



Origin and fate of particulate and dissolved organic matter in a naturally iron-fertilized region of the Southern Ocean

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Received: 3 September 2014 – Published in Biogeosciences Discuss.: 2 October 2014

Revised: 9 December 2014 – Accepted: 11 December 2014 – Published: 30 January 2015

Abstract. Natural iron fertilization of high-nutrient low-chlorophyll (HNLC) waters induces annually occurring spring phytoplankton blooms off the Kerguelen Islands (Southern Ocean). To examine the origin and fate of particulate and dissolved organic matter (POM and DOM), D- and L-amino acids (AA) were quantified at bloom and HNLC stations. Total hydrolyzable AA accounted for 21–25 % of surface particulate organic carbon (%POC_{AA}) at the bloom sites, but for 10 % at the HNLC site. A marked decrease in %POC_{AA} with depth was observed at the most productive stations leading to values between 3 and 5 % below 300 m depth. AA contributed to only 0.9–4.4 % of dissolved organic carbon (%DOC_{AA}) at all stations. The only consistent vertical trend was observed at the most productive station (A3-2) where %DOC_{AA} decreased from ~ 2 % in the surface waters to 0.9 % near 300 m. These AA yields revealed that POM and DOM were more rapidly altered or mineralized at the bloom sites compared to the HNLC site. Alteration state was also assessed by trends in C / N ratio, %D-AA and degradation index. Different molecular markers indicated that POM mostly originated from diatoms and bacteria. The estimated average proportion of POM from intact phytoplankton cells in surface waters was 45 % at the bloom station A3-2, but 14 % at the HNLC site. Estimates based on D-AA yields indicated that ~ 15 % of POM and ~ 30 % of DOM was of bacterial origin (cells and cell fragments) at all stations. Surprisingly, the DOM in HNLC waters appeared less altered than the DOM from the bloom, had slightly higher dissolved AA

concentrations, and showed no sign of alteration within the water column. Unfavorable conditions for bacterial degradation in HNLC regions can explain these findings. In contrast, large inputs of labile organic molecules and iron likely stimulate the degradation of organic matter (priming effect) and the production of more recalcitrant DOM (microbial carbon pump) during iron-fertilized blooms.

1 Introduction

The Southern Ocean plays a key role in the global carbon cycle because it represents an important physical and biological CO₂ sink (Le Quéré et al., 2007). The magnitude of the biological contribution to the CO₂ sink is driven by primary productivity and the fate of the organic matter (OM) produced. It was univocally shown that iron controls primary production in the Southern Ocean having the largest high-nutrient low-chlorophyll (HNLC) zone (Boyd et al., 2007). The extent to which biological degradation of particulate (POM) and dissolved organic matter (DOM) affect their potential export to depth and preservation is far from understood (Ebersbach and Trull, 2008).

Natural iron fertilization induces intense phytoplankton blooms in spring, especially around the Kerguelen Islands (Blain et al., 2007). Blooms induced by natural iron fertilization likely represent major events for heterotrophic bacteria which are reported to be limited by carbon in the HNLC

Southern Ocean (Church et al., 2000). Indeed, pronounced responses of heterotrophic bacteria are reported from natural iron-fertilization studies (Zubkov et al., 2007; Christaki et al., 2008, 2014). Rapid bacterial mineralization of ~ 45 % of the primary production was observed during the decline of the spring phytoplankton bloom above the Kerguelen Plateau (Obernosterer et al., 2008). Furthermore, the efficiency of the carbon export, defined as the ratio of particulate organic carbon (POC) in deep waters to primary production, was about 2 times lower above the Kerguelen Plateau than in HNLC waters (Savoie et al., 2008). These findings suggest that the POM produced during the bloom is relatively labile, and rapidly turned over. Although DOM production is an essential step during bacterial mineralization, the effect of iron fertilization on the concentration and reactivity of DOM has been far less studied.

To better understand the role of the Southern Ocean in the cycles of carbon and other vital elements, such as nitrogen, it is important to study the origin and fate of its OM in regions having contrasting conditions and productivities. To do so, molecular level analyses of POM and DOM provide unique and valuable pieces of information. Amino acid enantiomers (L- and D-AA) are among the few markers that can be analyzed in bulk seawater at ambient concentrations and provide numerous indicators on OM origin and fate (e.g., Kaiser and Benner, 2008; Dittmar et al., 2009). AA contain most of the nitrogen in living organisms and are important constituents of detrital POM and DOM (Cowie and Hedges, 1992). While L-AA are ubiquitous (e.g., all proteins), D-AA are only produced in important proportions in bacteria (Asano and Lübbenhüsen, 2000).

AA are generally selectively utilized by heterotrophic organisms compared to bulk OM. As a result, the proportions of bulk carbon and nitrogen in the form of AA, or AA yields, decrease with time and are useful indicators of POM and DOM alteration state (Cowie and Hedges, 1994; Davis et al., 2009). Other independent diagenetic markers are the molar percentage (mol%) of D-AA versus total hydrolyzable AA (THAA) (Tremblay and Benner, 2009) and the degradation index (DI) calculated based on the relative distribution of individual AA (Dauwe et al., 1999; Peter et al., 2012). An increase in mol% D-AA during degradation is attributed to the bacterial contribution of D-AA to the OM and to a lower degradation rate of the bacterial biomolecules rich in D-AA compared to proteins having only L-AA (Nagata et al., 2003). During degradation some AA are selectively degraded (the others are selectively preserved) and these changes reduce the DI value. AA yields are known to be more sensitive to the first alterations of the OM compared to mol% D-AA and DI which require more intense alterations before showing consistent changes (Davis et al., 2009; Bourgoin and Tremblay, 2010).

C- and N-normalized yields of individual D-AA (e.g., D-alanine) were shown to have a similar dynamic as bulk bacterial C and N, respectively (Tremblay and Benner, 2006;

Kaiser and Benner, 2008). Thus, these yields can be used to estimate the contribution of bacteria (from intact cells and cell fragments or detritus) to POM and DOM. The description of this approach and its limitations are presented in previous studies (Tremblay and Benner, 2006, 2009; Kaiser and Benner, 2008; Kawasaki et al., 2011). In addition, plots of the sum of serine and threonine versus the ratio aspartic acid / glycine have been previously utilized to discriminate different carbon sources, specifically diatoms, coccolithophores and bacteria (Ittekkot et al., 1984; Gupta and Kawahata, 2000; Salter et al., 2010).

The objective of this work was to evaluate the origin and fate of the OM produced during spring phytoplankton blooms in the naturally iron-fertilized Southern Ocean. To do so, bulk and AA enantiomeric analyses were done on unfiltered and filtered waters collected at different bloom sites and in HNLC waters during the KEOPS2 (Kerguelen Ocean and Plateau compared Study 2) expedition. This work provides a detailed description of OM characteristics and new evidences on the distinct fates of the OM from naturally iron-fertilized regions.

2 Materials and methods

2.1 Site description and sampling

Among the sampling sites visited during the KEOPS2 campaign in October and November 2011, five stations were selected for the present study (Table 1) (map available in Blain et al., 2014). Station R-2 (bottom depth = 2532 m) was located in a HNLC region outside the western boundary of the Kerguelen Plateau. In contrast, station A3-2 (bottom depth = 527 m) was in the center of the phytoplankton bloom that occurs annually above the Kerguelen Plateau. Stations E-1 (bottom depth = 2056 m) and E-5 (bottom depth = 1920 m) were located in offshore waters within a stationary meander south of the polar front and these stations were sampled in a quasi-Lagrangian manner before (E-1) and during the bloom (E-5). TEW-2 is a shallow station (total depth = 85 m) above the eastern continental shelf of Kerguelen Island and it is characterized by continental inputs and relatively high productivity. Sampling depths were selected to cover important features of water masses revealed by CTD (conductance, temperature, depth) profiling. The surface mixed layer depth was 105 m at station R-2, 153 m at station A3-2, 46 m at station E-5, 72 m at station E-1, and 40 m at station TEW-2. Because of water quantity and time limitations, only the first 350 m of the water column were sampled at E-5.

