 3.42 (Taylor, 1964). These methods for PIC and POC/particulate organic nitrogen (PON) were used for all subsequent years, with very slight modifications: (i) the wet cake method was replaced by drying prior to removing the material from the filter, (ii) in some years sieving and filtering was done at sea and the samples were frozen on the filters until dried upon returning to land. The silica methods varied more strongly over time: (i) for deployments beginning in 1998, 1999 and 2000, the use of HF in the digestion was replaced by high temperature combustion with lithium borate in a graphite crucible and HNO₃ digestion to determine total silicon and aluminium; (ii) biogenic silica for these years (and retroactively for 1997) was calculated using the updated estimate for the lithogenic
drogen peroxide and concentrated hydrochloric acid follow-

in 30 mL were treated with potassium permanganate, hy-

dygen carbonate (pH 8) and stored at 4°C in the dark

water to 40 mL, from which 10 mL were subsampled and

A total of 138 samples were processed for siliceous mi-

Al: Si mass ratio of 3.83 (Taylor and McLennan, 1985); (iii)

from 2001 onwards, total silica was not measured, instead

hot alkali digestion and colorimetry was used to estimate

biogenic silica directly (following the method of Quégüiner, 

3.4 Siliceous microplankton sample preparation

A total of 138 samples were processed for siliceous mi-

croplankton analysis. Each split was refilled with distilled

water to 40 mL, from which 10 mL were subsampled and

buffered with a solution of sodium carbonate and sodium

hydrogen carbonate (pH 8) and stored at 4°C in the dark

for future calcareous nannoplankton analysis. The remain-

ing 30 mL were treated with potassium permanganate, hy-

drogen peroxide and concentrated hydrochloric acid follow-

ing the methodology used by Romero et al. (1999). Three

slides per sample were prepared and mounted using the stan-

dard decantation method outlined by Bárcena and Abrantes 

(1998). This method produces random settling of the diatom

valves for quantitative microscopic purposes. Siliceous mi-

croplankton analysis was carried out on permanent slides 

(Norland optical adhesive 61 mounting medium; refractive 

index: 1.56) of acid-cleaned material. Qualitative and quan-

titative analysis were done at x1000 and x4000 magnifications 

using an Olympus BH-2 compound light optical microscope 

with phase-contrast illumination. In order to properly charac-

terise the diatom assemblages, a target of 400 diatom valves

was counted per sample. Owing to the strong seasonality 

diatom production, some cups collected very low numbers 

of diatom valves. For these samples a compromise between 

to number to be counted and time spent had to be reached but

the number of valves counted was never less than 100 with 

the exception of cup no. 6 of year 2000–2001, and cup no. 14 

of year 1999–2000 at the 47°S site which were not consid-

Table 2. Continued.

| Deployment | Cup | Sampling period mid-point | Length days | Diatoms 10^6 valves m^-2 d^-1 | Total mass mg m^-2 d^-1 | BSO2 mg m^-2 d^-1 | CaCO3 % | POC mg m^-2 d^-1 | POC % |
|------------|-----|---------------------------|-------------|-----------------------------|------------------------|-------------------|--------|-----------------|-------|
| 54°S 800m, 2002–2003 | 1 | 04/10/2003 | 14.0 | 0.352 | 15.1 | 8.1 | 54 | 4.5 | 30 | 0.5 | 3 |
| | 2 | 18/10/2003 | 14.0 | 1.143 | 23.4 | 12.6 | 54 | 7.0 | 30 | 0.8 | 3 |
| | 3 | 01/11/2003 | 14.0 | 2.888 | 34.8 | 18.6 | 54 | 10.8 | 31 | 1.1 | 3 |
| | 4 | 15/11/2003 | 14.0 | 11.519 | 121.0 | 54.6 | 45 | 19.9 | 61 | 2.0 | 2 |
| | 5 | 29/11/2003 | 14.0 | 7.748 | 75.5 | 35.9 | 48 | 14.6 | 69 | 1.6 | 2 |
| | 6 | 13/12/2003 | 14.0 | 19.892 | 178.6 | 111.6 | 62 | 40.2 | 35 | 3.6 | 2 |
| | 7 | 27/12/2003 | 14.0 | 10.320 | 232.6 | 134.8 | 58 | 43.6 | 19 | 7.2 | 3 |
| | 8 | 10/01/2004 | 14.0 | 30.998 | 182.0 | 114.5 | 63 | 33.7 | 19 | 6.2 | 3 |
| | 9 | 24/01/2004 | 14.0 | 16.786 | 121.9 | 81.7 | 67 | 21.0 | 17 | 3.3 | 3 |
| | 10 | 07/02/2004 | 14.0 | 1.142 | 63.9 | 37.0 | 58 | 10.8 | 17 | 2.4 | 2 |
| | 11 | 21/02/2004 | 14.0 | 5.982 | 28.5 | 16.6 | 58 | 7.0 | 24 | 1.0 | 4 |
| | 12 | 06/03/2004 | 14.0 | 6.189 | 31.0 | 18.4 | 59 | 7.7 | 25 | 0.9 | 3 |
| | 13 | 20/03/2004 | 14.0 | 9.824 | 66.2 | 45.3 | 68 | 12.8 | 54 | 0.9 | 4 |
| | 14 | 03/04/2004 | 14.0 | 65.376 | 363.9 | 203.6 | 56 | 81.0 | 22 | 12.3 | 3 |
| | 15 | 17/04/2004 | 14.0 | 1.549 | 19.3 | 11.3 | 59 | 4.5 | 23 | 1.1 | 6 |
| | 16 | 01/05/2004 | 14.0 | 3.336 | 39.3 | 32.6 | 62 | 16.9 | 20 | 0.8 | 1 |
| | 17 | 25/05/2004 | 14.0 | 3.431 | 19.6 | 13.7 | 59 | 3.5 | 18 | 0.4 | 2 |
| | 18 | 29/06/2004 | 14.0 | 1.263 | 14.5 | 6.0 | 41 | 2.8 | 20 | 2.0 | 14 |
| | 19 | 03/08/2004 | 14.0 | 1.088 | 10.3 | 5.0 | 49 | 3.8 | 17 | 0.5 | 3 |
| | 20 | 07/09/2004 | 14.0 | 0.458 | 7.3 | 3.6 | 49 | 2.5 | 34 | 0.5 | 7 |

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Table 2. Continued.

