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Water column distribution and carbon isotopic signal of cholesterol, brassicasterol and particulate organic carbon in the Atlantic sector of the Southern Ocean

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Abstract. The combination of concentrations and δ^{13} C signatures of Particulate Organic Carbon (POC) and sterols provides a powerful approach to study ecological and environmental changes in both the modern and ancient ocean. We applied this tool to study the biogeochemical changes in the modern ocean water column during the BONUS-GoodHope survey (February–March 2008) from Cape Basin to the northern part of the Weddell Gyre. Cholesterol and brassicasterol were chosen as ideal biomarkers of the heterotrophic and autotrophic carbon pools, respectively, because of their ubiquitous and relatively refractory nature.

We document depth distributions of concentrations (relative to bulk POC) and δ^{13} C signatures of cholesterol and brassicasterol combined with CO_2 aq. surface concentration variation. While the relationship between CO_2 aq. and $\delta^{13}C$ of bulk POC and biomarkers have been reported by others for the surface water, our data show that this persists in mesopelagic and deep waters, suggesting that δ^{13} C signatures of certain biomarkers in the water column could be applied as proxies for surface water CO₂ ag. We observed a general increase in sterol δ^{13} C signatures with depth, which is likely related to a combination of particle size effects, selective feeding on larger cells by zooplankton, and growth rate related effects. Our data suggest a key role of zooplankton fecal aggregates in carbon export for this part of the Southern Ocean (SO). Additionally, in the southern part of the transect south of the Polar Front (PF), the release of seaice algae during the ice demise in the Seasonal Ice Zone (SIZ) is hypothesized to influence the isotopic signature of sterols in the open ocean. Overall, the combined use of δ^{13} C values and concentrations measurements of both bulk organic C and specific sterols throughout the water column offers the promising potential to explore the recent history of plankton and the fate of organic matter in the SO.

1 Introduction

The intensity of organic matter (OM) export combined with the efficiency of deep water heterotrophic reprocessing of this material sets the sequestration's efficiency of the oceanic biological carbon pump (Honjo et al., 2008; Boyd and Trull, 2007; Battle et al., 2000). However, it appears difficult to balance the organic C demand by twilight zone (currently defined as the depth interval between 100-1000 m) heterotrophs with the export flux from the upper mixed layer

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(Burd et al., 2010; Steinberg et al., 2008; Reinthaler et al., 2006; Arístegui et al., 2005). Therefore it is essential to better understand the processes controlling the export and reprocessing of exported OM (Boyd and Trull, 2007). Gaining information about the sources and fate of sinking and suspended biogenic particles is necessary to improve our knowledge of the processes occurring below the euphotic layer, where the attenuation of the export flux is the strongest. One way to improve our understanding about mechanisms controlling interrelated biogeochemical processes involved in carbon export and reprocessing is to characterize the chemical composition of C-carrier phases, such as particles sinking rapidly through the water column as well as suspended fine particles with longer residence time (Wakeham et al., 2009).

As part of the Southern Ocean GEOTRACE program sampling opportunity, this study explores the fate of organic matter in the water column for the Atlantic sector of the Southern Ocean. Further dataset acquisition on the global ocean will help to reach a general picture using such type of information since, to the best of our knowledge, no dataset like this one is available on the ocean water column. In the following we focus our work on the Southern Ocean area.

While several studies have examined the δ^{13} C values of the bulk particulate organic carbon (POC) in the Southern Ocean (Lourey et al., 2004; Trull and Armand, 2001; Popp et al., 1999; Bentaleb et al., 1998; Dehairs et al., 1997; Rau et al., 1997; Kennedy and Robertson, 1995; François et al., 1993), fewer studies have focused on specific compounds. O'Leary et al. (2001), Tolosa et al. (1999) and Popp et al. (1999) report on the factors controlling the carbon isotopic composition of phytoplankton based on the study of δ^{13} C_{POC}, δ^{13} C_{sterols} and δ^{13} C_{phytol} values. These three studies, however, are mainly limited to an investigation of the surface ocean, so the distribution of these biomarkers in deeper waters remains unknown. To the best of our knowledge, no information is available on specific compounds and their isotopic composition for the intermediate and deep Southern Ocean. However, identifying the food web components controlling the C flux in the deeper water column and understanding compositional changes of suspended and sinking organic material is important to better constrain the fate of organic matter exported to the deep ocean. We investigated this by studying the variability in concentration of individual compounds and their ¹³C isotopic composition.

While cholesterol $(27\Delta^5;$ cholest-5-en-3 β -ol) is produced by some algae (Volkman, 2003, 1986) it is commonly considered as a biomarker for consumer organisms and a proxy for zooplanktonic herbivory (Grice et al., 1998). Schouten et al. (1998) propose cholesterol as a general biomarker for the eukaryotic marine community, thereby avoiding the complexity inherent to species-specific sterols that have different isotopic offsets relative to phytoplankton δ^{13} C value. A general biomarker averages out the effects due to variability of biosynthetic pathways, cell size, geometry, and growth rate between individual species. Grice et al. (1998) show via

controlled mesocosm experiments that grazers convert different algal precursor sterols into cholesterol without significant isotopic fractionation. Moreover, Chikaraishi (2006) in accordance with Grice et al. (1998) reports that no substantial carbon isotopic fractionation occurs during either heterotrophic sterol assimilation or de novo synthesis.

Schouten et al. (1998) state that the refractory nature and ubiquitous character of cholesterol as an eukaryotic marker should favor the integrity of the $\delta^{13}C_{cholesterol}$ signatures when particles sink to greater depth, thereby preserving the information acquired in surface waters. Brassicasterol (28 $\Delta^{5,22}$: 24-Methylcholesta-5,22E-dien-3 β -ol) is reported in a large number of algal classes (Volkman, 2003; 1986) and is considered as a strict phytosterol, i.e. it cannot be biosynthesized by zooplankton. For these reasons, it is commonly used as an indicator of marine algae and of diatoms, in particular. Recently, Rampen et al. (2010) stressed that when diatoms dominate the phytoplankton community, sterols, and brassicasterol in particular (abundant in the pennate diatoms), provide useful information on the type of diatoms that are present. Therefore, for the Southern Ocean where diatoms are dominant but where other phytoplankton groups also contribute to the primary production, we will consider brassicasterol as a strict phytoplankton indicator. Because of their ubiquitous and relative refractory nature (e.g. Volkman, 1986, 2003), cholesterol and brassicasterol were thus chosen in this study as ideal markers of the heterotrophic and autotrophic carbon pools that have potential for being sequestered in the deep ocean.

As a part of a broader study on the biogeochemistry and C fluxes along the Greenwich Meridian, we document variations in concentration (relative to bulk POC) and δ^{13} C compositions of cholesterol and brassicasterol over the whole water column from the Cape Basin to the northern Weddell Gyre. Combined with data on CO₂ aq. concentrations, our results furthermore enable us to verify earlier surface water observations about CO₂ substrate concentration and C isotopic composition of bulk POC and biomarkers. They also document to what extent this substrate-dependent C isotopic signature, acquired in the surface, is preserved deeper in the water column. This is important since particles may, or may not, become isotopically homogenous at depth as a result of mixing, depending on magnitude of sinking velocity relative to advection. This leads to answering the question whether the system is operating in 1-D (strong surface to deep links) or 3-D (strong mixing and homogenization in the deep ocean) configuration.