Water samples for AA, dissolved organic carbon (DOC) and nitrogen (DON) analyses were collected using 10 L Teflon-lined Niskin-1010X bottles mounted on a 1018 rosette system adapted for trace metal clean work (Bowie et al., 2014). Subsamples were transferred to 500 mL Schott glass bottles that were rinsed with acid (HCl, 10 %) and then ultrapure water before use. Part of each water sample

Table 1. Sampling sites, chlorophyll *a* (Chl *a*) concentrations, particulate and dissolved organic carbon (POC, DOC) and nitrogen (PON, DON) concentrations, and amino acid (AA) concentrations in particulate and dissolved (including free AA) phases.

Stations	Coordinates		Depth (m)	[Chl <i>a</i>] ($\mu\text{g L}^{-1}$) ^a	[POC] (μM) ^a	[DOC] (μM)	[PON] (μM) ^a	[DON] (μM) ^b	[AA] (nM)			Free AA ^c (%)
	Latitude ° S	Longitude ° E							partic. ^c	dissolv. ^d	free	
R-2 (HNLC) ^f 27 Oct 2011 ^g	50.383	66.683	40	0.32	7.8	48	0.8	6.0	162	335	38	11.3
			70	0.27	nd	47	nd	6.1	262	216	nd	
			100	0.23	5.2	48	0.7	6.9	498	276	nd	
			200	0.02	2.9	47	0.3	nd	45	173	nd	
			300	nd	nd	45	nd	nd	23	198	nd	
			500	nd	1.5	44	0.1	nd	77	186	nd	
			900	nd	1.6	45	0.1	nd	92	192	11	
1300	nd	nd	44	nd	nd	33	151	nd				
A3-2 17 Nov 2011 ^g	50.617	72.050	40	1.7	14.8	52	2.8	6.0	821	277	52	18.8
			70	2.1	14.3	51	2.6	5.1	716	292	69	
			110	2.4	13.9	50	2.6	3.9	656	237	nd	
			150	1.4	10.8	48	2.1	5.3	400	173	nd	
			210	0.15	3.5	47	0.4	3.9	44	144	10	
300	0.05	2.0	48	0.3	2.7	22	128	nd				
E-5 18 Nov 2011 ^g	48.400	71.884	25	1.2	11.0	48	2.2	6.5	539	208	28	13.5
			70	0.95	9.1	47	2.2	7.0	333	234	nd	
			110	0.36	nd	47	0.6	6.8	71	160	nd	
			200	0.12	3.5	46	0.3	6.9	45	191	6	
350	nd	3.0	45	0.2	nd	27	146	nd				
E-1 30 Oct 2011 ^g	48.450	72.183	20	0.99	7.7	49	1.3	4.8	223	406	nd	
			40	0.96	8.9	49	1.6	5.0	486	352	nd	
			70	0.86	4.5	47	1.0	5.2	124	495	nd	
			120	0.78	6.5	46	1.2	6.4	387	289	nd	
			200	0.07	5.1	45	1.0	nd	203	314	nd	
			500	nd	1.6	45	0.1	nd	nd	383	nd	
1300	nd	nd	45	nd	nd	44	199	nd				
TEW-2 (shelf) 31 Oct 2011 ^g	48.884	70.650	15	1.8	9.3	47	1.5	7.4	627	413	37	9.0
			30	1.4	5.7	46	1.0	7.1	338	489	nd	
			62	0.6	6.4	45	0.7	7.6	205	403	11	

^a Data are from Lasbleiz et al. (2014).^b Data are from Blain et al. (2014).^c Particulate AA = total hydrolyzable amino acids (THAA).^d Dissolved AA = dissolved combined and free amino acids.^e % free amino acids versus dissolved AA.^f High-nutrient low-chlorophyll.^g Dates of sampling.

nd: not determined.

was filtered through two combusted GF/F filters (glass fiber, 0.7 μm nominal pore size, Whatman) using a Hamilton glass syringe and PTFE (polytetrafluoroethylene) tubing. For AA analyses of the dissolved fraction, 20 mL of the GF/F filtered water were stored in HDPE (high-density polyethylene) bottles at -20°C . For DOC analyses, 15 mL of the GF/F filtered water were acidified (H_3PO_4 , final pH = 2) and stored in the dark in combusted and sealed glass ampoules. DON analyses were done with an aliquot of the GF/F filtered water that was stored poisoned (100 μL of HgCl_2 ; 4 g L^{-1} , working solution) in PTFE bottles (Blain et al., 2014). Total AA (dissolved and particulate) were determined in 20 mL of unfiltered water stored in HDPE bottles at -20°C . The concentration of particulate AA was calculated by subtracting the

AA concentration in filtered water from the total AA concentration.

Water samples for POC and PON analyses were collected with regular 12 L Niskin bottles. A total of 1 L of seawater was filtered on combusted GF/F filters, and these were stored in the dark until analyses (Lasbleiz et al., 2014). The sampling depths of these water samples did not always perfectly match the depths of the samples used for the other analyses. In these cases, the depths that were the closest to those shown on Table 1 were used for POC and PON.

2.2 Organic carbon and nitrogen analysis

DOC concentrations were measured on acidified aliquots using a Shimadzu TOC-VCP analyzer with a Pt catalyst at 680 °C (Benner and Strom, 1993). All DOC analyses were run in duplicate and 3–5 injections were done for each sample. Analytical precision was 0.2–1 % (CV). Reference materials (<http://www.rsmas.miami.edu/groups/biogeochem/CRM.html>) were injected every 15 samples to insure stable operating conditions. DON concentrations were determined after wet oxidation and calculated as the difference between total N concentration and the sum of inorganic N (i.e., nitrate, nitrite, ammonium) (Blain et al., 2014). Inorganic N was determined by the automated colorimetric technique on a Skalar autoanalyzer (Aminot et K erouel, 2007). POC and PON analyses of particles on filters were done on a PerkinElmer 2400 CHN analyzer (Lasbleiz et al., 2014). The filter blank values for POC and PON corresponded to $1.27 \mu\text{M} \pm 0.26$ and $0.06 \mu\text{M} \pm 0.02$, respectively.

2.3 Amino acid analysis

AA enantiomers (L and D) and achiral AA were measured after liquid-phase acid hydrolysis and reversed-phase high-performance liquid chromatography (HPLC) according to a newly developed procedure (Escoubeyrou and Tremblay, 2014). This procedure is similar to traditional ones, except that a high volume (100 μL) of undiluted samples is injected and a robust hybrid C-18 column is used. These changes increase the precision of the measurements while providing a more consistent separation.

Briefly, 366 μL of seawater (filtered or unfiltered) were transferred to an ampoule and mixed with 634 μL of HCl 30 % (Merck, Suprapur grade). Procedural blanks were also prepared with 634 μL of HCl 30 %. The ampoules were then vacuum sealed and placed at 110 °C for 20 h. After hydrolysis, 300 μL of hydrolyzate (190 μL for blanks) were evaporated on a Genevac Personal Evaporator (EZ-2 + HCl compatible). A total of 150 μL of ultrapure water were then added and evaporated to remove remaining traces of HCl. Samples consisting of 110 μL of nonhydrolyzed seawater were also evaporated for the analysis of free AA. Samples and blanks were then dissolved in 120 μL of borate buffer (0.4 M, Merck, Suprapur) with a pH adjusted to 10 with NaOH 6 M (Merck, Suprapur). Strong vortex agitation and sonication were necessary to completely dissolve the dried residue. The vials were then centrifuged (4300 rpm, 5 min) to separate the flocculate formed in borate buffer.

