Factors influencing the stable carbon isotopic composition of suspended and sinking organic matter in the coastal Antarctic sea ice environment


1 School of GeoSciences, The University of Edinburgh, West Mains Road, Edinburgh EH9 3JW, UK
2 Laboratory of Global Marine and Atmospheric Chemistry, School of Environmental Sciences, University of East Anglia, Norwich, NR4 7TJ, UK
3 UMR-CNRS 5805 EPOC, Universite Bordeaux 1, Av. Des Facultes, 33405 Talence, Cedex, France
4 Scottish Universities Environmental Research Centre, Scottish Enterprise Technology Park, Rankine Avenue, East Kilbride, Glasgow, G75 0QF, UK
5 British Antarctic Survey, High Cross, Madingley Road, Cambridge, CB3 0ET, UK

Correspondence to: S. F. Henley (s.f.henley@sms.ed.ac.uk)

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Abstract. A high resolution time-series analysis of stable carbon isotopic signatures in particulate organic carbon (δ13CPOC) and associated biogeochemical parameters in sea ice and surface waters provides an insight into the factors affecting δ13CPOC in the coastal western Antarctic Peninsula sea ice environment. The study covers two austral summer seasons in Ryder Bay, northern Marguerite Bay between 2004 and 2006. A shift in diatom species composition during the 2005/06 summer bloom to near-complete biomass dominance of Proboscia inermis is strongly correlated with a large ~10‰ negative isotopic shift in δ13CPOC that cannot be explained by a concurrent change in concentration or isotopic signature of CO2. We hypothesise that the δ13CPOC shift may be driven by the contrasting biochemical mechanisms and utilisation of carbon-concentrating mechanisms (CCMs) in different diatom species. Specifically, very low δ13CPOC in P. inermis may be caused by the lack of a CCM, whilst some diatom species abundant at times of higher δ13CPOC may employ CCMs. These short-lived yet pronounced negative δ13CPOC excursions drive a 4‰ decrease in the seasonal average δ13CPOC signal, which is transferred to sediment traps and core-top sediments and consequently has the potential for preservation in the sedimentary record. This 4‰ difference between seasons of contrasting sea ice conditions and upper water column stratification matches the full amplitude of glacial-interglacial Southern Ocean δ13CPOC variability and, as such, we invoke phytoplankton species changes as a potentially important factor influencing sedimentary δ13CPOC. We also find significantly higher δ13CPOC in sea ice than surface waters, consistent with autotrophic carbon fixation in a semi-closed environment and possible contributions from post-production degradation, biological utilisation of HCO3− and production of exopolymeric substances. This study demonstrates the importance of surface water diatom speciation effects and isotopically heavy sea ice-derived material for δ13CPOC in Antarctic coastal environments and underlying sediments, with consequences for the utility of diatom-based δ13CPOC in the sedimentary record.

1 Introduction

During photosynthetic uptake of aqueous carbon dioxide, marine phytoplankton preferentially assimilate the lighter isotope, carbon-12, thus increasing the stable carbon isotopic signature, δ13C, of the residual pool of dissolved inorganic carbon (DIC). As such, marine algae always display lower δ13CPOC than the inorganic carbon source they assimilate (Hayes, 1993). Several studies have demonstrated that on large oceanic scales, δ13C of the product organic carbon (δ13CPOC) is inversely correlated with the concentration of
dissolved molecular carbon dioxide ([CO$_2$(aq)]) in surface waters (Rau et al., 1989, 1991). This inverse relationship has been exploited to use $\delta^{13}$C$_{POC}$ in marine sediment cores as a proxy to reconstruct surface water [CO$_2$(aq)] and atmospheric $p$CO$_2$ in the past (Jasper and Hayes, 1990; Freeman and Hayes, 1992; Bentaleb and Fontugne, 1998).

However, several studies have demonstrated that this relationship cannot be applied universally and in high-southern latitudes particularly, the anti-correlation between $\delta^{13}$C$_{POC}$ and [CO$_2$(aq)] can be decoupled by physical and biological factors. Amongst these factors are phytoplankton growth rate and its regulation by temperature and light levels (O’Leary et al., 2001), cell size and shape (Popp et al., 1998; Burkhardt et al., 1999; Trull and Armand, 2001) and non-diffusive carbon uptake through carbon concentration mechanisms (Rau, 2001; Cassar et al., 2004).

Paleoceanographic studies of the Southern Ocean have observed that the $\delta^{13}$C of diatom-bound organic matter was depleted in $^{13}$C during glacial times relative to interglacials and the Holocene (Singer and Shemesh, 1995; Rosenthal et al., 2000; Crosta and Shemesh, 2002; Schneider-Mor et al., 2005). However, ice core records show that glacial $p$CO$_2$ was lower than during interglacials (Berner et al., 1980; Barnola et al., 1987; Masson-Delmotte et al., 2010), which would be expected to drive $\delta^{13}$C$_{POC}$ more positive. Definitive explanations for low glacial $\delta^{13}$C$_{POC}$ remain unclear but potential contributing factors include lower algal growth rates during glacial periods (Rosenthal et al., 2000), sea ice-triggered increase in [CO$_2$(aq)] (Crosta and Shemesh, 2002) and the effects of changes in diatom abundance or species composition (Crosta et al., 2005). Documenting the processes that decouple carbon isotopes from the classic $\delta^{13}$C$_{POC}$ versus $p$CO$_2$ relationship used for paleo-CO$_2$ reconstructions is important in understanding the role of the Southern Ocean in glacial-interglacial climate change. This study provides a detailed high-resolution time-series analysis of carbon isotopes and associated biogeochemical parameters in surface waters, sea ice, sediment traps and core-top sediments in order to elucidate the key factors influencing surface and sinking $\delta^{13}$C$_{POC}$ in the Antarctic sea ice zone on a seasonal timescale, as well as their potential for preservation in marine sediments.

2 Materials and methods

2.1 Study area

This study was conducted over two growing seasons and the intervening winter of full sea ice cover between 2004 and 2006 in Ryder Bay and Marguerite Bay, located south of Adelaide Island, west of the Antarctic Peninsula mainland (Fig. 1). Ryder Bay is a coastal, seasonally sea ice-covered Southern Ocean environment in which diatoms dominate the summer assemblages, with biomass of other phytoplankton such as prymnesiophytes and cryptophytes more than an order of magnitude lower (Garibotti et al., 2005). Ryder Bay adjoins Marguerite Bay and the principal study site is the Rothera Oceanographic and Biological Time-Series (RaTS) site at 67°34.02′ S, 68°14.02′ W (Clarke et al., 2008), situated in open water of depth 520 m. If access to the main RaTS site is prevented by weather or ice conditions, a secondary station at 67°34.85′ S, 68°09.34′ W of water depth $\sim$400 m is used as an alternative site also representative of prevailing oceanographic conditions in Ryder Bay. The Marguerite Bay site is located at 67°55.39′ S, 68°24.15′ W in open water of depth 840 m.

2.2 Sea ice sampling

Sea ice brine was sampled according to sea ice availability at three locations: the RaTS site, Hangar Cove and Lagoon Island (Fig. 1). Fifteen samples were taken over the course of the study: five Lagoon Island land-fast ice samples taken in December 2004, two winter sea ice samples taken at the RaTS site in September and October 2005 and eight early spring samples from Hangar Cove in November and December 2005.

Sea ice brine was sampled using a sack hole drilling method, with samples for the stable carbon isotopic composition of CO$_2$(δ$^{13}$CCO$_2$) and [CO$_2$(aq)] taken first to minimise atmospheric contamination. Samples for δ$^{13}$CCO$_2$ were taken using a 30 ml syringe and gently injected into a 12 ml glass exetainer vial preloaded with 50 µL of 35 gL$^{-1}$ copper (II) sulphate to suppress bacterial activity (Winslow et al., 2001).