2 Materials and methods

2.1 Sampling and hydrology

The "Biogeochemistry in the Southern Ocean: Interactions between Nutrients, Dynamics and Ecosystems Structures"

(BGH) expedition took place in the South East Atlantic and the Atlantic sector of the Southern Ocean during summer 2008 (February-March 2008) on board R/V Marion Dufresne (voyage track shown in Fig. 1). The BGH project in the region south of South Africa was built on two main topics related to (1) the large scale inter-ocean exchanges and (2) the characterization of the biogeochemical processes involved in the internal cycling of trace elements, isotopes and carbon. South of Africa, the Southern Ocean (SO) provides the export channel for North Atlantic Deep Water (NADW) to the global ocean and the passage for heat and salt water from the Indian and Pacific oceans to the Atlantic Ocean. The eastward flowing Antarctic Circumpolar Current (ACC), the South Atlantic Current and NADW meet with the westward flow of Indian waters carried by the Agulhas Current, leading to water mass exchanges through jets, meanders, vortices, and filament interactions. Indeed, the Agulhas Current is the major western boundary current of the Southern Hemisphere (Lutjeharms, 2006) and a key component of the global ocean "conveyor" circulation controlling the return flow to the Atlantic Ocean (Gordon, 2003, 1986). Unusual dynamics pervade the motion of this warm-water current: as it moves west around the Southern tip of Africa, it is reflected back east by the ACC. Not all waters are captured by this sudden diversion to the Indian Ocean and parts of the Agulhas Current leak away into the South Atlantic Ocean (BGH cruise report). However, the ACC fronts (Fig. 1) represent almost impermeable barriers delimiting zones with relatively constant hydrological and biogeochemical properties, and lower baroclinic transport (Sokolov and Rintoul, 2007). Cross frontal exchanges occur mainly locally and close to sharp topographic features. Water particles spend several years in each zone before being advected northward by Ekman pumping, and subsequently take part in several winter convective mixings (S. Speich, personal communication, 2009).

Amongst the sites studied during BGH, five sites had sufficient station time (about 48 h) to enable whole water column suspended matter sampling with large-volume in situ filtration systems (LVFS) because this operation is time consuming (6 to 8 h per deployment). Up to a maximum of 11 LVFS units were deployed covering the entire water column from the surface to the deep ocean. The five sites were selected to cover the major zonal systems framed by fronts (Fig. 1). The complete salinity and temperature sections during BGH are available in Fig. 1.

Suspended particulate organic matter was sampled using LVFS (Challenger Oceanics and McLane WTS6-1-142LV systems) fitted with 142 mm diameter filters holders. Water was pumped through two successive filters: (i) a 53 μ m mesh nylon screen (filter SEFAR-PETEX ; polyester) and (ii) a QM-A quartz fiber filter (\sim 1 μ m porosity, Pall Life). Total volume of seawater filtrated by LVFS ranged between 152 L (surface water) and 1983 L (deep water). Prior to use, PETEX screens were conditioned by soaking in HCl 5 %, rinsed

with Milli-Q grade water, dried at ambient temperature under a laminar flow hood and were stored in clean plastic zipbags till use. QMA filters were conditioned by combustion at $450\,^{\circ}\text{C}$ for $4\,\text{h}$ and were stored in clean plastic zip-bags till use.

Since the suspended matter samples were also intended for analysis of ²³⁴Th (Planchon et al., 2013), filters were partitioned among the different end-users, using sterile scalpels and a custom-build INOX steel support for the PETEX screens and a plexiglass punch of $\emptyset = 25.3$ mm for the QMA filters. These operations were conducted under a laminar flow hood. Whenever possible, both size fractions ($\sim 53 \,\mu m$ and 53–1 μm) were sampled for biomarker analysis. This was in general the case for the upper ocean mixed layer. Below the mixed layer, the large particle fraction (> 53 µm) could not be recovered because screens carried too little material. Aliquots dedicated to compound specific isotope analysis (CSIA) were packed in cryotubes and stored at -80 °C till processing in the home-laboratory. Aliquots dedicated to the analysis of $\delta^{13}C_{POC}$ were dried at 50 °C and stored in Petri dishes at ambient temperature till processing in the home laboratory.

2.2 $\delta^{13}C_{POC}$: sample preparation and analysis

POC concentrations and $\delta^{13}C_{POC}$ were analyzed via elemental analyzer - isotope ratio mass spectrometer (EA-IRMS). Prior to this, inorganic carbon (carbonates) was removed by exposing the filters to concentrated HCl vapor inside a closed-glass container for 4 h (Lorrain et al., 2003). After drying at 50 °C, the samples were packed in silver cups and analyzed with a Thermo Flash 112 elemental analyzer configured for C analysis and coupled on-line via a Con-Flo III interface to a Thermo-Finnigan Delta V IRMS. Acetanilide and IAEA-CH-6 reference materials were used for calibrating concentrations and isotopic composition, respectively. Using blank procedure, the detection and quantification limits for POC were measured as lower than 0.01 µM for 100 L. This is significantly lower than the concentrations measured from the surface to the deep waters for large and small particles.

2.3 $\delta^{13}C_{\text{sterols}}$: sample preparation and analysis

Samples were processed following the method described in Boschker (2004). Briefly, total lipids were extracted using a modified method from Bligh and Dyer (1959) with chloroform/methanol/Milli-Q water $(v:v,\ 15:15:6\ \text{mL})$ mixture. Usually, 70 to 80% of the chloroform phase containing the total lipid fraction was recovered, implying that true concentrations are 20 to 30% underestimated (Boschker, 2004). In the Supplement (Annex 1) concentrations for cholesterol and brassicasterol are not corrected from the 20 to 30% underestimation. This approach does not influence the observed sterol concentration ratios and the patterns we find

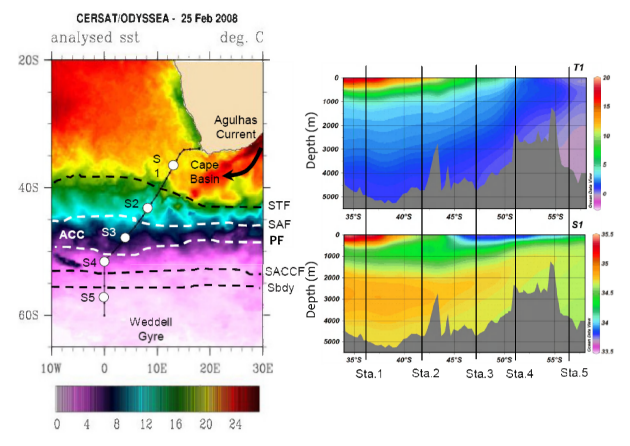


Fig. 1. Left part of panel = BONUS-GoodHope voyage track and station locations (S1 at $36^{\circ}31'$ S; S2 at $42^{\circ}28'$ S; S3 at $47^{\circ}33'$ S; S4 at $51^{\circ}52'$ S; S5 at $57^{\circ}33'$ S). The background is sea surface temperature SST (°C) from the CERSAT/ODYSSEA satellite. A SST scale is given at the bottom of the image. The SST north-south gradient is typical of the region in summer. Approximate location of ACC fronts in summer (STF = Sub-Tropical Front, SAF = Sub-Antarctic Front North, PF = Polar Front, SACCF = South Antarctic Circumpolar Current Front, Sbdy = southern boundary of the ACC) are shown. The area north of the STF is the Sub-Tropical Zone (STZ); the Sub-Antarctic Zone (SAZ) is located between STF and SAF; the Polar Front Zone (PFZ) is located between the SAF and PF; the Antarctic Zone (AZ) is located between the PF and Sbdy, and south of the Sbdy is identified as the northern part of the Weddell Gyre (WG). Right part of panel = Complete salinity S1 PSU (lower panel) and temperature T1 (°C, upper panel) sections during BGH (from BGH cruise report; courtesy from S. Speich). Station locations are shown.

in the ratios, if we assume that the degree of underestimation is similar for various sterols. Since the final standard error on the $\delta^{13} C$ -cholesterol and $\delta^{13} C$ -brassicasterol measurements is estimated to range between 1.1 and 2.0% (Supplement Annex 1), and because the total lipid extraction step implies permanent sample shaking to homogenize and optimize recovery of the extract, we assume that the isotopic bias due to incomplete recovery is negligible compared to the actual accuracy and precision of the measurement.