HPLC analyses were performed on an Ultimate 3000 Dionex system equipped with an autosampler and a fluorescence detector (excitation at 335 nm, emission at 450 nm). The separation was done with a Phenomenex Gemini C18 column (250 x 4.6 mm, 5 μm , 110 Å) and a SecurityGuard column (4 x 3 mm). Automated precolumn derivatization with o-phthalaldehyde (OPA) and N-isobutyryl-L-cysteine

(IBLC, L-run) or with OPA and N-isobutyryl-D-cysteine (IBDC, D-run) was performed. The reagents were prepared by dissolving 2 mg of OPA and 5 mg of IBLC or IBDC in 200 μL of methanol and diluting with 1.8 mL of ultrapure water. The derivatization mixture consisted of 100 μL of sample or blank, 10 μL of OPA/IBD(L)C reagent and 10 μL of S-methyl-L-cysteine (SMC, 0.74 μM) used as internal standard. After a reaction time of 2 min at 8 °C, 100 μL of this mixture were injected. OPA/IBLC (run L) and OPA/IBDC (run D) were used alternatively for every sample and blank. The elution order of L-AA and D-AA is reversed during the second run, which increases the accuracy of peak identification and quantification (Br uckner et al., 1994). The largest peak of the two runs was not considered because it may be caused by a coelution. The column temperature was maintained at 44 °C. A gradient of organic phase (93 % methanol, 7 % acetonitrile) and KH_2PO_4 aqueous solution (40 mM, pH 6.15) was used with a flow rate of 0.8 mL min^{-1} (Escoubeyrou and Tremblay, 2014).

Before injecting samples, blanks or standards, several analyses of borate buffer were performed to check the level of contamination from free AA. AA identification and quantification in unknown samples and blanks were achieved by comparing the retention times and peak areas with those of standard AA solutions. Coefficients of determination of calibration curves were always between 0.99 and 1.0. Standard solutions contained 23 AA: L- and D-aspartic acid (L/D-Asp), L- and D-glutamic acid (L/D-Glu), L- and D-serine (L/D-Ser), L- and D-alanine (L/D-Ala), L- and D-valine (L/D-Val), L- and D-leucine (L/D-Leu), L-threonine (L-Thr), glycine (Gly), L-arginine (L-Arg), L-isoleucine (L-Ileu), L-phenylalanine (L-Phe), L-lysine (L-Lys), L-histidine (L-His), L-tyrosine (L-Tyr), L-methionine (L-Met), γ -aminobutyric acid (Gaba), and β -alanine (BAIa).

Peak areas of AA measured in blanks were subtracted from those in samples. The borate buffer solution was used as the blank for the quantification of free AA. Individual AA concentrations in injected samples were then calculated using the slope of calibration curve of each AA. The average racemization rates of individual free and protein AA occurring during hydrolysis and measured by Kaiser and Benner (2005) were used to correct the concentrations of chiral AA. Asp and Glu peaks included the contribution of L-glutamine (Gln) and L-asparagine (Asn), respectively, which were deaminated during hydrolysis. They were thus identified as Asx and Glx.

Particulate THAA were quantified by the difference between the peak areas obtained with unfiltered and filtered hydrolyzed samples. The chromatograms of filtered samples were used to quantify dissolved AA, which are the sum of free and dissolved combined AA. Dissolved combined THAA can be determined by the difference between the peak areas obtained with filtered hydrolyzed and nonhydrolyzed samples. The limits of detection were 0.007–3.57 nM depending on the AA (Escoubeyrou and Tremblay, 2014). Most

particulate AA (determined by difference) were near their limit of detection in samples having low POC concentrations. This increases the uncertainty of particulate AA, POC and PON values in these samples.

2.4 Chlorophyll analysis

Concentrations of chlorophyll *a* were determined by HPLC following the protocol of Van Heukelem and Thomas (2001) modified by Ras et al. (2008). Seawater samples (1–2.23 L) were filtered on 25 mm GF/F filters (Whatman) which were then stored in cryotubes in liquid nitrogen until further processing in the home lab, as described in detail in Lasbleiz et al. (2014).

3 Results

3.1 Organic carbon and nitrogen

Striking differences in OM concentrations and compositions were observed between stations (Table 1). As expected, station R-2 located in the HNLC region exhibited the lowest chlorophyll *a* ($< 0.32 \mu\text{g L}^{-1}$) and POC concentrations (1.5–7.8 μM) compared to the other stations for similar depths. The highest concentrations of chlorophyll *a* (up to $2.4 \mu\text{g L}^{-1}$) and POC (up to $14.8 \mu\text{M}$) were observed in the mixed layer at station A3-2. Station E-1, sampled before the bloom, had relatively low POC concentrations in its surface waters (7.7 μM at 20 m). After 19 days, the POC concentrations increased to 11 μM at E-5. There was a marked decrease in POC concentrations with depth, especially at station A3-2 showing a 7.4-fold decrease. The lowest POC concentrations, 1.5–3 μM , were measured below 210 m.

DOC represented 78 (surface water of A3-2)–97 % (deep water of R-2) of total organic carbon (Table 1). Despite a slight decreasing trend with depth at all stations, DOC concentrations were much less variable than those of POC. The highest concentrations (51–52 μM) were measured in the surface waters of station A3-2, the bloom site above the plateau. The lowest concentration (44 μM) was found in the deep waters of the less-productive station R-2.

PON and DON concentrations also varied with location and depth (Table 1). They generally followed the same trends as their carbon counterparts, although not always to the same extent as indicated by the POC / PON and DOC / DON ratios in representative samples (Fig. 1). A preferential degradation of N-rich molecules in particles lead to a general increase in POC / PON ratios with depth (Fig. 1a). As expected, the particles in the surface waters of the bloom at stations A3-2 and E-5 were enriched in N. In contrast, the HNLC region (station R-2) had more N-poor particles (Fig. 1a) and 2–3.5 times lower PON concentrations (Table 1). DON concentrations varied from 7.6 μM at station TEW-2 to 2.7 μM in the deep waters of station A3-2. Station A3-2 was the only site where DON concentrations decreased (Table 1) and DOC / DON

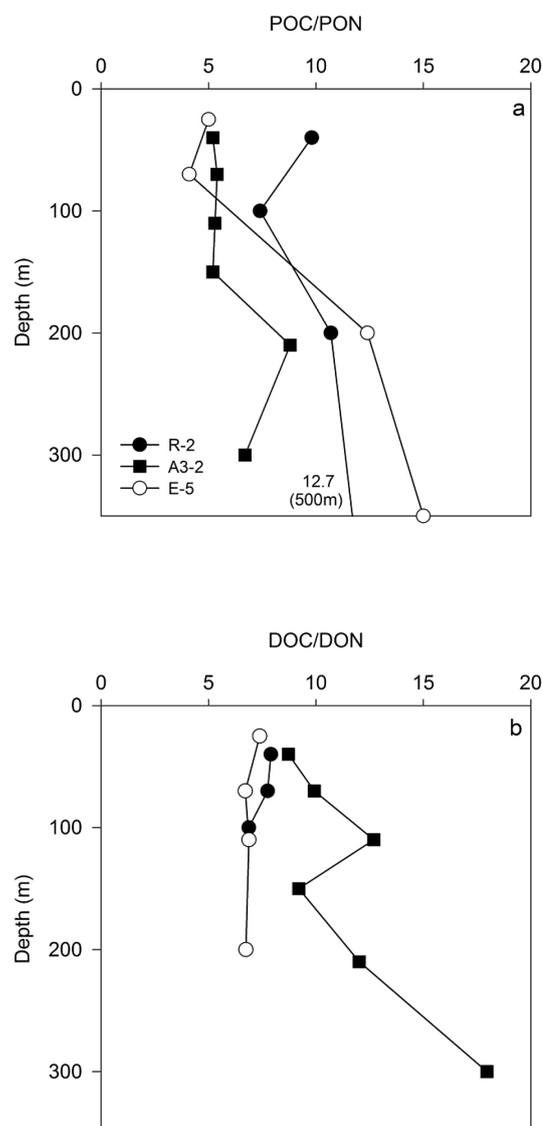


Figure 1. Carbon to nitrogen atomic ratios in the (a) particulate and (b) dissolved fractions for three representative stations (R-2, A3-2, E-5) at 0–350 m depth. The value close to the station R-2 profile is the next data point at 500 m. Ratios cannot be quantified at certain depths because elementary data were not available (Table 1).

ratios increased with depth (Fig. 1b). This indicates that N-rich dissolved organic molecules at station A3-2 are more easily or rapidly degraded than those from the other stations.