Samples for alkalinity and pH, for [CO$_2$(aq)] determination, were taken by immersing a 250 ml glass biological oxy-tern counter vial with a ground glass stopper. On return to the laboratory, samples were stored, unfiltered, in the dark overnight to allow them to reach room temperature and thus maintain a steady temperature throughout subsequent analysis on the following day.

For particulate organic carbon measurements, sea ice brine samples were filtered through muffle-furnaced (400°C) 47 mm diameter GF/F filters, of pore size $\sim$0.7 µm, within two hours of collection. The filters were then dried at 50°C overnight, and stored frozen until analysis. For diatom census counts, sea ice brine was filtered through 37 mm diameter polycarbonate filters, of pore size 0.45 µm. Filters were dried overnight at 50°C and stored in clean plastic Petri-slides until analysis.

2.3 Surface water sampling

A high resolution time series of surface water samples was taken in Ryder Bay, at the RaTS site during the austral spring and summer of 2004/05 and 2005/06. Low resolution time series sampling was conducted during winter 2005.
A normal sampling event consisted of collection of seawater samples from 15 m water depth, the average depth of the chlorophyll maximum in Ryder Bay since sampling began in 1997 (Clarke et al., 2008). Samples were taken for determination of chlorophyll $a$, $[\text{CO}_2\text{aq}]$, $\delta^{13}\text{C}_\text{CO}_2$, suspended POC, $\delta^{13}\text{C}_\text{POC}$ and diatom assemblages and measurements were taken for temperature and salinity using a YSI-500 multi-parameter meter. Each sampling event was accompanied by a full-depth Conductivity Temperature Depth (CTD) cast to monitor changes in mixed layer depth.

CTD casts were taken to 500 m depth using a Sea Bird 19+ CTD module with a WetLabs in-line fluorometer and LiCor PAR sensor. For measurement of temperature at the 15 m sampling depth, a Sensoren Instrumente Systeme GmbH reversible thermometer was lowered to 15 m and allowed to equilibrate for two minutes before a brass messenger was sent down to initiate temperature recording.

Surface water samples were taken using a 5 L Niskin bottle for chlorophyll $a$, $\delta^{13}\text{C}_\text{CO}_2$ and $[\text{CO}_2\text{aq}]$ measurements. For $\delta^{13}\text{C}_\text{CO}_2$, water was drawn from the Niskin bottle using a 50 ml syringe and gently injected into a 12 ml glass extainer vial preloaded with 50 µL of 35 g L$^{-1}$ copper sulphate to suppress bacterial activity (Winslow et al., 2001). Samples for alkalinity and pH, for $[\text{CO}_2\text{aq}]$ determination, were taken from the Niskin straight into a 250 ml glass BOD bottle, which was immediately sealed with a ground glass stopper whilst overflowing and ensuring that no air bubbles were present. These samples were left overnight, as for the equivalent sea ice samples. Chlorophyll samples were collected and treated as per Clarke et al. (2008). Particulate samples were retrieved using a 12 V whale pump and 15 m of silicone tubing weighted down at the end. Water from 15 m was pumped into 10 L HDPE carboys for transfer back to the laboratory. Surface water samples were prepared for particulate organic carbon measurements and diatom census counts in the same way as was sea ice brine.
2.4 Sediment trap and surface sediment sampling

Two sediment trap mooring arrays were deployed to catch sinking particles for \( \delta^{13} \)CPOC analysis and flux calculations, concurrent with the time series water sampling programme; one at the RaTS site and the other at the deeper Marguerite Bay site.

Each mooring consisted of two time-series sediment traps, at 200 m and 512 m for the RaTS mooring and 123 m and 745 m for the Marguerite Bay mooring. Each sediment trap consisted of 21 rotating cups programmed to rotate at predefined intervals. Cup turnover times were shorter giving higher resolution during periods of potential sea ice melt and the spring bloom, whilst lower resolution cup rotation was used during the low flux winter periods.

Both sediment trap mooring arrays were deployed from late-January 2005 to mid-February 2006. Upon recovery, all sediment trap bottles were removed and replaced, and the moorings redeployed.

Prior to deployment, each cup was filled with filtered seawater spiked with an extra 5 % NaCl in order to increase its density and prevent mixing with the overlying seawater. Finally, cups were spiked with formaldehyde to give an overall concentration of 2 % (v/v) to prevent bioturbation, by killing swimmers and stopping biological activity. Formaldehyde-preservation of sediment trap material for \( \delta^{13} \)CPOC analysis is widely used (Thunell et al., 2000; Struck et al., 2004; Mincks et al., 2008) and is deemed appropriate for the purposes of this study since formaldehyde preservative does not add sufficient organic carbon to sediment trap material to alter \( \delta^{13} \)CPOC (Altabet, 2001).

Box core samples were taken at the RaTS site and the Marguerite Bay site in January 2005 and December 2006 aboard R.R.S. James Clark Ross. In each case, the box core was taken and then four sub-cores of approximately 30 cm were taken by pushing plastic sleeves through the box core. Core-top samples were collected from the top two 0.5 cm intervals from each sub-core.

2.5 Surface water and sea ice [CO\(_2\)\(_{aq}\)] and \( \delta^{13} \)C\(_{CO_2}\) determination

[CO\(_2\)\(_{aq}\)] was determined using measurements of salinity and temperature, detailed above, with pH and alkalinity, both determined on the day following sampling. pH measurements were performed using a bench-top pH meter calibrated to buffer solutions of pH 4.01, 7.00 and 10.01. Maximum error on triplicate pH measurements across all samples was \( \pm 0.02 \). Alkalinity was determined by titration with 0.05 M HCl and the Gran plot method (Almgren et al., 1983). [CO\(_2\)\(_{aq}\)] was calculated using constants from Dickson and Millero (1987), Hannson (1973) and Mehrbach et al. (1973) using the CO2SYS programme (Lewis and Wallace, 1998). Maximum error on [CO\(_2\)\(_{aq}\)] calculations, taking into account the maximum error on all input parameters, is 11.0 %.

\( \delta^{13} \)DIC analysis was conducted by GC-IRMS using a method similar to Assayag et al. (2006). The 12 ml glass exetainer vial containing 12 ml of seawater sample spiked with CuSO\(_4\).5H\(_2\)O was split into two samples by inserting a closed syringe through the septum of the vial and injecting 6 ml of Helium gas into the sample vial using a separate needle and syringe. The 6 ml of sample forced into the closed syringe by the He injection was then injected into a clean 12 ml exetainer vial that had been under vacuum for 30 min. Each sample vial was then injected with 0.6 ml of concentrated H\(_3\)PO\(_4\) in order to convert the DIC into aqueous and gaseous CO\(_2\) for analysis. Three sets of isotopic standards were prepared (MAB2, CaCO\(_3\) and NaHCO\(_3\)) using a range of final DIC concentrations. The standards were weighed into 12 ml glass exetainer vials and then placed on a vacuum to remove all gases. 6 ml of 10 % H\(_3\)PO\(_4\) was then injected into each standard vial to reproduce the same conditions as in the sample vials. \( \delta^{13} \)C\(_{DIC}\) was analysed using a custom-built GC-IRMS system, from which raw \( \delta^{13} \)C values were corrected using the isotopic standards. Precision of \( \delta^{13} \)C\(_{DIC}\) values was generally better than 0.2 ‰. \( \delta^{13} \)C\(_{CO_2}\) was determined from \( \delta^{13} \)C\(_{DIC}\) and absolute temperature (\( T_K \)) in Kelvin) using Eq. (1) from Rau et al. (1996):

\[
\delta^{13} \text{C}_{CO_2} = \delta^{13} \text{C}_{DIC} + 23.644 - 9701.5/T_K
\]

2.6 POC, PN and \( \delta^{13} \)CPOC analysis

Bulk POC, particulate nitrogen (PN) and \( \delta^{13} \)CPOC analyses were conducted using a method similar to Lourey et al. (2004). Prior to analysis, the filters were decarbonated by wetting with Milli-Q water and fuming with HCl for 48 h and then drying at 50°C. Filters were cut in half and analysed for elemental POC, PN and \( \delta^{13} \)CPOC using a Carlo Erba NA 2500 elemental analyser in-line with a VG PRISM III isotope ratio mass spectrometer. The two halves were analysed separately and then data were summed, to achieve final representative values for the whole filters. All \( \delta^{13} \)C data are presented in the delta per mil notation versus V-PDB (%/vPDB).