The total lipid extract was then separated into neutral, glyco-, and polar lipids on silica chromatographic column (0.5 g Kieselgel 60; Merck): (i) the neutral phase was eluted with chloroform (7 mL), (ii) the glycolipidic phase with acetone (7 mL), and (iii) the polar phase with methanol (10 mL) (Boschker, 2004). The glycolipidic and the polar phases were stored at $-20\,^{\circ}\mathrm{C}$ while the neutral phase was immediately processed for analysis.

Neutral phases were completely dried under gentle inert N_2 flow (to avoid degradation) and a known quantity of squalane used as internal standard (IS, i.e., a stable compound which does not co-elute with natural compounds present in the samples) was added. To estimate compound concentrations in the samples, we used as reference the GC-c-IRMS peak area of squalane.

To increase volatility and enhance thermal stability of sterols for their separation by gas chromatography (e.g. Lagarda et al., 2006), dried neutral lipids were derivatized using bis-(trimethylsilyl)-trifluoroacetamide (BSTFA, 99 %)/toluene ($v:v,\ 1:1$). Trimethylsilyl (TMS) groups were substituted to the hydrogen of the hydroxyl groups of the compounds to form trimethylsyliloxy groups [-O-Si-(CH₃)₃]. In practice, derivatization was achieved by adding 50 µL of BSTFA-toluene to the dried samples (+ IS) and heating during 60 min at 60 °C.

All the TMS-derivatized neutral fractions (+ IS) were analyzed using a Trace GC Ultra coupled to Trace Plus MS (Interscience) for compound identification and a Thermo Finnigan GC equipped with a combustion furnace (CuO/NiO/Pt reactor at 940 °C) coupled to a Delta*Plus* XL isotope ratio mass spectrometer (IRMS) for δ^{13} C and concentration measurements. Compound identification was achieved by (i) retention time matching between standard mix and sample, and (ii) GC-MS mass spectrometry characterization (matching with Goad and Akihisa, 1997).

Both gas chromatographs were equipped with similar capillary columns (DB-5-type (J&W Scientific), $30\,\mathrm{m}\times0.32\,\mathrm{mm}$ i.d. with $0.25\,\mu\mathrm{m}$ film thickness). Identical temperature programming was used for both GC ovens starting from $50\,^\circ\mathrm{C}$ (2 min) to $300\,^\circ\mathrm{C}$ (15 min) with a ramp of $4\,^\circ\mathrm{C}$ min $^{-1}$. Carrier gas flow (He) was $2\,\mathrm{mL\,min}^{-1}$, and injection occurred in splitless mode at $300\,^\circ\mathrm{C}$. Blanks (hexane) and mixtures of standards were regularly measured in between sample analyses (standard bracketing) to check the stability of the systems and absence of potential contaminants.

The carbon isotopic composition of squalane (internal standard), treated similarly as the samples (extracted neutral phase), was used to check reproducibility and accuracy of the GC-c-IRMS system. Aliquots of 0.5 mg squalane powder were weighed into tin capsules (IVA, 3.3×5 mm) and analyzed via EA-IRMS (Thermo) using IAEA-CH₆ as reference material. A good agreement was observed between the mean $\delta^{13}C_{\text{squalane}}$ obtained via EA-IRMS (δ^{13} C_{squalane} = $-20.2 \pm 0.1 \%$, 1σ , n = 3) and GC-c-IRMS (δ^{13} C_{squalane} = -21.2 ± 0.9 %, 1σ , n = 69). Various amounts of squalane were measured to evaluate the impact of signal size or linearity of the GC-c-IRMS. From a range of signal intensity being 177-3406 mV, no significant trend is observed ($y = -0.0002 \times -20.917$; $R^2 = 0.0356$; n = 69; p > 0.05; not shown). It can thus be concluded that, in case of this specific IRMS system and tuning, low signal intensity does not impact the measured δ^{13} C. For this study, the cutoff limit considered adequate to accept or reject δ^{13} C-sterols results was set at 100 mV.

Due to the addition of a trimethylsilyl group to each individual compound (TMS-derivatization), the δ^{13} C of cholesterol ($27\Delta^5$; cholest-5-en-3 β -ol) and brassicasterol ($28\Delta^{5,22}$; 24-Methylcholesta-5,22E-dien-3 β -ol) obtained from GC-c-IRMS needs to be corrected. Four standards – cholesterin- α 24, stigmasterol- α 22, β -sitosterol, and 7-dehydrocholesterol – are used for this purpose. 5 α -cholestane is used as a fifth standard to test GC-c-IRMS accuracy. Comparison between mean δ^{13} C measured by GC-c-IRMS on derivatized (n=3) and non-derivatized (n=3) standards' mix is achieved using a functional relationship estimation by maximum likelihood (FREML – Ripley and Thompson, 1987; AMC technical brief report no. 10, 2002) (Supplement Annex 2) via Eq. (1):

$$\delta^{13}C_{\text{sterol}} = \frac{(n+3) \times \delta^{13}C_{\text{TMS-sterol}} - 3 \times \delta^{13}C_{\text{BSTFA}}}{n}, \quad (1)$$

where $\delta^{13}\mathrm{C}_{\mathrm{sterol}}$ is the measured carbon isotopic signature of non-derivatized standard sterol, n is the number of carbon atoms of the sterols without derivatization, $\delta^{13}\mathrm{C}_{\mathrm{TMS-sterol}}$ is the measured carbon isotopic signature of derivatized standard sterol and $\delta^{13}\mathrm{C}_{\mathrm{BSTFA}}$ is the measured carbon isotopic signature of BSTFA-toluene used to derivatize sterols.

Furthermore, since no certified reference materials for compound-specific stable isotope measurements on sterols exist, we compare mean $\delta^{13}\mathrm{C}$ measured by GC-c-IRMS and EA-IRMS of the same non TMS-derivatized compounds (n=3) and include it in the $\delta^{13}\mathrm{C}_{\text{sterol}}$ correction following a FREML equation as described above (Eq. 1 applied for non-derivatized standard sterols analyzed either via EA-IRMS or via GC-c-IRMS) (Supplement Annex 2).

The final δ^{13} C value for each compound is thereby corrected for isotopic deviation from TMS-derivatization, as well as possible isotopic deviation from GC-c-IRMS compared to EA-IRMS. The uncertainty on final δ^{13} C of cholesterol (cho) and brassicasterol (bra) ($\pm 1\sigma$; Supplement Annex 1) is calculated by propagating standard deviations from triplicate measurements (i.e. 1σ) and correction from derivatization.

3 Results

3.1 Contents of cholesterol, brassicasterol and POC-bulk

Results for POC (large and small particles), brassicasterol and cholesterol concentrations in the small (1–53 $\mu m)$ and large (> 53 $\mu m)$ particle size fractions are available in the Supplement (Annex 1). Brassicasterol and cholesterol contents are of similar magnitude (10 \times 10 1 ng L^{-1}). Cholesterol was usually slightly less abundant than brassicasterol in the upper 200 m (Fig. 2), but was detectable over the entire water column, while brassicasterol was undetectable in deeper water at the most northward stations S1 and S2 (i.e. peak amplitude below the GC-c-IRMS/GC-MS detection limit fixed at 100 mV).

Vertical profiles of large particle POC (LP-POC) are shown relative to total POC (T-POC = LP-POC + SP-POC, where SP-POC is small particle POC) to highlight specific features of the profiles (Fig. 2, upper panel). We also plotted the parameters vs. density rather than vs. depth, which allows for a better discernment of the variations in the surface and mesopelagic waters. Vertical profiles of small particle cholesterol (SP-cholesterol) and brassicasterol (SP-brassicasterol) are shown relative to SP-POC (Fig. 2, lower panel) following the same rationale.