3.2 Amino acid concentrations

Table 1 presents AA concentrations measured in the particulate and dissolved fractions. As expected, the most productive stations (A3-2, E-5, and TEW-2) had the highest concentrations of particulate THAA. The water collected at the surface of these stations contained between 539 and 821 nM of particulate THAA, while concentrations were 162–223 nM

in the surface waters of stations R-2 and E-1. Though deep waters contained less particulate THAA than surface waters at all stations, the decrease with depth was much faster at the bloom sites. These concentrations dropped by a factor of 20 and 37 at stations E-5 and A3-2, respectively, but only by a factor of ~ 5 at stations R-2 and E-1. At stations R-2 and E-1, particulate THAA concentrations were maximal in the subsurface layer (40–100 m), and not at the surface like for bloom locations. This suggests a production of particulate THAA in the subsurface or a localized removal mechanism (e.g., photo-oxidation) in the first meters of the water column at stations R-2 and E-1.

In contrast to particulate THAA, concentrations of dissolved AA (combined and free AA) were generally higher at the less productive locations (R-2 and E-1; up to 495 nM) than at the bloom sites (A3-2 and E-5; 128–292 nM) for similar depths (Table 1). The shallow station TEW-2 also had high concentrations (403–489 nM). However, a greater proportion of dissolved AA were free AA at the surface of stations A3-2 and E-5 compared to station R-2 (13.5–23.6 % vs. 11.3 %; Table 1). Station A3-2 exhibited the highest concentrations of free AA (52–69 nM; Table 1), but lower dissolved combined AA (e.g., in proteins and peptides) concentrations than station R-2.

Concentrations of total dissolved AA decreased with depth, by a factor 1.4–2.2 (Table 1), except in the shallow water column at station TEW-2. The decrease in free AA was more important (factor 3.4–5.2). These trends were more pronounced at the bloom station A3-2, but were not as marked as for particulate THAA.

3.3 Amino acid yields and diagenetic markers

AA are the most abundant organic N-molecules in nature. We thus expect to see similar trends in the proportions of N and of THAA in POM as indicated by POC/PON (Fig. 1a) and %POC_{AA} (particulate organic carbon quantified as amino acids) (Fig. 2a), respectively. Both parameters indicated higher proportions of N-molecules in POM at stations A3-2 and E-5 and a decrease in these proportions with depth at all stations. The decrease in %POC_{AA} was particularly marked at stations A3-2 and E-5 where THAA accounted for 22–25 % of POC in the surface waters, but only for 4.2–4.9 % below 300 m (Fig. 2a). Particulate THAA accounted for 30–51 % of PON in the surface of the more productive stations and for 27 % at stations R-2, and these proportions decreased with depth at all stations, especially at station A3-2 (not shown). The degradation of particulate THAA was thus more rapid than that of other N molecules.

Dissolved AA contributed to only 0.86–2.7 % of DOC at stations R-2, A3-2 and E-5 (Fig. 2b). %DOC_{AA} (dissolved organic carbon quantified as amino acids) were slightly higher in the surface sample of stations E-1 and TEW-2 (2.4–4.4 %, E-1 and TEW-2 data not shown). The only consistent vertical trend was observed at station A3-2 where %DOC_{AA}

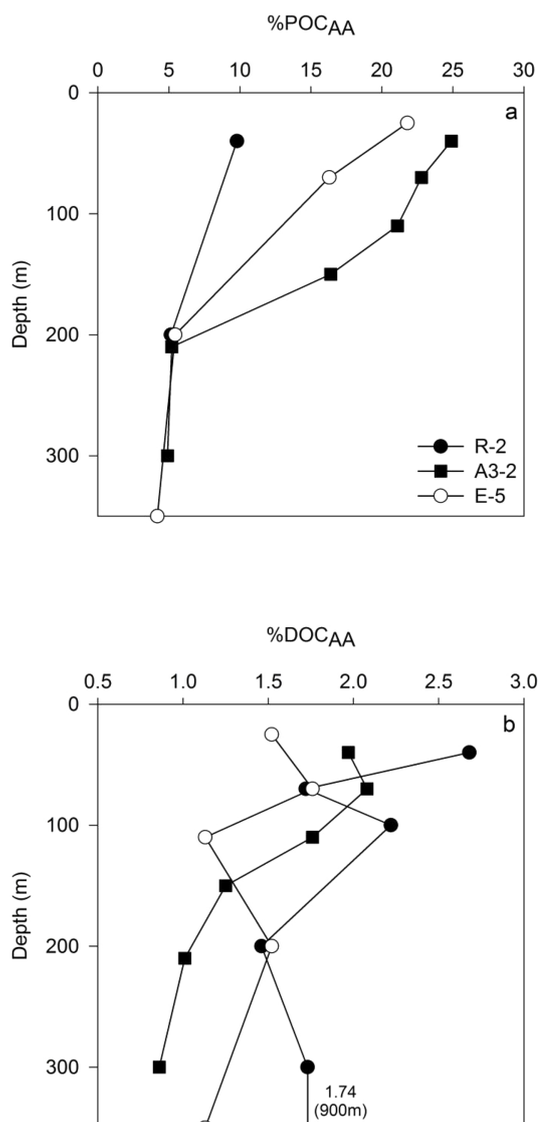


Figure 2. Proportion of (a) particulate and (b) dissolved organic carbon quantified as amino acids (%POC_{AA} and %DOC_{AA}) for three representative stations (R-2, A3-2, E-5) at 0–350 m depth. The value close to the station R-2 profile is the next data point at 900 m.

decreased with depth from ~ 2 to 0.86 %. This suggests a preferential removal of dissolved AA with depth at this bloom station. This trend is not observed or is less marked at the other stations. Dissolved AA contributed to 2.3–8 % of DON with no trend according to station or depth (not shown).

During degradation, the mol% D-AA should increase and the DI should decrease. These markers indicated that POM from station R-2 was more altered than the POM from the other stations (Figs. 3a, 4a), in agreement with its lower N and THAA contents. However, mol% D-AA and DI did not show consistent trends with depth, except for mol% D-AA at station R-2 (Fig. 3a) and for DI in deep waters only (Fig. 4a). Clearly, these markers were not as sensitive, or effective, as

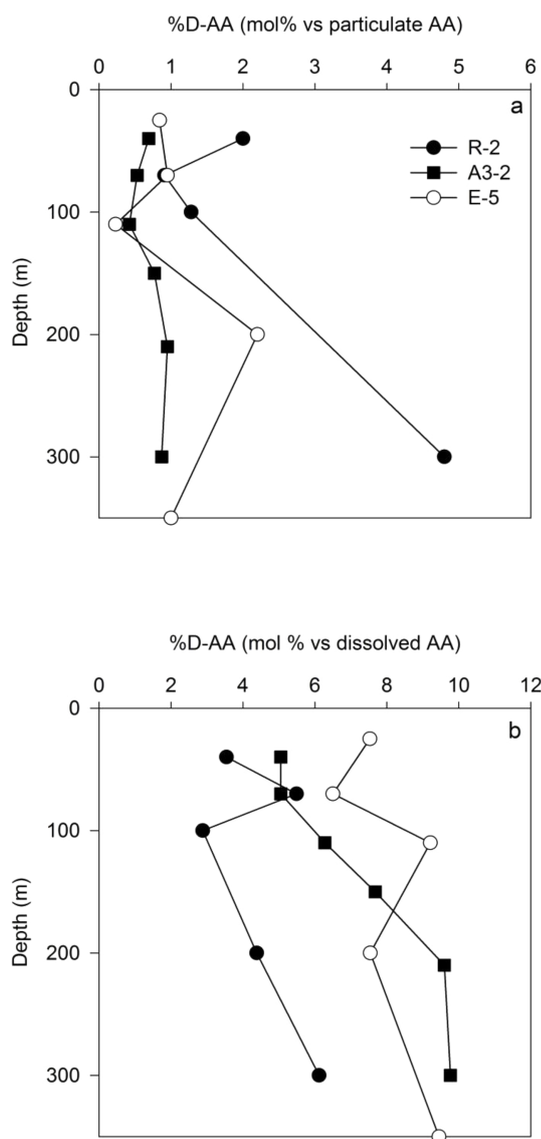


Figure 3. Proportion of D-amino acids (%D-AA) versus total amino acids (AA) in the (a) particulate and (b) dissolved fractions for three representative stations (R-2, A3-2, E-5) at 0–350 m depth.