| Deployment | Cup | Sampling period | Length | Diatoms \(10^6\) m\(^{-2}\) d\(^{-1}\) | Total mass | BSiO\(_2\) | CaCO\(_3\) | POC |
|------------|-----|----------------|-------|---------------------------------|------------|----------|--------|-----|
| 54° 800 m, 2005–2006 | 1 | 22/12/2005 | 17.0 | 24.184 | 197.9 | 120.0 | 61 | 43.5 | 22 | 4.6 | 2 |
| | 2 | 08/03/2006 | 17.0 | 19.228 | 193.6 | 115.2 | 59 | 47.5 | 25 | 3.2 | 2 |
| | 3 | 25/01/2006 | 17.0 | 47.505 | 418.1 | 275.3 | 66 | 65.1 | 16 | 12.2 | 3 |
| | 4 | 11/02/2006 | 17.0 | 11.571 | 178.0 | 60.8 | 34 | 92.1 | 52 | 3.9 | 2 |
| | 5 | 28/02/2006 | 17.0 | 4.840 | 116.7 | 31.7 | 27 | 66.9 | 57 | 2.4 | 2 |
| | 6 | 17/03/2006 | 17.0 | 1.180 | 44.9 | 7.9 | 17 | 30.1 | 67 | 1.5 | 3 |
| | 7 | 03/04/2006 | 17.0 | 8.853 | 110.6 | 46.0 | 42 | 50.2 | 45 | 2.6 | 2 |
| | 8 | 20/04/2006 | 17.0 | 11.216 | 142.8 | 86.5 | 61 | 33.0 | 23 | 3.1 | 2 |
| | 9 | 07/05/2006 | 17.0 | 9.477 | 107.0 | 63.5 | 59 | 23.3 | 22 | 2.6 | 2 |
| | 10 | 07/06/2006 | 45.0 | 0.904 | 18.6 | 7.6 | 41 | 8.0 | 43 | 0.6 | 3 |
| | 11 | 22/07/2006 | 45.0 | 0.072 | 3.6 | 1.6 | 45 | 1.2 | 33 | 0.4 | 10 |
| | 12 | 05/09/2006 | 45.0 | 0.778 | 9.1 | 4.6 | 50 | 2.8 | 31 | 0.3 | 3 |
| | 13 | 18/10/2006 | 16.0 | 0.291 | 6.4 | 1.9 | 29 | 1.0 | 15 | 0.4 | 7 |
| | 14 | 03/11/2006 | 16.0 | 1.065 | 23.9 | 6.9 | 29 | 3.4 | 14 | 1.2 | 5 |
| | 15 | 19/11/2006 | 16.0 | 4.461 | 77.6 | 25.1 | 32 | 13.6 | 18 | 3.4 | 4 |
| | 16 | 05/12/2006 | 16.0 | 23.929 | 131.5 | 71.5 | 54 | 28.7 | 22 | 4.5 | 3 |
| | 17 | 21/12/2006 | 16.0 | 72.099 | 333.9 | 220.9 | 66 | 30.8 | 9 | 10.5 | 3 |
| | 18 | 03/01/2006 | 16.0 | 100.622 | 335.5 | 243.5 | 73 | 18.4 | 5 | 9.5 | 3 |
| | 19 | 18/10/2006 | 16.0 | 0.012 | 0.6 | 0.4 | 72 | 0.1 | 11 | 0.0 | 2 |
| | 20 | 23/02/2006 | 16.0 | 0.016 | 0.6 | 0.4 | 72 | 0.1 | 11 | 0.0 | 2 |
| | 21 | 11/03/2006 | 16.0 | 0.021 | 0.6 | 0.4 | 72 | 0.1 | 11 | 0.0 | 2 |
| | 22 | 27/03/2007 | 16.0 | 7.500 | 33.0 | 22.8 | 69 | 3.8 | 12 | 1.1 | 3 |
| | 23 | 12/04/2006 | 16.0 | 0.030 | 1.8 | 1.2 | 69 | 0.8 | 43 | 0.1 | 4 |
| | 24 | 28/04/2006 | 16.0 | 0.106 | 6.2 | 4.3 | 69 | 4.1 | 67 | 0.1 | 2 |
| | 25 | 14/05/2006 | 16.0 | 0.010 | 0.7 | 0.5 | 69 | 0.3 | 51 | 0.0 | 3 |
| | 26 | 30/05/2006 | 16.0 | 0.013 | 0.5 | 0.4 | 69 | 0.3 | 51 | 0.0 | 3 |
| | 27 | 18/06/2006 | 16.0 | 0.420 | 9.0 | 6.2 | 69 | 3.9 | 43 | 0.3 | 3 |
| | 28 | 11/07/2006 | 16.0 | 0.005 | 0.3 | 0.2 | 69 | 0.3 | 43 | 0.0 | 3 |
| | 29 | 03/08/2006 | 16.0 | 0.001 | 0.2 | 0.2 | 69 | 0.1 | 43 | 0.0 | 3 |
| | 30 | 23/08/2006 | 16.0 | 0.001 | 0.1 | 0.1 | 69 | 0.0 | 43 | 0.0 | 3 |
| | 31 | 08/09/2006 | 16.0 | 0.140 | 4.2 | 2.9 | 69 | 1.9 | 45 | 0.1 | 3 |
| | 32 | 24/09/2006 | 16.0 | 0.087 | 3.1 | 2.2 | 69 | 1.3 | 42 | 0.2 | 6 |

* Component fluxes representing intervals for which insufficient material was available for component measurement and were estimated.

3.5 Taxonomic identifications

All diatom and silicoflagellate specimens were identified to the lowest taxonomic level possible. Radiolarians were only identified to group level. Scanning electron microscope imagery was used on selected samples to verify taxonomic identifications made with the light microscope. Taxonomy followed modern concepts in Hasle and Syvertsen (1997). The resting spores of members of the subgenus *Hyalochaete* of the genus *Chaetoceros* were identified only at group level due to a lack of morphological criteria. The differentiation between *Pseudo-nitzschia lineola* and *Pseudo-nitzschia turgiduloides* was often difficult due to their state of preservation in the samples; therefore, they were grouped under the category *Pseudo-nitzschia cf. lineola* in this study. A species or group of species of the genus *Thalassiosira* larger than 20 μm, highly dissolved and with radial to fasciculated areolation were grouped together under the name *Thalassiosira* sp. 1. Several small *Thalassiosira* species with similar morphological features were assembled together under *Thalassiosira trifulta* group following Shiono and Koizumi (2000). Due to the gradational nature of the morphology between *Thalassiosira gracilis* var. *gracilis* and *T. gracilis* var. *expecta*, both varieties were grouped together under the name *T. gracilis* group following the recommendations of Crosta et al. (2005).

3.6 Statistical analysis

In order to enable comparison with other sites, annual flux estimates are provided in Table 3. These were obtained by assuming that total mass flux outside of the sampling period was constant and by linearly interpolating values for the small gaps (i.e. 8.5–17 days intervals) during the productive season. No attempt was made to annualise the relative contribution of the diatom taxa, and therefore average values of the integrated diatom assemblage for whole sampling interval are provided in Table 3.

In order to investigate the covariability between the main diatom taxa along our sediment trap records, we conducted separate principal component analyses (PCA) for each site using of Statistica 7.0® software. PCA analysis is a statis-
Table 3. Estimated annual export fluxes of total mass flux, biogenic silica, calcium carbonate, POC and diatom valves for < 1 mm fraction at the 47, 54 and 61° S sites.

(a) Annual fluxes of biogeochemical components (g m\(^{-2}\) yr\(^{-1}\)) and diatom valves (valves 10\(^8\) m\(^{-2}\) yr\(^{-1}\)).

| Trap | Year           | Total mass flux | SiO\(_2\)-biogenic PIC as CaCO\(_3\) | POC | Diatom valves (x10\(^8\)) |
|------|----------------|----------------|--------------------------------------|-----|--------------------------|
| 47_1000 | 1999–2000   | 12             | 1                                    | 8   | 1.1                      | 0.3                 |
|       | 2000–2001   | 15             | 1                                    | 12  | 0.9                      | 0.8                 |
|       | Average ± SD| 14 ± 2         | 1 ± 0                                | 10 ± 3 | 1.0 ± 0.1         | 0.5 ± 0.4            |
| 54_800 | 1997–1998   | 19             | 10                                   | 4   | 0.6                      | 30.2                |
|       | 1999–2000   | 52             | 31                                   | 9   | 1.4                      | 29.1                |
|       | 2002–2003   | 30             | 16                                   | 8   | 1.1                      | 39.0                |
|       | 2003–2004   | 20             | 12                                   | 4   | 0.6                      | 22.5                |
|       | 2005–2006   | 29             | 15                                   | 9   | 0.7                      | 30.8                |
|       | 2006–2007   | 16             | 10                                   | 2   | 0.5                      | 34.5                |
|       | Average ± SD| 24 ± 13        | 12 ± 9                               | 7 ± 3 | 0.8 ± 0.4         | 31.0 ± 5.5            |
| 61_2000 | 2001–2002   | 85             | 65                                   | 6   | 1.2                      | 242.9               |

(b) Proportion of biogeochemical components (wt %).

| Trap | Year           | SiO\(_2\)-biogenic PIC as CaCO\(_3\) | POC |
|------|----------------|--------------------------------------|-----|
| 47_1000 | 1999–2000   | 8                                    | 65  | 9.0                 |
|       | 2000–2001   | 7                                    | 76  | 5.6                 |
|       | Average ± SD| 7 ± 1                                | 70 ± 8 | 7.3 ± 2.4       |
| 54_800 | 1997–1998   | 53                                   | 23  | 2.9                 |
|       | 1999–2000   | 60                                   | 17  | 2.7                 |
|       | 2002–2003   | 52                                   | 25  | 3.5                 |
|       | 2003–2004   | 59                                   | 20  | 3.1                 |
|       | 2005–2006   | 53                                   | 30  | 2.5                 |
|       | 2006–2007   | 63                                   | 12  | 3.3                 |
|       | Average ± SD| 57 ± 4                               | 21 ± 7 | 3.0 ± 0.4       |
| 61_2000 | 2001–2002   | 76                                   | 7   | 1.4                 |

(c) Annual fluxes of biogeochemical elements, mmol m\(^{-2}\) yr\(^{-1}\) and mole ratios.