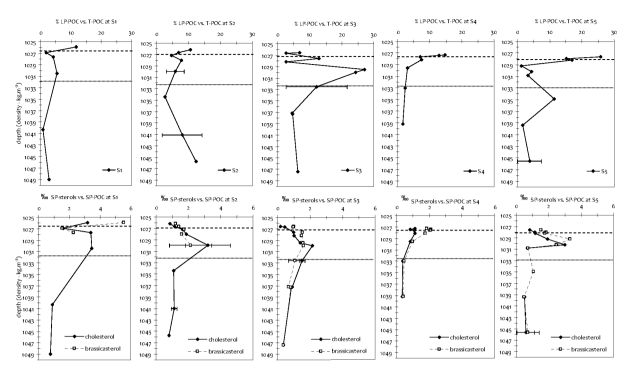


Fig. 2. Upper panel: profiles of large particle (> $53 \,\mu m$) POC content over total POC content (in %). Lower panel: profiles of small particle (1– $53 \,\mu m$) cholesterol and brassicasterol content over small particle POC content (in %). Y-axis depth is shown as seawater density; the 100 m (dashed line) and 1000 m (dotted line) depth horizons are shown. The standard error is estimated as \pm 30 % for brassicasterol and cholesterol. Error bars shown represent the average standard deviation ($\pm 1\sigma_{SD}$) for two close-by samples (see also Supplement Annex 1).

3.1.1 General features

Relative contents of LP-POC and SP-sterols changed considerably in the surface waters along the section. Overall, the contributions of LP-POC ranged from 0.8 to 27.0 % of total POC, with highest values at S3 (27.0%) in the Polar Frontal Zone (PFZ), and S5 (25.8%) in the northern Weddell Gyre. LP-POC values often displayed a maximum in mesopelagic to deep waters. This maximum was clearly visible at S1 (750 m) and S3 (450 m) but less prominent at S2 (250 m) and absent at S4 (Fig. 2, upper panel). At S5, an LP-POC maximum was present at 1500 m, below the mesopelagic layer. With the exception of station S4, a systematic increase of LP-POC over T-POC ratio was apparent near the seafloor (possibly indicating the nepheloïd layer), ranging between 2.7% (station S1) and 12.3% (station S2). The large error bars (up to 100%) in Fig. 2 reflected the large variability between close-by samples (see also Supplement Annex 1), reflecting heterogeneity of the system (close-by depths were often sampled with an interval of 24 h) and variability due to sampling method.

Profiles of relative SP-cholesterol and SP-brassicasterol contents (Fig. 2, lower panel) were also characterized by mesopelagic maxima, except at S1 where brassicasterol below 750 m was below detection limit and at S4 where no mesopelagic maximum was present. The depths of

mesopelagic maxima for cholesterol and brassicasterol were rather similar between sites, and ranged between circa 200 and 800 m (Fig. 2 lower panel; see Supplement Annex 1 for depth meter vs. density correspondence). The highest brassicasterol contents were observed in surface water at S1 and S5, in good accordance with the presence of the diatom pigment fucoxanthin (J. Ras, personal communication, 2008). Below the mesopelagic maximum the relative contents of cholesterol and brassicasterol decreased toward the seafloor (S1, S2, S3; Fig. 2 lower panel), or remained rather unchanged (S4, S5). In contrast to relative contents of LP-POC, no increase of brassicasterol and cholesterol contents was observed near the seafloor.

3.1.2 Site specific features

The surface waters at station S1 showed the highest relative SP-brassicasterol content (5.5%; Fig. 2 lower panel) and also the highest concentration (65.3 ng $\rm L^{-1}$; Supplement Annex 1) of the whole dataset. This is in accordance with the observed high concentrations of Chl a and phaeopigments (Le Moigne et al., 2013). The profiles of relative SP-cholesterol and LP-POC contents at S1 were quite similar, with pronounced surface and broad mesopelagic maxima; the latter reached circa 100% of the surface content for cholesterol, and circa 40% for LP-POC.

Station S2, also located in the area of high Chl a and phaeopigment concentrations north of the Sub-Antarctic Front (SAF) (Le Moigne et al., 2013), showed LP-POC contents similar to S1. Here, surface water brassicasterol reached its second highest value (46.8 ng L $^{-1}$; see Supplement Annex 1) for the entire cruise, in agreement with the high chloropigments concentrations (J. Ras, personal communication, 2008). The mesopelagic maximum of relative cholesterol and brassicasterol contents, located at about 600 m, was pronounced and sharp, whereas relative content of LP-POC showed a weak mesopelagic maximum expanding from 200 to 1000 m (Fig. 2). At this site the increase of LP-POC towards the seafloor was the strongest (reaching up to 12 % of total POC) of the entire survey.

At S3 the mesopelagic LP-POC content was the highest of the entire section (Fig. 2). A clear broad mesopelagic maximum of LP-POC and SP-cholesterol content was present between 500 and 600 m. SP-brassicasterol showed a weak maximum expanding from the near-surface water to 600 m. Again, profiles of LP-POC and SP-cholesterol relative contents were quite similar.

Station S4 was located above the Atlantic-Indian ridge in an area with lowest Chl *a* and phaeopigment concentrations (Le Moigne et al., 2013). At this site, no mesopelagic maximum was observed and depth profiles of SP-sterols and LP-POC were quite similar. The maximum relative content of SP-sterols and LP-POC was meanwhile located closer to the surface, at 80 m.

Station S5 was characterized by deep reaching Chl $\it a$ concentrations (circa 150 m) and small sub-surface (100–125 m) phaeopigment concentration (Le Moigne et al., 2013). Surface waters showed the highest relative content of LP-POC (Fig. 2) and also the highest total POC concentrations (up to 25.8 μ g L $^{-1}$; see Supplement Annex 1) of the entire cruise. A second LP-POC maximum was present at 1500 m (Fig. 2). The relative contents of brassicasterol and cholesterol showed a sharp mesopelagic maximum, ranging between 300 to 500 m. Brassicasterol showed a second smaller maximum at 1500 m, coinciding with the LP-POC maximum.

3.2 δ^{13} C composition of POC, cholesterol and brassicasterol

The vertical profiles of $\delta^{13}C_{LP\text{-POC}}$, $\delta^{13}C_{SP\text{-POC}}$, $\delta^{13}C_{SP\text{-POC}}$, $\delta^{13}C_{SP\text{-cholesterol}}$ and $\delta^{13}C_{SP\text{-brassicasterol}}$ are shown in Figure 3 (for complete dataset, see Supplement Annex 1). Note that for S1 and S2 SP-brassicasterol data are limited to the upper water column, since concentrations fell below detection limit for the deeper water column. The range of $\delta^{13}C$ variation through the water column can reach up to 10 % (see for example the $\delta^{13}C_{SP\text{-cholesterol}}$ profile at station S2; Fig. 3). At stations S1 and S2, $\delta^{13}C$ profiles of sterols and bulk POC do cross over at depth, while at stations S3 to

S5 these remain clearly separated from each other over the entire water column.

At S1 and to a lesser extent at S2, $\delta^{13}C_{LP-POC}$ values decreased below 2750 m and converged toward the δ^{13} C_{SP-POC} values (Fig. 3). At these depths the water mass consisted of diluted North Atlantic Deep Water (NADW) (S. Speich, personal communication, 2008). This could be an indication of NADW transporting highly degraded organic matter constituted by isotopically light refractory lipidic material since it has been shown that proteins and carbohydrates (higher δ^{13} C isotopic signals than lipids, see Galimov, 2006) are more labile compared to lipids and are then removed preferentially from the particulate organic matter (POM) (Degens, 1969). Also, intense bacterial degradation of the more labile carbohydrates and proteins could explain this strong decrease in δ^{13} C-POC observed for large particles at S1 and S2. Stations S1 and S2 are defined by mixed phytoplanktonic community composed of dinoflagellates, chromophytes, nanoflagellates and cyanobacteria, and are systems supported to a large degree by regenerated production (Joubert et al., 2011). Observed variations of $\delta^{13}C_{SP-POC}$ and $\delta^{13}C_{LP-POC}$ along the S1 and S2 water columns could thus be linked to high degradation processes occurring in the surface water.