2.7 Diatom species counts

Diatom assemblages were determined by analysing a sub-sample of each polycarbonate filter by scanning electron microscopy. Counting methods, surface area, volume and biomass determinations and species identification in surface samples are detailed in Annett et al. (2010). Sea ice samples were analysed following identical protocols. Diatom census counts were also conducted on sediment trap material, according to the methods of Laws (1983) and Schrader and Gersonde (1978). Full details on slide preparation and diatom identification are as per Crosta et al. (2004).
2.8 Sediment trap and core-top sediment $\delta^{13}$C$_{POC}$ analysis

After recovery of the sediment trap mooring arrays, the solution in each sample cup was allowed to settle, the supernatant siphoned off and the swimmers removed manually using HCl-cleaned plastic forceps and a x10 dissecting binocular microscope. Each sample cup was then split into 10 fractions using a rotary splitter at the National Oceanography Centre (NOC), Southampton, UK.

One fraction from each sediment collection cup was washed, freeze-dried and ground for analysis of $\delta^{13}$C$_{POC}$. Duplicate 10 mg aliquots of this dried sediment were weighed into silver capsules, acidified with 5% HCl to remove carbonates and then dried at 60°C overnight. Decarbonated samples were then analysed for $\delta^{13}$C$_{POC}$ using a VG PRISM III isotope ratio mass spectrometer. One sub-core of each box core was prepared and analysed for $\delta^{13}$C$_{POC}$ in the same way as the sediment trap cup fraction.

2.9 Data analysis and statistics

All statistical analyses were performed using R computing software. Relevant information for each analysis is summarised in Appendix Table 1, in the order in which results appear in the text, and given due consideration in the discussion that follows.

3 Results

3.1 Seasonal sea ice cover and productivity

Sea ice cover, mixed layer depth and chlorophyll $a$ data from the austral summer growing seasons of 2004/05 and 2005/06 are presented in Fig. 2. Total sea ice cover was variable between the two seasons at the RaTS site, with full cover occurring for 138 days from 16 June to 1 November during winter 2004 and 198 days from 10 June to 25 December 2005. The mixed layer depth data show typical seasonality for Ryder Bay, with a deep winter mixed layer and a shallow surface layer in summer, influenced heavily by sea ice and surrounding glaciers (Meredith et al., 2004). Mixed layer depth is defined as the depth at which $\sigma_0 = \sigma_0$ (surface) + 0.05 (Barth et al., 2001), where $\sigma_0$ is the potential density anomaly = $\rho - 1000$, and $\rho$ is density in kg m$^{-3}$. A stratified surface ocean during summer reduces wind-induced vertical mixing and provides a stable environment for proliferation of diatom blooms and resultant seasonal drawdown of macro- and micromutrients (Clarke et al., 2008). In this study, both growing seasons lasted ~4 months, but the 2005/06 phytoplankton bloom occurred around 6 weeks later in accordance with later sea ice retreat. Summer surface water conditions were also much more stable in 2005/06 with a longer period characterised by a shallow mixed layer, in agreement with more persistent sea ice cover during the preceding winter.
3.2 Dissolved carbon dioxide and $\delta^{13}$C$_{CO_2}$ in surface waters and sea ice

The concentration of CO$_2$ and $\delta^{13}$C$_{CO_2}$ in surface waters show a general trend of [CO$_2$(aq)] decrease and $^{13}$C enrichment during spring and summer during both summer seasons (Fig. 3). During the 2004/05 season, [CO$_2$(aq)] decreased from values as high as 54.2 ± 6.0 µM to 5.1 ± 0.6 µM whilst $\delta^{13}$C$_{CO_2}$ values rose from –11.2 to –9.0‰. Similarly during the 2005/06 season, [CO$_2$(aq)] decreased from a high winter value of 33.7 ± 3.7 µM to 13.6 ± 1.5 µM and $\delta^{13}$C$_{CO_2}$ values rose from –11.1 to –9.2‰. Important to note however, is that the 2004/05 season was characterised by rapid and large fluctuations in [CO$_2$(aq)]; in fact, season maximum concentration occurs after the first chlorophyll peak in the middle of the growing season. Conversely, [CO$_2$(aq)] shows a much more systematic reduction over the duration of the 2005/06 growing season, albeit with a lesser overall drawdown. Similarly, $\delta^{13}$C$_{CO_2}$ shows much more variability in 2004/05 than the gradual increase seen in summer 2005/06. The greater variability seen in the 2004/05 season depicts regular inputs of CO$_2$, which resulted in small negative shifts in $\delta^{13}$C$_{CO_2}$ to values as low as –10.5‰ during January and February 2005. The absence of such fluctuations in $\delta^{13}$C$_{CO_2}$ during summer 2005/06 shows that there is no regular mid-season input of CO$_2$. However, there is one marked increase in [CO$_2$(aq)] and simultaneous decrease in $\delta^{13}$C$_{CO_2}$, which provides evidence for a one-off mid-season input of CO$_2$, coincident with a mid-season chlorophyll reduction between two periods of greater phytoplankton productivity.

The $\delta^{13}$C$_{CO_2}$ in sea ice is generally enriched relative to surface waters and exhibits greater temporal variability, with values ranging from –10.7 to –4.8‰ (Fig. 3). CO$_2$ concentrations in sea ice brine are lower than in surface waters, consistent with higher $\delta^{13}$C$_{CO_2}$. In addition to temporal variability in [CO$_2$(aq)] and $\delta^{13}$C$_{CO_2}$, the greater variability seen here than in surface water samples is partly spatial, as is common in sea ice brine (Rau et al., 1992; Kennedy et al., 2002) since samples were taken from different locations in the study area according to availability of ice.
3.3 Particulate organic carbon in surface waters and sea ice

Concentrations of POC in Ryder Bay surface waters mimic levels of chlorophyll \( a \) and show similar variability over summer 2004/05 and gradual trends in 2005/06 as do \([\text{CO}_2]_{(aq)}\) and \(\delta^{13}\text{C}_{\text{CO}_2}\) (Fig. 3). However, surface water \(\delta^{13}\text{C}_{\text{POC}}\) shows high inter-annual variability between the two growing seasons. During the 2004/05 season, \(\delta^{13}\text{C}_{\text{POC}}\) increases gradually over the course of the phytoplankton bloom from \(-21.2\) to \(-17.9\) ‰, with a small \(-1\)‰ decrease in \(\delta^{13}\text{C}_{\text{POC}}\) during late December 2004 when chlorophyll \( a \) declined and \([\text{CO}_2]_{(aq)}\) increased. In February 2005, when chlorophyll \( a \) began to decline at the end of the growing season, a large yet short-lived \(-9\)‰ negative shift is observed in \(\delta^{13}\text{C}_{\text{POC}}\) to a season-low of \(-26.7\) ‰. This occurs in concert with an increase in \([\text{CO}_2]_{(aq)}\) of \(-8\) \(\mu\text{M}\) and a decrease in \(\delta^{13}\text{C}_{\text{CO}_2}\) of \(-1.2\)‰. At the end of the growing season, \(\delta^{13}\text{C}_{\text{POC}}\) returned to a near winter value of \(-23\)‰. During the 2005/06 growing season, \(\delta^{13}\text{C}_{\text{POC}}\) increased from a winter low of \(-25\)‰ in September 2005 to a season high of \(-18.8\)‰ in December 2005 just prior to sea ice retreat. Once the open water spring phytoplankton bloom was underway, \(\delta^{13}\text{C}_{\text{POC}}\) was consistently around \(-21\)‰ until there was an injection of \(-7\) \(\mu\text{M}\) \(\text{CO}_2\) into the system during late January and concomitant decreases in \(\delta^{13}\text{C}_{\text{CO}_2}\) and \(\delta^{13}\text{C}_{\text{POC}}\) of \(0.7\)‰ and \(2\)‰, respectively. In late January and early February 2006, at the commencement of the second chlorophyll \( a \) peak, there was a large negative shift in \(\delta^{13}\text{C}_{\text{POC}}\) of \(-10\)‰ to values as low as \(-28.7\)‰ (Fig. 3). This negative shift in \(\delta^{13}\text{C}_{\text{POC}}\) was maintained throughout the second chlorophyll \( a \) peak and once chlorophyll had declined at the end of the growing season, towards the end of March 2006, the \(\delta^{13}\text{C}_{\text{POC}}\) returned to a typical winter value of \(-25\)‰. In agreement with this large and prolonged negative isotopic transition, a seasonal POC concentration-weighted average \(\delta^{13}\text{C}_{\text{POC}}\) of \(-24.5\)‰ was significantly lower for 2005/06 than the 2004/05 season average of \(-20.0\)‰ (2-sample t-test \(p < 0.001\)).