| Trap | Year           | BSi     | PIC     | POC     | BSi : PIC | POC : BSi |
|------|----------------|---------|---------|---------|-----------|-----------|
| 47_1000 | 1999–2000   | 15      | 76      | 89      | 0.2       | 5.7       |
|       | 2000–2001   | 17      | 116     | 72      | 0.1       | 4.2       |
|       | Average ± SD| 16 ± 1  | 96 ± 28 | 80 ± 12 | 0.2 ± 0.0 | 4.9 ± 1.1 |
| 54_800 | 1997–1998   | 168     | 43      | 46      | 3.9       | 0.3       |
|       | 1999–2000   | 513     | 87      | 117     | 5.9       | 0.2       |
|       | 2002–2003   | 264     | 77      | 89      | 3.4       | 0.3       |
|       | 2003–2004   | 192     | 40      | 50      | 4.8       | 0.3       |
|       | 2005–2006   | 252     | 87      | 59      | 2.9       | 0.2       |
|       | 2006–2007   | 168     | 19      | 43      | 8.9       | 0.3       |
|       | Average ± SD| 259 ± 131 | 59 ± 29 | 67 ± 29 | 5.0 ± 2.2 | 0.3 ± 0.0 |
| 61_2000 | 2001–2002   | 1081    | 63      | 102     | 17.3      | 0.1       |
tical technique that reduces the information brought by a high number of independent variables into a smaller set of dimensions (factors) with a minimum loss of information. Only species and taxonomic groups with relative contributions > 1 % for the entire sampling period were considered in the analysis, i.e. thirteen taxa from site 47° S and nine taxa from site 54° S. The relative contribution of these groups of species was recalculated for each sample and then a log transformation (log x +1) was applied in order to normalise the distribution of the data. Diatom groups were then determined using a Q-mode factor analysis of the samples with a maximised variance (VARIMAX) rotation.

The Shannon’s diversity index (Shannon, 1949) was used to document latitudinal diversity trends across sites (Table 4).

3.7 Environmental variables

Weekly SSTs for the decade 1997–2007 were derived from the IGOSS NMC (the Integrated Global Ocean Services System Products Bulletin, National Meteorological Center; Reynolds et al., 2002) database, each value is a weekly composite of data collected within the area 48.5–45.5° S × 130–150° E for the 47° S site and 55.5–52.5° S × 130–150° E for the 54° S site (Figs. 2a and 3a). Sea-viewing Wide Field-of-View Sensor (SeaWiFS) satellite-derived chlorophyll a and photosynthetically available radiation (PAR) estimates were obtained from NASA’s Giovanni online data system (Acker and Leptoukh, 2007) for the same area used for the SST estimates (Figs. 2a and 3a).

Primary productivity values (mg C m⁻² d⁻¹) for all the sites were obtained from the Ocean Productivity website (www.science.oregonstate.edu/ocean.productivity/index.php), which provides estimates of net primary productivity derived from SeaWiFS satellite data by the standard vertically generalized production model (VGPM; Behrenfeld and Falkowski, 1997) and the carbon-based production model (CbPM; Behrenfeld et al., 2005).

4 Results

Here, we present the chemical (total mass, biogenic silica, carbonate and POC) and biological (diatom species) compositions of the particle fluxes registered at ~1000 m at the 47° S site during 2 years (July 1999–October 2001) and at 54° S site during 6 years (September 1997–February 1998, July 1999–August 2000, November 2002–October 2004 and December 2005–October 2007; Table 2). A description of the total particle flux and its chemical signature at stations 47 and 54° S for the first year deployment of the sediment traps (1997–1998) can be found in Trull et al. (2001).

4.1 Biogeochemical fluxes

The total mass and bulk component (biogenic silica, carbonate and POC) fluxes for both traps are shown in Figs. 2b and 3b and listed in Table 2. Annual total mass flux at ~1 km depth was the lowest at station 47° S (14±2 g m⁻² yr⁻¹; 2-year average ± standard deviation) and the highest at station 54° S (24±13 g m⁻² yr⁻¹; 6-year average ± standard deviation; Table 3). BSi flux followed a similar latitudinal trend with lower fluxes at 47° S (1±0 g m⁻² yr⁻¹) compared to 54° S (12±9 g m⁻² yr⁻¹). Carbonate export exhibited less variability between sites, with values somewhat higher at 47° S (10±3 g m⁻² yr⁻¹) than those measured at 54° S (7±3 g m⁻² yr⁻¹, respectively). Interestingly, despite the strong latitudinal differences in the magnitude of the mass fluxes, POC export was very similar at both stations (1.0±0.1 and 0.8±0.4 g m⁻² yr⁻¹, for 47 and 54° S, respectively).

In terms of relative abundance, the biogenic silica fraction represented 57 % of the mass flux at the 54° S site, whereas its contribution dramatically dropped to 7 % at the 47° S station. Calcium carbonate and POC accounted for 70 and 7.3 % at the 47° S site, respectively, and 21 and 3 % at the 54° S station (Table 3). These differences were primarily driven by the northward decrease in the biogenic silica fluxes. The BSI : PIC molar ratios decreased northward mirroring the latitudinal variations of the particle composition, from 5.0 at station 54° S to 0.2 at station 47° S (Table 3; Fig. S1 in the Supplement). The POC : BSi followed an opposite pattern with 0.3 at 54 and 4.9 at 47° S.

The seasonality of the total mass flux at station 47° S during the 2-year record showed a period of enhanced particle export in spring and secondary peaks in summer and autumn (Fig. 2b). The highest fluxes were registered in November–December 2000 (92–176 mg m⁻² d⁻¹), March 2001 (105 mg m⁻² d⁻¹) and October 1999 (90 mg m⁻² d⁻¹). Total mass flux at the 54° S site was strongly seasonal with maximum values occurring during the late spring–summer and very low export prevailing through the autumn and winter months. The late spring–summer export maxima were as short as 3 months and often showed a bimodal distribution (e.g. 1997–1998, 1999–2000, 2002–2003; Fig. 3b). The highest total mass fluxes at this site were collected during December–January 1999 (511–724 mg m⁻² d⁻¹), January 2006 (418 mg m⁻² d⁻¹), February 2003 (397 mg m⁻² d⁻¹) and January 1998 (396 mg m⁻² d⁻¹).

4.2 Diatom fluxes

The biogenic silica flux at the 47 and 54° S site was composed of diatoms, silicoflagellates, radiolarians and a handful of skeletons of the dinoflagellate Actiniscus pentasterias. Diatom fluxes were 1 order of magnitude higher than those of silicoflagellates and radiolarians at the 47° S site, and 1 and 3 orders of magnitude higher, respectively, at the 54° S site. Consistent with the biogenic silica flux, diatoms were most numerous in the 54° S site with an annual flux of 31±5.5×10⁸ valves m⁻² yr⁻¹ (6-year average ± standard deviation).
Table 4. List of diatom species recorded in the sediment traps of the 47, 54 and 61° S (2000 m) sites along the 140° E. Relative abundances < 0.1 are represented by an asterisk (*), whereas the absence of a taxon in a given site is represented by an empty circle (○).