At S2 all $\delta^{13}C$ values in the upper 600 m decreased with depth, whereas below 600 m they increased (note that SP-brassicasterol was not detected below 600 m). Below 1500 m LP- $\delta^{13}C_{POC}$ decreased again, as observed at S1 (see above). At S3 below the surface waters, $\delta^{13}C_{LP-POC}$ and $\delta^{13}C_{SP-cholesterol}$ showed a gradual increase with depth, while $\delta^{13}C_{SP-DOC}$ and $\delta^{13}C_{SP-brassicasterol}$ remained unchanged over the entire water column (Fig. 3). At S4 within the upper 100 m, $\delta^{13}C_{LP-POC}$ and $\delta^{13}C_{SP-brassicasterol}$ decreased with depth. At greater depth, $\delta^{13}C_{SP-brassicasterol}$ decreased with depth. At greater depth, $\delta^{13}C_{LP-POC}$ and $\delta^{13}C_{SP-cholesterol}$ increased uniformly with depth while $\delta^{13}C_{SP-DOC}$ barely changed with depth. $\delta^{13}C_{SP-brassicasterol}$ strongly increased from the surface water to 1000 m. At S5 all components showed a gradual $\delta^{13}C$ increase with depth (Fig. 3).

4 Discussion

This study presents the first complete water column profiles of $\delta^{13}C$ data on cholesterol and brassicasterol in the Southern Ocean from the Cape Basin to the northern Weddell Gyre. We focus the discussion on the observed variation below the surface layers, with the aim of using these proxies to provide insight into the fate of organic matter sinking out of the surface water where the photosynthetic signal is acquired.

4.1 Variation of brassicasterol and cholesterol content with depth

The increase of SP-sterol content relative to SP-POC and (Fig. 2, lower panel) in the mesopelagic layer (100–1000 m)

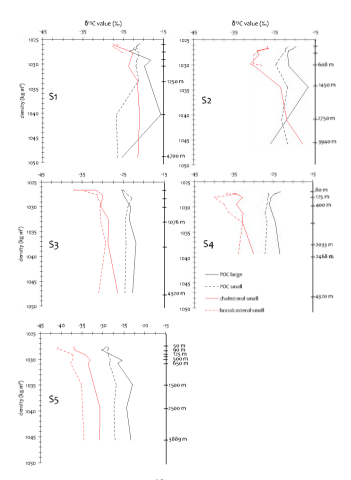


Fig. 3. Vertical profiles of $\delta^{13}C_{POC}$ (‰) for large (> 53 µm) and small (1–53 µm) particles. $\delta^{13}C_{cholesterol}$ (‰) and $\delta^{13}C_{brassicasterol}$ (‰) for small particles vs. seawater density. As an indicator, right Y-axis shows depth meter scale.

overlaps with the domain of excess ²³⁴Th activity and excess particulate barium during the BONUS-GoodHope expedition, discussed by Planchon et al. (2013). In that paper it is reported that excess ²³⁴Th and particulate barium reflect the processes of particle break-up and organic matter remineralization. The observation that in some cases the contribution of large particles to total POC (Fig. 2, upper panel) also increases in mesopelagic waters may reflect local zooplanktonic activity, possibly feeding on the vertical particle flux, thereby synthesizing cholesterol. Our present data do confirm the significance of mesopelagic waters as a region where the flux of material exported from the surface undergoes major changes.

Figure 4a shows the depth variation of brassicasterol over cholesterol ratios (bra/cho) with depth (see also Supplement Annex 1). Two clear observations emerge from these data: (i) first, bra/cho ratios are highly variable and mainly > 1 in the upper 500 m waters (surface water and upper part of the mesopelagic layer); (ii) secondly, bra/cho ratios stabilize at values of ~ 1 in the lower 500 m waters (lower mesopelagic

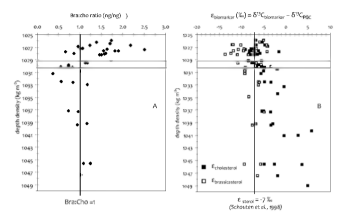


Fig. 4. (A) = vertical profile of small particle brassicasterol vs. cholesterol concentration ratio (ng ng $^{-1}$; from Supplement Annex 1) for the whole dataset. (B) = vertical profile of $\varepsilon_{cholesterol}$ and $\varepsilon_{brassicasterol}$ for the whole dataset (ε_{sterol} (%) = $\delta^{13}C_{sterol}$ – $\delta^{13}C_{sp.POC}$); standard error propagation range between 1.1 and 1.8 %. For each panel, the space located between the two horizontal lines represents the range of 500 m depth horizons along the transect.

layer and deep water). This reflects the relative decrease of autotrophic organic material (brassicasterol – phytoplankton) vs. heterotrophic material (cholesterol – zooplankton) with increasing depth. Furthermore, the lowest values of bra/cho (0.5 at station S2, 608 m; 0.4 at station S4, 749 m; Supplement Annex 1) are observed at intermediate depths, between 500 and 1000 m. Our results suggest that in the upper 500 m the major contributor to the sterol pool is phytoplankton, whereas between 500 and 1000 m it is zooplankton. Finally, these two components reach equivalent proportions below 1000 m depth. These results suggest a key role of fecal aggregates in routing carbon to the deep ocean, as highlighted by Ebersbach et al. (2011) for the Australian sector of the Southern Ocean.

From Fig. 4a observations, it is assumed that the entire dataset can be split into two groups (from surface to 500 m and from 500 m to the deep ocean). We obtain a general picture for the BONUS-GoodHope section covering 5 complete water column sampling stations located in five different Southern Ocean regions. Regression slopes are 0.5 ± 0.1 ($R^2 = 0.8$; p value < 0.05) and 1.5 ± 0.2 ($R^2 = 0.8$; p value < 0.05) for the upper 500 m and the water column below 500 m, respectively. Not shown.

4.2 Variation of δ^{13} C signatures with depth

In general, the $\delta^{13}C_{SP\text{-sterols}}$ but also $\delta^{13}C_{LP\text{-POC}}$ (S1, S2 excepted; see above section part 3.2. and Fig. 3) increase with depth, whereas $\delta^{13}C_{SP\text{-POC}}$ remains stable throughout the water column profile. Figure 4b shows depth variations of calculated values of $\varepsilon_{cholesterol}$ and $\varepsilon_{brassicasterol}$ for the complete dataset. This ε is based on a simple calculation

 $(\varepsilon_{\text{sterol}}) = \delta^{13} C_{\text{SP-sterol}} - \delta^{13} C_{\text{SP-POC}}$, allowing us to estimate the apparent offset between the ¹³C isotopic signal of SP-POC and studied sterols. Indeed, $\varepsilon_{brassicasterol}$ ranges between -13.6% and -4.1% in the upper 500 m and then stabilizes around -7% below 500 m. On the other hand, $\varepsilon_{\text{cholesterol}}$ shows a pronounced increase from the surface water ($\varepsilon_{\text{cholesterol}}$ ranges between -12.3% and -5.2%) to the deep ocean where it reaches a value of $\sim +5$ %. The stabilization of $\varepsilon_{brassicasterol}$ around $-7\,\%$ below $500\,\mathrm{m}$ is in accordance with previous laboratory experiments estimating an $\varepsilon_{\text{sterol}}$ of -7% (Schouten et al., 1998; see also Popp et al., 1999 for an application to Southern Ocean surface water data analysis). From batch cultures experiments, Riebesell et al. (2000) estimate the mean isotopic difference between POC and sterols at 8.5 ± 1.1 %, which is in accordance with Schouten et al. (1998). Summarized in Hayes (2001), it appears common, but far from universal, that sterols are depleted relative to biomass by 5-8 %. Previous studies were performed for laboratory experiments on growth rate of various phytoplankton groups (Riebesell et al., 2000; Schouten et al., 1998) or surface water samples (Popp et al., 1999), while the present observation concerns samples below the euphotic layer where no further biosynthesis of brassicasterol occurs since it is only synthesized by phytoplankton. Since we observe that $\delta^{13}C_{SP-POC}$ is stable throughout the water column, it is thus hypothesized that the stable $\varepsilon_{\text{brassicasterol}}$ of -7 %below 500 m could indicate that this sterol is biosynthesized in the surface water and not below. On the other hand, the increase of $\varepsilon_{\text{cholesterol}}$ with depth could reflect several factors which are discussed below.