POC:PN ratios of suspended material averaged 5.8 indicating a wholly marine origin, as would be expected at a site like Ryder Bay, due to the relative paucity of terrestrial organic matter in the vicinity (Fig. 4a). The dominant marine phytoplankton source of Ryder Bay organic matter is confirmed by POC:chl \( a < 200\) in the vast majority of suspended samples.

Sea ice POC:PN is highly variable throughout the season of sea ice coverage, but an average value of 10 is higher than in surface waters (Fig. 4b). Most of our values for POC:chl \( a \) in sea ice range from 83.7 to 497.0 (Fig. 4b) and therefore fall within the range of POC:chl \( a \) values found in previous studies of marine biota (Eppley et al., 1973; Pollehne et al., 1993) and sea ice algal assemblages (Gleitz and Thomas, 1993). We also observe significantly higher POC:chl \( a \) values of 1250 to 1750 in late-December 2004. Conversely, at
3.4 Diatom assemblages and size classes

The surface water phytoplankton bloom in Ryder Bay is typically dominated by the microplankton fraction (>20 µm), so large solitary or chain-forming diatoms dominate over the smaller nanoplankton and picoplankton (Clarke et al., 2008). Diatom assemblages show distinct changes throughout the two growing seasons in surface waters and sea ice (Fig. 5; Annett et al., 2010). Briefly, diatom biomass in 2004/05 was initially relatively diverse, with substantial contributions from *Minidiscus chilensis* and *Chaetoceros* (*Hyalochaeta* subgenus) species. Mid-season assemblages were dominated by *Odontella weissflogii*, accounting for up to 80% of the estimated diatom community. Late-season assemblages returned to a more diverse composition. The early part of the 2005/06 season showed a mixed diatom assemblage, consisting largely of *Fragilariopsis cylindrus*, large and medium centrics (>50 µm and 20 to 50 µm, respectively) and a small contribution from *Proboscia inermis*. A shift towards the almost complete dominance of *P. inermis* occurred at the time of the late-season negative excursion in δ13CPOC. In both seasons, surface area to volume ratios (SA:V) estimated for the diatom community are initially high (~0.76 µm²:µm³) and decline thereafter (0.2 to 0.3 µm²:µm³). More variability is seen in SA:V in 2004/05 than in 2005/06, in accordance with the more diverse assemblages in the earlier season.
In sea ice, we observe less species variability than in surface waters (Fig. 5). In December 2004, sea ice biomass is made up of medium centric groups, such as *Porosira* and *Thalassiosira* species. In the 2005/06 season, sea ice diatom assemblages were dominated initially by *Chaetoceros simplex* and very small centric species (<10 µm). In early December 2005, the main contribution comes from the *Fragilariopsis curta* group, primarily *F. cylindrus* but also *F. curta*.

Diatom census counts in the sediment traps show that species composition at depth is broadly similar to that in surface waters, but with different proportions of each species (Fig. 6), presumably a result of differential dissolution during sinking through the water column. Shallow traps are dominated by cold water species such as the *F. curta* and *Chaetoceros* groups, although higher resolution sampling in Marguerite Bay shows minor contributions from the *F. obliquecostata* group and *Thalassiosira* spp. throughout the sampling period. Both deeper traps are overwhelmingly dominated by *Chaetoceros* with minor contributions from *F. curta* and *T. antarctica* groups. *P. inermis* abundance was low in all sediment traps throughout the study period.

### 3.5 Sinking particulate organic carbon

Sediment trap carbon flux and $\delta^{13}$C$_{POC}$ data from Ryder Bay and Marguerite Bay are shown in Fig. 7, and Table 1 provides a summary of sediment trap isotopic data. Carbon fluxes were highly variable between traps and between seasons, but the Ryder Bay traps show greater overall export than the Marguerite Bay traps (Fig. 7). There is no significant change in seasonal flux-weighted average $\delta^{13}$C$_{POC}$ with water depth at either mooring site in either growing season.

Seasonal variability in surface water $\delta^{13}$C$_{POC}$ is, to some extent, reflected in the sediment trap data (Fig. 7). During the 2004/05 season, surface water $\delta^{13}$C$_{POC}$ ranged from $-21.2\%e$ to $-17.5\%e$ from early December to the beginning of February (Fig. 3). Whilst this $4\%e$ enrichment is not fully expressed in the sediment trap data, we do see a slight $\delta^{13}$C$_{POC}$ increase in the RaTS 200 m sediment trap from $-20.7\%e$ to $-19.7\%e$ between late January and late February. We see a more pronounced negative shift to $-22.5\%e$ in sediment trap $\delta^{13}$C$_{POC}$ at the beginning of March 2005, approximately one month after the negative shift in surface $\delta^{13}$C$_{POC}$ to $-26.7\%e$ observed at the beginning of February 2005. A similar response is not observed in the Marguerite Bay traps, possibly due to different diatom assemblages in the more open

![Fig. 6. Time-series plots of diatom species composition in sediment traps between December 2004 and March 2006, location and depth of traps as described on each plot.](image-url)
Fig. 7. $\delta^{13}\text{C}_{\text{POC}}$ and carbon (C) flux data from time-series sediment traps from December 2004 until March 2006, as per legend. RaTS is the trap mooring at the routine sampling site in Ryder Bay. MB is the deeper trap mooring in Marguerite Bay. The top panel shows sea ice coverage as measured daily in Ryder Bay (data from the British Antarctic Survey). Maximum error on sediment trap $\delta^{13}\text{C}_{\text{POC}}$ values is 1.0 ‰ associated with formaldehyde preservation (Mincks et al., 2008) since this vastly exceeds analytical error.

Table 1. Sediment trap seasonal flux-weighted average $\delta^{13}\text{C}_{\text{POC}}$ presented in ‰ versus VPDB; maximum error quoted as the 1.0‰ uncertainty associated with formaldehyde preservation (Mincks et al., 2008), as this vastly exceeds analytical error.