%POC_{AA} (or POC/PON) for tracking POM degradation with depth in the present study. In the dissolved fraction, mol% D-AA and DI values suggest, once again, that DOM from station R-2 was less altered (Figs. 3b, 4b). However, no sign of alteration with depth was observed at this station. In fact, the only station showing a clear increase of DOM alteration state with depth was, once again, A3-2 (Figs. 3b, 4b). From 40 to 300 m, mol%D-AA went from 5 to 10% and the DI from 0.5 to -1.7 in DOM. The increase in mol%D-AA in total OM (POM and DOM) with depth at station A3-2 was even more important (from ~ 2 to 8%, not shown).

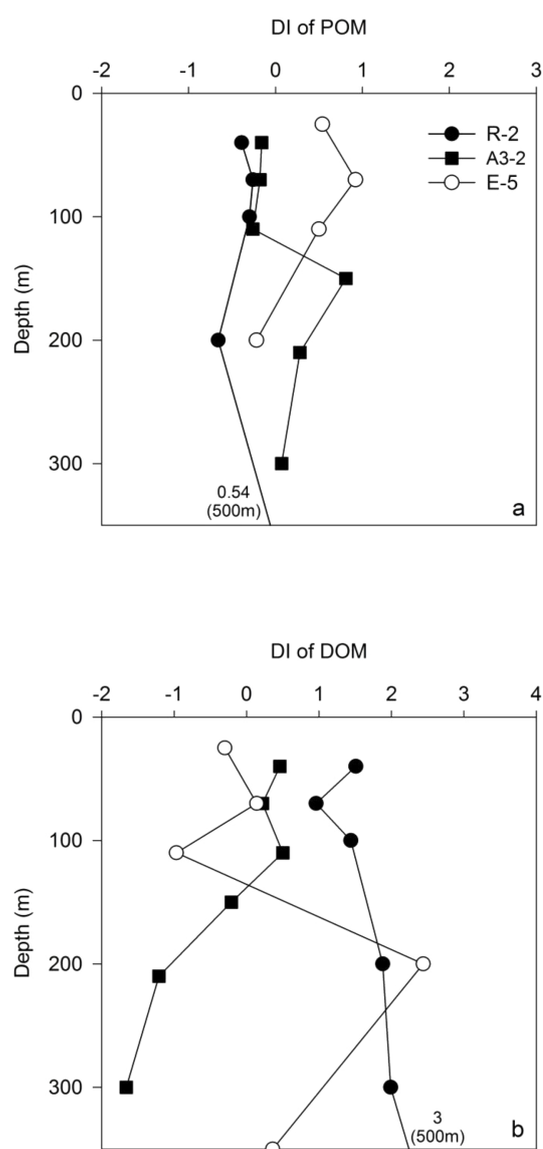


Figure 4. DI of (a) POM (Dauwe et al., 1999) and (b) DOM (Peter et al., 2012) calculated for three representative stations (R-2, A3-2, E-5) at 0–350 m depth. Values close to the station R-2 profiles are the next data point at 500 m.

3.4 Markers of organic matter origin

The average values of Ser + Thr and Asp / Gly found in diatoms, coccolithophores and bacteria (Muller et al., 1986; Cowie and Hedges, 1992) are presented in Fig. 5, along with the values measured here at stations A3-2 and E-5. All the data points for POM (open circles) are near average values (open boxes) found in diatoms or between those of diatoms and bacteria, suggesting that the POM mostly originated from diatoms and bacteria. Except for one data point at each of the other stations having an Asp / Gly ratio of over 1.5, the same conclusion can be stated for the other studied

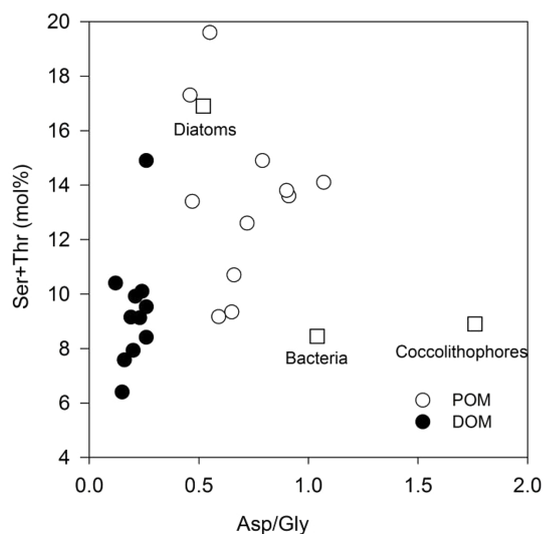


Figure 5. Molar contribution of serine + threonine (Ser + Thr) versus the molar ratio of aspartic acid and glycine (Asp / Gly) in the particulate and dissolved organic matter (POM and DOM) of stations A3-2 and E-5. Average values for diatoms, coccolithophores, and bacteria are based on Muller et al. (1986) and Cowie and Hedges (1992).

locations (not shown). No trend with depth was observed for Ser + Thr, but Asp / Gly generally decreased with depth (not shown). Gly is known to be selectively preserved during POM degradation (Dauwe et al., 1999). Changes in the composition of AA during degradation (e.g., microbial) are certainly responsible for the lower Ser + Thr and Asp + Gly values measured in DOM (Fig. 5).

Chlorophyll *a* and D-AA can be used to estimate the proportions of organic matter associated with phytoplankton (cells) and bacteria (cells and detritus), respectively. The ratios of chlorophyll *a* to POC concentrations (concentrations in Table 1) at stations A3-2 and E-5 are more than twice those measured in the HNLC station R-2. This difference can be due to a higher proportion of phytoplanktonic cells in the POM from the bloom and/or to lower intracellular chlorophyll content in the phytoplankton of the HNLC region. Assuming that diatom-dominated phytoplankton communities have 40 times more carbon than chlorophyll *a* on a per cell basis (Sathyendranath et al., 2009), calculations indicated that phytoplankton biomass could account for 14 % of POC in the surface water of station R-2, but for 35–63 % in the surface waters of the other stations (not shown), except at 30 m of TEW-2 where 81 % was calculated. Estimates were much lower in the deep waters, with 2.3 % calculated at station R-2 (200 m) and 8.3 % at station A3-2 (300 m). This approach takes mostly into account phytoplankton cells (vs. altered fragments) considering that a rapid decrease in chlorophyll *a* content is expected in phytoplankton detritus.

D-AA can be used to estimate the proportions of OM coming from bacterial cells and altered cell fragments or detri-

tus. D-Asx yields varied from < 2 to 31.7 nmol mg⁻¹ POC and D-Glx yields varied from < 2 to 18.1 nmol mg⁻¹ POC (Table 2). In both cases, the highest values were measured above 200 m in the water columns. D-Ala yields were highly variable, < 2–68.8 nmol mg⁻¹ POC with no clear trend with depth. The sample collected at 200 m at station R-2 exhibited relatively high values for each D-Asx, D-Glx and D-Ala (31.7, 18.1 and 53.7 nmol mg⁻¹ POC, respectively), suggesting a greater contribution of bacteria to POC in this sample. Average yields reported in cultured marine bacteria (Kaiser and Benner, 2008) represent 15, 48 and 50 nmol mg⁻¹ POC for D-Asx, D-Glx and D-Ala, respectively (Table 2). Comparisons between these values and those measured here in POM suggest that a significant, although variable, fraction of POC can be from bacteria. For instance, at station A3-2, where higher POC concentrations increase estimation accuracy and precision, D-Glx and D-Ala yields suggest that ~ 5–20 % of the POC was from bacteria. However, the lack of correlations, or consistent trends, between the D-AA biomarkers and the large ranges of D-AA yields in POM (and bacteria) makes it difficult to provide estimates of the percentages of POM coming from bacteria based on this approach.