| Species                              | SAZ (47° S) | PFZ (54° S) | AZ (61° S) |
|--------------------------------------|-------------|-------------|------------|
| Actinocyclus actinochilus (Ehrenberg) Simonsen | *           | ○           | *          |
| Actinocyclus curvatulus Janisch       | 0.3         | *           | ○          |
| Actinocyclus exigius Fryxell and Semina | *           | ○           | ○          |
| Actinocyclus octonarius Ehrenberg     | *           | ○           | ○          |
| Actinocyclus spp.                     | 0.6         | *           | *          |
| Alveus marinus (Grunow) Kaczmarska and Fryxell | *           | ○           | ○          |
| Asteromphalus hookeri Ehrenberg       | *           | 0.5         | 0.2        |
| A. hyalinus Karsten                   | *           | 0.2         | 0.2        |
| A. parvulus Karsten                   | *           | 0.1         | 0.2        |
| Asteromphalus spp.                    | ○           | ○           | ○          |
| Azpeitia tabularis (Grunow) Fryxell and Sims | 10.8       | 0.8         | 0.7        |
| Chaetoceros aequatorialis var. antarcticus Manguin | ○           | 0.1         | *          |
| Ch. atlanticus Cleve                  | 0.1         | 0.5         | 0.2        |
| Ch. dichaeta Ehrenberg                | ○           | 0.6         | 0.1        |
| Ch. peruvianus Brightwell             | 0.1         | *           | ○          |
| Chaetoceros subgenus Hyalochaete spp. | 0.2         | 0.4         | 0.2        |
| Chaetoceros subgenus Phaeoceros spp.  | ○           | 0.1         | 0.2        |
| Chaetoceros resting spores            | 2.3         | 0.5         | 0.1        |
| Cocconeis spp.                        | 0.2         | ○           | ○          |
| Corethron sp.                         | *           | *           | *          |
| Cyclotella spp.                       | *           | ○           | ○          |
| Dactyliosolen antarcticus Castracane   |             |             |            |
| Diploneis bombus (Ehrenberg) Ehrenberg | *           | ○           | ○          |
| Eucamnia antarctica (Castracanee) Mangin (summer form) | ○           | 0.3         | *          |
| E. antarctica (Castracane) Mangin (winter form) | *           | *           | 0.1        |
| Fragilaropsis curta (Van Heurck) Hustedt | ○           | *           | 0.6        |
| F. cylindrus (Grunow) Krieger         | ○           | *           | 0.2        |
| F. doliolus (Wallich) Medlin and Sims | 0.5         | ○           | ○          |
| F. kerguelensis (O’Meara) Hustedt     | 43          | 59.3        | 79.9       |
| F. obliquecostata (van Heurck) Heiden  | ○           | *           | *          |
| F. pseudonana (Hasle) Hasle           | *           | 3.1         | 2          |
| F. rhombica (O’Meara) Hustedt         | 0.4         | 2.3         | 0.9        |
| F. ritscheri Hustedt                  | 0.4         | 0.1         | 0.1        |
| F. separanda Hustedt                  | ○           | 0.1         | 2.1        |
| F. cf. sublineata (Van Heurck) Heiden  | ○           | ○           | *          |
| Fragilariosis spp.                    | ○           | *           | ○          |
| Gyrosigma spp.                        | ○           | *           | ○          |
| Haslea trompii (Cleve) Simonsen       | ○           | 0.1         | *          |
| Hemidiscus cuneiformis Wallich        | 3.7         | ○           | ○          |
| Navicula directa (Smith) Ralfs in Pritchard | ○          | 4.6         | 0.3        |
| Nitzschia bicapitata Cleve            | 2.8         | *           | ○          |
| N. braarudii (Hasle)                  | 0.2         | ○           | ○          |
| N. kolaczekii Grunow                  | 0.6         | *           | ○          |
| N. sicula (Castracane) Hustedt var. bicuneata Grunow | 2           | *           | 0.1        |
| N. sicula (Castracane) Hustedt var. rostrata Hustedt | *           | *           | ○          |
| Nitzschia spp.                        | *           | *           | ○          |
| Paralia spp.                          | *           | ○           | ○          |
| Pleurosigma spp.                      | 0.2         | 0.1         | *          |
| Porosira pseudodenticulata (Hustedt) Jousé | 0.1         | *           | *          |
| Proboscia alata (Brightwell) Sundström | 0.1         | *           | ○          |
| P. inermis (Castracane) Jordan Ligowski | ○           | *           | ○          |
| Proboscia spp.                        | *           | *           | ○          |
deviation) compared to $0.5 \pm 0.4 \times 10^8$ valves m$^{-2}$ yr$^{-1}$ (2-year average ± standard deviation) of the 47° S site.

Total diatom-valve flux at the 47° S site (Fig. 4a) showed a less pronounced seasonality than that observed at 54° S (Fig. 5a) and exhibited a weak correlation with the total mass ($r = 0.37$, $n = 30$) and BSi ($r = 0.42$, $n = 29$) fluxes. Diatoms occurred in the greatest numbers during November 2000 ($1.6 \times 10^6$ valves m$^{-2}$ d$^{-1}$), February–

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**Table 4. Continued.**

| Species | SAZ (47° S) | PFZ (54° S) | AZ (61° S) |
|---------|-------------|-------------|------------|
| *Psammodiction panduriforme* (Gregory) Mann | 0.1 | ☐ | ☐ |
| *Pseudo-nitzschia* cf. *lineola* | * | 8.1 | 0.4 |
| *P. heimii* Manguin | * | 4.6 | * |
| *Pseudo-nitzschia* spp. | * | ☐ | 0.1 |
| *Rhizosolenia antennata* (Ehrenberg) Brown f. *antennata* | * | * | ☐ |
| *R. antennata* (Ehrenberg) Brown f. *semispina* Sundström | 0.1 | * | ☐ |
| *R. bergonii* Peragallo | 1.8 | * | ☐ |
| *Rhizosolenia* cf. *costata* | ☐ | * | ☐ |
| *R. curvata* Zacharias | ☐ | * | ☐ |
| *R. polydactyla* Castracane f. *polydactyla* | * | * | ☐ |
| *Rhizosolenia* sp. f. 1A (Armand et Zielinski) | ☐ | * | * |
| *Rhizosolenia* spp. | 0.2 | * | 0.1 |
| *Roperia tesselata* (Roper) Grunow | 3.2 | * | ☐ |
| *Stellarima stellaris* (Roper) Hasle et Sims | 0.5 | * | ☐ |
| *Thalassionema nitzschioides* var. *capitulata* (Castracane) Moreno-Ruiz | * | 0.2 | 0.1 |
| *T. nitzschioides* var. *lanceolata* (Grunow) Pergallo et Pergallo | 0.2 | 0.9 | 0.1 |
| *T. nitzschioides* var. *parvum* (Zielinski et Gersonde) | 0.2 | 0.4 | ☐ |
| *Thalassiosira eccentrica* (Ehrenberg) Cleve | 0.9 | 0.1 | 0.2 |
| *T. ferelineata* Hasle et Fryxell | 0.3 | ☐ | ☐ |
| *T. gracilis* var. *expecta* (Van Landingham) Fryxell et Hasle | 0.1 | 0.7 | 0.4 |
| *T. gracilis* var. *gracilis* (Karsten) Hustedt | 0.3 | 3.9 | 3.6 |
| *T. gracilis* group | 0.3 | 4.6 | 4.1 |
| *T. gravida* Cleve | ☐ | * | ☐ |
| *T. lentiginosa* (Janisch) Fryxell | 2.1 | 2.1 | 5 |
| *T. leptopus* (Grunow ex Van Heurck) Hasle et G.Fryxell | ☐ | ☐ | * |
| *T. lineata* Jousé | 2.2 | ☐ | ☐ |
| *T. maculata* Fryxell et Johans. | 0.2 | 0.1 | * |
| *T. oestrupii* (Ostenfeld) Hasle var. *oestrupii* Fryxell and Hasle | 3.6 | 0.3 | * |
| *T. oestrupii* (Ostenfeld) Hasle var. *venrickiae* Fryxell and Hasle | 0.6 | 0.1 | ☐ |
| *T. oliveriana* (O' Meara) Makarova et Nikolaev | 0.1 | 0.6 | 0.7 |
| *T. symmetrica* Fryxell et Hasle | 0.3 | * | ☐ |
| *T. trifulta* Fryxell | 0.1 | ☐ | ☐ |
| *T. tumida* (Janisch) Hasle | 0.2 | 0.3 | 0.1 |
| *Thalassiosira* sp. 1 | 5.4 | 0.1 | * |
| *Thalassiosira* sp. 2 | 0.2 | 0.1 | ☐ |
| *Thalassiosira* sp. 3 | ☐ | 0.2 | ☐ |
| *Thalassiosira* eccentric group | 0.1 | ☐ | ☐ |
| *T. linear* group | 0.2 | * | 0.1 |
| *T. trifulta* group | 1.9 | 0.1 | ☐ |
| *Thalassiosira* spp. < 20μm | 0.9 | 1.4 | 0.4 |
| *Thalassiosira* spp. > 20μm | 1.7 | 0.1 | * |
| *Thalassiothrix* antarctica Schimper ex Karsten | 2.2 | 0.4 | 0.2 |
| *Trachyneis aspera* (Ehrenberg) Cleve | ☐ | ☐ | ☐ |
| *Trichotoxon reinboldii* (Van Heurck) Reid et Round | ☐ | * | ☐ |
| *Tropidoneis* group | * | 0.4 | * |
| Other centrics | 1.2 | 0.1 | * |
| Other pennates | 0.2 | * | 0.1 |
| Shannon’s diversity index | 2.48 | 1.86 | 1.04 |
March 2001 (0.4–0.8 \times 10^6 \text{ valves m}^{-2} \text{ d}^{-1}) and October 1999 (0.4 \times 10^6 \text{ valves m}^{-2} \text{ d}^{-1}).