<table>
<thead>
<tr>
<th>Trap</th>
<th>$\delta^{13}\text{C}_{\text{POC}}$ (2004/05)</th>
<th>$\delta^{13}\text{C}_{\text{POC}}$ (2005/06)</th>
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</thead>
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<tr>
<td>RaTS 200 m</td>
<td>$-20.3 \pm 1.0$</td>
<td>$-23.3 \pm 1.0$</td>
</tr>
<tr>
<td>RaTS 512 m</td>
<td>$-20.4 \pm 1.0$</td>
<td>$-24.7 \pm 1.0$</td>
</tr>
<tr>
<td>MB 123 m</td>
<td>$-19.1 \pm 1.0$</td>
<td>$-20.5 \pm 1.0$</td>
</tr>
<tr>
<td>MB 745 m</td>
<td>$-19.1 \pm 1.0$</td>
<td>$-22.3 \pm 1.0$</td>
</tr>
</tbody>
</table>

As in surface waters, 2005/06 sediment trap data show very different $\delta^{13}\text{C}_{\text{POC}}$ characteristics to those from 2004/05. Although data are somewhat sparse for the Ryder Bay trap, they do highlight a similar negative transition to very low $\delta^{13}\text{C}_{\text{POC}}$ values at the end of the 2005/06 growing season as seen in surface values. In surface waters, $\delta^{13}\text{C}_{\text{POC}}$ undergoes a large negative shift at the end of January to $-28.7\%e$. Although there is a time lag, the same negative shift is seen in late February in the 200 m RaTS trap ($\delta^{13}\text{C}_{\text{POC}} = -28.7\%e$) and the 512 m trap, where $\delta^{13}\text{C}_{\text{POC}}$ drops from $-21.8\%e$ in early February to $-27.8\%e$ in late February (Fig. 7).

4 Discussion

4.1 $\delta^{13}\text{C}_{\text{POC}}$ and [CO$_2$(aq)] in surface waters and sea ice

According to the classic CO$_2$–$\delta^{13}\text{C}_{\text{POC}}$ relationship (François et al., 1993), we would expect $\delta^{13}\text{C}_{\text{POC}}$ to vary depending on the balance between supply and demand of the photosynthetic carbon source. This supply and demand model is regulated by two steps in the photosynthetic process: transport of the inorganic carbon reactant into the internal cell carbon pool and subsequent fixation to organic carbon (Popp et al., 1999; Trull and Armand, 2001). According to this model, an increase in external [CO$_2$(aq)] would increase the fractionation factor ($\varepsilon_p$) of inorganic carbon assimilation and decrease $\delta^{13}\text{C}_{\text{POC}}$, independent of the initial $\delta^{13}\text{C}_{\text{CO}_2}$ (François et al., 1993; Rau et al., 1996; Burkhardt et al., 1999; Lourey et al., 2004).

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We investigate the influence of changing $[\text{CO}_2]_{\text{aq}}$ on $\delta_{p}$ and $\delta^{13}\text{C}_{\text{POC}}$ in Fig. 8, which presents $\delta^{13}\text{C}_{\text{POC}}$ data from this study plotted against concurrent $[\text{CO}_2]_{\text{aq}}$ in sea water and sea ice brine, as well as theoretical relationships between $\delta^{13}\text{C}_{\text{POC}}$ and $[\text{CO}_2]_{\text{aq}}$. These theoretical relationships are based on $\delta_{p} = 25\%$ which is within the commonly accepted range of maximum $\delta_{p}$ values, 25 to 28$\%$ (Raven and Johnston, 1991; Goericke et al., 1994), and closed and open system isotope fractionation approaches. Open system isotopic evolution is calculated using Eq. (2), where $\delta^{13}\text{C}_{\text{CO}_2}$ ini is the initial isotopic value of reactant CO$_2$, for which we use $-11\%$ for surface water and sea ice. $f$ is the fraction of CO$_2$ utilised compared to maximum values of 60$\mu$M in sea water and 10$\mu$M in sea ice (Fig. 3).

$$\delta^{13}\text{C}_{\text{POC}} = \delta^{13}\text{C}_{\text{CO}_2}\text{ini} + \delta_{p} \times f$$

Closed system isotopic evolution is calculated using the Rayleigh accumulated product equation (Eq. 3).

$$\delta^{13}\text{C}_{\text{POC}} = \delta^{13}\text{C}_{\text{CO}_2}\text{ini} - \delta_{p} \left( \frac{f \ln(f)}{1-f} \right)$$

Unlike the strong negative correlation observed between $\delta^{13}\text{C}_{\text{POC}}$ and $[\text{CO}_2]_{\text{aq}}$ in open-ocean studies in the Southern Ocean (Lourey et al., 2004), we demonstrate no significant relationship in coastal sea ice or surface waters, assuming either closed or open system dynamics (Fig. 8). We see no correlation between observed $\delta^{13}\text{C}_{\text{POC}}$ and $[\text{CO}_2]_{\text{aq}}$ in surface water ($r^2 = 0.247$; $p = 0.120$) or sea ice ($r^2 = 0.200$; $p = 0.511$) and the data clearly do not fit a modelled isotopic evolution at any value of $\delta_{p}$ assuming closed or open system dynamics. $\delta^{13}\text{C}_{\text{CO}_2}$ can be excluded as a possible driver of the observed negative shifts in $\delta^{13}\text{C}_{\text{POC}}$, since a $\delta^{13}\text{C}_{\text{CO}_2}$ shift of only $-2.2\%$ in surface waters cannot account for the large shift in $\delta^{13}\text{C}_{\text{POC}}$ of $\sim -10\%$ (Fig. 3). Therefore, although some more subtle changes in $\delta^{13}\text{C}_{\text{POC}}$ are consistent with minor changes in $[\text{CO}_2]_{\text{aq}}$ and $\delta^{13}\text{C}_{\text{CO}_2}$ during biological uptake and/or nutrient injections, inorganic carbon availability is not the primary control on $\delta^{13}\text{C}_{\text{POC}}$ in sea ice or surface waters in Ryder Bay.

### 4.2 Factors influencing $\delta^{13}\text{C}_{\text{POC}}$ in surface waters

Phytoplankton growth rates typically show a positive relationship with $\delta^{13}\text{C}_{\text{POC}}$, since higher growth rates increase carbon demand and restrict $\delta_{p}$ (Laws et al., 1995; Rau et al., 1996; Popp et al., 1998; Burkhardt et al., 1999; Villinski et al., 2000; Trull et al., 2008). While growth rate data are not available for the study period, a role for algal growth rate cannot be ruled out. However, this would more likely be significant at the onset of the phytoplankton bloom and growth rate-related $\delta^{13}\text{C}_{\text{POC}}$ changes in iron-replete Southern Ocean environments typically account for up to 2$\%$ (Trull et al., 2008). Growth rate cannot therefore explain the full extent of the rapid late-season isotopic shifts seen in Ryder Bay. Since Ryder Bay surface waters do not conform to the widely published growth rate/[CO$_2$]$_{\text{aq}}$ vs. $\delta^{13}\text{C}_{\text{POC}}$ relationship.

![Diagram](https://www.biogeosciences.net/9/1137/2012/fig8.png)
(e.g. Jasper et al., 1994), an alternative mechanism must be driving the large decreases in δ13CPOC observed in the latter part of the Ryder Bay growing seasons.

In this study, the largest demonstrated determinant on seasonal δ13CPOC changes in surface waters is diatom assemblage. However, least-squares linear regression analysis shows that there is only a weak negative relationship between diatom SA:V and δ13CPOC, even when early season samples likely dominated by sea ice-derived organics (those plotting on the same line as sea ice samples) are excluded ($r^2 = 0.472; p = 0.017$, Fig. 9). Time-series data (Fig. 9) clearly show that whilst the negative excursions in δ13CPOC are accompanied by slight changes in SA:V, there is no significant SA:V control on δ13CPOC. We deduce therefore that in contrast to Popp et al. (1998, 1999), Burkhardt et al. (1999) and Trull and Armand (2001), diatom species control of δ13CPOC is not related to cell size or SA:V in this study, but rather that these parameters are all responding to an additional factor.