4.2.1 Growth rate related effect

Fry and Wainright (1991) report that fast-growing diatoms are ¹³C-enriched compared to more slowly growing cells. Highest growth rates occur during the early season when there is no nutrient limitation, while later in the season growth rates decrease due to the onset of nutrient limitation conditions (see, e.g. Arrigo, 2002), the latter leading to more pronounced isotope fractionation and thus more ¹³Cdepleted phytoplankton (see, e.g. Popp et al., 1999). Indeed, numerous parameters such as light irradiance, supply of nutrients such as nitrogen and phosphorus, trace metals such as iron and copper (see, e.g. Boyd, 2002) can induce phytoplankton growth limitation. During the end of the growth season, we would thus expect relatively ¹³C-depleted phytoplankton in the upper water column, while the ¹³C-enriched cells from the start of the growing season would have reached the deeper water column. Such a mechanism would impact the vertical distribution of $\delta^{13}C_{brassicasterol}$ and $\delta^{13}C_{cholesterol}$. Though $\delta^{13}C_{brassicasterol}$ vertical variation was not measurable in the deep water at stations 1 and 2, the observed enrichment in $\delta^{13}C_{cholesterol}$ with depth might be explained by the growth rate related effect.

4.2.2 Particle size related effects

Pancost et al. (1997) report a particle size-dependency for $\delta^{13}C_{brassicasterol}$, with a ^{13}C enrichment in suspended particles $>20\,\mu\mathrm{m}$ relative to smaller ones. This suggests that surface-area to volume ratios control the fractionation of carbon isotopes by diatoms, as confirmed by Popp et al. (1998). Therefore, a relatively increased contribution of larger phytoplankton to the export flux could possibly explain the observed $\delta^{13}C$ enrichment with depth. Selective feeding of zooplankton on larger cells and active transport of this material to deeper layers during zooplankton migration (Steinberg et al., 2008) might cause the translocation of isotopically heavier material to deeper waters, while leaving isotopically lighter material – associated with smaller particles – in the surface layers.

4.2.3 Sea-ice algae related effect

A further possible process leading to ¹³C-enriched organic material is related to the activity of phytoplankton thriving in sea-ice brines. Sea-ice phytoplankton is significantly enriched in ¹³C compared to pelagic phytoplankton in adjacent open waters because of carbon limitation in the brine pockets and due to physiological properties such as the presence of carbon concentrating mechanisms and/or the uptake of bicarbonate (HCO₃) (Gleitz et al., 1996; Gibson et al., 1999; Villinski et al., 2000). Melting of sea-ice with release of seaice phytoplankton occurs during the growth season, so these isotopically heavy particles, if sinking out of the surface waters, can be expected to be found deeper in the water column. For the most southern stations (S4, S5), which are influenced by the Seasonal Ice Zone (SIZ) in winter, sinking of sea-ice algae could contribute to ¹³C enrichment of organic C at depth.

4.2.4 Other processes potentially affecting variations of $\delta^{13}C_{POC}$

Several further processes different from those listed above could affect the isotopic composition of deep ocean POC. These processes include (i) preferential degradation of the more labile carbohydrate and protein material, which is 13 C-enriched compared to lipids (Degens, 1969; Galimov, 2006); (ii) 13 C enrichment of sinking particles due to selective grazing on the larger 13 C-enriched cells (see discussion above) followed by fecal pellet production; (iii) loss of 13 C-depleted carbon associated with methane production by prokaryotes within zooplankton digestive tracts (Freeman, 2001; Hayes, 1993); and (iv) increase in the contribution of heterotrophic POC to the total POC pool due to transfer of organic carbon to higher trophic levels ($\sim 1{\text -}2$ % enrichment per increasing trophic level due to respiratory effects; DeNiro and Epstein, 1978; Fry, 1988; Fry and Sherr, 1984).

We note, however, that $\delta^{13}C_{SP\text{-POC}}$ did not show an increase with depth. We hypothesize that in this case there may be a balance between a $\delta^{\hat{1}\hat{3}}C_{SP-POC}$ decrease due to preferential loss of ¹³C-enriched, more labile carbon phases (carbohydrates, proteins; see above) and a $\delta^{13}C_{SP-POC}$ increase due to the trophic level succession effect, keeping $\delta^{13}C_{SP-POC}$ rather unchanged throughout the deep water column. Alternatively, it might simply be that small particles are indeed representative of the organic matter biosynthesized in the surface water. This would imply a fast transfer mechanism for these particles. Recent work by McDonnell and Buesseler (2010) did reveal that sinking velocity is greatest not only for the largest particles (up to 1 mm), but also for the small sized particles (< 50 μm). Clearly, any realistic scenario for ¹³C enrichment of biomarkers with depth needs also to comply with the observation that POC of the < 53 µm size class does not show a significant ¹³C enrichment with depth.

In summary, the observed isotopic increase for SPcholesterol with depth would need a cell size or growth rate related impact on isotopic fractionation and a mechanism separating isotopically heavy from light cells. This mechanism could be (1) a separation in time between sinking of isotopically heavier cells (early season) and light cells (late season) and/or (2) selective zooplankton grazing on larger isotopically heavier cells and their transfer to the deep sea via fecal pellets. The first hypothesis implies an age difference between deep and surface particles, while the second one does not. However, the absence of a significant δ^{13} C variation of SP-POC with depth requires that the above processes affecting $\delta^{13}C_{SP\text{-cholesterol}}$ are somehow neutralized for SP-POC (via a balance between preferential degradation of isotopically heavy compounds such as carbohydrates and proteins and isotopic enrichment due to trophic translocation of organic matter).

As an additional hypothesis, we also stress a possible effect of high pressure on cholesterol biosynthesis occurring below the surface water: cholesterol is a component of the cell membrane and the idea is that the $^{13}C^{-13}C$ covalent bond is stronger than the $^{13}C^{-12}C$, which is itself stronger than the $^{12}C^{-12}C$ covalent bond. Such specificity could be of great interest to resist better to high pressure effect. This would imply an increase of $\delta^{13}C_{cholesterol}$ with depth.

Our observations stress the interest of studying both $\delta^{13}C_{POC}$ and $\delta^{13}C_{sterol}$ in sinking and suspended particles since it allows gaining information on the fate of organic matter in the whole water column via cross-comparison.

4.3 Variation of whole water column δ^{13} C signatures with [CO₂] aq.

In this section we discuss for the first time variation of whole water column $\delta^{13} \text{C}$ signatures (surface water, mesopelagic layer and deep ocean) with surface CO_2 aq. concentration in the context of previous studies mainly limited to the surface water.

Figure 5 shows the latitudinal variation of $\delta^{13}C_{LP\text{-POC}}$ (Fig. 5a), $\delta^{13}C_{SP\text{-POC}}$ (Fig. 4b), $\delta^{13}C_{SP\text{-cholesterol}}$ (Fig. 5c), $\delta^{13}C_{SP\text{-brassicasterol}}$ (Fig. 5d) vs. surface CO_2 aq. concentration (µmol kg⁻¹) (CO_2 aq. data from Gonzáles-Dávila et al., 2011) for the arbitrarily defined surface water (0–100 m), mesopelagic layer (100–1000 m), deep ocean (> 1000 m) and bottom waters. In parallel, we obtain a good correlation between $\delta^{13}C_{SP\text{-brassicasterol}}$ and $\delta^{13}C_{SP\text{-cholesterol}}$ full dataset ($\delta^{13}C_{SP\text{-cholesterol}} = (0.7 \times \delta^{13}C_{SP\text{-brassicasterol}}) - 8.0$; $R^2 = 0.8$; not shown).