D-AA yields in the dissolved fraction were much more consistent (Table 2). Figure 6 shows average values measured in DOM produced by bacteria (Kaiser and Benner, 2008) and in DOM from surface (0–120 m) and deep waters (> 120 m) at stations impacted (A3-2 + E-5) or not (R-2 + E-1) by the bloom. D-Asx and D-Ala yields suggest that 16.5–35 % of this DOC is from bacteria, with the lowest estimates found in the deep waters of stations R-2 and E-1. Similar trends, but lower estimates (10.4–17.5 %) were obtained when D-Glx yields are used. D-Glx yields in bacterial DOM were highly variable (Kaiser and Benner, 2008), and the reference average value used here might be higher than that of the bacterial DOM produced in the studied waters.

4 Discussion

4.1 Amino acids in the Southern Ocean

This study presents the first detailed quantification of particulate and dissolved AA in the Southern Ocean. The measured concentrations, yields, and relative distributions of AA in POM and DOM were in the range of values previously found in the open equatorial Pacific (Lee et al., 2000), the coastal waters of the Arctic (Dittmar et al., 2001; Shen et al., 2012) and North Atlantic Ocean (Bourgoin and Tremblay, 2010), and in the polar front zone of the Southern Ocean (Panagiotopoulos et al., 2002). THAA represented between 21 and 25 % of POC in the surface waters of bloom sites, which is close to the 27 % measured in suspended particles at the chlorophyll maximum depth in the equatorial Pacific (Lee et al., 2000). Such high values are expected when protein-rich plankton represents an important part of POM (see Sect. 4.2

Table 2. Yields of D-AA in particulate and dissolved fractions along with average yields measured in cultured bacteria and bacterial DOM.

Stations	Depth (m)	D-Asx ^a	D-Glx ^b	D-Ala ^c	D-Asx	D-Glx	D-Ala
		nmol (mg POC) ⁻¹			nmol (mg DOC) ⁻¹		
R-2 (HNLC)	40	10.2	2.83	13.9	5.31	1.77	10.6
	70	nd	nd	nd	6.09	1.59	9.75
27 Oct 2011 [§]	100	7.06	6.36	68.8	6.52	1.81	4.70
	200	31.7	18.1	53.7	4.27	0.81	7.17
	300	nd	nd	nd	6.58	1.63	11.3
	500	nd	nd	nd	3.80	1.06	7.69
	900	5.48	8.22	nd	4.85	1.84	10.5
	1300	nd	nd	nd	5.30	1.82	9.53
A3-2 17 Nov 2011 [§]	40	13.5	9.33	5.34	6.36	2.61	11.0
	70	9.80	8.99	2.17	6.88	2.56	12.1
	110	6.12	7.65	2.62	7.29	2.63	11.9
	150	7.30	6.25	6.77	6.59	2.62	11.0
	210	nd	2.40	7.51	6.88	2.61	11.6
	300	nd	6.57	nd	6.80	2.09	10.5
E-5 18 Nov 2011 [§]	25	14.8	10.5	4.25	7.44	3.24	13.5
	70	9.67	4.37	10.4	8.14	3.38	12.9
	110	nd	nd	nd	7.32	2.87	13.0
	200	7.79	1.91	10.8	6.95	2.50	13.0
	350	2.13	0.55	nd	6.82	2.73	14.1
E-1 30 Oct 2011 [§]	20	1.70	7.13	nd	5.64	2.45	12.1
	40	5.27	4.62	14.0	5.39	2.47	11.2
	70	nd	2.70	4.34	7.47	2.92	11.4
	120	nd	7.09	nd	6.51	2.44	11.3
	200	nd	2.17	nd	2.69	2.44	9.94
	500	nd	nd	nd	1.94	1.88	5.63
	1300	8.39	nd	nd	2.63	2.25	8.17
TEW-2 31 Oct 2011 [§]	15	nd	2.85	nd	8.39	4.13	14.7
	30	21.3	14.3	nd	8.46	4.24	15.8
	62	9.26	2.74	nd	8.12	4.10	16.0
Bacteria ^d		15 (22)	48 (26)	50 (19)	24 (3)	17 (7)	35 (7)

^{a, b, c} D-aspartic acid or D-asparagine, D-glutamic acid or D-glutamine, and D-alanine, respectively.

^d From Kaiser and Benner (2008). POC-normalized values are averages from three phototrophic and five heterotrophic marine bacteria assuming 20 % phototrophic and 80 % heterotrophic. DOC-normalized values are averages measured in three DOM samples freshly produced by bacteria. Values in parentheses are standard deviations.
nd: not determined or below detection (for POC or D-AA measurements).

below) (Cowie and Hedges, 1992; 1994). Concentrations of chlorophyll *a* and POM were highest in the surface water of the bloom station A3-2, consistent with a greater production of biomass induced by iron inputs. In contrast, THAA represented 10 % of POC at station R-2, which is expected in more-altered POM (Cowie and Hedges, 1994).

A particularity of the Southern Ocean surface waters are the lower DOC concentrations compared to other oceans (i.e., ~ 50 vs. 70–80 μM) (Hansell, 2013). Here, AA accounted for only 0.9–4.4 % of DOC, which is in the range of values found in other regions (Benner, 2002; Bourgoin and Tremblay, 2010; Shen et al., 2012). However, AA yields in the DOC of HNLC surface water (2.7 %) were about twice

that measured in the oligotrophic North Pacific near Hawaii (Kaiser and Benner, 2008). This suggests more favorable conditions for AA accumulation/preservation in the HNLC of the Southern Ocean.

4.2 Origin and fate of POM

Molecular AA markers (i.e., Ser + Thr, Asp / Gly and D-AA yields) indicated, at least qualitatively, that the POM at the bloom sites mostly originated from diatoms and bacteria. Different approaches can be used to estimate the proportion of each of these components of the plankton community. Pigment analysis and microscopic observations confirmed that diatoms were abundant in surface waters of the bloom sites

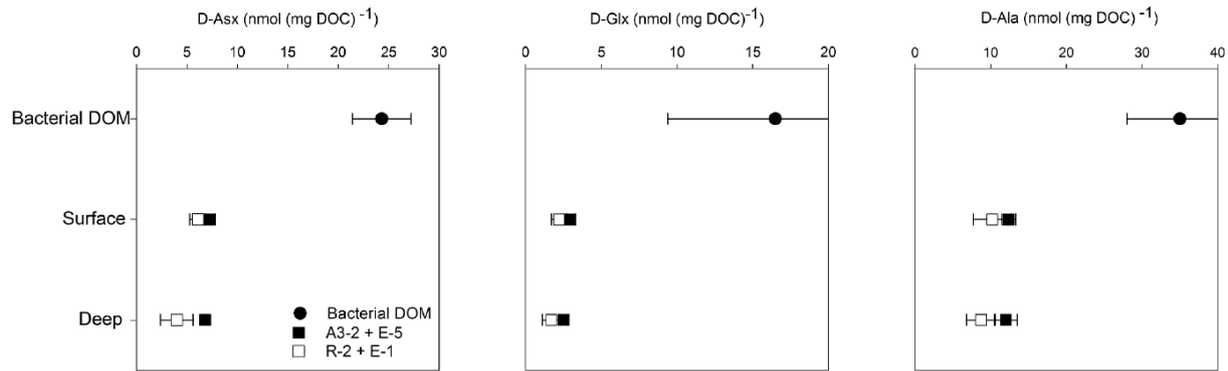


Figure 6. Average C-normalized yields of D-aspartic acid or D-asparagine (D-Asx), D-glutamic acid or D-glutamine (D-Glx), and D-alanine (D-Ala) in bacterial DOM (Kaiser and Benner 2008) and DOM from surface (0–120 m) and deep waters (> 120 m) at stations impacted (A3-2 + E-5) or not (R-2 + E-1) by the bloom. Error bars represent standard deviations.