The most striking change in surface water diatom assemblages concurrent with the large negative shift in δ13CPOC is the shift to near total dominance of Proboscia species, particularly P. inermis, during the second chlorophyll peak of summer 2005/06 (Fig. 10). Least-squares linear regression analysis reveals an extremely significant relationship ($r^2 = 0.968; p = 0.000243$) between the percentage of total biomass made up of P. inermis and δ13CPOC for 2005/06. Although diatom species changes appear to have a much greater effect on δ13CPOC during the 2005/06 season when P. inermis makes a significantly greater contribution to total diatom biomass, the relatively high abundance (21%) of P. inermis in one sample of the 2004/05 growing season also seems related to isotopically lighter POC (–22.4 ‰). However, presence of P. inermis cannot explain the lowest value...
Fig. 10. Comparison of the fraction of surface water diatom biomass made up of *P. inermis* (triangles), δ¹³C_POC (circles; solid line) and ε_p (crosses; dashed line) for 2004/05 and 2005/06. Time-series data is shown in (a), note the reversal of the right hand y-axis (% *P. inermis*) to better illustrate the coupling between *P. inermis* biomass and δ¹³C_POC. Variation of δ¹³C_POC and ε_p vs. *P. inermis* biomass for all 2005/6 samples and the only sample from 2004/5 where *P. inermis* accounts for >2% of total biomass are shown in (b).

Least-squares linear regression analysis on all samples where *P. inermis* is expected to exert a significant control on δ¹³C_POC (>2% of total biomass) shows a very strong relationship between *P. inermis* abundance and δ¹³C_POC (r² = 0.918; p = 0.000423, Fig. 10) and therefore suggests that the majority of δ¹³C_POC variability can be attributed to species effects. Similar biomass dominance by *O. weissflogii* in 2004/5 corresponds with no appreciable shift in δ¹³C_POC to either lighter or heavier values, so we argue that species-specific effects on δ¹³C_POC are not exerted by all diatom species in Ryder Bay. Instead, it is the unusual biochemistry of *P. inermis* that drives distinct negative shifts to δ¹³C_POC values as low as −29‰ in this study.

Proboscia diatoms are known to synthesise a unique set of long-chain 1,14-diols and 12-hydroxy methyl alkanoates with strongly depleted carbon isotope signatures of <−32‰ and <−34‰, respectively (Sinninghe Damsté et al., 2003). While these *Proboscia* lipids can contribute up to 35% of total lipid flux in sediment traps (Wakeham et al., 2002), it is unlikely that they alone account for the low δ¹³C_POC signatures seen in this study, as they typically make up less than 1% of total organic carbon (raw data from Wakeham et al. (2002), published online at http://usigofs.whoi.edu/PI-NOTES/arabian/Wakeham/sedtrap_lipid_raw.html). However, the ¹³C depletion of these lipids is greater than those of alkenones from haptophytes or dinosterol from dinoflagellates and points to a substantially depleted pool of intracellular carbon in...
Proboscia spp. not readily explained by factors such as SA:V (Sinninghe Damsté et al., 2003). Sinninghe Damsté et al. (2003) found that all Proboscia species analysed synthesised these “Proboscia lipids”, thus it is likely that the potential influence on δ\(^{13}\)C\(_{\text{POC}}\) is not limited to P. inermis. Accordingly, samples in this study where other Proboscia species were observed (Proboscia truncata), albeit at low abundances, fall on the same trend presented in Fig. 10 and least-squares linear regression analysis yields an almost identical \(r^2\) value as for P. inermis alone (\(r^2 = 0.922, p = 0.00037\); data not shown). However, the Proboscia bloom presented here was overwhelmingly dominated by P. inermis, so we restrict our conclusions regarding species effects to this species.

A detailed examination of why the internal carbon pool of P. inermis is so depleted is beyond the scope of this study. However, we suggest that carbon concentrating mechanisms (CCMs) employed by many diatom species to buffer the impacts of variability in [CO\(_2\)\(_{\text{a}}\)] may play an important role in the species-related differences in δ\(^{13}\)C\(_{\text{POC}}\) seen here (Sharkey and Berry, 1985; Descolas-Gros and Fontugne, 1985; Goericke et al., 1994; Laws et al., 1995).

Phytoplankton employing typical C\(_3\) biochemistry in the absence of CCMs, i.e. diffusive CO\(_2\) transfer to the internal cell carbon pool and eukaryotic Rubisco carboxylation (Kerby and Raven, 1985), fractionate CO\(_2\) by \(\sim 29\%\)e and produce organic carbon of \(-25\) to \(-30\%\)e (Raven et al., 1994). We propose therefore that P. inermis utilises a simple C\(_3\) photosynthetic pathway with no employment of CCMs. An additional contribution to low δ\(^{13}\)C\(_{\text{POC}}\) values associated with P. inermis may arise from the production of isotopically light “Proboscia lipids”.

Why a more mixed phytoplankton assemblage prior to the bloom of P. inermis coincides with higher δ\(^{13}\)C\(_{\text{POC}}\) (\(-25\) to \(-15\%\)e) remains unclear. However, direct active HCO\(_3\) uptake, which is significant in the marine environment (Tortell et al., 1997, 2006; Cassar et al., 2004), can drive δ\(^{13}\)C\(_{\text{POC}}\) to values \(\gtrsim 10\%\)e (Raven, 1997 and references therein), since HCO\(_3\) is isotopically enriched relative to CO\(_2\) by approximately \(12\%\)e at 0 °C (Deines et al., 1974; Mook et al., 1974). HCO\(_3\) utilisation mediated by phosphoenolpyruvate carboxylase (PEPC) would also drive δ\(^{13}\)C\(_{\text{POC}}\) relatively high due to a much smaller \(\varepsilon_p\) than the initial HCO\(_3\) enrichment relative to CO\(_2\) (O’Leary, 1981; O’Leary et al., 1992). We speculate therefore that higher δ\(^{13}\)C\(_{\text{POC}}\) prior to the P. inermis bloom may have been driven by some diatom species in the mixed assemblage employing CCMs and/or utilising HCO\(_3\) as a carbon substrate.

In summary, we find a striking relationship between diatom species composition and δ\(^{13}\)C\(_{\text{POC}}\) in surface waters such that large and rapid increases in abundance of P. inermis appear to explain the large negative shifts in surface water δ\(^{13}\)C\(_{\text{POC}}\). Although the exact nature of the biochemical mechanisms employed by Ryder Bay diatom species remain unknown, we speculate that the unusual biochemistry of P. inermis and its lack of a CCM during photosynthetic uptake give it a characteristically light isotopic signature. The large negative shift in surface water δ\(^{13}\)C\(_{\text{POC}}\) accompanying the large and rapid increase in P. inermis abundance can then be explained by the low δ\(^{13}\)C\(_{\text{POC}}\) signature of the large proportion of diatom biomass attributable to this species.

4.3 Factors influencing δ\(^{13}\)C\(_{\text{POC}}\) in sea ice

Carbon isotopic signatures in sea ice are important because sea ice-derived organic material is released to the underlying water column during brine drainage events with implications for the sinking flux of δ\(^{13}\)C\(_{\text{POC}}\). Input from sea ice melting is thought to be less significant in Ryder Bay than in other coastal Antarctic environments as sea ice tends to blow out of the bay rather than undergoing extensive melting in situ (Clarke et al., 2008). However based on POC:chl \(a\) and POC:PN ratios, we estimate that the maximum contribution to suspended organic carbon from sea ice material is 10 to 34 % in 2004/05 and 18 to 53 % in 2005/06. These estimates are based on minimum POC:chl \(a\) of 84 for sea ice material and 22 for surface waters and minimum POC:PN of 5.35 in surface waters and 6.94 and 6.26 for sea ice in 2004/5 and 2005/6 respectively. These end-member ratios were used to calculate the relative proportions of sea ice and surface water material required to produce the ratios measured in surface waters on 13 December 2004 and 23 December 2005, the two samples where sea ice material is likely to have had the strongest impact on δ\(^{13}\)C\(_{\text{POC}}\). Although approximate, these estimations suggest that the input of sea ice-derived organics does exert a control on surface water δ\(^{13}\)C\(_{\text{POC}}\) and support observations of increased surface water δ\(^{13}\)C\(_{\text{POC}}\) when sea ice was present in Ryder Bay.