At station 54° S, total diatom-valve flux was highly seasonal and followed a similar pattern to that of the total mass ($r = 0.66$, $n = 108$) and BSi fluxes ($r = 0.68$, $n = 108$). These correlations are high despite the biases associated with our diatom-valve counting technique which does not allow for quantification of small valve fragments. In particular the high diatom-valve fragmentation observed during the productive period of 1999–2000 reduced the correlations between diatom-valve flux and total mass and BSi fluxes. In fact, the latter correlations increased significantly after excluding the 1999–2000 data ($r = 0.85$, $n = 88$ and $r = 0.87$, $n = 88$, respectively).

The spring–summer diatom bloom often exhibited two peaks of enhanced export separated by a period of lower flux (e.g. 1997–1998, 1999–2000, 2002–2003; Figs. 5a and 7). During the productive period of 2006–2007, the diatom bloom exhibited one single peak during which the largest diatom fluxes of the record were registered (up to $100 \times 10^6 \text{ valves m}^{-2} \text{ d}^{-1}$ in January 2007; Fig. 5a). Secondary diatom flux maxima were registered in January 1998 ($71 \times 10^6 \text{ valves m}^{-2} \text{ d}^{-1}$), December 2002 ($65 \times 10^6 \text{ valves m}^{-2} \text{ d}^{-1}$) and December 1999 ($52 \times 10^6 \text{ valves m}^{-2} \text{ d}^{-1}$). We noticed that during the 1999–2000, summer bloom the high BSi fluxes were not coupled with a proportional increase of the diatom valves (Fig. 5a). The higher degree of fragmentation observed on these samples could be attributed to either a more intense grazing pressure by the zooplankton community that year or by a higher fragmentation of the valves during the sample preparation due to the presence of abundant numbers of weakly silicified diatoms (e.g. species of the genus \textit{Pseudo-nitzschia}) which are more prone to break during the sample processing (Rembauville et al., 2015).

In terms of diatom assemblage composition, the occurrence and fractional contributions of all the diatom taxa found at the 47 and 54° S study sites, as well as at 61° S (Rigual-Hernández et al., 2015), are provided in Table 4. The diatom sinking assemblage at station 47° S was more.
Figure 5. Temporal variability of (a) total diatom flux and biogenic silica and (b) flux and relative contribution of the main diatom species at 800 m at the 54° S site for the periods July 1999–August 2000, November 2002–October 2004 and December 2005–October 2007.
Table 5. Varimax loadings matrix (rotation: varimax normalised). Factor loadings for analysis on sediment trap samples using species and group of species with abundances higher than 1 % of the total integrated diatom assemblage for the whole sampling period at the 47° S (a) and 54° S site (b). Bold print indicates the dominant species or group of species within each factor (loadings > 0.70).

(a) 47° S site.

| Diatom species or groups of species | Factor 1 | Factor 2 | Factor 3 | Factor 4 |
|------------------------------------|----------|----------|----------|----------|
| Fragilariopsis kerguelensis        | 0.05     | -0.73    | -0.46    | -0.35    |
| Azpeitia tabularis                 | 0.59     | 0.38     | 0.06     | 0.55     |
| Thalassiosira sp. 1                | -0.11    | 0.91     | -0.17    | -0.21    |
| Nitzschia bicapitata               | -0.82    | 0.41     | 0.09     | 0.19     |
| Chaetoceros resting spores         | -0.59    | 0.12     | 0.56     | 0.08     |
| Thalassiosira oestrupii var. oestrupii | -0.06   | -0.75   | 0.20     | 0.13     |
| Hemidiscus cuneiformis             | 0.63     | 0.47     | 0.03     | -0.09    |
| Roperia tesselata                  | 0.04     | -0.29    | 0.73     | 0.11     |
| Thalassiothrix antarctica          | 0.06     | -0.05    | 0.05     | 0.61     |
| Nitzschia sicula var. bicuneata    | -0.64    | -0.09    | -0.03    |          |
| Thalassiosira lineata              | 0.15     | 0.02     | 0.78     | -0.39    |
| Rhizosolenia bergonii              | 0.08     | 0.14     | 0.17     | -0.57    |
| Thalassiosira lentiginosa           | 0.47     | -0.16    | 0.15     | 0.29     |
| Thalassiosira trifulta group       | -0.53    | -0.04    | -0.06    | -0.57    |
| Variance (%)                       | 19       | 19       | 13       | 13       |
| Cumulative variance                | 19       | 38       | 51       | 64       |

(b) 54° S site.

| Diatom species or groups of species | Factor 1 | Factor 2 | Factor 3 |
|------------------------------------|----------|----------|----------|
| Fragilariopsis kerguelensis        | -0.91    | -0.20    | -0.11    |
| Navicula directa                   | 0.84     | -0.14    | 0.17     |
| Pseudo-nitzschia heimii            | 0.35     | 0.70     | 0.36     |
| Pseudo-nitzschia cf. lineola       | 0.77     | 0.25     | 0.41     |
| Thalassiosira gracilis group       | 0.81     | -0.19    | 0.19     |
| Fragilariopsis pseudonana          | 0.79     | 0.09     | -0.08    |
| Fragilariopsis rhombica            | 0.85     | 0.29     | 0.14     |
| Thalassiosira lentiginosa           | 0.09     | -0.89    | 0.00     |
| Chaetoceros group                  | 0.12     | 0.12     | 0.93     |
| Variance (%)                       | 48       | 17       | 14       |
| Cumulative total variance (%)      | 48       | 65       | 79       |

Bold values highlight the taxa that define each factor.

diversified ($H'$ for the entire sampling period = 2.48) than those found south the SAF ($H' = 1.86$ at the 54° S; $H' = 1.04$ at the 61° S) consisting of 79 species or groups of species. The most abundant species was *Fragilariopsis kerguelensis*, which represented 43 % of the integrated assemblage for the entire sampling period (Fig. 4). Subordinate contributions to the diatom assemblage were made by *Azpeitia tabularis* (10 %), *Thalassiosira* sp. 1 (4 %), *Nitzschia bicapitata* (4 %), resting spores of *Chaetoceros* spp. (subgenus *Hyalochaetae*: 3 %), *Thalassiosira oestrupii* var. *oestrupii* (3 %), *Hemidiscus cuneiformis* (3 %) and *Roperia tesselata* (3 %; Fig. 4). A total of 77 taxa were identified at the 54° S site (Table 4). *F. kerguelensis* was also the dominant species, contributing up to 59 % of the diatom assemblage for the whole sampling period (Fig. 5). Secondary contributors correspond to *Pseudo-nitzschia* cf. *lineola* (8 %), *Pseudo-nitzschia heimii* (5 %), *Thalassiosira gracilis* group (4 %), *Fragilariopsis pseudonana* (3 %), *Fragilariopsis rhombica* (2 %) and *Thalassiosira lentiginosa* (2 %; Fig. 5).

4.3 Principal component analysis of diatom assemblages

The PCA for the 47° S site identified four components containing 64 % of the total variance, whereas that of the 54° S site required three components to describe 79 % of the information of diatom data (Table 5). Figure 6 shows the position of the species on the first two PCA axes for the 47 and 54° S sites. Together with the species, we plotted total and major components mass fluxes. The first component of the PCA for the 47° S site accounted for 19 % of the variance. The centric species *A. tabu-
ulate and H. cuneiformis (Fig. 6) had a positive loading on factor 1 and exhibited their highest relative abundance during spring and summer (Fig. 4). Factor 2 explained 19% of the variance and was dominated by F. kerguelensis, T. oestrupii var. oestrupii and Thalassiosira sp. 1. F. kerguelensis maintained a relatively constant contribution to the diatom assemblages during the whole sampling interval with a tendency to peak in late summer and autumn together with T. oestrupii var. oestrupii. None of the factors of the PCA of the 47° S site were significantly correlated with the biogenic particle fluxes (Fig. 6a and Table 6a).