(Lasbleiz et al., 2014). Species-specific biovolume determination in the euphotic layer (between the surface and 0.01 % PAR – photosynthetically active radiation) revealed that diatoms accounted, on average, for 44 % of POC at station A3-2, while their contribution to POC was only 5 % at the HNLC site (Lasbleiz, 2014) (Table 3). These observations are consistent with the present estimates based on chlorophyll *a* concentrations, indicating that phytoplankton cells could account on average for 45 % of POC in the surface waters of station A3-2, but 14 % of POC in the surface waters of station R-2 (Table 3). The lower proportion of diatoms among the phytoplankton community at station R-2 is in agreement with direct observations (Lasbleiz et al., 2014). These estimates were 5–7 times lower below 200 m at both locations (Table 3). In deep waters, the phytoplankton contribution to POC is thus likely dominated by detrital material (vs. intact cells). Laurenceau et al. (2014) used microscopic observations of material collected in gel traps and estimated that phytodetrital aggregates represented 41 and 34 % of POC at stations A3-2 and R-2, respectively, in sinking particles collected at 210 m (Table 3). Both types of estimates show the same trend of a higher phytoplankton contribution at station A3-2. The higher percentage contribution obtained for the gel traps is most likely due to the fact that this approach considers both living and dead phytoplankton cells, and that particles defined as phytodetrital aggregates can contain POC from other origins (e.g., bacteria).

D-AA yields were used to estimate the proportion of POC from bacteria (cells and altered debris of bacteria). However, some extreme and inconsistent values were measured in the present dataset. Extreme values or the disagreement between D-AA yields are probably caused by the fact that yields in POM were calculated by the difference between the chromatographic peaks measured in unfiltered and filtered water samples. In many samples this difference was very small because of the very low POM concentrations, increasing the uncertainty of these yields. Different dynamics between D-

Asx, D-Glx et D-Ala in the studied POM could also explain inconsistent values. This appears to be the case at station A3-2, where D-Glx and D-Ala markers were not always in agreement, despite the increased estimation accuracy and precision due to the higher POC concentrations. For instance, at 70 m, at this site, estimates based on D-Glx and D-Ala were 19 and 5 %, respectively (not shown). Averages suggest that ~ 15 % of the POC originated from bacteria at all depths at station A3-2 (Table 3, Bacteria total). Using the same approach and markers, Kaiser and Benner (2008) estimated that about 25 % of POC was of bacterial origin in oligotrophic areas of the North Pacific and North Atlantic.

An independent approach, based on cell counts of autotrophic (unpublished data) and heterotrophic bacteria (Christaki et al., 2014) and conversion of cell counts to carbon units using $250 \text{ fg C cell}^{-1}$ for autotrophic (Campbell et al., 1994) and $12.4 \text{ fg C cell}^{-1}$ for heterotrophic bacteria (Fukuda et al., 1998), revealed that only 2–3 % of POC are accounted for by bacterial cells as determined by flow cytometry in the surface waters at station A3-2 (Table 3, Bacteria biomass). However, there was a steady increase of this proportion with depth at station A3-2, reaching 10 % at 300 m. When compared to the D-AA estimate of ~ 15 % of POC originating from bacteria, it appears that intact bacterial cells represent a small proportion (< 20 %) of the total bacterial contribution (living + detritus) in the surface water, which is in agreement with previous studies (e.g., White and Howes, 1994; Bourgoin and Tremblay, 2010). However, intact bacterial cells appear to dominate (67 %) the bacterial contribution in the deep waters at the bloom site. This increase in bacterial cell contribution with depth was not observed at the HNLC site where proportions fluctuated between 4 and 7 % of POC. Different trends have been reported in previous studies. As observed here at the bloom site, the proportion of POC as bacterial biomass increased with depth in the St. Lawrence Gulf (Bourgoin and Tremblay, 2010). This trend was caused by a depletion of bacterial POM with depth

Table 3. Summary of the characteristics of POM and DOM in HNLC waters (station R-2) and at the bloom site above the Kerguelen Plateau (station A3-2). Surface refers to 10–90 m depth layer and deep refers to 210–300 m depth. Contributions to POM and DOM are average values given as percent of bulk organic carbon.

Depth	Parameters	POM		DOM	
		HNLC	Bloom	HNLC	Bloom
Surface					
	Phytoplankton				
	diatom cells ^a	5 %	44 %	nd	nd
	total cells ^b	14 %	45 %	nd	nd
	Bacteria				
	cells ^c	5 %	3 %	nd	nd
	total ^d	> 15 % ^g	15 %	30 %	30 %
	Alteration state	high	low	low	med
	Alterations with depth ^e	med	high	low	high
Deep					
	Phytoplankton				
	total cells ^b	2 %	8 %	nd	nd
	phytodetritus ^f	34 %	41 %	nd	nd
	Bacteria				
	cells ^c	7 %	10 %	nd	nd
	total ^d	> 15 % ^g	15 %	25 %	30 %
	Alteration state	high	high	med	high

^a From microscopic counts and biovolume determination (Lasbleiz, 2014).

^b Total cells assuming that phytoplankton cells have 40 times more carbon than chlorophyll *a* (Sathyendranath et al., 2009).

^c From cell counts of autotrophic and heterotrophic bacteria and conversion factors (Christaki et al., 2014).

^d Total bacterial contribution estimated from D-AA yields (see text), for POC it includes living cells and cell fragments or detritus.

^e Alterations promoted by different organic matter inputs or environmental conditions (see text).

^f Estimated based on direct observation of sinking particles defined as phytodetrital aggregates (Laurenceau et al., 2014).

^g Estimates were inconsistent but generally > 15 % (see text).

nd: not determined.

med: medium.

that was more intense than the decrease of bacterial biomass with depth. In contrast, Kawasaki et al. (2011) estimated that bacterial detritus represented a greater proportion of POC (vs. living bacterial biomass) at 100 m (~ 7 %) than at 5 m (~ 4.5 %) in an oligotrophic area of the North Pacific gyre. However, autotrophic bacteria accounted for ~ 13 % of their POC which was about 50 % of the total bacterial contribution (living + detritus). In the present study, autotrophic bacteria (*Synechococcus*) had low cell numbers ($2\text{--}10 \times 10^5 \text{ cells L}^{-1}$; data not shown). It thus appears that a high lability of bacterial detritus and a low proportion of autotrophic bacteria increase the relative proportion of living bacteria versus bacterial detritus in POC in deep waters as in the present study (see below).

A large fraction of the POM remains of unknown origin, especially in the HNLC site. This fraction could be mostly from detritus of various origins including zooplankton. Laurenceau et al. (2014) estimated that zooplankton fecal pellets represented on average $56 \pm 19 \%$ of POC flux below 110 m

at the studied locations. These pellets contain materials from different origins including phytoplankton (Silver and Gowing, 1991).

The reactivity of POM was markedly higher at the bloom site A3-2 compared to the HNLC site, as revealed by a much more rapid decrease of POM and particulate THAA concentrations with depth. In addition, THAA yields in POM (%POC_{AA} and %PON_{AA}), reliable indicators of POM alteration or diagenetic state (Cowie and Hedges, 1994; Tremblay and Benner, 2009), indicated that the POM from the bloom was relatively fresh or unaltered, but was more rapidly degraded during sinking than the POM from the HNLC site (Table 3). These findings could explain the lower efficiency of carbon export at 100 m at the bloom sites A3-2 (5 %) and E-5 (11 %) than in HNLC waters (station R-2, 34 %) (Planchon et al., 2014). Laurenceau et al. (2014) also found that export efficiencies (at 200 m) were negatively correlated to net primary productivity during the same sampling campaign.