In general, sea ice δ\(^{13}\)C\(_{\text{POC}}\) was enriched relative to surface waters, consistent with generally higher δ\(^{13}\)C\(_{\text{CO}}\) in particular in December 2005, and lower CO\(_2\) concentration in the sea ice brine (Fig. 3). This is in agreement with autotrophic carbon fixation by phytoplankton in a closed or semi-closed system and, as such, a higher degree of CO\(_2\) utilisation than in the open surface water system (Gibson et al., 1999; Villinski et al., 2000). Consequently, Ryder Bay sea ice is characterised by seasonal deficits of all major nutrients, higher dissolved oxygen concentrations and lower total alkalinity compared to surface waters (Carson, 2008). Sea ice thickness in Ryder Bay rarely exceeds 0.5 m (Meredith et al., 2008), so we assume that the relatively thin first year ice was undergoing some exchange with surrounding sea water. Although sea ice porosity data are not available for Ryder Bay specifically, we know that sea ice in Marguerite Bay is comparatively porous due to relatively warm conditions (Fritsen et al., 2008), ice formation through the pancake ice cycle (Eicken, 1992; Thomas and Dieckmann, 2002) and subsequent deformation and snow-ice formation (Perovich et al., 2004). Our observations agree with the expected occasional
nutrient inputs from, and brine drainage to, surrounding sea water, and consequently less extreme environmental conditions in Ryder Bay sea ice than in more permanent sea ice environments (Gleitz et al., 1995; Kattner et al., 2004; Papadimitriou et al., 2007).

In comparison to organic carbon synthesised under closed system dynamics in multiyear ice (Papadimitriou et al., 2009), δ\(^{13}\)C\(_{\text{POC}}\) values found here in 2005/06 are somewhat lower, as would be expected for a semi-closed system setting. Conversely, sea ice δ\(^{13}\)C\(_{\text{POC}}\) in 2004/05 is more 13C-enriched than those of Papadimitriou et al. (2009) whilst [CO\(_{2}\text{(aq)}\)] was lower and significantly higher POC:chl \(a\) ratios compared well with values of 1262 ± 2276 found in intact ice floes (Kennedy et al., 2002). This suggests that 2004/05 sea ice, albeit short-lived, was closer to a closed system, but this is contested by lower δ\(^{13}\)C\(_{\text{CO}_2}\) and is not thought to be representative of prevailing sea ice conditions in Ryder Bay.

The prevailing semi-closed system dynamics at work in Ryder Bay sea ice explain higher δ\(^{13}\)C\(_{\text{POC}}\) than in surface waters, since photosynthesis is partially carbon-limited. We find no good correlation between δ\(^{13}\)C\(_{\text{POC}}\) and ambient [CO\(_{2}\text{(aq)}\)] \(r^2 = 0.200; p = 0.511\) (Fig. 8) due to occasional exchange with seawater and therefore ambient \(\text{CO}_2\) is not necessarily representative of \(\text{CO}_2\) assimilated during POC synthesis. δ\(^{13}\)C\(_{\text{CO}_2}\) is also not necessarily in equilibrium with δ\(^{13}\)C\(_{\text{POC}}\) sampled at the same time and may instead reflect recent replenishment of isotopically light \(\text{CO}_2\). This would explain why sea ice δ\(^{13}\)C\(_{\text{CO}_2}\) shows no response to the preferential biological uptake of \(\text{C}^{13}\), which drives enrichment of δ\(^{13}\)C\(_{\text{POC}}\), and its utility for describing sea ice processes is therefore limited.

In addition to biological production, nutrient drawdown and isotopic enrichment in the semi-closed sea ice ecosystem, relatively high sea ice δ\(^{13}\)C\(_{\text{POC}}\) may be influenced by some species utilising HCO\(_3^-\) as \(\text{CO}_2\) becomes limiting (Papadimitriou et al., 2009). This would increase δ\(^{13}\)C\(_{\text{POC}}\) in the same way as discussed for surface waters and would impact on δ\(^{13}\)C\(_{\text{DIC}}\) rather than specifically δ\(^{13}\)C\(_{\text{CO}_2}\). \(\text{CO}_2\) degassing and carbonate mineral precipitation due to \(\text{CO}_2\) saturation or supersaturation in brine inclusions upon sea ice formation may further affect δ\(^{13}\)C\(_{\text{CO}_2}\) (Romanek et al., 1992; Papadimitriou et al., 2003, 2007) and therefore δ\(^{13}\)C\(_{\text{POC}}\). However, low \(\text{CO}_2\) concentrations in Ryder Bay sea ice make this scenario unlikely and occasional flushing by seawater would overprint any small contribution of these processes to δ\(^{13}\)C\(_{\text{CO}_2}\).

Post-production degradation processes may also contribute to higher δ\(^{13}\)C\(_{\text{POC}}\) in sea ice since \(\text{C}^{12}\) is preferentially degraded, leaving the remaining organic carbon enriched in \(\text{C}^{13}\). This is in agreement with higher POC:chl \(a\) ratios found here than in previous studies (Gosselin et al., 1990; Lizotte and Sullivan, 1992), due to additional detrital and non-algal carbon from grazing activity and high retention (Daly, 1990; Gleitz and Thomas, 1993; Bentaleb et al., 1998). Active degradation in sea ice is also consistent with the high POC:P\(_N\) ratios (Fig. 4), as organic nitrogen is preferentially degraded over carbon-bearing compounds (Rosenfeld, 1981; Hedges et al., 1986; Ganeshram et al., 1999). However, POC:P\(_N\) > 10 is common for sea ice microalgae and often implies nitrate-deprived algal metabolism (Gleitz and Thomas, 1993 and references therein), as we would expect from a semi-closed system setting. High POC:P\(_N\) could also be explained by the influence of exopolymeric substances produced by diatoms and bacteria, which are abundant in the Antarctic marine environment, especially in sea ice (Meiners et al., 2004; Mancuso Nichols et al., 2005), and so may not be diagnostic of post-production degradation alone. We cannot use δ\(^{13}\)C\(_{\text{CO}_2}\) to confirm whether in situ degradation is an important influence on sea ice δ\(^{13}\)C\(_{\text{POC}}\), because the aforementioned exchange with isotopically light seawater \(\text{CO}_2\) would mask any δ\(^{13}\)C\(_{\text{CO}_2}\) depletion that would accompany preferential degradation of organic \(\text{C}^{12}\). Proboscia species were found in only one sea ice sample, but abundance was negligible, therefore we observe no P. inermis control on sea ice δ\(^{13}\)C\(_{\text{POC}}\) such as we demonstrate in surface waters. However, unlike surface waters, least-squares linear regression analysis of sea ice brine samples shows a good relationship between SA:V and δ\(^{13}\)C\(_{\text{POC}}\) \(r^2 = 0.713, p = 0.101, n = 4\); Fig. 9), which becomes statistically significant when we include early season surface water samples thought to be dominated by sea ice material \(r^2 = 0.761, p = 0.0146, n = 6\). However, given the difference in δ\(^{13}\)C\(_{\text{POC}}\) between sea ice and water samples with similar SA:V ratios, as well as different SA:V in sea ice versus water samples with similar δ\(^{13}\)C\(_{\text{POC}}\) values, the effect of diatom SA:V on \(\varepsilon\)\(_p\) alone is unable to account for higher δ\(^{13}\)C\(_{\text{POC}}\) in sea ice than surface waters.

In summary, higher δ\(^{13}\)C\(_{\text{POC}}\) in sea ice than surface waters is likely attributable to a higher degree of \(\text{CO}_2\) utilisation due to the semi-closed nature of the sea ice ecosystem. Post-production degradation of organic material, direct HCO\(_3^-\) uptake by some sea ice diatoms and possible production of exopolymeric substances may further contribute to isotopically heavy sea ice-derived organic material.