For the global ocean (Goericke and Fry, 1994) and the Southern Ocean (Lourev et al., 2004; O'Learv et al., 2001; Popp et al., 1999; Dehairs et al., 1997), a number of studies highlight a close inverse relationship between sea surface temperature (SST), CO₂ aq. and δ^{13} C_{POC}. Using models from O'Leary (1981) and Farguhar et al. (1982), François et al. (1993) demonstrate that the carbon isotopic composition of phytoplankton is primarily determined by (i) the isotopic composition of the source of inorganic carbon; (ii) the isotopic fractionation during transport into and out of the cell; and (iii) isotopic discrimination during carboxylation and the degree of leakage of intracellular inorganic carbon. Looking at the isotopic composition of the source (marine algae take up either CO₂ aq. or bicarbonate HCO₃, or both), François et al. (1993) stress that δ^{13} C-bicarbonate varies little from warm to cold surface water ($\sim +1$ to +2 %; Kroopnick, 1985) and in cases where phytoplankton incorporate bicarbonate, this would tend to increase $\delta^{13}C_{POC}$. In contrast, $\delta^{13}C_{CO_2}$ and systematically decreases from $\sim -7.5\,\%$ in warm water to \sim -10% in the cold Southern Ocean surface water. François et al. (1993) conclude that changes in the isotopic composition of CO₂ aq. can contribute significantly to the observed latitudinal trend in $\delta^{13}C_{POC}$ (a circa 2.5% decrease). For $\delta^{13}C_{\text{sterols}}$ in surface water, Popp et al. (1999) and Tolosa et al. (1999) demonstrate similar behavior as observed for $\delta^{13}C_{POC}$ in Southern Ocean and in Indian Ocean surface waters, respectively.

Several factors such as substrate concentration, growth rate, cell geometry, plankton species, and nutrient availability reviewed in François et al. (1993) and Popp et al. (1999) may account for the covariance between $\delta^{13} C$ of phytoplankton or specific compounds and CO_2 aq., and for the differences between regressions for the different C pools. However, the significant correlation between $\delta^{13} C_{SP\text{-brassicasterol}}$ and $\delta^{13} C_{SP\text{-cholesterol}}$ full dataset (see discussion above) mainly reflects the key role of change in the isotopic composition of CO_2 aq. with latitude.

4.3.1 Surface water variations

 $\delta^{13}C_{LP\text{-POC}}$ and $\delta^{13}C_{SP\text{-POC}}$ values decrease from -21.2 ± 0.7 (S1, $36^{\circ}31'$ S) to -29.4 ± 0.5 % (S5, $57^{\circ}33'$ S) and from -20.9 ± 0.1 (S1) to -28.7 ± 0.2 % (S5), respectively (Table 1). In contrast, CO₂ aq. concentrations increase between 36° S and 57° S from $10.7\pm0.3~\mu\mathrm{mol~kg^{-1}}$

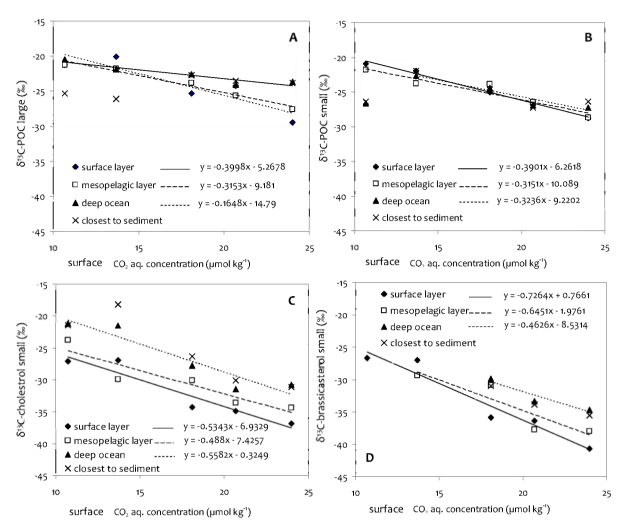


Fig. 5. Upper panel: relationship between surface water (0–100 m) CO₂ aq. concentration (μmol kg $^{-1}$. Table 1) and δ^{13} CpOC (‰) in large (> 53 μm) (A) and small (1–53 μm) (B) particles in surface water (0–100 m, closed diamonds), mesopelagic layer (100–1000 m, open squares), deep ocean (> 1000 m, closed triangles), and bottom waters (crosses). Lower panel: relationship between surface water CO₂ aq. concentration and small particles (1–53 μm) δ^{13} Ccholesterol (C) and δ^{13} Cbrassicasterol (D) in surface water (0–100 m, closed diamonds), mesopelagic layer (100–1000 m, open square), deep ocean (> 1000 m, closed triangles), and bottom water (crosses). Averages (±1 σ) are given in Table 2. Standard errors are not shown here for the sake of clarity.

to $24.0\pm0.3\,\mu\mathrm{mol\,kg^{-1}}$ (Table 1, Gonzáles-Dávila et al., 2011). This results in significant inverse correlations between $\delta^{13}\mathrm{C}_{POC}$ and surface CO_2 aq. (Fig. 5, Tables 1 and 3). The $\delta^{13}\mathrm{C}$ decrease is similar between small and large particles reaching on average $8.0\pm0.2\%$ from S1 to S5 (from Table 1). Such a decrease is similar to what has been reported by Goericke and Fry (1994) for Southern Ocean waters, but larger than the value of approximately $5\,\%$ reported by François et al. (1993) and Dehairs et al. (1997).

As observed for $\delta^{13}C_{POC}$, the decrease in $\delta^{13}C_{LP}$ and SP-cholesterol and $\delta^{13}C_{LP}$ and SP-brassicasterol in the surface water coincides with an increase in CO_2 aq. content (Table 1). However, the average (LP and SP) decrease reaches -11.1 ± 1.9 % and -14.7 ± 0.9 % for cholesterol and brassicasterol, respectively (from Table 1),

which is larger than for small and large particle POC. This observation is in agreement with Popp et al. (1999) who report that the individual sterol isotopic compositions in surface waters (WOCE SR3 transect, 145° E, between 45°55′ S to 65°24′ S) generally decrease southward and exceed the variations in bulk δ^{13} Cpoc.

Our data reveal that for large particles the difference in $\delta^{13}\mathrm{C}$ composition between LP-POC and LP-cholesterol is between -6.0 and $-13.0\,\%$ (Table 1), while the difference between LP-POC and LP-brassicasterol $\delta^{13}\mathrm{C}$ ranges between -6.8 and $-13.9\,\%$. For small particles the difference between POC and cholesterol ranges between -5.0 to $-9.3\,\%$ and the difference between POC and brassicasterol between -5.8 to $-12.0\,\%$. Such offsets appear to increase southward. Tolosa et al. (1999) estimate the $\delta^{13}\mathrm{C}$ isotopic

Table 1. Mixed layer depth MLD (m), sea surface temperature (SST), surface CO_2 aq. concentration (μmol kg $^{-1}$), surface $\delta^{13}C$ (‰) for LP and SP-POC, LP and SP-cholesterol, LP and SP-brassicasterol. Standard deviation ($\pm 1\sigma$) is the average standard deviation with one exception: (*) = analytical standard error ($\pm 1\sigma$). STZ = Sub-Tropical Zone, SAZ = Sub-Antarctic Zone, PFZ = Polar Frontal Zone, IPFZ = Intermediate Polar Frontal Zone (also named Southern Antarctic Circumpolar Current Front or SACCF), and WG = Weddell Gyre. N.D. means no data.