In contrast to THAA yields, no consistent trend in mol% D-AA and DI values were observed with depth, probably because more intense degradation of bulk POM is required to be detectable with these markers (Davis et al., 2009; Bourgoïn and Tremblay, 2010). Our data show that a large fraction of particulate AA is rapidly utilized in surface waters of the bloom. The AA that remain in the POM below the surface layer are probably in more recalcitrant structures. These structures may be too recalcitrant to show significant change in AA composition (i.e., mol% D-AA and DI values) with depth. In addition, an apparent mismatch between DI values and the expected trend during degradation has been previously observed in the Southern Ocean and was attributed to the unusual AA composition of diatoms, the dominant source of POM (Ingalls et al., 2003).

4.3 Origin and fate of DOM

Although an important fraction of the primary production is transferred to the DOM pool (e.g., 30–50 % as short lived labile DOM: Pomeroy et al., 2007; Obernosterer et al., 2008; Hansell, 2013), it was not possible to estimate the phytoplankton contribution to bulk DOC with the markers used in the present study. By contrast, the bacterial contribution to DOC was estimated using D-AA yields, indicating that 16.5–35 % of bulk DOC was of bacterial origin with no consistent trend with depth or location. These contributions were near 30 % in surface waters, and similar between Fe-fertilized waters and the HNLC site (Table 3). Similarly, DOC concentrations were not substantially different among sites (i.e., 3–4 μM greater in the bloom). Taken together, these results suggest that DOM production was counterbalanced by bacterial consumption (Christaki et al. 2014) of DOM from bacteria and other sources. The relatively constant bacterial contribution to DOC measured here ($\sim 30\%$) or in the North Atlantic and North Pacific ($\sim 25\%$, Kaiser and Benner, 2008) indicates that bacterial DOM has an average reactivity similar to bulk DOM and contributes to all DOM fractions, from labile to refractory DOM.

This study shows for the first time the contrasting fates of the DOM produced during natural iron fertilization and the DOM from an HNLC region (Table 3). The bloom site A3-2 clearly shows intense DOM alteration within the water column. At station A3-2, all the molecular diagenetic markers (%DOC_{AA}, mol% D-AA, DI values) indicated that surface DOM was less altered than deep water DOM. In contrast, DOM in HNLC surface waters revealed a lower alteration state that did not change with depth (Table 3). The DOM produced during the bloom was thus more rapidly degraded and mineralized, and this pattern reflects well that observed for POM. Mixing of deep and surface water layers at station A3-2 could explain the gradual character of the trends observed in profiles of DOM molecular markers at this location. Despite the fact that a water-column stratification was present during sampling, with a surface mixed layer of 153 m, station

A3-2 is located above the Kerguelen Plateau which is surrounded by deeper areas and thus a more intense or frequent vertical mixing may occur. However, this probable mixing cannot explain why station A3-2 had more altered DOM (and lower dissolved AA) than in the HNLC area even in deep waters.

A surprising finding of the present study is the lower total dissolved AA concentrations and yields in surface water DOM in the bloom sites as compared to HNLC waters. These differences were due to $\sim 30\%$ more dissolved combined AA in HNLC waters. By contrast, free AA had higher concentrations and represented a greater proportion of dissolved AA at the bloom sites (13.5–23.6 % vs. 11.3 % at HNLC) which likely reflects the higher phytoplankton activity and associated DOM release mechanisms (Fuhrman, 1987; Sarmiento et al., 2013).

We propose four possibilities that could explain the lower alteration state of DOM and the higher concentration of dissolved combined AA in less productive sites (i.e., stations R-2 and E-1) compared to the bloom sites (i.e., stations A3-2 and E-5). First, the higher quantities of labile DOM produced at the bloom sites could induce a priming effect (Bianchi, 2011). These large quantities of labile DOM (e.g., free AA) induce a strong microbial response (Christaki et al., 2014) and the energy gained through labile DOM may allow for the degradation of more recalcitrant DOM including dissolved combined AA. This degradation was particularly marked within the water column of the bloom site A3-2. As a consequence, the DOM at the bloom sites appears more altered than the DOM from the HNLC site (Table 3). The higher concentration of more altered DOM in the bloom sites is consistent with the microbial carbon pump concept in which marine microbes transform labile DOM into refractory DOM (Jiao et al., 2011). Important inputs of labile DOM likely stimulate bacterial activity leading to more intense DOM degradation and production of more recalcitrant DOM. The more rapid degradation of the DOM from the bloom is consistent with a recent study in Arctic fjords (Osterholz et al., 2014). Their findings suggest that DOM produced during the spring/summer bloom is degraded in a few weeks by a specialized resident microbial community. Second, another reason for the slower DOM degradation and the accumulation of combined dissolved AA in HNLC waters is an iron limitation for heterotrophic bacteria (Obernosterer et al., 2014). Third, the DOM in HNLC waters can be involved in physicochemical processes (e.g., inter-molecular interactions, partial photooxidation) that protect DOM such as dissolved combined AA from microbial attack (Keil and Kirchman, 1994). Protection mechanisms may be too slow to significantly reduce the microbial degradation rate of the labile DOM produced during the bloom. Finally, a fourth explanation could be that HNLC waters produce DOM molecules, including ones that contain combined AA, which are more recalcitrant to degradation.

The first three explanations are associated with unfavorable conditions for DOM degradation in HNLC areas, namely low quantity of labile or priming DOM, iron limitation, and DOM protection. The last explanation implies that the DOM produced in the HNLC region is intrinsically more recalcitrant than the DOM produced during the bloom. Though the DOM from the HNLC site appears less degraded, the relative lability or recalcitrance of this DOM remains unknown. This notion of substrate quality “may only be valid in the context of the ambient environment” or conditions (Bianchi, 2011). Freshly produced and highly altered or old molecules can persist in unfavorable conditions, but be degraded in another environment.

5 Conclusions

Important differences in POM and DOM origins, compositions, and alterations were observed between the HNLC region and the areas where a phytoplankton bloom was induced by natural iron fertilization in the Southern Ocean. Phytoplankton cells represent an important fraction of POC (~ 45 %) in the surface waters at the bloom sites, while the bacterial total (cells and detritus) contribution was much lower (~ 15 %). Molecular indicators demonstrate that POM originating from the bloom was more rapidly degraded during sinking than the POM from the HNLC site. The fraction of the bloom POM that was transferred to DOM is unknown, but ~ 30 % of the DOM appears to come from bacteria. As for POM, the DOM from the bloom sites was involved in more intense degradation processes. In contrast, the DOM in HNLC waters was less altered and showed no significant alteration throughout the water column. The bloom induced by natural iron fertilization probably leads to favorable conditions (e.g., large quantities of labile OM) supporting high microbial activity (Christaki et al., 2014) and enhanced POM and DOM degradation. The more intense degradation in bloom sites reduces the carbon export efficiency and the net impact of such blooms on the global carbon budget. However, a higher microbial activity appears to also lead to a greater production of more recalcitrant DOM or a more active microbial carbon pump (Jiao et al., 2011).

Acknowledgements. We thank Louise Oriol and Karine Escoubeyrou for their help during sampling and analyses. We also thank our many colleagues who participated in the collection of various data sets, the KEOPS coordinator S. Blain, the chief on board, B. Quéguiner, and the crew aboard the R/V *Marion Dufresne* for their help in the successful completion of the cruise. C. Panagiotopoulos and an anonymous reviewer provided valuable comments that improved the manuscript. This research was supported by the project KEOPS2 (ANR-10-Blanc-614, LEFE INSU, CNES, IPEV) and the Natural Science and Engineering Research Council of Canada (NSERC). L. Tremblay received support from the UPMC during his sabbatical stay at the Laboratoire

d’Océanographie Microbienne. We are grateful to the BIO2MAR platform (<http://bio2mar.obs-banyuls.fr>) for providing access to instrumentation.

Edited by: B. Quéguiner

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