At the 54° S site, the first component (48% of the total variance) was highly correlated with the bulk components of the flux (Fig. 6b and Table 6b) and individualises two groups of diatom species. High-positive factor loadings characterise the bloom-forming *Pseudo-nitzschia* cf. *lineola*, *F. rhombica*, *F. pseudonana* and *N. directa* and the cool-open-ocean diatom *T. gracilis* group. The relative contribution of these species peaked during the productive season (Fig. 5) and showed a strong positive correlation with all the components of the flux (Fig. 6b and Table 6b). Therefore, diatom species characterized by a high-positive first factor loading can be defined as the “high-export group”. In contrast, a high-negative factor loading on the first PCA axis was attributed to *F. kerguelensis*, which peaked during winter and autumn, coinciding with very low particle fluxes. *Pseudo-nitzschia heimii* was the only species with a high positive factor loading on the second PCA axis (Fig. 6b and Table 5) and its relative abundance peaked mainly from mid-summer to autumn. With the exception of year 2002-2003, a consistent diatom species succession was consistently observed over the growth season at the 54° S site (Figs. 5b and 7). During those years with a double peak diatom sedimentation bloom, the first maximum (November to early December) was always dominated by *F. kerguelensis* and by other large and heavily silicified diatoms, such as *T. lentiginosa*. During the second peak (in late December to early February), the relative contribution of *Pseudo-nitzschia* cf. *lineola* and small *Fragilariaopsis* species increased sharply, representing together up to 50% of the diatom assemblage in January 2000 (Figs. 5b and 7). Even during year 2006–2007, when the diatom sedimentation bloom exhibited a single maximum, a similar succession can be discerned within the peak.

5 Discussion

5.1 Latitudinal trend of biogenic particle fluxes to the ocean interior

The contrasting latitudinal variations in the composition and magnitude of the particle fluxes along the 140°E transect reflect the physicochemical and biological characteristics of the different zonal systems sampled by the traps. Relatively low BSi and diatom export measured in the mesopelagic wa-

| Table 6. Correlation coefficients between the PCA factors and the fluxes of bulk components (total mass, carbonate, biogenic silica and POC) for the 47° S (a) and 54° S (b) sites. |
|----------------------------------|------------------|------------------|------------------|------------------|
|                                 | Factor 1 | Factor 2 | Factor 3 | Factor 4 |
| (a) 47° S site.                 |          |          |          |          |
| Total mass                      | -0.10    | -0.14    | 0.01     | -0.19   |
| Biogenic silica                 | -0.09    | -0.10    | 0.12     | -0.11   |
| Carbonate                       | -0.07    | -0.18    | -0.03    | -0.20   |
| POC                             | -0.28    | 0.06     | 0.14     | -0.12   |
| (b) 54° S site.                 |          |          |          |          |
| Total mass                      | 0.73     | 0.07     | 0.03     |          |
| Biogenic silica                 | 0.71     | 0.08     | -0.01    |          |
| Carbonate                       | 0.66     | 0.06     | 0.15     |          |
| POC                             | 0.75     | 0.12     | 0.06     |          |

Bold values are significant at \( p < 0.05 \).

ters of the SAZ (Fig. 8a and Table 3) are consistent with the low-to-moderate diatom biomass accumulation in the surface layer of this region (Kopczynska et al., 2001; de Salas et al., 2011). Low silicic acid (Bowie et al., 2011a) and iron levels (Sedwick et al., 2008; Bowie et al., 2009; Mongin et al., 2011), together with light limitation, as a result of cloudiness (Bishop and Rossow, 1991) and deep summer mixed layers (70–100 m; Rintoul and Trull, 2001), are considered the main factors responsible for the reduced diatom production in the SAZ. Moreover, the low BSi : PIC mole ratios measured by the traps (< < 1; Table 3) illustrate the relatively low contribution of diatoms to the particle flux export to the ocean interior. Low diatom export fluxes and BSi : PIC mole ratios are characteristic of carbonate-dominated and low-productivity regimes (Honjo et al., 2008) and typical of much of the circumpolar SAZ (Honjo et al., 2000; Trull et al., 2001a).

The higher diatom-valve fluxes and BSi export at the 54° S site (Table 3; Fig. 8a) agrees well with previous studies of the PFZ surface waters south of Tasmania, which reported relatively large and heavily silicified diatoms as major contributors to the phytoplankton biomass (Kopczynska et al., 2001; de Salas et al., 2011). Higher levels of silicic acid (Smith Jr et al., 2000), colder summer surface waters and shallower mixed winter layers than those of the SAZ (Rintoul and Trull, 2001) are most likely the main factors responsible for the greater prevalence of diatoms in this region. As a result of the enhanced diatom production and the drop in the abundance of calcifying phytoplankton (Findlay and Giraudau, 2000; Honjo et al., 2000), BSi : PIC mole ratios of the settling material at this site shift to > 1 (Table 3).

Further south, at station 61° S in the southern AZ, Rigual-Hernández et al. (2015) documented an annual diatom flux 1 order of magnitude greater than that measured at the 54° S site (243 × 10^8 valves m^-2 d^-1 at 2000 m; Table 3 and...
5.2 Latitudinal diatom species distribution

The species occurrence observed along the 140°E transect is consistent with previous reports on diatom assemblage composition in the surface waters (Kopczynska et al., 1986; Kopczynska et al., 2001; de Salas et al., 2011) and sediments (Armand et al., 2005; Crosta et al., 2005; Romero et al., 2005) of the Australian sector of the Southern Ocean, and provide evidence, once again, that the frontal systems represent natural physical boundaries for phytoplankton species distribution (Boyd, 2002).

Overall, the diatom assemblage registered at the 47°S site is typical of the SAZ and differs significantly from those found in the PFZ and AZ (Table 4). The SAZ represents a “buffer zone” between the subtropical gyres to the north and the polar waters to the south which results in a highly diverse diatom community as highlighted by the highest $H'$ (2.48; Table 4) of the study transect. The occurrence of the warm water taxa *H. cuneiformis*, *Fragilariopsis doliolus*, *Nitzschia kolaczeckii* and *T. lineata* (Romero et al., 2005; Venrick et al., 2008) is restricted to this station, and therefore, these species appear as good indicators for the southward migration of the warmer, saltier and nutrient-poor water masses of the SAZ into the ACC. Moreover, the stark increase in the abundance of the open-ocean diatoms *A. tabularis*, *N. bicapitata*, *R. tesselata* and *Thalassiosira oestrupii* north of the SAF suggest the preference of these species for warmer waters (Hasle and Syvertsen, 1997; Romero et al., 2005).

The sinking diatom assemblage registered at the 54°S site is characteristic of the ACC waters and largely defined by the dominance of *F. kerguelensis*. The relative abundance of *F. kerguelensis* at the PFZ (59%) represents a transitional value between that of the AZ (80%) and that of the SAZ (43%). This strong latitudinal gradient mirrors its distribution in the surface sediments, which has been previously tied to summer SST (Crosta et al., 2005; Esper et al., 2010). However, other potentially important influences such as mixed layer depth, seasonality, and iron and silicate abundance also exhibit latitudinal gradients, and therefore may also influence the distribution of this species. Peak abundances of *Pseudo-nitzschia* species along the 140°E transect are observed in the PFZ (Table 4) and are consistent with previous studies that described this genus as a major contributor to the bulk phytoplankton biomass in the ACC waters (e.g. Kopczynska et al., 2001; Smetacek et al., 2002; de Salas et al., 2011). Moreover, it is worth noting that *P-n. heimii*, together with
other large diatoms (e.g. Thalassiothrix and Proboscia), have been reported to be major contributors of a SCM consistently observed between 53 and 58° S along 140° E (Kopczynska et al., 2001; Parslow et al., 2001). Navicula directa also showed maximum abundances at the PFZ site with values ~5%. This species has been traditionally described as a benthic-dwelling species (Scott and Marchant, 2005 and references therein) with an affinity for sea ice conditions (Armand, 1997). However, its persistent presence throughout the 6-year record and similar seasonal flux pattern to that of other well-known open-ocean species of the ACC, such as Thalassiosira gracilis group (r = 0.8, n = 108; Fig. 5b), point to a pelagic distribution of this species. This concept agrees well with Kopczynska et al. (1986) and Waite and Nodder (2001), who documented Navicula populations of considerable abundance in areas remote from coastal and sea ice influence in the Australian sector.

Although in many aspects the composition of the diatom assemblage at the 61° S site was similar to that of station 54° S, there were some qualitative and quantitative differences. As a result of the southward increase in the relative abundance of F. kerguelensis, the diversity (H’) and the relative contribution of most of the secondary constituents of the diatom assemblage at 61° S exhibited lower values than at 54° S (Table 4). For example, Pseudo-nitzchia species that represented cumulatively 13% of the integrated assemblage, dropped to <1% at the 61° S site. Navicula directa followed a similar pattern with maximum abundances at 54° S (5%) and negligible fluxes at 61° S. It is possible, however, that other factors, such as selective grazing or ecological constraints, may also account for the lower contribution of these species in the AZ.