### 4.4 Sinking particulate organic carbon

Sinking particulate δ\(^{13}\)C\(_{\text{POC}}\) time-series data (Fig. 7) show similar features to the surface water time-series (Fig. 3), suggesting that although P. inermis was not dominant in sediment traps, the associated δ\(^{13}\)C\(_{\text{POC}}\) signatures produced in surface waters are transferred to depth. Down-depth trends in seasonal average δ\(^{13}\)C\(_{\text{POC}}\) through the water column to sediment core-top material (Fig. 11) show that Ryder Bay sinking particulate matter is more depleted in δ\(^{13}\)C\(_{\text{POC}}\) in 2005/06 than 2004/05, consistent with much lower season-average surface water δ\(^{13}\)C\(_{\text{POC}}\) in 2005/06 (~24.5 ‰ vs. ~20.0 ‰). This is in response to the large and prolonged late-season negative δ\(^{13}\)C\(_{\text{POC}}\) shift, which is observed in both the
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200 m and 512 m sediment traps, albeit approximately one month later (Fig. 7). Although Marguerite Bay sediment trap δ13CPOC is also lower in 2005/06 than 2004/05, the signal is much more pronounced in Ryder Bay, suggesting that low δ13CPOC related to *P. inermis* dominance is a localised phenomenon.

Sinking δ13CPOC is always higher in Marguerite Bay than Ryder Bay at any given time. However, within each season at both sites, δ13CPOC in the deepest trap is within 0.4‰ of its surface water (Ryder Bay) or shallow trap (Marguerite Bay) counterpart. The only exception is Marguerite Bay in 2005/06, where shallow and deep trap values fall within 2‰. This clear relationship between δ13CPOC in surface waters and sediment traps provides evidence that surface ocean δ13CPOC signatures are faithfully exported to depth in the water column, even despite the loss of key diatom species during sinking.

Sediment core-top δ13CPOC in Ryder Bay is slightly higher than in the deepest sediment trap (Fig. 11), likely because of minor sedimentary remineralisation or due to the fact that surface sediments integrate δ13CPOC signatures over longer time scales. However, enrichment of core-top δ13CPOC relative to deep trap δ13CPOC is close to error and so we suggest that δ13CPOC of sinking particles is reliably transferred to marine sediments.

### 4.5 Potential implications for δ13CPOC in Southern Ocean sediments as a paleoceanographic proxy

Results presented in this study hold important implications for the use of sedimentary δ13CPOC as a proxy for past environmental conditions in the coastal Southern Ocean. Marine sedimentary records show glacial Southern Ocean δ13CPOC to be approximately 4‰ depleted relative to interglacial epochs (Singer and Shemesh, 1995; Rosenthal et al., 2000; Crosta and Shemesh, 2002; Schneider-Mor et al., 2005). Traditionally, low glacial δ13CPOC was explained by higher [CO2(aq)] due to strengthening of the thermohaline circulation and wide-spread enhanced upwelling (Rau et al., 1992; Singer and Shemesh, 1995). However, later studies contradict this upwelling theory by using other proxy records such as δ15Norg & Ba/Al to infer a stratified glacial Southern Ocean and reduced productivity (François et al., 1997).

The anti-correlation of low glacial δ13CPOC and high glacial δ15Norg can be reconciled by increased stratification restricting nitrate supply and increasing δ15Norg and sea ice cover preventing ocean-atmosphere gas exchange so that [CO2(aq)] remains high and δ13CPOC low (Crosta and Shemesh, 2002).

We have shown that seasonal changes in diatom assemblages can drive short-lived yet large isotopic transitions in coastal Antarctic surface waters and have a profound impact on seasonal average δ13CPOC exported to depth and underlying sediments. Most importantly, seasonal average δ13CPOC for a season of well-mixed conditions is 4‰ higher than a much more stratified season preceded by a heavy sea ice winter, such as may have been typical of glacial times. The 4‰ difference matches the full amplitude of the glacial-interglacial offset in δ13CPOC from Southern Ocean sediment cores.

We hypothesise therefore that diatom species shifts may be an important driver of lower glacial δ13CPOC in the Southern Ocean, in agreement with Jacot Des Combes et al. (2008). Whilst it is *P. inermis* that appears to be driving large isotopic shifts in this study, we do not specifically invoke this species as a driver of δ13CPOC over glacial-interglacial cycles. Other diatom species employing similar unusual biochemistry may be important contributors to low glacial δ13CPOC. Although sedimentary diatom assemblages do not show such drastic changes as witnessed in this study (Gersonde and Zielinski, 2000; Bianchi and Gersonde, 2004) and
there is no evidence for significant changes in Proboscia species in the open ocean on glacial-interglacial timescales (Crosta et al., 2004), this does not preclude a species control on low glacial δ¹³C_POC. Instead, the species responsible for low glacial δ¹³C_POC may not be well preserved in sediments, whilst its isotopic signature is preserved, as is demonstrated here for P. inermis.

With these caveats in mind, we demonstrate that changes in surface water diatom assemblages can drive shifts in seasonal average δ¹³C_POC of equal amplitude to the 4‰ glacial-interglacial δ¹³C_POC offset observed in marine sedimentary records. We show that these surface water isotopic shifts are transferred to marine sediments and we propose therefore that at least part of the lower glacial δ¹³C_POC signal may be due to changes in diatom assemblages. If the more glacial-type conditions of heavier winter sea ice and upper water stratification in 2005/06 were responsible for driving a shift to diatom species characterised by lower isotopic signatures, in this case P. inermis, then it follows that species compositional shifts may be a significant influence on δ¹³C_POC on glacial-interglacial timescales. Further studies are required to elucidate the processes underlying this relationship.

5 Conclusions

This study presents a unique insight into the factors affecting δ¹³C_POC in the coastal Antarctic sea ice environment. In agreement with previous studies, we find higher δ¹³C_POC in sea ice brine relative to surface waters, consistent with autotrophic carbon fixation in a semi-closed environment. Possible secondary effects on sea ice δ¹³C_POC may result from biological utilisation of HCO₃⁻ in addition to CO₂ as a carbon substrate, production of exopolymeric substances and/or post-production degradation of organic matter within the ice matrix. Sea ice-derived organics exert a short-lived impact on surface water δ¹³C_POC in Ryder Bay due to brine drainage processes whilst sea ice is present. Isotopically heavy sea ice material tends to sink quickly, so may be preserved more effectively in the sedimentary record and may consequently bias the overall δ¹³C_POC signal in marine sediments.

We demonstrate that [CO₂(aq)] and δ¹³C CO₂ are not the primary factors controlling variations in δ¹³C_POC in surface waters in the Antarctic sea ice environment. Instead, we argue that ~10‰ negative excursions in surface water δ¹³C_POC are driven by seasonal shifts in diatom assemblages, in this case specifically to dominance of P. inermis. While the exact mechanisms remain unknown, we postulate that P. inermis may modify δ¹³C_POC through its internal cell biochemistry and lack of a CCM, whilst other species present at different times in the growing seasons do employ CCMs. Consequently, seasonal species-related changes in ε_p further complicate the relationship between δ¹³C_POC and [CO₂(aq)].

Finally, sediment trap data indicate that although much of the surface suspended material, including certain diatom species, undergoes recycling in the upper ocean and is not exported to depth, the δ¹³C_POC signal is transferred to depth in the water column by sinking particles. Further, we show how isotopic signatures in these sinking particles are transferred to marine sediments unaltered. This study therefore identifies the importance of seasonal changes in surface water diatom speciation and isotopically heavy sea ice-derived material for δ¹³C_POC signatures in Antarctic coastal environments and underlying sediments, and thus highlights the need for analysis of species-specific or diatom-bound δ¹³C_POC in order to reliably interpret sedimentary δ¹³C records.
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References


Crosta, X., Crespin, J., Billy, I., and Ther, O.: Major factors con-