station	latitude	longitude	SO region	mixed layer depth (m)	surface water SST (°C)	(I) surface water $[CO_2]$ aq. $(\mu mol kg^{-1})$	δ ¹³ C cholesterol large particles	δ ¹³ C cholesterol small particles	δ ¹³ C brassicasterol large particles	δ ¹³ C brassicasterol small particles	δ ¹³ C POC large particles	δ ¹³ C POC small particles
S1	36°31′ S	13°07′ E	STZ	40	20.2 ± 0.7	10.7 ± 0.3	-27.2 ± 0.0	-27.1 ± 0.7	-28.0 ± 1.2	-26.7 ± 1.3	-21.2 ± 0.7	-20.9 ± 0.1 (*)
S2	42°28′ S	$08^{\circ}55'$ E	SAZ	50	13.2 ± 0.3	13.7 ± 0.5	-29.7 ± 0.7	-26.9 ± 0.1	-29.5 ± 0.6	-27.0 ± 0.6	-20.1 ± 0.6	-21.9 ± 0.6
S3	47°33′S	04°22′ E	PFZ	100	6.4 ± 0.1	18.1 ± 0.2	N.D.	-34.3 ± 4.0	N.D.	-35.9 ± 1.4	-25.3 ± 0.2	-25.0 ± 0.1
S4	51°52′S	00°00′ E	IPFZ	100	2.4 ± 0.3	20.7 ± 0.2	-37.2 ± 1.2	-34.9 ± 0.8	-47.0 ± 1.9	-36.4 ± 0.8	-24.2 ± 0.8	-26.4 ± 0.1
S5	57°33′ S	00°03′ E	WG	80	0.39 ± 0.01	24.0 ± 0.3	-39.7 ± 3.8	-36.9 ± 0.1	-43.3 ± 2.0	-40.7 ± 0.2	-29.4 ± 0.5	-28.7 ± 0.2

⁽I) data from Gonzáles-Dávila et al. (2011)

Table 2. Slope, standard error, p value and R^2 for variation of δ^{13} C (‰), POC large particles, small particles, cholesterol and brassicasterol in small particles vs. surface CO_2 aq. content, for the arbitrarily defined surface water (0–100 m), mesopelagic layer (100–1000 m) and deep ocean (1000-seafloor).

	slope	p value	R^2					
δ^{13} C-POC large partic	³ C-POC large particles							
surface water	-0.62 ± 0.12	< 0.05	0.81					
mesopelagic layer	-0.48 ± 0.04	< 0.05	0.97					
deep ocean	-0.25 ± 0.04	< 0.05	0.88					
δ^{13} C-POC small parti	δ^{13} C-POC small particles							
surface water	-0.60 ± 0.04	< 0.05	0.99					
mesopelagic layer	-0.47 ± 0.05	< 0.05	0.90					
deep ocean	-0.48 ± 0.08	=0.05	0.92					
δ^{13} C-cholesterol smal	ll particles							
surface water	-0.84 ± 0.13	< 0.05	0.91					
mesopelagic layer	-0.73 ± 0.10	< 0.05	0.86					
deep ocean	-0.88 ± 0.14	< 0.05	0.89					
δ^{13} C-brassicasterol sr	³ C-brassicasterol small particles							
surface water	-1.13 ± 0.14	< 0.05	0.94					
mesopelagic layer	-0.94 ± 0.20	> 0.05	0.83					
deep ocean	-0.79 ± 0.17	> 0.05	0.90					

difference between bulk organic matter and cholesterol to be 5%, and stress that such difference is typically observed between whole plant material and extractable lipids (Hayes, 1993). While for the northern part of the BGH transect, our data are close to the observations of Tolosa et al. (1999), in general we observe a larger δ^{13} C difference between POC and cholesterol, and the discrepancy is enhanced southward. This enhanced discrepancy could be related to the contribution of larger cells (diatoms).

4.3.2 Mesopelagic and deep ocean

For the $\delta^{13}C_{LP\text{-POC}}$ regressions against CO₂ aq., the slopes are quite variable (Fig. 5a, Table 2) without a clear trend with depth. On the other hand, the slopes of the $\delta^{13}C_{SP\text{-POC}}$

vs. CO_2 aq. regressions are quite conservative throughout the water column (Fig. 5b, Table 2). Thus, observations of $\delta^{13}C_{SP\text{-}POC}$ signal conservation with depth suggest that this proxy would be useful for paleoclimatology. On the other hand and in accordance with Hedges et al. (2001), this proxy is confirmed as useless to trace fate of organic matter in the water column since it shows stable signal from the surface to the deep ocean.

While the slopes of the regressions of $\delta^{13} C_{SP\text{-cholesterol}}$ vs. CO_2 aq. are also significantly conserved with depth (Fig. 5c, Table 2), the regression intercepts do evolve with depth, reflecting the tendency for higher $\delta^{13} C_{\text{cholesterol}}$ values in the deep ocean and in proximity of the seafloor (discussed in Sects. 4.1.1 and 4.1.2). $\delta^{13} C_{SP\text{-brassicasterol}}$ shows a significant decrease of regression slopes between surface and deep ocean and from the north to the south part of the Southern Ocean (Fig. 5d, Table 2). This could reflect an enhanced contribution of sea-ice diatoms south of the BGH transect (S4 and S5, see above Sect. 4.1.3), although this cannot be unambiguously demonstrated by our data. These two proxies thus appear useful to trace fate of organic matter in the whole water column.

5 Conclusions

Although the BONUS-GoodHope section crossed several frontal structures and specific biogeochemical zones, our study reveals general patterns in brassicasterol and cholesterol concentrations, bra/cho ratios, and $\delta^{13} \rm C$ signatures of POC, cholesterol and brassicasterol along with depth and latitudinal gradients. First, the bra/cho ratio, which reflects the relative importance of autotrophs vs. heterotrophs, shows a general trend relevant for each of the five studied stations: in the upper 500 m depth the major component of organic material is phytoplankton (bra/cho ratios highly variable and > 1); however, between 500 and 1000 m depth the major component becomes zooplankton (bra/cho ratios reaching values < 1 in the mesopelagic layer). These two components reach a stable ratio below 1000 m depth (bra/cho ~ 1). Our

observations support a key role of fecal aggregates in the fate of organic matter and export to deep ocean.

Second, variations of $\varepsilon_{brassicasterol}$ and $\varepsilon_{cholesterol}$ with depth also show general trends valuable for the five stations. This indeed highlights (i) that brassicasterol is only biosynthesized in the surface water, thus showing a stable value close to -7% below 500 m depth in accordance with Schouten et al. (1998) and Popp et al. (1999) application for the Southern Ocean; and (ii) an increase of $\varepsilon_{cholesterol}$ with depth which is mainly defined to be related to growth rate related effect and/or particle size related effect and selective feeding by zooplankton. A role of sea-ice algae release south of the section is also proposed to act on latitudinal and depth variations of $\delta^{13}C_{brassicasterol}$ and $\delta^{13}C_{cholesterol}$. As an additional hypothesis, we also suggest that a high pressure effect on cholesterol biosynthesis below the surface water might induce an increase of $\delta^{13}C_{cholesterol}$ in deeper waters.

Third, the relationship between CO₂ aq. concentration and $\delta^{13}C_{SP-sterols}$ is generally maintained throughout the deep ocean. This observation was not expected a priori, since circulation of deep waters carrying suspended matter formed in different regions may possibly dilute the isotopic signatures resulting from biological activity in local surface waters. It thus seems that export of organic matter and fate of organic compounds in the deep sea closely fit to a 1-D scheme (surface to deep ocean). If further studies confirm this observation, the fact that regressions of $\delta^{13}C_{cholesterol}$ and $\delta^{13}C_{SP-POC}$ versus CO_2 ag. are quite conservative with depth highlights the potential of these proxies for surface water CO₂ aq. concentration. Future field work is necessary to confirm variability and trends observed during this study, and to adequately constrain the application of such potential proxy as presented here.

While not conclusive, this first dataset on combined δ^{13} C value and concentration measurements of both bulk organic C and specific sterol markers throughout the water column shows the promising potential of analyzing δ^{13} C signatures of individual marine sterols to explore the recent history of plankton and the fate of organic matter in the SO.

Supplementary material related to this article is available online at: http://www.biogeosciences.net/10/2787/2013/bg-10-2787-2013-supplement.zip.

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