5.3 Seasonal variability of diatom assemblages

Taking into account that diatoms are, by far, the main contributors to the BSi production at the 54° S site, and that the BSi fraction, in turn, dominated the total mass flux, the strong correlation between diatom-valve and mass fluxes \( r = 0.85; n = 88 \) suggests that the particle export at the PFZ is mainly mediated by diatoms. In contrast, at the 47° S site, the silica-poor content of the particles and the low correlation between diatom-valve and mass fluxes \( r = 0.37; n = 30 \) indicates a minor role for diatoms in regulating the export in the SAZ. These results underscore the contrasting role that diatoms play in the controls on the flux north and south of the SAF (Trull et al., 2001a; Ebersbach et al., 2011).

The less defined seasonal pattern and lower amplitude of the diatom fluxes observed at the 47° S site (Fig. 4) are a reflection of the different algal community north of the
SAF, dominated by non-siliceous phytoplankton (Odate and Fukuchi, 1995; Kopczynska et al., 2001; de Salas et al., 2011). For both years of our study, the highest annual diatom export events coincided with the onset of the biomass accumulation in the surface waters, indicating that diatoms responded rapidly to the enhanced light levels (Fig. 2a) and to the formation of a stable and shallow mixed layer (Rintoul and Trull, 2001). However, unlike the chlorophyll a concentration that gradually increased throughout the spring, diatom export rapidly returned to winter values most likely caused by the depletion of the winter silicate and/or iron stocks (Lannuzel et al., 2011). This seasonal pattern is characteristic of the SAZ and other silicate-poor environments, where diatoms typically bloom at the beginning of the successional sequence and then are replaced by other functional groups (Margalef, 1978; Balch, 2004; Alvain et al., 2008; Rigual-Hernández et al., 2013). The increase in the diatom and BSi fluxes from January to early March 2001 suggests the export of a second diatom bloom that year. South of Tasmania the SAZ exhibits a complex physical structure with frequent wind mixing events (Yuan, 2004) and fronts meandering and forming eddies that can reach the trap location (Rintoul and Trull, 2001; Herreraiz-Borreguero and Rintoul, 2011). Thus, it is likely that one of these mechanisms injected nutrients into the surface layer of the 47° S site fuelling diatom production and allowing the “reset” of phytoplankton succession. In terms of population dynamics, the seasonal succession of species at the 47° S site was not as clearly expressed as in station 54° S and none of the diatom species seem to play an important role in the export controls of any of the components of the flux as indicated by the results of the PCA (Fig. 6a). *F. kerguelensis* exhibited fairly constant relative abundances throughout the record suggesting little competition for resources with other diatom species. The temperate-to-warm water species *H. cuneiformis* and *A. tabularis* showed their maximum contribution at times of maximum diatom export which suggests that these species are the first to respond to nutrient supply in the surface waters in this region.

The annual export maxima of total mass and diatom-valve flux at 54° S were separated into two peaks for most of the years (Figs. 5a and 7). A similar double peak feature of the particle bloom has been previously reported in the AZ of the Pacific (Honjo et al., 2000; Grigorov et al., 2014) and Atlantic (Fischer et al., 2002). Honjo (2004) speculated that such a double peak structure may be due to a break in primary production caused by a temporary depletion of a limiting nutrient, while Grigorov et al. (2014) attributed the drop in the diatom flux between two periods of enhanced export to a storm event that mixed the diatom biomass out of the surface layer. The lack of accompanying in situ measurements of nutrient concentration and mixed layer depth precludes the direct assessment of these possibilities.

The initial diatom population size, species-specific physiological traits and selective grazing pressure are crucial factors determining which diatom species dominates or co-dominates an individual bloom (Assmy et al., 2007; Assmy et al., 2013; Boyd, 2013). The chain-forming *F. kerguelensis* is one of the most abundant diatom species in ACC waters (e.g. Laubscher et al., 1993; Bathmann et al., 1997; Smetacek et al., 2002) and has been reported to represent up to 90% of the summer diatom populations in the AZ (Gall et al., 2001). The high relative contribution of *F. kerguelensis* throughout our record is consistent with these latter studies and suggests the presence of a large seeding population of this species before the onset of the bloom. These large initial seed stocks, together with the effective mechanical protection of its robust frustule (Hamm et al., 2003) against the heavy copepod grazing pressure of the ACC (Pollard et al., 2002; McLeod...
et al., 2010) are most likely the main factors determining the dominance of *F. kerguelensis* during the growth season. The increase in the relative abundance of the lightly silicified *Pseudo-nitzschia* cf. *lineola* and small *Fragilariopsis* species during the second part of the bloom (Figs. 5b and 7) is consistent with the observations of Kopczynska et al. (2001) who reported *F. pseudonana* and *P. n. lineola* dominating the diatom assemblages in the PFZ waters south of Tasmania during late summer. Assmy et al. (2007) reported large numbers of *P. n. lineola* during the last stages of the fertilisation experiment EisenEx, indicating the capacity of this diatom to outcompete other taxa under iron-limiting conditions. Moreover, small *Fragilariopsis* and *Pseudo-nitzschia* species are known to produce an iron-storage protein (ferritin) that allow them to undergo more cell divisions than other open-ocean diatoms under low iron concentrations (Marchetti et al., 2009). We speculate, that due to these particular physiological traits *Pseudo-nitzschia* and small *Fragilariopsis* species may gain a competitive advantage under the environmental conditions during the last stages of the diatom bloom (i.e. low silica and iron concentrations, and enhanced PAR), enabling such species to escape grazing and/or outcompete other diatoms. However, this scenario does not account for our observations in 2002–2003, when *Pseudo-nitzschia* and small *Fragilariopsis* species exhibited higher relative contribution in the first seasonal export peak (Figs. 5b and 7). This exceptional seasonal flux peak remained unexplained and likely due to other environmental conditions not captured by our study.

5.4 Ecological flux vectors in the PFZ

The short and vigorous summer particle export, consistently observed during our 6 year record at the 54°S trap is characteristic of high latitude systems (e.g. Honjo et al., 2000; Fischer et al., 2002; Pilskañ et al., 2004) and can contribute up to 66% of the annual POC export to 800 m in just 2 months (e.g. 1999–2000). Therefore, these large summer pulses of POC are responsible for a major proportion of the variability in carbon sequestration from the atmosphere in the PFZ. The mechanism is primarily through the increase in the overall flux, because the fractional POC content was not observed to increase during high flux periods. For example, % POC for the year 1999–2000 ranged between 1.2% and 3.7%, and maximum relative abundances occurred at times of relatively low fluxes (Fig. 3b).

The strong positive correlation between factor 1 and POC fluxes at the 54°S site (Table 6) indicates an intimate association between high relative abundances of the high-export group species and pulses of POC export. As a specific example of this, during January 1999 and December 2000, when the highest contribution of the high-export group was noted (55–60% of the total diatom flux; Fig 7), the PFZ sediment trap registered the largest POC fluxes of the record (up to 23 mg m$^{-2}$ d$^{-1}$; Fig. 7). Interestingly, these observations of elevated POC flux coincide with significantly lower summer SSTs than other years.

All the members of the high-export group have been previously reported as important components of both natural and iron-fertilised blooms in the Southern Ocean (Bathmann et al., 1997; Waite and Nodder, 2001; Smetacek et al., 2002; Assmy et al., 2007; Quéguiner, 2013; Grigorov et al., 2014; Rigual-Hernández et al., 2015). The increase in the relative abundance and fluxes of these species during the growth season indicates that they respond opportunistically to the enhanced light levels, most likely undergoing cycles of rapid biomass build-up followed by mass mortality and sinking in the form of aggregates (Smetacek et al., 2004; Green and Sambrotto, 2006; Assmy et al., 2013). This concept is supported by the recent findings of Closett et al. (2015) who documented an increase in the particle sinking speeds at the 54°S site during the summer 1999–2000 of up to at least 35 m d$^{-1}$, a value that falls within the range of previous estimates for marine snow sinking rates (Turner, 2002; Trull et al., 2008; Laurenceau-Cornec et al., 2015). Moreover, other regionally relevant PFZ studies (Ebersbach et al., 2011; Grigorov et al., 2014) concluded that aggregates are the principal form of particle export during the growth season. Taken together, our data and these studies strongly suggest that aggregate formation is a widespread mechanism of the summer bloom in the open-ocean waters of the ACC.

We speculate that the massive development of high-export group diatoms during the growth season facilitates the formation of aggregates in the upper water column, which results in an increase in sinking rates and POC fluxes. Aggregates, and particularly diatom flocs, are rich in exopolymers that increase their effectiveness at scavenging particles they have collided with (Allredge and McGillivary, 1991; Passow and De La Rocha, 2006). Therefore, it is possible that the formation of aggregates during the diatom bloom facilitated the scavenging of other particles (including phytoplankton chains and cells, biominerals and detritus), leading to the co-sedimentation of the major components of the flux (i.e. calcium carbonate, silica and organic carbon). This scavenging mechanism is consistent with previous laboratory observations made by Passow and De La Rocha (2006) and can explain the increase of the sinking rates during the growth season as well as the positive correlation between factor 1 and all bulk components of the flux (Fig. 6b, Table 6b).

Since most of the members of the high-export group are of relatively small in size and weakly silicified, it is unlikely that these species accounted for the major fraction of the BSi export during the summer bloom for most of the years. In contrast, the thick-shelled *F. kerguelensis* is a more compelling candidate to be responsible for the bulk of the BSi export, because despite the fact that its relative abundance exhibited the lowest values of the record during summer, its valve fluxes always were the highest during this season. In terms of carbonate export, the correlation between factor 1 and carbonate flux is not as strong as with the rest of the components of the
of the SCM shade flora (e.g. *Thalassiothrix* and some rhi-
zosolenids) in autumn and winter (the so-called fall dump) has been hypothesised to contribute to a substantial fraction of the annual carbon export in the PFZ (Kemp et al., 2006; Kemp and Villareal, 2013; Quégúiner, 2013). At the 54° S site, the highest fluxes and relative contribution of the deep dwellers *Thalassiothrix antarctica* and *Proboscia* were recorded between the end of the productive period and winter (Fig. 5); however their contribution to the total diatom assemblage was always low (<3.6%) and their flux pulses were not coupled with significant increases in POC export (Fig. 7). Thus, our data do not provide evidence that the sedimentation of these species was associated with a “fall dump” in the PFZ south of Tasmania. However, it has been suggested that sediment traps do not act as good samplers of large mats formed by these long diatoms (Kemp et al., 2006), and therefore their mass sedimentation during autumn and/or winter could have been missed by our sampling technique.

## 5.5 Relative importance of the SAZ and PFZ to carbon export

The depth at which the organic carbon is re-mineralised to CO$_2$ by zooplankton and bacteria determines the timescales during which carbon is sequestered from the atmosphere (Yamanaka and Tajika, 1996; Smetacek et al., 2012). In the SAZ and the PFZ, the fraction of organic carbon recycled within the winter mixed layer (>400 m in the SAZ and between 150 and 200 m in the PFZ; Rintoul and Bullister, 1999; Rintoul and Trull, 2001) would re-equilibrate with the atmosphere within months, whereas only the comparatively smaller portion that reaches deeper layers will remain in the ocean interior for centuries or longer timescales (Trull et al., 2001a). Thus, from the perspective of carbon sequestration, the POC fluxes measured by the traps reported in this study are probably of greater importance than those re-mineralised at mid-depths.

Despite the fact that total mass fluxes in the PFZ at ~1 km were twofold larger than those of the SAZ, the annual POC export was almost identical in both regions (Fig. 8a), implying that particles sinking out of the mixed layer in the SAZ were relatively POC rich (Trull et al., 2001a; Ebersbach et al., 2011). Taking into account that gross primary production is similar in the two zones, or perhaps somewhat lower in the PFZ (Fig. 8b; Lourey and Trull, 2001; Cavagna et al., 2011; Westwood et al., 2011), our results challenge the notion that for a given similar level of production, diatom-dominated ecosystems export greater amounts of carbon to the deep ocean than ecosystems dominated by smaller, non-siliceous phytoplankton (Buesseler, 1998; Boyd and Newton, 1999; Laws et al., 2000). Trull et al. (2001) hypothesised that the similar POC export at both sites could be due to either (1) a more efficient repackaging of carbon for deep transport by the zooplankton community in the SAZ than in the PFZ or (2) to the fact that the silicate-rich particles exported in the PFZ may experience stronger losses of organic carbon at mesopelagic depths than do the carbonate-rich particles of the SAZ. Results from of the SAZ-sense programme (Bowie et al., 2011b) taken together with the data presented in this study provide key information to assess these hypotheses.

Analysis of the flux size spectra at the 47 and 54° S sites by Ebersbach et al. (2011) during January and February 2007 revealed that the vertical export at both stations was dominated by heavily processed particles, mainly faecal aggregates with a slight shift towards smaller particles within the PFZ due to abundant chains of diatoms sinking individually or as part of unconsolidated aggregates. Although the latter study was limited to a short observational period, the results of Ebersbach et al. (2011) suggest that zooplankton grazing had a similar impact on the control of particle export at both sites, and therefore the first hypothesis seems unlikely.

On the other hand, our data show that only a few diatom species, particularly *F. kerguelensis*, dominate the particle export in the silicate-rich and iron-limited waters of the PFZ and AZ. Most of these species are known to significantly increase their BSi : PON and BSi : POC ratios under iron deficiency resulting in the thickening of its already robust frustule (Takeda, 1998; Hoffmann et al., 2007). Furthermore, recent findings from the European Iron Fertilisation Experiment (EIFEX; Smetacek et al., 2012) illustrated that the cellular content of a large fraction of the *F. kerguelensis* stock outside and inside the patch was recycled in the surface layer, resulting in the disproportional sinking of empty frustules to the deep ocean (Assmy et al., 2013). Assmy et al. (2013) concluded that due to these particular traits, *F. kerguelensis* and other exceptionally robust diatoms, such as *Thalassiosira lentiginosa* and *Thalassionema nitzschioides*, preferentially sequester silicon relative to carbon in the iron-limited waters of the ACC. This concept is consistent with our findings in the open waters of the Australian sector south of the SAF, and would help to explain the low POC content and POC : BSi ratios of the particles registered at meso- and bathypelagic depths by our PFZ and AZ traps. Significant, comparisons of our results (Fig. 8a) with satellite and in situ measurements of primary production (Fig. 8b) suggest that high BSi sedimentation rates should be interpreted as a proxy for iron-limited diatom assemblages.
(Hutchins and Bruland, 1998; Takeda, 1998; Assmy et al., 2013) rather than for high primary production. This conclusion raises corresponding caution to previous studies that suggest that higher BSi fluxes in the past refer to a stronger biological carbon pump (Anderson et al., 2009; Sigman et al., 2010).

6 Conclusions

This study reports on the chemical (biogenic silica, carbonate and POC) and biological (diatoms) composition of material exported at ∼ 1 km depth at two sites representative of two major hydrological regions of the Australian sector of the Southern Ocean, the SAZ and PFZ. As a result of different algal communities, the composition and magnitude of the sinking particle fluxes was very different between sites, with higher and BSi-dominated fluxes in the PFZ versus lower and carbonate-dominated fluxes in the SAZ. Despite these differences, the POC export reaching the traps was indistinguishable between sites (∼ 1 g m⁻² yr⁻¹). Seasonality and flux magnitude was more pronounced in the PFZ. The vigorous settling of biogenic particles during summer in the PFZ accounted for a large fraction of the annual POC export. These summer pulses are a major factor responsible for the variability in carbon sequestration from the atmosphere in this region. Our results suggest that the development of a group of bloom-forming diatom species during the growth season probably led to the formation of algal and/or faecal aggregates. The production and sinking of these aggregates most likely facilitated the scavenging of other particles in the water column, and thus co-sedimentation of all the components of the flux. Fragilariopsis kerguelensis dominated the diatom sinking assemblage at both sites and was considered the major biological vector decoupling the carbon and silicon cycles in the waters south the SAF. Comparisons of our data with in situ and satellite primary production estimates were made available by the Ocean Productivity website (http://www.science.oregonstate.edu/ocean.productivity/). Diana M. Davies is thanked for performing biogeochemical analyses on the trap material. Anne-Marie Ballegeer is acknowledged for her technical support in the preparation of samples and comments on an early draft of the manuscript. Thanks to Jan Lieser and John Baumann for their assistance in the remote sensing analysis. The authors are thankful to Jessica Wilks and Kelly Lawler for their help in the microscopy analysis and taxonomic identifications. The authors acknowledge the assistance and support of Nicole Vella and Debra Birch from the Macquarie University Microscopy Unit in the scanning electron microscopy analysis.

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